# Mitochondrial dynamics underlying thermal plasticity of cuttlefish (*Sepia officinalis*) hearts

Key words: Temperature sensitivity, cephalopod, evolutionary adaptation, thermal acclimation, proton leak, lactate and octopine dehydrogenase, respiration, systemic and branchial hearts, cardiac fibres

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# 1 ABSTRACT

In the eurythermal cuttlefish Sepia officinalis, performance depends on hearts that ensure 2 3 systemic oxygen supply over a broad range of temperatures. We therefore aimed to identify adjustments in energetic cardiac capacity and underlying mitochondrial function supporting 4 thermal acclimation and adaptation that could be critical for the cuttlefish's competitive 5 success in variable environments. Two genetically distinct cuttlefish populations were 6 acclimated to 11°C, 16°C and 21°C, respectively. Subsequently, skinned and permeabilised 7 heart fibres were used to assess mitochondrial functioning by means of high-resolution 8 9 respirometry and a substrate-inhibitor protocol, followed by measurements of cardiac citrate synthase and cytosolic enzyme activities. Temperate English Channel cuttlefish had lower 10 mitochondrial capacities but larger hearts than subtropical Adriatic cuttlefish. Warm 11 12 acclimation to 21°C decreased mitochondrial complex I activity in Adriatic cuttlefish and 13 increased complex IV activity in English Channel cuttlefish. However, compensation of mitochondrial capacities did not occur during cold acclimation to 11°C. In systemic hearts, 14 thermal sensitivity of mitochondrial substrate oxidation was high for proline and pyruvate but 15 low for succinate. Oxygen efficiency of catabolism rose from 11°C to 21°C via shifts to 16 17 oxygen-conserving oxidation of proline and pyruvate and via reduced relative proton leak. 18 The changes observed for substrate oxidation, mitochondrial complexes, relative proton leak 19 or heart weights improve energetic efficiency and essentially seem to extend tolerance to high temperatures and reduce associated tissue hypoxia. We conclude that cuttlefish sustain cardiac 20 21 performance and thus, systemic oxygen delivery over short and long-term changes of 22 temperature and environmental conditions by multiple adjustments in cellular and 23 mitochondrial energetics.

#### 24 INTRODUCTION

25 The evolution of modern cephalopods was shaped by the rising competition with marine 26 vertebrates and led to innovations that advance high levels of performance. Concomitant 27 increases of energy demands required extant cephalopods to optimise supply and use of oxygen (O'Dor and Webber, 1986; O'Dor and Webber, 1991). As a result, cephalopods 28 29 evolved high concentrations of blood pigment optimised for oxygen transport and a closed 30 circulatory system driven by two branchial and one powerful systemic heart (Schipp, 1987; Wells and Smith, 1987; Wells, 1992; Pörtner and Zielinski, 1998). However, design 31 32 constraints were involved, leading to locomotion by jet propulsion and comparatively low 33 blood oxygen carrying capacities. Optimisation of performance led at least some cephalopod 34 species to operate at their functional limits (O'Dor and Webber, 1986; Pörtner, 2002b). 35 Environmental stressors such as fluctuations in ambient temperature or ambient hypoxia may be particularly challenging for the highly oxygen dependent cephalopods that face high 36 competitive pressure (Rosa and Seibel, 2008). 37

Body functions of animals operate only within a certain thermal range and are set by 38 ambient temperature for ectotherms. According to recent evidence, oxygen supply becomes 39 40 limiting at high temperatures when oxygen demand increases due to limited functional 41 capacities of the circulation and ventilation system to deliver oxygen (Pörtner and Knust, 42 2007; Pörtner and Farrell, 2008). Adaptive adjustments of e.g. ventilatory musculature and 43 heart rate or stroke volume are suitable to compensate for temperature induced impairments of oxygen supply (Wells, 1992; Frederich and Pörtner, 2000). These adjustments, however, 44 45 involve changes in the capacity of mitochondria to provide sufficient aerobic energy to vital tissues like the heart (Pörtner, 2002a). Low temperatures also cause capacity limitations, 46 47 which may involve insufficient release of oxygen to tissues by the blood pigment 48 haemocyanin (Melzner et al., 2007b) and limited mitochondrial energy provision as required 49 to power circulation or ventilation (Pörtner, 2002a).

Ectothermic hearts play a major role in defining limits of aerobic performance and thermal tolerance, as demonstrated by impaired cardiac function close to extreme temperatures or exercise levels (Farrell, 2002; Somero, 2010). In cephalopods, systemic hearts cover increased metabolic demands during either exercise or rising temperatures by a 2-3 fold increase of stroke volume or heartbeat frequency (Wells, 1992), but show limited performance at high critical temperatures, as indicated in *Sepia officinalis*, where blood perfusion fails to increase further beyond 23°C (Melzner et al., 2007a; Melzner et al., 2007b).

Yet, unlike in fishes, cephalopod systemic hearts receive well oxygenated blood and gain 57 support by two accessory hearts, contractile blood vessels and mantle pressure oscillations 58 (Schipp, 1987; Melzner et al., 2007a). Adversely, high blood viscosity, 2-3 fold lower blood 59 oxygen carrying capacities and higher resting metabolic rates compared to haemoglobin 60 bearing fish with a similar lifestyle (Wells, 1992; Pörtner, 1994) demand higher performance of the systemic heart. While cephalopod hearts are able to meet this workload within limits during acute rises of metabolic demand, it is unknown whether they adjust to seasonal (i.e. acclimation) or long term environmental changes over multiple generations (i.e. evolutionary adaptation). Such temperature acclimation or adaptation has been shown for fish hearts. Here, changes comprise increases in heart size (Goolish, 1987; Kent et al., 1988), increased mitochondrial content (Kleckner and Sidell, 1985; Johnston and Harrison, 1987; Kolok, 1991), shifts from carbohydrate to fatty acid oxidation (Sephton and Driedzic, 1991; Sidell et 68 al., 1995) or enhanced activities of enzymes essential in aerobic metabolism (Crockett and 70 Sidell, 1990; Podrabsky et al., 2000), following cold-exposure. Yet, evidence for such temperature related adjustments is still lacking for cephalopod hearts.

The eurythermal common cuttlefish *Sepia officinalis* (Linnaeus 1758) lives on the continental shelf from the cold-temperate eastern North Atlantic to warm-subtropical Mediterranean and Atlantic waters off the Senegalese coast and follows a bottom dwelling, migratory life style on rocky to sandy grounds down to 200 m depth. Cuttlefish are rather sluggish, grow fast (up to two kg) and spawn after one or two years in shallow waters and die thereafter (Jereb and Roper, 2005). In this study, we compared a cuttlefish population from the English Channel living between 9°C and 17.5°C with a genetically distinct Mediterranean population (Wolfram et al., 2006) facing a range from 10°C to 25°C, respectively, between winter and summer (Boucaud-Camou and Boismery, 1991; Artegiani et al., 1997; Wang et al., 2003).

In this study, we aimed to understand how temperature changes affect the heart of the common cuttlefish *Sepia officinalis* by (1) exploring cardiac adjustments following long term genetic isolation in a temperate and subtropical habitat as well as (2) following thermal acclimation and (3) by investigating the thermal sensitivity of heart mitochondria.

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#### 86 MATERIAL AND METHODS

#### 87 Experimental animals

88 Wild laid eggs of the European cuttlefish (Sepia officinalis) were collected from the temperate Oosterschelde lagoon, Netherlands (3°56'E, 51°35'N) and the subtropical Venetian 89 lagoon close to Chioggia, Adria, Italy (12°18'E, 45°13'N). Populations from these localities 90 91 form geographically distanced and genetically distinct clades without evidence of on-going genetic exchange (Wolfram et al., 2006; Perez-Losada et al., 2007). Thus, inter-population 92 93 differences will most likely reflect evolutionary adaptation or genetic drift. About 300 eggs from five to six different egg masses (50 to 60 eggs per mass) were collected from a 94 95 frequently visited spawning ground (>100 individuals per day) in the Oosterschelde lagoon and 200 eggs from more than ten egg masses were collected by scuba diving and snorkelling 96 97 in the Venetian lagoon in 2008 and 2009. After transport to our institute individuals from each 98 population were hatched and raised in separate tanks, connected to a re-circulating 99 aquaculture system at constant temperature (mean  $\pm$  s.d., 15.9°C  $\pm$  0.1) at suitable stocking densities (Hanley et al., 1998) under constant 12 h dark:12 h light cycle. Hatchlings were fed 100 101 daily with live shrimps (Neomysis integer, Palaemonetes varians and Crangon crangon) and 102 after reaching a bigger size, exclusively with frozen brown shrimp (Crangon crangon). Water 103 quality parameters were monitored weekly and kept at levels appropriate for cuttlefish culture (Hanley et al., 1998) (mean  $\pm$  s.d.,  $O_2 = 10.07 \pm 0.43$  mg l<sup>-1</sup>, pH = 8.05  $\pm$  0.02, salinity = 32.6 104  $psu \pm 0.5$ ,  $NH_4^+ < 0.2 \text{ mg l}^{-1}$ ,  $NO_2^- < 0.2 \text{ mg l}^{-1}$ ,  $NO_3^- < 80 \text{ mg l}^{-1}$ ) by means of water 105 treatment (protein skimmers, mechanical and biological filters, UV sterilisation) or water 106 107 replacement.

After animal size had reached 30 - 40 g, each population was divided into three groups 108 109 that were acclimated to 11°C, 16°C and 21°C from 8 to 22 weeks. These acclimation 110 temperatures were selected to expose European cuttlefish to their seasonal and depth 111 dependent temperature range, 10°C to 17.5°C for the English Channel population and 10°C to 25°C for the Adriatic Sea population. Each group was kept in separate tanks connected to 112 independent re-circulating systems for each temperature. Only animals showing normal 113 114 behaviour without signs of illness (skin infections mostly due to jetting against tank walls) 115 were selected for experiments. Although Adriatic cuttlefish weighed significantly more in the 21°C group (Supplementary table S2) at the end of the acclimation period (possibly due to 116

- 117 enhanced growth rates), correlation analysis did not show clear evidence that weight
- 118 differences obscured our data.

#### 119 Mitochondrial respiration

# 120 Dissection and muscle fibre preparation

121 Animals were anesthetized in 3% ethanol until non-responsive before culling. Subsequently, total weight, mantle length, total length and sex were recorded. After opening 122 123 the mantle cavity, samples of blood, gills and ink were taken and frozen for further experiments. The three hearts were excised starting with the branchial hearts and placed 124 125 immediately into 1 ml ice-cold biopsy buffer (modified after (Kuznetsov et al., 2008)) to preserve mitochondrial function. The biopsy buffer contained (in mmol 1<sup>-1</sup>, 2.77 CaK<sub>2</sub>EGTA, 126 127 7.23 EGTA, 14.46 KOH, 5.77 Na<sub>2</sub>ATP, 6.56 MgCl<sub>2</sub> 20 taurine, 20 imidazole, 0.5 dithiothreitol (DTT), 50 MES, 588 sucrose, 252 glycine, pH 7.4 at 26°C, 1000 mosmol l<sup>-1</sup>). 128 129 After removal of non-cardiac tissues and weighing of all three hearts, approximately 50 130 mg of systemic heart tissue were placed into a drop of biopsy buffer on a Petri dish on ice and 131 then coarsely torn apart and dissected into small fibre bundles using two tweezers. The 132 remainder of heart tissue and the branchial hearts were snap-frozen in liquid nitrogen and 133 stored at -80°C for subsequent enzyme assays. Fibre bundles were immediately transferred to 1 ml biopsy buffer in a 12 well multiwell culture plate and permeabilised with 50 µg ml<sup>-1</sup> 134 135 saponin (Note: preliminary testing confirmed appropriate saponin concentration) by gentle mixing (115 rpm) on ice for 30 min. The fibres were then removed and washed three times for 136 137 10 min in 2 ml ice-cold mitochondrial respiration medium (in mmol 1<sup>-1</sup>, 50 HEPES, 25 KH<sub>2</sub>PO<sub>4</sub>, 0.5 EGTA, 50 KCl, 50 NaCl, 10 MgCl<sub>2</sub>, 20 taurine, 50 lactobionate, 350 sucrose, 138 150 glycine, 1 g l<sup>-1</sup> freshly added fatty acid free BSA, pH 7.4 at 22°C, 1000 mosmol l<sup>-1</sup>, 139 140 modified after (Mommsen and Hochachka, 1981; Agnisola et al., 1991; Kuznetsov et al., 141 2008)) and stored in ice-cold respiration medium until use (protocol modified after (Saks et 142 al., 1998)).

# 143 Measurements of mitochondrial respiration

Before each assay, fibres were blotted dry on chilled Whatman® paper, divided into two 2-6 mg bundles and transferred to 2 ml duplicate chambers of an Oxygraph-2k respirometer (Oroboros Instruments, Innsbruck, Austria) containing air saturated respiration medium at the experimental temperature. Mitochondrial respiration was then measured online as background corrected weight specific oxygen consumption rate (pmol  $O_2$  s<sup>-1</sup> mg<sup>-1</sup>, i.e. negative time 150 Innsbruck, Austria). Assays were performed consecutively at 11°C, 16°C and 21°C in

151 randomised order. Chambers were washed with 96% ethanol and Milli-Q water to thoroughly

152 remove inhibitors and substrates after each assay. To protect fibres from extreme oxygen

153 levels, oxygen concentrations in chambers were kept between 80 nmol  $O_2$  ml<sup>-1</sup> (maximum

154 stimulated respiration remained stable down to 80 nmol  $O_2 ml^{-1}$ ) and air saturation (270-340

155  $\text{nmol } O_2 \text{ ml}^{-1}$ ) by re-oxygenation with pure oxygen gas.

To assess mitochondrial function, respirometry on permeabilised skinned heart fibres was 156 157 combined with a substrate-inhibitor protocol as follows. Mitochondria were first fuelled successively and in excess with the amino acid proline (5 mmol  $1^{-1}$ ), ADP (2.5 mmol  $1^{-1}$ ) as 158 well as pyruvate (5 mmol  $l^{-1}$ ) and succinate (10 mmol  $l^{-1}$ ) in order to reach maximum coupled 159 oxidative phosphorylation (i.e. state 3 respiration tied to ATP production). The choice of 160 161 substrates corresponds to own tests and to previous studies that showed high rates of 162 oxidation for proline, pyruvate and succinate in squid hearts (Ballantyne et al., 1981; 163 Mommsen and Hochachka, 1981). In situ, cephalopod hearts derive pyruvate most evidently 164 from blood glucose or oxidised octopine and proline from high intracellular stores (> 12 µmol g<sup>-1</sup>). Succinate can originate from both pyruvate derivatives and amino acids (e.g. ornithine, 165 arginine, glutamate and proline) with the latter feeding into the Krebs cycle at the level of 166 167 alpha-ketoglutarate (Ballantyne et al., 1981; Mommsen and Hochachka, 1981; Hochachka 168 and Fields, 1982).

169 Integrity of mitochondrial membranes was tested by the addition of cytochrome c (0.01 mmol l<sup>-1</sup>) that would stimulate respiration in case of damaged outer membranes. Proline 170 171 stimulated respiration without ADP (state 2) provided an estimate of mitochondrial proton 172 leak (Iftikar et al., 2010). Uncoupling of respiration from ATP production by 173 carbonylcvanide-p-(trifluoromethyl) phenylhydrazone (FCCP, up to 2.5 umol 1<sup>-1</sup>) denoted maximum capacity of the electron transport chain. The loss of activity with rotenone (2.5 174 umol l<sup>-1</sup>) indicated NADH dehydrogenase (complex I) activity. The loss of activity with 175 antimycin A (2.5  $\mu$ mol l<sup>-1</sup>) quantified the non-mitochondrial background respiration. 176 177 Cytochrome c oxidase (complex IV) activity was tested using the redox pair ascorbate (2 mmol 1<sup>-1</sup>) and N,N,N',N'- tetramethyl-p-phenylenediamine dihydrochloride (TMPD, 0.5 178 mmol l<sup>-1</sup>, see list in supplementary table S1). Background auto-oxidation of the redox pair 179 180 was determined for all experimental temperatures and subtracted from final results. Suitability 181 and concentrations of substrates were tested beforehand. All chemicals were purchased from

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Sigma-Aldrich (Schnelldorf, Germany)

#### 183 Enzyme assays

#### 184 Protein extraction and sample preparation

185 To extract the protein, frozen heart tissue was ground by hand in a mortar filled with liquid nitrogen. The frozen tissue powder was then weighed and suspended in 10 volumes 186 (w:v) of ice cold extraction buffer (50 mmol 1<sup>-1</sup> Tris-HCl (pH 7.4 at 16°C), 1 mmol 1<sup>-1</sup> EDTA, 187 188 0.1% Triton X-100) and sonicated for 90 s at 0°C in a Branson Sonifier 450 (output control 8, 189 Duty cycle 50%). Following two centrifugations for 10 min at 6000g and 4°C, supernatants 190 were used for protein and enzyme assays. Pellets remaining after centrifugation were 191 resuspended and re-extracted in ice cold extraction buffer to test for residual protein and 192 citrate synthase activity.

For enzyme assays, samples were diluted 1:10 (v:v) with 75 mmol l<sup>-1</sup> Tris-HCl (pH 8.1 at 16°C) and equilibrated to approximately 16°C using a temperature controlled metal block connected to a thermostat (Haake C25, Thermo Scientific, Karlsruhe, Germany) prior to each measurement. Absorbance was measured in triplicates using 96 well F-bottom microplates (Nunc GmbH & Co. KG, Wiesbaden, Germany), a multiplate reader (Fluostar Galaxy, BMG Labtechnologies, Offenburg, Germany) and analyzing software (FLUO 32, version 4.31 R5).

# 199 Enzyme activities and protein content

Citrate synthase (EC 4.1.3.7) activity was determined after Sidell et al. (1987) as the increase of absorbance at 412 nm by means of DTNB (5,5'-dithio-bis-(2-nitrobenzoic acid)). First, background deacylase activity was measured with 20  $\mu$ l of diluted sample supernatant and 160  $\mu$ l reaction mixture (in mmol l<sup>-1</sup>, 75 Tris-HCl, 0.25 DTNB, 0.4 acetyl~CoA, pH 8.1) followed by the addition of 20  $\mu$ l 0.5 mmol l<sup>-1</sup> oxaloacetate to start the reaction. As DTNB reacts with sulphur groups (SH), dithiothreitol (DTT) was used as standard for the calculation of SH group turn-over.

Activities of octopine dehydrogenase (ODH, EC 1.5.1.11) and lactate dehydrogenase 207 208 (LDH, EC 1.1.1.27) indicate capacities to form or recycle anaerobic end products in 209 cephalopods (Grieshaber and Gäde, 1976; Storey, 1977; Storey and Storey, 1979). ODH and 210 LDH activities were determined after Storey (1977) and Driedzic et al. (1990) as the decrease of absorbance at 340 nm. First, background activity was measured by combining 20 µl of 211 diluted sample supernatant and 160 µl reaction mixture (in mmol l<sup>-1</sup>, 100 Tris-HCl (pH 7.0), 212 0.45 NADH, 1 KCN, 10 arginine were added for quantifying the ODH reaction) followed by 213 the addition of 20  $\mu$ l 4 mmol l<sup>-1</sup> pyruvate to start the reaction. Final ODH activity was 214

determined by subtracting the LDH activity, which was measured simultaneously. NADHwas used as a standard.

Enzyme activities were standardised to protein content determined after Bradford (1976). Sample supernatant and pellet homogenate were diluted 1:10 (v:v) with 0.9% NaCl. 5  $\mu$ l were transferred to 250  $\mu$ l Bradford dye reagent (0.1 mg ml<sup>-1</sup> Coomassie Brilliant Blue G-250, 5% ethanol, 8.5% H<sub>3</sub>PO<sub>4</sub>). After mixing and incubation for 10 min at room temperature (21°C), absorbance was recorded at 595 nm. Bovine albumin serum (BSA) was used as protein standard.

#### 223 Statistical analysis

224 Statistical analysis was performed to spot significant differences (P < 0.05) using SPSS 225 (SPSS© Inc., version 14.0.1) by employing tests as followed: analysis of variance (ANOVA) or non-parametric tests; additional post-hoc Tukey or Hochberg's GT2 test for equal or 226 227 unequal sample sizes respectively to compare acclimation treatments or assay temperatures; 228 Pearson's correlation analysis to test for allometric effects; Kolmogorov-Smirnov and 229 Levene's test assessed normality and homogeneity of variances, respectively. Data were expressed as means and the range of their 95% confidence interval if not stated otherwise. 230 231 Outliers were detected using Nalimov's test (P < 0.01) and excluded if justified. The 232 temperature coefficient Q<sub>10</sub> was calculated as follows:

$$Q_{10} = \left(\frac{R_2}{R_1}\right)^{\frac{10}{(T_2 - T_1)}}$$

233 *R* denotes the respiratory rate at a higher  $(T_2)$  or lower temperature  $(T_1)$ . Mitochondrial 234 respiration was expressed per mg wet weight of blotted heart fibres.

#### 235 **RESULTS**

#### 236 Evolutionary Adaptation

237 Mitochondrial respiration differed clearly between the two genetically distinct cuttlefish populations. On average, cardiac mitochondria, fully fuelled by substrates and ADP (state 3), 238 showed significantly higher respiration rates in Adriatic cuttlefish with 41.1 pmol O<sub>2</sub> s<sup>-1</sup> mg<sup>-1</sup> 239 (95% CI range from 37.3 to 44.9 pmol  $O_2$  s<sup>-1</sup> mg<sup>-1</sup>) compared to only 32.9 pmol  $O_2$  s<sup>-1</sup> mg<sup>-1</sup> 240 241 (30.0-35.9) as seen in English Channel cuttlefish (ANOVA, F(1, 128) = 11.92, P < 0.01, 242 calculations based on entire pooled data set). These 20% higher cardiac aerobic capacities in 243 cuttlefish from warmer waters were most pronounced at lower assay temperatures, and were 244 mainly due to a higher contribution of pyruvate to overall respiration (average contribution of 15.2 pmol  $O_2$  s<sup>-1</sup> mg<sup>-1</sup>, (13.1-17.3)) for Adriatic cuttlefish compared to 9.0 pmol  $O_2$  s<sup>-1</sup> mg<sup>-1</sup>, 245 246 (7.8-10.2) for English Channel cuttlefish, Figure 1). Consequently, fractions of pyruvate dependent respiration constituted 35.4% (32.8-38.0) in systemic hearts of Adriatic Sea but 247 248 only 26.2% (24.1-28.3) in English Channel cuttlefish. Conversely, proline dependent fractions 249 were higher (45.2% (42.5-47.8)) in English Channel- than in Adriatic Sea cuttlefish (35.2% 250 (32.4-38.0)). Succinate dependent fractions were similar in the two populations. 251 While body weights did not differ between populations (ANOVA, F(1, 64) = 2.36, P =252 0.13). English Channel cuttlefish contained 43% heavier systemic and 38% heavier branchial 253 hearts than Adriatic cuttlefish (systemic hearts: Kruskal-Wallis, H(1) = 13.69, P < 0.01; 254 branchial hearts: Kruskal-Wallis, H(1) = 10.22, P < 0.01, for raw data see supplementary 255 table S2). Relative weights of systemic hearts constituted 0.083% (0.078-0.088) in English

256 Channel and 0.059% (0.056-0.062) in Adriatic cuttlefish. Similarly, relative weights of

branchial hearts constituted 0.047% (0.044-0.050) in English Channel and 0.039% (0.036-

258 0.042) in Adriatic cuttlefish. As a result, respiration calculated for the whole systemic heart

259 (state 3 respiration x heart weight) did not differ between populations (ANOVA, F(1, 41) = 0.03, P = 0.86), due to the larger hearts in English Channel cuttlefish.

Further, systemic hearts of Adriatic cuttlefish showed partly higher protein specific enzyme activities compared to English Channel cuttlefish, which was most apparent for octopine dehydrogenase (Table 1). Enzyme activities of branchial hearts were similar between populations except for enhanced lactate dehydrogenase activities in English Channel cuttlefish acclimated to 11°C and 16°C (Table 1).

#### 266 Thermal acclimation

Acclimation of cuttlefish to 11°C, 16°C and 21°C did not affect overall maximum state 3 267 268 respiration in the English Channel (ANOVA, F(2, 68) = 0.04, P = 0.96) and in the Adriatic Sea populations (Kruskal-Wallis, H(2) = 0.94, P = 0.63) but caused minor shifts in substrate 269 270 dependent fractions (Figure 1) and slight changes in cardiac enzyme activities of English 271 Channel cuttlefish, shown by enhanced citrate synthase activities at 11°C and lower lactate 272 dehydrogenase activities at 21°C compared to animals acclimated to 16°C (Table 1). Also, 273 relative heart weights did not change with thermal acclimation for systemic- (ANOVA, 274 English Channel F (2, 29) = 0.50, P = 0.61, Adriatic Sea F (2, 30) = 0.95, P = 0.40) and 275 branchial hearts (ANOVA, English Channel F(2, 29) = 0.46, P = 0.64, Adriatic Sea F(2, 30)276 = 3.0, P = 0.07).

277 Thermal acclimation affected the activity of mitochondrial complexes in systemic heart 278 fibres of cuttlefish. In this regard, Adriatic cuttlefish acclimated to 21°C showed between 10-19% lower complex I activity compared to cuttlefish acclimated to 11°C and 16°C whereas 279 280 complex I activity in English Channel cuttlefish did not respond to thermal acclimation 281 (Figure 2A). In contrast, systemic hearts of English Channel cuttlefish contained 282 mitochondria that displayed increased complex IV activity following acclimation to 21°C, at assay temperatures of 11°C and 21°C (Figure 2B). In Adriatic cuttlefish hearts, however, 283 284 complex IV activity remained unaffected by thermal acclimation but showed between 8-30% 285 higher enzyme activity at lower acclimation temperatures compared to English Channel 286 cuttlefish (Figure 2B).

#### 287 Thermal sensitivity of cardiac mitochondria

288 In both cuttlefish populations cardiac mitochondria responded similarly to acute 289 temperature changes ranging from 11°C to 21°C. State 3 respiration increased from 20.4 (18.1-22.8) to 46.4 pmol O<sub>2</sub> s<sup>-1</sup> mg<sup>-1</sup> (42.9-49.9) in English Channel and from 27.8 (24.8-30.8) 290 to 51.2 pmol  $O_2$  s<sup>-1</sup> mg<sup>-1</sup> (45.3-57.1) in Adriatic cuttlefish, with average  $O_{10}$  of 2.3 in English 291 Channel and 2.0 in Adriatic animals (Figure 3). Respiration resolved for substrates revealed a 292 293 high thermal sensitivity of proline- and pyruvate stimulated respiration but a low thermal 294 sensitivity of succinate stimulated respiration (Figure 3). This caused succinate to contribute 295 most to overall state 3 respiration at 11°C assay temperature (English Channel 40.4% (37.8-296 43.0); Adriatic Sea 43.4% (39.6-47.2)) but to be a minor substrate at 21°C (English Channel 297 18.7% (15.9-21.5); Adriatic Sea 19.0% (16.0-22.0)). Conversely, proline and pyruvate

- prevailed as oxidative substrates at 21°C assay temperature, whereas their fractions declined
  upon cooling towards 11°C (Figure 1).
- 300 Mitochondrial complexes displayed different temperature dependencies upon acute
- 301 exposure. Similar to succinate stimulated respiration (which indicates activity of complex II)
- 302 the activity of complex IV (indicated by respiration fuelled with ascorbate and TMPD)
- 303 showed low thermal sensitivity (Figure 3).
- 304 Interestingly, although absolute leak rates increased with warmer assay temperatures up to 305 21°C, mitochondrial proton leak relative to ADP stimulated respiration decreased in systemic
- 306 hearts of English Channel cuttlefish and by trend in Adriatic cuttlefish upon warming (Figure
- 307 4A, B). Outer membrane integrity, tested by cytochrome c addition, decreased with increasing
- 308 assay temperatures in both cuttlefish populations (Figure 5). Further, English Channel
- 309 cuttlefish showed more defective outer membranes and higher relative proton leak at  $11^{\circ}C$
- 310 assay temperature compared to Adriatic cuttlefish (Figure 5).

#### 311 **DISCUSSION**

#### 312 **Evolutionary adaptation**

313 There is strong evidence for evolutionary changes of heart function between the genetically distinct temperate and subtropical cuttlefish populations. Systemic hearts of 314 315 temperate (English Channel) cuttlefish contained lower aerobic and partly lower anaerobic 316 capacities (Figure 1 & Table 1) that may reduce their cardiac output and hence lower their 317 ability to increase heart rates during exercise or elevated temperatures. Whether mitochondrial 318 density parameters, enzyme numbers or specific enzyme activities accounted for the 319 differential aerobic capacities remains to be resolved. Nevertheless, similar total heart 320 respiration between populations showed that English Channel cuttlefish compensate for 321 reduced respiratory capacities by larger hearts. Compensatory increases of heart weight are 322 also common in teleost fish living at cooler temperatures (Foster et al., 1993; Driedzic et al., 323 1996). Larger hearts support pumping of larger blood volumes per stroke, thereby, they may 324 compensate for low energetic capacities but also for rising blood viscosities at colder 325 temperatures (Goolish, 1987; Driedzic et al., 1996). Thus, cuttlefish heart function in the cold 326 may rather be sustained through an increase in organ size, than through an increase of cellular 327 energetic capacities.

328 Furthermore, the findings of predominant oxidation of proline in systemic hearts of 329 temperate cuttlefish but enhanced pyruvate oxidation in subtropical cuttlefish are well in line 330 with findings for cold adapted fish that show suppressed carbohydrate metabolism but 331 enhanced lipid oxidation (Crockett and Sidell, 1990; Sidell et al., 1995). Fuels like lipids and 332 proline are less oxygen efficient than carbohydrates, which are generally favoured in 333 cephalopods (Hochachka, 1994). Carbohydrate oxidation reduces oxygen consumption of the 334 heart and threats from tissue hypoxia due to higher ATP yields per molecule of oxygen 335 consumed (Higgins et al., 1980; Kahles et al., 1982; Hochachka, 1994). In cephalopod hearts, 336 a mole of proline requires half a mole of dioxygen to oxidize it to glutamate before entering 337 the Krebs cycle, thus decreasing its oxygen efficiency (Mommsen and Hochachka, 1981). 338 Therefore, impaired oxygen supply at higher temperatures or environmental hypoxia, which 339 cuttlefish face often in northern Adriatic lagoons (Diaz, 2001; Sorokin and Dallocchio, 2008), 340 may favour carbohydrates and thus, pyruvate as the more oxygen efficient substrate.

#### 341 Thermal acclimation

#### 342 Substrate oxidation

In cuttlefish hearts, thermal acclimation did not affect aerobic capacities (i.e. state 3 343 344 respiration, Figure 1) and caused only minor changes of citrate synthase and lactate dehydrogenase activities in temperate cuttlefish (Table 1). At first glance, this contrasts with 345 346 the thermal compensation found for English Channel cuttlefish. Here, routine metabolic rates 347 of cuttlefish acclimated to 20°C fell below routine metabolic rates of individuals acclimated 348 to 15°C, once above 20°C experimental temperature, thereby supporting an upward shift of 349 limiting temperatures beyond 23°C (Melzner, 2005). This pattern of thermal acclimation may 350 well be explained by a suppression of oxygen consumption rates in organs other than the 351 hearts (e.g. hepatopancreas, mantle). Such a one-sided reduction of oxygen consumption in 352 some tissues but concomitant maintenance of cardiac capacities may then free aerobic scope 353 necessary to shift thermal tolerance upwards. On the other hand, in parallel to cardiac 354 capacities, routine metabolic rates decline by 2-3.5 times from  $20^{\circ}$ C to  $8^{\circ}$ C to equally low 355 levels irrespective of the acclimation mode (Melzner, 2005). As a result, cardiac as well as 356 whole animal energetic capacities decline with decreasing temperatures and thus match a 357 reduction of energy turnover during the cold season, marked by reduced growth rates of 358 cuttlefish during winter (Le Goff and Daguzan, 1991).

#### 359 Mitochondrial complexes

360 Effects of warm acclimation on mitochondrial complexes I and IV in systemic hearts of 361 cuttlefish suggest modifications that delay heat-induced tissue hypoxia and formation of 362 reactive oxygen species (ROS). Due to constraints on their systemic (i.e. low blood oxygen carrying capacity) and intracellular oxygen transport system (e.g. lack of intracellular oxygen 363 364 delivering protein like myoglobin, (Hochachka, 1994)), cephalopods are chronically prone to facing critically low intracellular oxygen concentrations, particularly when high temperatures 365 366 induce tissue hypoxia (Pörtner, 2001). Gnaiger et al. (1998) argued that an excess capacity of 367 complex IV sustains high affinities for oxygen in mitochondria. Therefore, the increase of 368 complex IV capacity in systemic hearts of English Channel cuttlefish following warm acclimation to 21°C (Figure 2B) likely enhanced oxygen affinity, thus supporting oxygen 369 370 diffusion to mitochondria at high and hypoxia inducing temperatures. This response is in 371 agreement with fishes that are capable to increase cardiac complex IV activity following 372 warm-acclimation as seen in the highly hypoxia tolerant carp (Cai and Adelman, 1990) but 373 also in cod (Foster et al., 1993) or in the liver of Antarctic eelpout (Windisch et al., 2011).

374 ROS formation in response to heat stress and environmental hypoxia is common among 375 marine ectotherms (Abele et al., 2007), and may occur more often in the warm and often 376 hypoxic lagoons of the northern Adriatic Sea during summer (Diaz, 2001; Sorokin et al., 377 2002) where cuttlefish are common (Rossetto, 2001). Complex I is one of the major sites of 378 ROS production and has been related to cardiac failure (Ide et al., 1999; Sorescu and 379 Griendling, 2002), therefore, a suppression of complex I capacity following warm-380 acclimation in Adriatic cuttlefish (Figure 2A) may reduce heat and hypoxia related ROS 381 formation to preserve mitochondrial function. As a corollary, changed activities of 382 mitochondrial complexes following warm acclimation may serve cuttlefish to reduce 383 temperature induced hypoxia or harmful oxygen stress in cardiac tissues.

# 384 Thermal sensitivity of cardiac mitochondria

# 385 Substrate oxidation

386 During summer, cuttlefish pass a steep thermocline during their daily vertical migration 387 with possible temperature changes of up to 2°C per m (e.g. Adriatic Sea: 23°C to 14°C down 388 to 80 m; English Channel: 19°C to 11°C down to 40 m) (Artegiani et al., 1997; Sharples et al., 389 2001). Our findings demonstrate that noticeable increases of mitochondrial respiration (state 390 3) from 11°C to 21°C support systemic hearts of cuttlefish to operate aerobically over this 391 range of temperatures. Interestingly, various substrates contributed differently to this thermal 392 response (Figure 3). The substrate dependent effect on thermal sensitivity of oxidative 393 pathways may allow mitochondria to produce aerobic energy over a broader range of 394 temperatures. While less temperature sensitive pathways (e.g. that augment succinate) 395 attenuate a rapid decline of mitochondrial respiration (i.e. ATP provision) at low 396 temperatures, thermally more sensitive pathways (e.g. fuelled by pyruvate or proline) 397 complement less responsive pathways at higher temperatures to match enhanced ATP 398 demands of the heart. Although pyruvate and proline oxidation declines at low temperatures, 399 in situ supply of succinate may be sustained by amino acids such as glutamate, ornithine or 400 arginine that are well oxidized and readily available from the blood or intracellular stores 401 (Ballantyne et al., 1981; Mommsen and Hochachka, 1981; Hochachka and Fields, 1982; 402 Mommsen et al., 1983). Observations in fish hearts support this pattern, as fatty acid 403 oxidation has mostly lower or at least different thermal sensitivities than carbohydrate 404 oxidation, which kicks in in the warmth (Sephton et al., 1990; Sephton and Driedzic, 1991). 405 The predominance of pyruvate and proline oxidation at higher temperatures (Figure 1) 406 improves oxygen efficient production of ATP and likely reduces the threat of tissue hypoxia.

407 After entering mitochondria, proline is oxidized to glutamate and then enters the Krebs cycle 408 via alpha-ketoglutarate and yields 4-5 NADH and 1 FADH<sub>2</sub> (i.e. assuming no diffusive loss of 409 products), similar to pyruvate that yields 4 NADH and 1 FADH<sub>2</sub> during its oxidation via the 410 citric acid cycle (Storey and Storey, 1983). As a consequence, proline and pyruvate feed about 411 80% of their electrons into complex I and thus produce more ATP per mole of oxygen 412 consumed. In contrast, succinate yields 1 NADH and 1 FADH<sub>2</sub> and thus, diverts only 50% of 413 its electrons to complex I when fully catabolized to oxaloacetate. Oxygen efficient ATP 414 production may be advantageous at high temperatures when oxygen becomes limiting 415 ((Pörtner, 2010) for recent review) and is well in line with the general trend of modifications 416 in substrate use by mitochondria that enhance oxygen efficiency in cephalopods (Hochachka, 417 1994).

# 418 Membrane leakiness and integrity

419 Surprisingly, in cuttlefish hearts, relative proton leak decreased and outer membrane 420 integrity increased (i.e. shown by decreased stimulation of respiration by cytochrome c) up to 421 a temperature close to the whole animal critical temperature of 23°C ((Melzner et al., 2007b), 422 Figure 4, 5). In contrast to the present observation, relative proton leak increases with 423 temperature in mitochondria isolated from hearts, red muscle and liver of fish (Hardewig et al., 1999; Fangue et al., 2009; Hilton et al., 2010), gills of Antarctic bivalves (Pörtner et al., 424 1999) or the body wall of a lugworm (Sommer and Pörtner, 2002). Our finding thus contrasts 425 426 with the general view that higher temperatures increase membrane fluidity and thus proton 427 leak (Hazel, 1995; Pörtner, 2001). This could be explained by a decrease of mitochondrial 428 membrane potential via increased ATP synthase activity relative to electron flux through the 429 electron transport system, which would induce reduced proton gradients over the membrane 430 leading to reduced proton leak (Nicholls, 2004). Alternatively, structural changes of mitochondrial membranes that reduce membrane permeability may have occurred, as 431 432 indicated by reduced respiratory stimulation by cytochrome c (Figure 5), which passes the 433 outer membrane by diffusion only (Gellerich et al., 2000). Such evident changes of the outer 434 membrane permeability likely affected the inner mitochondrial membrane too and hence 435 supported the observed decline of relative proton leak with rising temperatures. Even though 436 the underlying mechanisms remain unclear, decreasing relative proton leak with acutely rising 437 temperature aids cuttlefish hearts to be more oxygen efficient and thereby shift temperatures that entail oxygen deficiency to higher tolerated values. These functional characteristics may 438 439 be adaptive in supporting eurythermy of this species.

Interestingly, temperate English Channel cuttlefish mitochondria displayed lower
respiratory capacities and contained more permeable and therefore less efficient membranes at
11°C assay temperature than the subtropical Adriatic cuttlefish (Figure 4, 5). Their
mitochondria operate thus less efficient and more "costly" at cool temperatures. Conversely, a
higher "futile cycling" through proton leak would make them more responsive to sudden
increases in workloads. This may be needed less in warm acclimated hearts.

# 446 Conclusion

Our study about cuttlefish heart fibres and their mitochondria revealed inherent potential 447 448 to cope with thermal challenges faced during an individual's lifetime, but also genetic plasticity between populations relevant for adaptation to long-term environmental change. 449 450 Most modifications improve cardiac efficiency and extend tolerance to high temperatures and 451 associated hypoxemic conditions. Cold compensation on evolutionary time scales occurs via a 452 shift in substrate and enhanced proton leak as a sign of enhanced futile cycling in 453 mitochondrial metabolism. Lack of cold compensation by adjusting mitochondrial or enzyme 454 capacities during cold acclimation, on the other hand, suggest decreased cardiac energetic 455 capacities during the cold season. Overall, we conclude that the observed flexibility of cardiac 456 function, based on specific cephalopod-type characteristics, assures cardiac power output and 457 systemic oxygen delivery at various temperatures. It thus conforms to the capacity of Sepia 458 officinalis to tolerate a broad range of temperatures and supports their ability to contend with 459 fishes - their prime competitors - in rapidly changing environments.

460	LIST OF SYM	BOLS AND ABBREVIATIONS
461	ATP	Adenosin-5'-triphosphate
462	BSA	Bovine Serum Albumin
463	Complex I	NADH dehydrogenase
464	Complex IV	Cytochrome c oxidase
465	DTNB	5,5'-dithio-bis-(2-nitro-benzoic acid)
466	DTT	Dithiothreitol
467	EDTA	Ethylenediaminetetraacetic acid
468	EGTA	Ethylene glycol tetraacetic acid
469	HEPES	N-(2-Hydroxyethyl)piperazine-N'-(2-ethanesulfonic acid
470	LDH	Lactate dehydrogenase
471	ODH	Octopine dehydrogenase
472	Q <sub>10</sub>	Temperature coefficient
473	ROS	Reactive oxygen species
474	TMPD	N,N,N',N'-tetramethyl-p-phenylenediamine dihydrochloride

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#### 486 FIGURES LEGENDS

Figure 1: Maximum state 3 respiration (pmol  $O_2 \text{ s}^{-1} \text{ mg}^{-1}$ ) resolved for each substrate, after addition of ADP, at 11°C, 16°C and 21°C assay temperature in comparison between acclimation treatments (*N* = 5-11) as well as between cuttlefish from (A) the English Channel and (B) Adriatic Sea. Significant differences (*P* < 0.05) between acclimation groups are marked by asterisks and for total state 3 respiration between populations by roofs. Values are means ± 95% C.I..

Figure 2: Effects of thermal acclimation on mitochondrial complex I / NADH dehydrogenase (A, relative decrease (%) of uncoupled respiration following rotenone addition) and complex IV / cytochrome c oxidase activity (B, respiration (pmol  $O_2 s^{-1}$ mg<sup>-1</sup>) after addition of ascorbate and TMPD) in systemic hearts of cuttlefish from the English Channel and Adriatic Sea. Significant differences (*P* < 0.05) between acclimation temperatures (*N*=5-11) are marked by asterisks and between populations by roofs.

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Figure 3: Temperature coefficients ( $Q_{10}$ , means ± 95% C.I.) of cardiac mitochondrial respiration from 11°C to 21°C, resolved for each substrate and for total state 3 and complex IV respiration. As there were no differences between acclimations, data were pooled for each case.  $Q_{10}$  of pyruvate and succinate are based on stimulating respiration.

Figure 4: Change of proline stimulated respiration without ADP (denoted as proton 508 leak) and with ADP (pmol  $O_2$  s<sup>-1</sup> mg<sup>-1</sup>) and relative proton leak (%, calculated as 509 fraction of leak rates relative to coupled respiration with ADP) from 11°C to 21°C 510 511 assay temperature in systemic hearts, in comparison between (A) English Channel 512 (N=24) and (B) Adriatic (N=19-20) cuttlefish. Due to the lack of differences between 513 acclimation treatments, data were pooled for each assay temperature and 514 population. Values are means  $\pm$  95% C.I. Significant differences (P < 0.05) between 515 assay temperatures for relative proton leak are marked by unequal letters of the 516 same case and between populations by roofs.

517

- 518 Figure 5: Relative change of maximum state 3 respiration (%) following cytochrome c
- addition, from 11°C to 21°C assay temperature in systemic hearts, in comparison
- 520 between English Channel (N=23-24) and Adriatic (N=19-20) cuttlefish. Due to the
- 521 lack of differences between acclimation treatments, data were pooled for each assay
- 522 temperature and population. Values are means ± 95% C.I.. Significant differences (*P*
- 523 < 0.05) between assay temperatures are marked by unequal letters of the same case
- 524 and between populations by roofs.
- 525
- 526 Table 1: Enzyme activities (µmol min<sup>-1</sup> mg protein<sup>-1</sup>, at 16°C) of (A) citrate synthase,
- 527 (B) lactate dehydrogenase and (C) octopine dehydrogenase of cuttlefish systemic
- 528 and branchial hearts, in comparison between thermal acclimations, as well as
- 529 between English Channel and Adriatic Sea populations.

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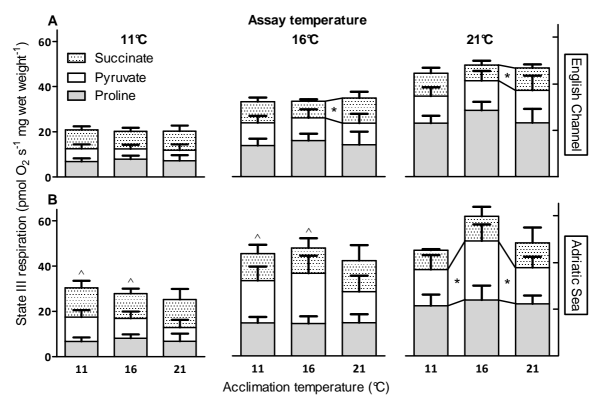
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- 741 cuttlefish (*Sepia officinalis* L.) populations in the English Channel and the Bay of
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Figure 1



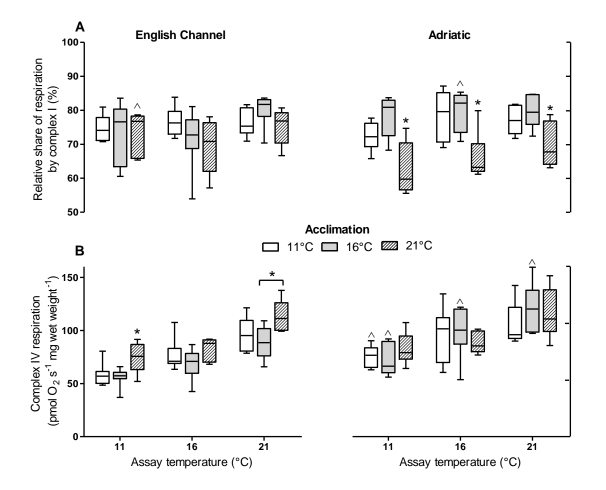
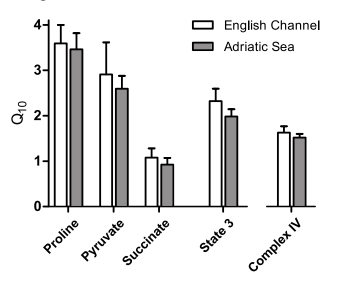


Figure 3





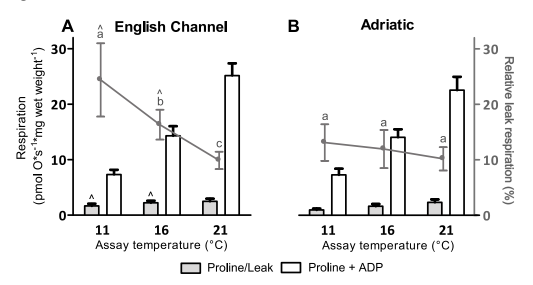


Figure 5

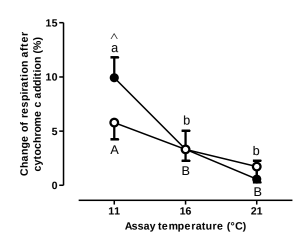


Table	1

			Systemic heart		Branchial heart			
Enzyme	Heart type							
		11	16	21	11	16	21	
Citrate synthase	English Channel	0.89 (0.82-0.96) <sup>a</sup>	0.73 (0.65-0.82) <sup>b</sup>	0.74 (0.60-0.88)	0.64 (0.51-0.77)	0.78 (0.67-0.89)	0.81 (0.74-0.88)	
	Adriatic Sea	1.07 (0.53-1.63)	▲ 1.01 (0.80-1.22)	0.81 (0.67-0.96)	0.63 (0.53-0.74)	0.64 (0.51-0.76)	0.78 (0.63-0.93)	
Lactate dehydrogenase	English Channel	0.10 (0.07-0.14)	0.11 (0.09-0.12) <sup>a</sup>	0.05 (0.04-0.07) <sup>b</sup>	0.12 (0.10-0.14)	0.10 (0.09-0.12)	0.08 (0.06-0.10)	
	Adriatic Sea	0.13 (0.08-0.19)	0.09 (0.05-0.13)	0.08 (0.07-0.10)	0.07 (0.05-0.10)	0.07 (0.06-0.09)	0.06 (0.06-0.07)	
Octopine dehydrogenase	English Channel	0.40 (0.34-0.46)	0.48 (0.43-0.53)	0.39 (0.29-0.48)	0.22 (0.17-0.27)	0.24 (0.20-0.27)	0.23 (0.20-0.26)	
	Adriatic Sea	0.56 (0.46-0.66)	0.57 (0.35-0.80)	0.62 (0.49-0.75)	0.26 (0.18-0.33)	0.23 (0.18-0.29)	0.28 (0.23-0.34)	

Agent	Action	Final concentration
Substrates		
Proline	Amino acid substrate	5 mmol l <sup>-1</sup>
Pyruvate	Carbohydrate substrate	5 mmol l <sup>−1</sup>
Succinate	Carbohydrate substrate	10 mmol I <sup>-1</sup>
ADP	Substrate for ATP generation	2.5 mmol I <sup>-1</sup>
Cytochrome c	Electron carrier, tests for outer membrane integrity	0.01 mmol l <sup>-1</sup>
TMPD	Complex IV substrate	5 mmol l <sup>−1</sup>
Ascorbate	Complex IV substrate	20 mmol l <sup>-1</sup>
Inhibitors		
Antimycin	Inhibits complex III, to test non-mitochondrial respiration	2.5 μmol l <sup>−1</sup>
Rotenone	Complex I inhibitor	2.5 μmol l <sup>−1</sup>
Oligomycin	Inhibits ATP synthase, to test proton leak	2 µg ml <sup>−1</sup>
Uncoupler		
FCCP	Uncouples the electron transport system electron flow from	2.5 μmol l <sup>−1</sup>
	ATP generation by relieving the proton gradient	

Table S1. Action and concentrations of the agents used to assay mitochondrial respiration of
permeabilised heart fibres from the European cuttlefish

Table S2. Measurements of the Europ	pean cuttlefish <i>Sepia officinali</i> s	s used in acclimation experiments

	0.1.1	Acclimation	0	Date of	<b>-</b>	Mantle length	Systemic heart	Branchial heart	Branchial heart	
dentifier	Origin	temperature (°C)	Sex	sampling	Total mass (g)	(cm)	mass (mg)	#1 mass (mg)	#2 mass (mg)	Comments
.06	Adriatic	11	Male	7/07/2009	90.22	8.8	58.3	38.5	33.5	
10	Adriatic	11	Male	14/07/2009	63.34	8.0	39.3	25.4	28.9	
.13	Adriatic	11	Male	16/07/2009	83.75	8.5	49.3	37.0	34.3	Animal was inking during capture
30	Adriatic	11	Female	21/09/2009	99.20	9.1	58.6	38.0	37.4	Animal was inking during capture
.63	Adriatic	11	Female	14/07/2010	110.53	9.3	58.3	39.7	39.2	Animal was inking during capture
64	Adriatic	11	Male	19/07/2010	78.47	8.2	40.1	27.8	30.9	Animal was inking during capture
.65	Adriatic	11	Female	20/07/2010	86.40	8.0	43.1	34.9	35.3	Animal was slightly inking during capture
69*	Adriatic	11	Male	22/11/2010	48.20	-	24.6	19.9	18.4	
.07	Adriatic	16	Female	8/07/2009	46.90	6.8	30.5	23.7	22.4	
.09	Adriatic	16	Male	13/07/2009	40.57	7.0	27.0	11.6	19.5	Animal was inking during capture
14	Adriatic	16	Male	20/07/2009	72.52	8.2	44.9	24.7	27.5	Animal was inking during capture
19	Adriatic	16	Female	30/07/2009	77.88	7.9	46.4	42.2	38.5	Animal was inking during capture, very well developed gonads
24	Adriatic	16	Female	10/08/2009	59.11	7.4	37.1	41.6	42.6	
46*	Adriatic	16	Female	20/04/2010	50.10	7.1	27.3	18.3	18.7	Animal was inking during capture
47*	Adriatic	16	Male	21/04/2010	49.87	7.0	27.1	15.0	16.5	Animal was inking during capture
48*	Adriatic	16	Male	26/04/2010	59.67	7.5	33.7	24.5	21.8	Animal was inking during capture
49*	Adriatic	16	Female	27/04/2010	62.66	7.2	33.0	20.9	24.7	Animal was inking during capture
50*	Adriatic	16	Male	28/04/2010	52.05	6.9	29.1	22.1	24.3	Animal was inking during capture
51*	Adriatic	16	Female	29/04/2010	41.34	6.2	25.6	13.6	16.7	Animal was inking during capture
51	Adriatic	16	Male	12/07/2010	109.93	9.5	77.4	42.1	44.5	Animal was inking during capture
62	Adriatic	16	Male	13/07/2010	144.00	10.5	101.1	56.3	56.2	5 · · · 5 · · · · ·
52*	Adriatic	21	Female	26/05/2010	148.82	10.5	74.7	40.3	43.8	
53*	Adriatic	21	Male	31/05/2010	128.71	10.0	68.7	40.0	39.8	
54*	Adriatic	21	Female	2/06/2010	228.17	11.2	113.2	65.3	78.8	Female had proper developed gonads with yellowish transparent eggs
55*	Adriatic	21	Male	3/06/2010	252.92	12.7	135.6	74.7	74.3	)
56*	Adriatic	21	Male	7/06/2010	149.30	10.4	75.7	55.3	47.6	
57*	Adriatic	21	Male	9/06/2010	150.37	10.1	75.6	41.5	45.3	
58	Adriatic	21	Female	7/07/2010	147.75	10.0	86.7	58.7	56.9	Female had eggs
59	Adriatic	21	Female	8/07/2010	226.54	11.3	140.7	82.9	79.9	Animal was inking during capture, female had ego
50 60	Adriatic	21	Male	9/07/2010	198.63	11.6	109.0	69.4	74.8	Animal was inking during capture
66	Adriatic	21	Female	21/07/2010	184.21	10.8	145.4	87.7	87.2	Female already mated as there were sucker mark on the head, had hardly ink in ink sac
67	Adriatic	21	Female	23/07/2010	144.90	9.5	108.5	57.4	59.1	Female already mated as there were sucker marks on the head, had hardly ink in ink sac
68	Adriatic	21	Female	26/07/2010	248.32	12.2	161.3	96.0	101.3	····, ··· · , ····
.08	Oosterschelde	11	Male	10/07/2009	59.09	7.3	42.8	23.9	27.5	
.11	Oosterschelde	11	Male	15/07/2009	79.46	7.9	63.5	41.2	38.3	
12	Oosterschelde	11	Female	21/07/2009	89.19	8.3	63.2	37.5	39.1	
23	Oosterschelde	11	Male	6/08/2009	96.64	8.9	81.2	51.3	48.5	Animal was inking during capture
25	Oosterschelde	11	Male	12/08/2009	112.12	9.5	70.1	47.0	42.0	
26	Oosterschelde	11	Male	13/08/2009	77.84	8.5	60.1	37.0	38.5	
27	Oosterschelde	11	Male	18/08/2009	92.99	8.6	73.0	43.4	45.8	Animal was inking during capture
28	Oosterschelde	11	Male	8/09/2009	79.98	8.1	65.1	38.1	36.0	
20 29	Oosterschelde	11	Male	17/09/2009	83.31	8.7	83.6	53.9	50.6	
29 04	Oosterschelde	16	Female	4/06/2009	72.46	8.5	-	-	-	

A05	Oosterschelde	16	Female	17/06/2009	84.50	8.6	57.3	31.6	31.4	Animal was inking during capture
A15	Oosterschelde	16	Female	22/07/2009	97.70	8.9	61.6	38.4		
A18	Oosterschelde	16	Male	29/07/2009	108.29	9.1	111.4	46.4	43.8	Animal was inking during capture
A31	Oosterschelde	16	Male	22/09/2009	125.10	9.3	114.5	50.4	59.3	
A32	Oosterschelde	16	Male	23/09/2009	79.75	7.7	73.1	34.4	33.8	
A33	Oosterschelde	16	Female	24/09/2009	253.83	12.5	315.7	130.0	134.3	Female had proper developed gonads with
										yellowish transparent and white eggs
A34*	Oosterschelde	16	Female	28/08/2009	164.90	9.3	133.6	93.5	86.7	
A35	Oosterschelde	16	Male	29/09/2009	155.89	10.2	131.5	65.4	68.0	Animal was inking during capture
A36	Oosterschelde	16	Female	30/09/2009	128.38	9.1	97.3	54.7	56.1	Animal was inking during capture
A37	Oosterschelde	16	Female	1/10/2009	165.94	10.3	183.0	90.1	87.0	
A38	Oosterschelde	16	Male	5/10/2009	164.39	11.4	167.9	79.0	70.0	
A39*	Oosterschelde	16	Male	6/10/2009	278.93	12.6	214.8	127.8	133.0	
A40*	Oosterschelde	16	Male	7/10/2009	177.30	11.0	142.2	87.4	80.5	
A41*	Oosterschelde	16	Male	8/10/2009	178.59	10.5	123.9	80.8	89.8	Animal was slightly inking during capture
A42*	Oosterschelde	16	Female	12/10/2009	225.04	10.9	183.3	108.6	116.9	
A43*	Oosterschelde	16	Male	13/10/2009	258.39	12.3	210.0	131.8	138.1	Animal was slightly inking during capture
A44*	Oosterschelde	16	Female	14/10/2009	299.15	12.8	194.7	161.2	133.2	Female had proper developed gonads with yellowish transparent eggs
A45*	Oosterschelde	16	Female	15/10/2009	276.24	12.3	184.6	146.7	155.6	Female had proper developed gonads with yellowish transparent eggs
A16	Oosterschelde	21	Female	27/07/2009	71.33	7.9	56.8	36.2	32.7	
A17	Oosterschelde	21	Male	28/07/2009	83.59	8.1	77.7	41.4	39.5	Heart muscle was relatively tough
A20	Oosterschelde	21	Female	3/08/2009	102.13	8.5	75.5	45.7	44.5	
A21	Oosterschelde	21	Male	4/08/2009	124.66	9.9	106.3	62.3	67.9	
A22	Oosterschelde	21	Male	5/08/2009	123.77	8.9	110.7	65.3	60.4	
Individua	als marked with asterisks	were used for	morphometric st	atistics but not for	measurement of r	nitochondrial resp	iration.			