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Identification of the alternative oxidase gene and its expression in the copepod Tigriopus californicus

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| 1 | Title: Identification of the alternative oxidase gene and its expression in the copepod |
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21 Abstract

22 In addition to the typical electron transport system (ETS) in animal mitochondria 23 responsible for oxidative phosphorylation, in some species there exists an alternative 24 oxidase (AOX) pathway capable of catalyzing the oxidation of ubiquinol and the 25 reduction of oxygen to water. The discovery of AOX in animals is recent and further 26 investigations into its expression, regulation, and physiological role have been 27 hampered by the lack of a tractable experimental model organism. Our recent DNA 28 database searches using bioinformatics revealed an AOX sequence in several marine 29 copepods including *Tigriopus californicus*. This species lives in tidepools along the west 30 coast of North America and is subject to a wide variety of daily environmental stresses. 31 Here we verify the presence of the AOX gene in *T. californicus* and the expression of 32 AOX mRNA and AOX protein in various life stages of the animal. We demonstrate that 33 levels of the AOX protein increase in T. californicus in response to cold and heat stress 34 compared to normal rearing temperature. We predict that a functional AOX pathway is 35 present in *T. californicus*, propose that this species will be a useful model organism for 36 the study of AOX in animals, and discuss future directions for animal AOX research.

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Key Words: arthropod, electron transport system, environmental stress, mitochondria,
 protein isolation, respiration, tide-pool

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42 Introduction

43 <u>The Alternative Oxidase (AOX) Pathway</u>

44 Cellular respiration in animal cells takes place in the mitochondria via the 45 electron transport system (ETS). The ETS is comprised of multi-subunit complexes and 46 mobile electron carriers in the inner membrane of the mitochondria (Genova, 2014). 47 Four complexes, (I, II, III, and IV), and the mobile carriers ubiquinone/ubiquinol and 48 cytochrome c, shuttle electrons through the system to the final electron acceptor oxygen 49 (Genova, 2014). At the same time, several of these complexes move protons from the 50 matrix to the intermitochondrial membrane space and result in a proton motive force 51 (PMF) across the inner mitochondrial membrane (Genova, 2014). This PMF is used by 52 complex V (ATP synthase), and as protons flow back into the matrix through this 53 enzyme, ATP is synthesized through the process of oxidative phosphorylation (Genova, 54 2014).

55 Many animals also contain additional enzymes capable of putting electrons into 56 and/or removing electrons from the ETS (McDonald and Gospodaryov, 2018). One such 57 enzyme is the alternative oxidase (AOX), a terminal quinol oxidase that catalyzes the 58 oxidation of ubiguinol and the reduction of oxygen to water (McDonald and 59 Vanlerberghe, 2004; McDonald, 2008). In contrast to cytochrome c oxidase, which is 60 comprised of nuclear and mitochondria encoded subunits, the AOX is encoded by a 61 single nuclear gene (McDonald, 2008). The AOX protein is composed of four helices 62 and several conserved amino acid residues that are important for its enzymatic function 63 (McDonald, 2008). Use of the AOX pathway causes electrons to bypass complexes III

and IV and results in cyanide-resistant respiration (Rogov et al., 2014). As AOX is nonproton pumping (in contrast to complexes I, III, and IV), it does not contribute directly to
the proton motive force (PMF), and the energy associated with electron transport is
dissipated as heat (McDonald, 2008). Therefore, AOX is viewed as energetically
inefficient.

69 AOX in Animals

70 AOX has been identified in several animal species due to the presence of AOX 71 DNA or mRNA sequences in public molecular databases (McDonald and Vanlerberghe, 72 2004; McDonald et al., 2009). Recent database searches have revealed the presence 73 of AOX DNA or mRNA in the phyla Ctenophora, Platyhelminthes, Arthropoda, 74 Tardigrada, Scalidophora, Brachiopoda, and Rotifera for the first time (McDonald and 75 Gospodaryov, 2018). Experimental evidence for the expression of AOX mRNA exists for 76 the sponge *Ephydatia muelleri*, and the molluscs *Anadara ovalis*, *Crassostrea gigas*, 77 Crassostrea virginica, and Mercenaria mercenaria (McDonald et al., 2009; Liu and Guo, 78 2017). Thus far, the only investigation of changes in AOX expression due to 79 environmental conditions has been the use of quantitative PCR (qPCR) to investigate 80 AOX transcript levels in the bivalves *Crassostrea gigas* and *Diplodon chilensis* which 81 increased in response to hypoxia and anoxia (Sussarellu et al., 2012; Yusseppone et 82 al., 2018).

The bulk of information about the AOX of animals exists at the level of genes or transcripts. The only reports of naturally occurring AOX proteins present in animals are from the brine shrimp *Artemia franciscana* and the Pacific oyster *Crassostrea gigas*;

86 AOX proteins in these species are ~35 kDa in size (Rodriguez-Armenta et al., 2018). 87 Knowledge of other animal AOX proteins is limited to studies that have taken advantage 88 of heterologous expression systems, therefore it is difficult to determine whether the 89 molecular masses of AOX reported reflect reality in vivo. For example, the expression of 90 the AOX from the sea squirt *Ciona intestinalis* produced a functional enzyme capable of 91 oxygen consumption of 42 kDa that was targeted to mitochondria in human kidney cells, 92 mice, and fruit flies (Hakkaart et al., 2006; El-Khoury et al., 2013; Kemppainen et al., 93 2014).). Heterologous expression of the Pacific oyster AOX has also occurred in the 94 yeast Saccharomyces cerevisiae (Robertson et al., 2016). It is clear that there is a need 95 for an animal model that naturally expresses the AOX protein. The importance of finding 96 a suitable organism for the study of animal AOX is important as typically used model 97 systems (i.e. mouse and fruit fly) do not contain a naturally occurring AOX and therefore 98 cannot be studied (McDonald et al., 2009). No information exists regarding the post-99 translational regulation of an animal AOX. In contrast to Complexes I-IV of the ETS, 100 there is no evidence that AOX exists as part of any respiratory supercomplex (Schertl 101 and Braun, 2014). The activation and inhibition kinetics of animal AOX proteins is also 102 unknown. Information on the characteristics of animal AOX proteins is needed, both due 103 to the desire for basic biological knowledge, and the realization that this knowledge is 104 required to assess the efficacy and safety of proposals to use AOX as a gene therapy 105 tool in humans for the treatment of mitochondrial diseases (El-Khoury et al., 2014). This 106 gap in our knowledge exists because to date animals known to contain AOX gene 107 sequences are often expensive to grow and culture in the lab, take up large amounts of 108 laboratory space, and do not have rapid life cycles which makes them impractical and

109 challenging for scientific study. In addition, research efforts have been hampered by the 110 existence of few genetic, molecular biology, and biochemical protocols and tools that 111 are available for use in these animals. It is clear that a worthy endeavor is to find a 112 useful AOX containing animal that is inexpensive to culture, has a rapid generation time, 113 takes up little space, and for which protocols and tool development is possible. In this 114 paper we assert that the intertidal copepod *Tigriopus californicus* meets all of these 115 criteria and can be used as a model system for the investigation of an animal AOX 116 protein.

117 <u>Tigriopus californicus</u>

118 T. californicus is an intertidal species of copepod found on the Pacific coast of 119 North America that inhabits rock pools (Burton and Lee, 1994). Due to T. californicus' 120 intertidal habitat, these animals are exposed to ever changing environmental stressors 121 including temperature, salinity, and oxygen levels (Burton and Lee, 1994) and 122 presumably have mechanisms in place to deal with such challenges. T. californicus has 123 a short generation time (~2-4 weeks depending on rearing temperature)(Hong and 124 Shurin, 2015), small space needs, and genetically divergent populations that can be 125 cross-bred in the laboratory (Burton & Feldman, 1981). A strong argument for using T. 126 californicus as a model organism in ecotoxicology and environmental genomics has 127 been made based on the above characteristics (Raisuddin et al., 2007). In addition, a 128 protocol and tool exists for the suppression of gene transcription using RNA interference 129 in this species (Barreto et al., 2015b).

130 T. californicus has been used to investigate the mechanisms of speciation and 131 incompatibilities in hybrid animals derived from mating different geographical 132 populations (Barreto et al., 2015a). Hybrid incompatibilities have been demonstrated to 133 have a negative impact on oxidative phosphorylation and the interaction between the 134 nuclear and mitochondrial genome encoded processes (Barreto et al., 2015a). This 135 incompatible hybridization causes mitochondrial dysfunction caused including inefficient 136 OXPHOS, a decrease in ATP biosynthesis, and increased ROS production (Barreto et 137 al., 2015a). An investigation into the bioenergetics of T. californicus is therefore 138 warranted at the level of the ETS by examining complexes I-IV and AOX.

In this study we confirm the presence of the AOX gene, the expression of AOX mRNA, and demonstrate that the AOX protein is present in the copepod *T. californicus*. We believe this is the first step in establishing this copepod as a model species for animal AOX research. By demonstrating that *T. californicus* is an effective model species with experimental tools that are available, future research can examine the physiological role of AOX in *T. californicus* and will explore questions about the expression and regulation of AOX proteins in an animal system.

146 Materials and Methods

147 In silico identification of copepod AOX sequences

Molecular database searches using various animal AOX sequences in BLAST (Basic Local Alignment Search Tool) using default settings at the National Center for Biotechnology Information (NCBI) (<u>http://www.ncbi.nlm.nih.gov/</u>) (this was an iterative process that used novel AOX sequences as they were discovered; full procedure as

described in McDonald et al., 2009) identified several putative copepod AOX

153 sequences. DNA sequences were translated into protein sequences using the ExPASy

- 154 Translate Tool (<u>http://web.expasy.org/translate/</u>).
- 155 Laboratory Culturing Conditions of *T. californicus*

156 Animals were obtained from J&L Aquatics Canada, Vancouver, British Columbia. 157 The animals arrived in a ~150mL bottle containing ~1000-2000 individuals at various life 158 stages. Animals used for DNA and RNA isolation were maintained in cultures of 20-200 159 individuals in 600 mL beakers containing 400mL of seawater (37.5 g/L) with a specific 160 gravity of 0.026 and grown at 21°C (room temperature) (Edmands and Deimler, 2004). 161 The salt water was made with Instant Ocean Salt and the specific gravity was tested 162 with the Instant Ocean Saltwater Aquarium Hydrometer (Instant Ocean, Blacksburg, VA, 163 USA). Each beaker was covered with the top plate of a petri dish to slow the rate of 164 evaporation. Beakers were placed in an incubator programmed to run on a 12 hour light 165 and 12 hour dark cycle. The animals were fed 4 drops of Reef Nutrition Phyto-Feast 166 Live Premium Phytoplankton (J&L Aquatics Canada, Vancouver, British Columbia) 167 every other day for the entire culturing period. Preliminary experiments demonstrated 168 that our animals completed their life cycle faster at 15.0°C, so a change in rearing 169 protocol occurred about 2 years into culturing *T. californicus* in the lab. Animals used for 170 protein isolations were cultured in 400mL beakers filled with 250mL of salt water (41.7 g 171 of Instant Ocean salt per 1L of MilliQ filtered water) at 15°C using a 12 hour light and 12 172 hour dark cycle. The copepods were fed once a week with 0.01 g of Tetramin fish food 173 (Nutrafin Basix) and 0.005 g of Spirulina powder (EarthRise) in 50mL of salt water 174 added to each habitat. The water in each habitat was changed weekly in order to

175 prevent the build-up of moulted exosketetons and waste in the beakers. Animals used in

temperature experiments were acclimated to 15°C (normal growth temperature) for 2-4

177 weeks before being exposed to 6°C (cold) or 28°C (hot) for 24 hours (acute treatment)

178 or 1 week (chronic treatment) before collection for protein isolation.

179 DNA and RNA isolation and AOX Gene and Transcript analysis

180 Primer Design

181 Gene specific primers for the *T. californicus* 551 bp AOX cDNA fragment

182 (JW502496) identified in the NCBI database were designed using the Primer3 program

183 (http://frodo.wi.mit.edu/primer3). Custom primers were then synthesized (Invitrogen Life

184 Technologies, Carlsbad, CA, USA) for use in PCR and RT-PCR protocols (Table 1).

185 Three different forward and reverse primer sets were created in order to generate cDNA

186 products ranging from 200- 500bp; multiple primer sets were designed in case one or

187 more sets proved unsuccessful in producing a product.

188 <u>Collection and preparation of copepods</u>

189 In order to collect the copepods from the cultures for the isolation of DNA and RNA 190 the animals were strained out from the water using Miracloth (pore size: 22-25µm) 191 (Merck KGaA, Darmstadt, Germany). Once the water was completely drained from the 192 beaker, the copepods were washed several times with clean salt water and quickly 193 transferred to a pre-weighed Eppendorf tube, the mass of the animals was determined 194 (~200 mg), and the tube was frozen in liquid nitrogen. The frozen copepods were then 195 removed from the Eppendorf tube and ground into a fine powder using a previously 196 autoclaved and chilled pestle and mortar. The powdered sample was kept cold by 197 grinding under liquid nitrogen.

198 Isolation of nucleic acids and amplification parameters

199 For the extraction of DNA and RNA from *T. californicus* Trizol reagent proved 200 most effective (Invitrogen Life Technologies, Carlsbad, CA, USA) and the protocol was 201 conducted as per the manufacturer's instructions. Once the nucleic acids were 202 successfully isolated from T. californicus and quantified using spectrophotometry, PCR 203 and reverse transcriptase (RT)-PCR were conducted in order to amplify the targeted 204 AOX sequence. PCR was conducted with 5 µg of *T. californicus* DNA using 0.5 µL iTaq 205 DNA polymerase (Bio-Rad Laboratories Ltd., Mississauga, ON, CA), 2 µL of each 10 206 μ M forward and reverse primer, 2 μ L of 10x PCR buffer, 2 μ L of dNTP mix, and 1.5 μ L 207 of nuclease-free water in 20 µL total volume. The thermal cycler was set for 1 cycle of 2 208 minutes at 94°C for initial denaturation, 40 cycles consisting of: denaturation for 20 209 seconds at 94°C, annealing for 10 seconds at 58°C, and extension for 30 seconds at 210 68°C, followed by 1 cycle of final extension for 10 minutes at 68°C and the final soak 211 cycle at 4°C held indefinitely, until the sample was removed from the thermocycler (Bio-212 Rad Laboratories Ltd., Mississauga, ON, CA). 213 RT-PCR was conducted using the Access RT-PCR Introductory System (Promega 214 Corporation, Madison, WI, USA) with 2 µg of total DNase treated RNA in 50 µL

reactions containing 10 μL AMV/Tfl 5x reaction buffer, 1 μL dNTP mix, 2 μL of each 10

 μ M forward and reverse primer, 2 μ L of 25 mM MgSO₄, 1 μ L AMV RTase, 1 μ L Tfl DNA

217 polymerase, and 29.5 µL nuclease-free water. The thermal cycler was set for first strand

cDNA synthesis consisting of 45 minutes at 45°C for reverse transcription and 2

219 minutes at 94°C for AMV RT inactivation. This was followed by 40 cycles of

denaturation for 30 seconds at 94°C, annealing for 1 minute at 60°C, and extension for

2 minutes at 68°C, followed by 1 cycle of final extension for 7 minutes at 68°C and the
final soak cycle held at 4°C, until the sample was removed from the thermocycler (BioRad Laboratories Ltd., Mississauga, ON, CA).

224 The amplification products were separated using DNA gel electrophoresis using a 225 1.2% agarose gel containing 1mL 50x Tris-acetate-EDTA (TAE) buffer, 49 mL nuclease-226 free water (dH_2O), and 0.60 g agarose powder run in a tank with 1x TAE running buffer. 227 5 µL 1 kb or 100 bp DNA ladder (GeneDirex, USA) was loaded in the first lane of the 228 gel, followed by 5-10 µL of each PCR or RT-PCR reaction combined with 6X DNA 229 loading buffer in subsequent lanes. Gels were run at 80-120V for 45 minutes and 230 stained with ethidium bromide in order to visualize the DNA under UV light (VersaDoc 231 400) (Bio-Rad Laboratories Ltd., Mississauga, ON, CA).

232 Isolation of Proteins from Copepods

233 For each protein extraction, approximately 0.05-0.10 g of copepods were isolated 234 from their habitats by filtration using a 100 µm filter (Fisherbrand) and placed in an 235 Eppendorf tube. Various combinations of different volumes of SDS-PAGE sample buffer 236 $(350-500 \ \mu\text{L})$ and 20 μL of β -mercaptoethanol solution were pipetted into each 237 Eppendorf tube. Each sample was sonicated (Omni International Inc.) for 10 seconds at 238 a frequency of 6 kHz and/ or homogenized to extract the proteins from the copepods. 239 The Precellys 24 lysis and homogenization machine (Bertin Technologies) was used to 240 perform the homogenization of copepod tissue. The homogenization processes 241 consisted of adding beads into each sample, loading the samples into the machine, and 242 3D bead-beating for 30 seconds at 5000rpm. These various combinations were used in

order to identify the optimal conditions for protein isolation from our laboratory growncopepods.

245 Protein Analysis: Gel Electrophoresis

246 Protein samples were analyzed using reducing SDS-PAGE and Western blotting. 247 For gel electrophoresis, a 15 well mini-PROTEAN TGX Stain-Free gel (Bio-rad), was 248 loaded with 15 µL of each copepod sample. In order to determine the molecular weight 249 of the proteins in our samples, two wells were loaded with 5 µL of the Precision Plus 250 Protein WesternC Standards (Bio-rad). A yeast sample expressing the Pacific oyster 251 alternative oxidase (Robertson et al., 2006) served as a positive control in order to verify 252 the cross-reactivity of the AOX antibody with our copepod samples. A negative control 253 blot was run using proteins isolated from the algal food source to rule out the possibility 254 of an algal AOX being detected in our copepod samples. Following the loading of the 255 wells the gel was run at a constant voltage (200V) for 35 minutes. After the run was 256 complete, the gel was visualized using a VersaDoc (Bio-rad, USA) so an image could 257 be taken to visualize protein loading. For protein gels using our temperature experiment 258 samples, proteins in each sample were quantified using Quick Start Bradford Protein 259 Assay using bovine serum albumin standards and a spectrophotometre. Each sample 260 consisted of 20 µg of protein added to each lane.

261 Protein Analysis: Western Blot

The Trans-Blot Turbo Transfer System (Bio-rad) was used to transfer the
proteins onto a nitrocellulose membrane in 3 minutes using Trans-Blot Transfer Medium
(Bio-rad). Following protein transfer, the membrane was placed in 5% dry milk in 10 mL

265 of 1X TBS-T on a shaking platform for 1 hour. The membrane was then subsequently 266 washed three times in TBS-T. 4µL of the primary AOX antibody (Plant AOX1/2, Cat. # 267 ABIN3197483, Agrisera Antibodies) was then added to 20 mL of 1X TBS-T for 1 hour 268 (dilution 1: 10,000). Following this, the membrane was again washed three times (15 269 minutes, 10 minutes, and 10 minutes) with TBS-T. The membrane was then incubated 270 in 2 µL secondary antibody, a goat anti rabbit IgG (Cat. # ABIN101988, Agrisera 271 Antibodies, dilution 1:25,000), and 2 µL of the Precision Protein StrepTactin HRP 272 Conjugate (Bio-rad) in 20mL of 1x TBS-T for 1 hour followed by three washes in 1X 273 TBS-T. It was subsequently placed in a mixture of luminol/enhancer and peroxide buffer 274 solution from the Immun Star Western C Chemiluminescent kit (Bio-rad) in a 1:1 ratio 275 for 5 minutes. The membrane was then placed in the VersaDoc, and the 276 chemiluminescent signal was detected and imaged.

277 **Results**

278 Identification of putative AOX sequences in copepods

A molecular database search using a BLAST (Basic Local Alignment Search Tool) at the National Center for Biotechnology Information (NCBI) revealed the presence of putative AOX sequences in 14 different copepod species (Table 2). The copepod species are representatives of 3 different copepod orders and 12 different families (Table 2). The predicted AOX proteins all contained at least 2 or more of the 4 iron-binding motifs (Table 2) containing conserved glutamate and histidine residues required for the enzyme to be functional (McDonald et al., 2009). AOX DNA and mRNA

in copepods are present in multiple developmental stages, both sexes, and varioushabitats around the world (Table 2).

288 In silico analyses of the Tigriopus californicus AOX sequence

289 A molecular database search revealed a putative T. californicus AOX DNA 290 sequence of 551 bp (Figure A.1). The predicted protein from T. californicus was a 291 partial sequence, however 3 out of the 4 iron binding motifs containing conserved 292 glutamate and histidine residues required for the enzyme to be functional (McDonald et 293 al., 2009) were present (Figure A.1). From previous studies, the C-terminal motif N-P-294 [YF]-X-P-G-[KQE], was determined to be diagnostic for animal AOX proteins (McDonald 295 et al., 2009). T. californicus demonstrates a high level of motif conservation in the region 296 of the protein mentioned above (i.e. N- P-F-E-K-G-K) (Figure A.1).

297 Molecular analyses of the *T. californicus* AOX

298 Trizol reagent proved to be effective in isolating RNA from the copepods and 299 allowed us to proceed with RT-PCR. All three AOX primer sets yielded cDNA fragments 300 of the expected sizes (Figure 1). Proteins were successfully isolated from T. californicus 301 using 0.05 g of copepods and 350 or 500 μ L of SDS-PAGE sample buffer containing β -302 mercaptoethanol and using various combinations of sonication and homogenization 303 (Figure A.2). The *T. californicus* AOX was recognized by a plant AOX antibody and was 304 ~ 50 kDa in size (Figure 2). The antibody failed to cross-react with the AOX protein 305 likely present in the algae provided to the copepods as food (Figure A.3). Previous 306 studies have demonstrated that algal AOXs contain features that differ from those of 307 other organisms (e.g. extended N-terminus and other insertion/deletions) (Nemanis et

al., 2013). We therefore expected that a plant AOX antibody would not cross-react withthe algal AOX.

310 Response of AOX Protein Levels to Temperature Treatments

Copepods were acclimated to 15°C for 2-4 weeks and then subjected to one of three different temperatures (6, 15, or 28°C) for 24 hours or 1 week and then sampled. Protein gels demonstrated that equal protein loading had occurred (Figure A.4 and A.5). Western blots indicate that AOX protein levels are low under control conditions (24 hours at 15°C), but AOX levels are higher after 24 hours of exposure to 6°C and 28°C (Figure 3). Similarly, AOX protein levels are low under control conditions (1 week at 15°C), but AOX levels are higher after 1 week of exposure to 6°C and 28°C (Figure 4).

318 **Discussion**

319 Discovery of AOX in Copepods and the Phylum Arthropoda

320 We have shown that AOX is in the genomes of multiple species of copepods 321 from around the world that inhabit a wide variety of ecological niches (Table 2.) The first 322 reports of AOX from arthropods were in the brine shrimp Artemia franciscana and 323 putative sequences from other members of the Chelicerata, Hexapoda, and Crustacea 324 (Rodriguez-Armenta et al., 2018; McDonald and Gospodaryov, 2018). It was previously 325 hypothesized that AOX was not present in arthropods due to a gene loss event 326 (McDonald et al., 2009), however, the above data refute this hypothesis. 327 Based on information gathered using bioinformatics tools, the copepod T.

328 *californicus* contains an AOX sequence (Figure A.1, Table 2). Our next goal was to

determine whether the AOX gene of *T. californicus* is expressed in the animal. RT-PCR
using RNA isolated from pooled developmental stages of the organism verified that
AOX mRNA is expressed in the copepod (Figure 1). This represents the first AOX
mRNA detected experimentally in any arthropod using gene specific primers for AOX.

In order to verify that the AOX sequence recovered from the arthropod *T. californicus* was from this animal and not a contaminant (e.g. from a microbial symbiont or pathogen) the animal's 551 bp cDNA sequence was translated to its predicted protein sequence and was found to possess the C-terminal region N-P-F-E-K-G-K (Figure A.1). This C-terminal motif is of particular interest because it is characteristic of only animal AOXs and is different from the C-terminal regions observed in plants and fungal species (McDonald et al., 2009).

340 We wished to confirm that AOX protein was detectable in *T. californicus*. Reducing 341 SDS-PAGE followed by Western blotting with a commercial plant AOX1/2 antibody 342 detected a protein of ~ 50 kDa in our copepod samples (Figure 2). Heterologously 343 expressed AOX proteins from *Ciona intestinalis* and *Crassostrea gigas* are 35-42 kDa in 344 size (Hakkaart et al., 2006; Robertson et al., 2016). The AOX protein of Artemia 345 *franciscana* is between 25 and 37 kDa on Western blots using the anti-AOX 346 Sauromatum guttatum monoclonal antibody (Rodriguez-Armenta et al., 2018). It is 347 expected that the food source (Spirulina) would contain AOX as it is an algae, however, 348 the AOX antibody did not recognize a protein in this sample (Figure A.3). 349 Finally, we wished to determine if AOX protein levels would change in response to

an environmental stressor. The copepods are normally reared at 15°C and have been

351 cultured at this temperature for several years and have likely adapted to this growth 352 temperature over many generations in our lab. We therefore selected 15°C as our 353 control temperature and exposed treatment groups to either 6°C (cold) or 28°C (hot) for 354 an acute time period of 24 hours or a chronic time period of 1 week prior to sampling the 355 animals. Animals grown at 15°C exhibited very low levels of detectable AOX protein 356 (Figures 3 and 4). In contrast, animals grown at 6°C or 28°C exhibited higher levels of 357 AOX protein after 24 hours of exposure (Figure 3) or 1 week of exposure (Figure 4). 358 Similar to what we report here, an increase in AOX protein levels in response to cold 359 stress has been reported in a variety of plants (McDonald, 2008). In addition, AOX1 360 protein levels sharply increase in response to heat stress in the green alga 361 Chlamydomonas reinhardtii (Zalutskaya et al., 2015). Unfortunately, no other studies 362 have examined the effects of biotic or abiotic stressors on AOX protein levels in any 363 animal that naturally possesses the enzyme. The only other investigation of a native 364 AOX from an animal is a Western blot confirming its presence in the brine shrimp 365 Artemia franciscana (Rodriguez-Armenta et al., 2018). Our finding is therefore the first 366 to demonstrate that the levels of a native AOX protein in an animal change in response 367 to an environmental stressor.

368 <u>The Power of *T. californicus* as an Experimental System to Examine AOX</u>

T. californicus represents an emerging model organism for research because of its abundance, wide geographic distribution, ease of manipulation in laboratory settings, and its ecological relevance due to its position in marine food webs (Raisuddin et al., 2007). Lab grown cultures of *T. californicus* can be maintained for many generations and are relatively inexpensive to house. We were able to isolate copepod DNA and

RNA using Trizol reagent and present a protocol here for successfully isolating proteins
from whole animals. Coupled with newer techniques such as RNA interference (RNAi)
(Barreto et al., 2015), it is clear that many experimental tools can be used to answer
questions about AOX in this organism.

378 Based on our results, we can begin to ask questions about the genes involved in 379 regulatory pathways that control transcript expression levels in response to 380 environmental changes, developmental stages, and gender. Analysis of AOX 381 expression levels can be conducted by subjecting T. californicus to a variety of 382 environmental stressors such as alterations in salinity and temperature (Lauritano et al., 383 2012; Zhang et al., 2013) in order to observe patterns of AOX gene expression under 384 varying environmental conditions. The environmental stress of fluctuating salinity levels 385 is a key stressor to T. californicus and has been noted to affect the expression of a 386 variety of genes in the copepod (Burton and Lee, 1994; Van Aken et al., 2009). We 387 have demonstrated that AOX protein levels increase in *T. californicus* in response to 388 temperature stress, however, at this time it is not known why T. californicus has the 389 gene for and produces AOX protein. It is thought that the AOX pathway provides 390 metabolic flexibility and gives the organism the ability to survive under a multitude of 391 environmental stressors (Vanlerberghe, 2013). In addition, this system also represents 392 an opportunity to study the post-translational regulation of the AOX protein in T. 393 californicus.

394 Future Direction and Applications

395 Knowledge about animal AOX also has applications in human and animal 396 medicine. Comparative research may aid in the treatment of diseases caused by 397 parasitic protists, where AOX is a current target of drug design (May et al., 2017). AOX 398 research could lead to the development of anti-parasitic drugs that can be used to kill 399 parasitic copepods that live on the skin of economically valuable fish species. Humans 400 do not contain AOX, and the information gathered from our research may eventually 401 contribute to the treatment of mitochondrial dysfunctions and disorders in humans using 402 AOX gene therapy (El-Khoury et al., 2013; Fernandez-Ayala et al., 2009; Hakkaart et 403 al., 2006; Kemppainen et al., 2014). Heterologous expression systems provide a 404 potential route of expression to rescue electron flow and test hypotheses in order to 405 mitigate the deleterious complications involved with respiratory chain dysfunctions (such 406 as Parkinson's, diabetes mellitus, and deafness) (Hakkaart et al, 2006; Kemppainen et 407 al., 2014), but this requires extensive knowledge about the regulation of AOX gene 408 expression and the post-translational regulation of the AOX protein.

We believe that *T. californicus* is an excellent model system in which to study AOX. These animals can be used to study the physiological role of AOX and to test functional hypotheses about the enzyme. This will lead to an increase in our knowledge of AOX in general and contribute to the assessment its future applications in healthcare and aquaculture.

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- 419 The authors report no conflict of interest.

420 **Figure Captions**

Figure 1. Reverse-transcriptase PCR products using *T. californicus* AOX gene specific
primers with DNase treated RNA. Lane 1, 5 μL 100bp DNA ladder; lane 2, 10 μL AOX1
primer set product (232bp product size); lane 3, 10 μL AOX2 primer set product (344bp
product size); lane 4, 10 μL AOX3 primer set product (503bp product size); lane 5, 10
μL positive control from kit (323bp product size).

426 Figure 2. Detection of *T. californicus* AOX protein by reducing SDS-PAGE and Western 427 blotting using a plant AOX antibody. All T. californicus samples per well are 14µL of 428 protein samples derived from 0.05 g of copepods of mixed developmental stages 429 subjected to either sonication (S), homogenization (H), or a combination of the two (H & 430 S). The protein sample buffer volume used (in microliters) is indicated by the number 431 below the isolation method. Ladder lanes contain 5µL of the Precision Plus Protein 432 Western C Standards. The positive control is 14 µL of a protein sample from isolated 433 mitochondria from Saccharomyces cerevisiae overexpressing the Crassostrea gigas 434 AOX.

Figure 3. Detection of *T. californicus* AOX protein by reducing SDS-PAGE and Western
blotting using a plant AOX antibody under 24 hours exposure to different temperatures.
All *T. californicus* samples per well are 20 µg of protein sample derived from 0.05-0.10 g
of copepods of mixed developmental stages. Ladder lanes contain 5µL of the Precision

Plus Protein Western C Standards. Temperature treatments are indicated above eachsample well.

Figure 4. Detection of *T. californicus* AOX protein by reducing SDS-PAGE and Western
blotting using a plant AOX antibody under 1 week exposure to different temperatures.
All *T. californicus* samples per well are 20 µg of protein sample derived from 0.05-0.10 g
of copepods of mixed developmental stages. Ladder lanes contain 5µL of the Precision
Plus Protein Western C Standards. Temperature treatments are indicated above each
sample well.

Figure A1. cDNA and predicted protein sequences of the AOX from *Tigriopus*

448 *californicus*. The highlighted region contains the C-terminal motif that is highly

449 conserved in animal AOX proteins.

450 Figure A2. Protein samples run on reducing SDS-PAGE. Ladder lanes contain 5µL of

451 $\,$ the Precision Plus Protein Western C Standards. The positive control is 14 μL of a

452 protein sample from isolated mitochondria from *Saccharomyces cerevisiae*

453 overexpressing the Crassostrea gigas AOX. All T. californicus samples per well are

454 14μL of protein samples derived from 0.05 g of copepods of mixed developmental

455 stages subjected to either sonication (S), homogenization (H), or a combination of the

456 two (H & S). The protein sample buffer volume used (in microliters) is indicated by the

457 number below the isolation method.

458 Figure A3. Protein samples run on reducing SDS-PAGE and Western blotted using a

459 plant AOX antibody. The ladder lane contains 5µL of the Precision Plus Protein Western

460 C Standards. The algae lane contains a sample derived from the *Spirulina* used to feed

| 461 | the copepods. This sample was sonicated in the same manner as the copepod samples |
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| 462 | and 14µL were loaded into the well. |

- Figure A4. Protein samples run on reducing SDS-PAGE. Ladder lanes contain 5µL of
- the Precision Plus Protein Western C Standards. All *T. californicus* samples per well are
- 465 20 µg of protein samples derived from 0.05-0.010 g of copepods of mixed
- 466 developmental stages subjected to 24 hours at the temperatures indicated above the
- 467 wells of the gel.
- Figure A5. Protein samples run on reducing SDS-PAGE. Ladder lanes contain 5µL of
- the Precision Plus Protein Western C Standards. All *T. californicus* samples per well are
- 470 20 μg of protein samples derived from 0.05-0.010 g of copepods of mixed
- 471 developmental stages subjected to 1 week at the temperatures indicated above the
- wells of the gel.
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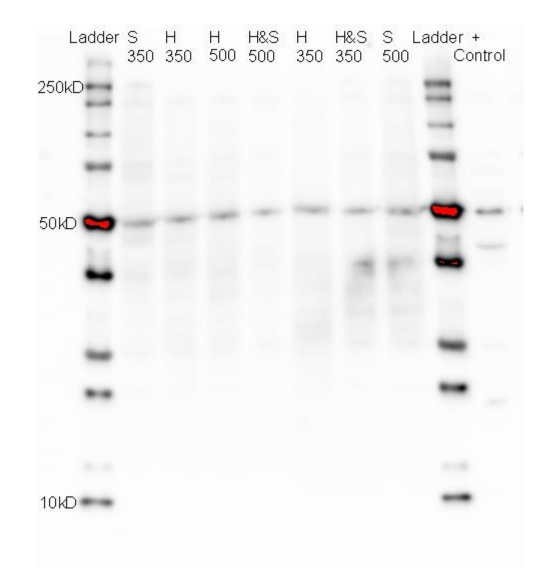
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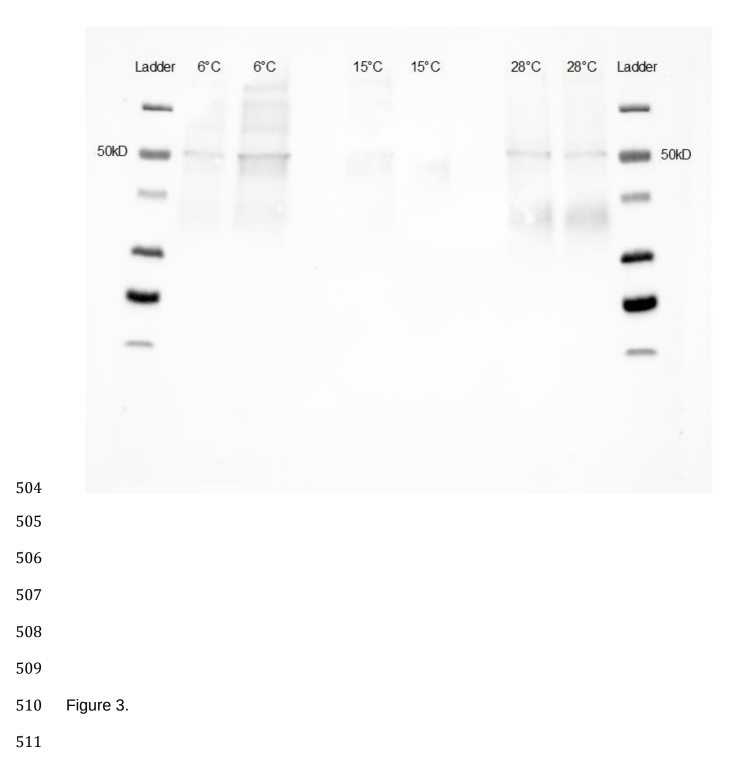
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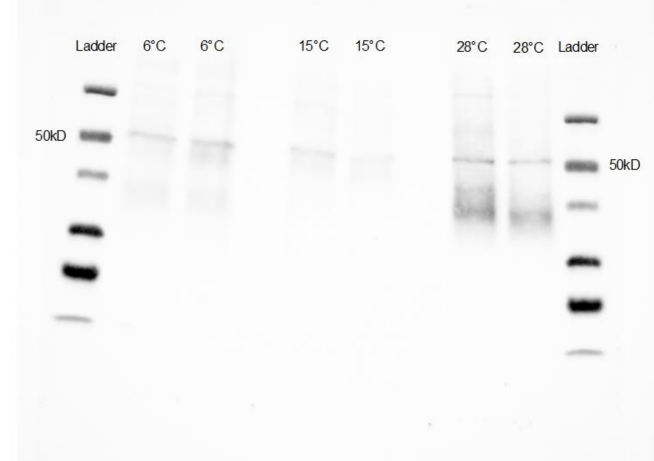
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- 497 Figure 1.



502 Figure 2.

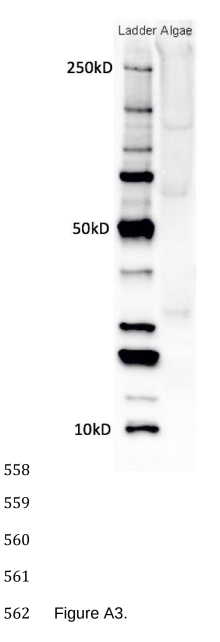


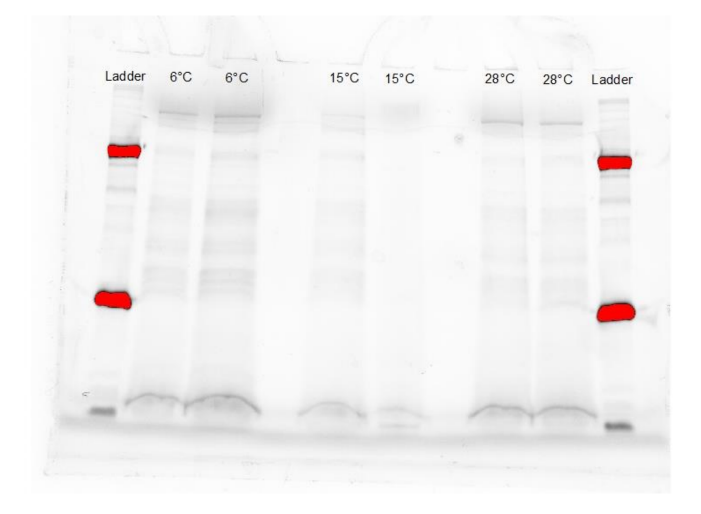


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- 523 <u>Tigriopus californicus 551bp cDNA sequence</u>
- 524 GAATGCACGTGATCATGGATGGATCCACACTCTCCTGGAAGAAGCGGAGAATGAA
- 525 AGGATGCATCTGATGACCTTCATGCGACTCCGAAGACCCGGGCCCATTTTCCGAG
- 526 GCACCGTGATCCTGACTCAATGGCTGTTCACATTTACGTTCTCATTCGCTTACATAC
- 528 ACTTACACTCACATCCTGGAAGAAATCGACGCAGGACGATTGCCCATGTGGAAGA
- 529 CCTTGCCAGCTCCGGAATTGGCCATCAAGTATTGGAGATTGCCCGAAGACGCCAA
- 530 GATGCGGGAAGTCATTTTGGCAATCCGAGCCGATGAAGCTCATCATCGGCTTGTG
- 531 AATCACACCCTTGGATCGATGGACCTCAAATCAGACAATCCTTTTGAGAAAGGGAA
- 532 ATAACTTTTGTCTTCGGCTCAGATACAAATTTAATGGTCAATAACAAGTATTCCATG
- 533 GTAAAAACGAACGGAGGTTGAGAAATATAAATAAATTGTCGAGTCTTTTC
- 534
- 535 <u>Tigriopus californicus predicted protein sequence</u>
- 536 NERMHLMTFMRLRRPGPIFRGTVILTQWLFTFTFSFAYILSPNFCHRFVGYLEEQAVVT
- 537 YTHILEEIDAGRLPMWKTLPAPELAIKYWRLPEDAKMREVILAIRADEAHHRLVNHTLGS
- 538 MDLKDNPFEKGK
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- 540 Figure A1.
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| Ladder | S, 500 | H, 350 | H, 500 | H&S 500 | H, 350 | H&S, 350 | S, 500 ⁻ | Ladder | Positive control |
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Table 1. Gene-specific primers designed for *Tigriopus californicus* AOX amplification.

| Primer Name | Forward Primer Sequence (5' → 3') | Reverse Primer Sequence (5' → 3') | Expected cDNA Product Size (bp) |
|----------------|--------------------------------------|--------------------------------------|--|
| AOX1 | CCTGACTCAATGGCTGTTCA | CAAAATGACTTCCCGCATCT | 232 |
| AOX2 | TGCCACAGATTTGTTGGGTA | TTCTCAACCTCCGTTCGTTT | 344 |
| AOX3 | GGATCCACACTCTCCTGGAA | CTCAACCTCCGTTCGTTTT | 503 |

Table 2. Putative AOX sequences in copepods recovered using bioinformatics searchesof public databases.

| Order | Family | Species | Representative Accession Number | Iron- binding sites present in translated AOX protein | Information on Source Material |
|------------|---------------|--------------------------|---------------------------------------|---|---|
| Calanoida | Acartiidae | Acartia tonsa | HAGX01001020 | All 4 | All developmental stages |
| | Calanidae | Calanus finmarchicus | GAXK01135432 | All 4 | C5 copepodite, Trondheim, Norway |
| | | Calanus glacialis | HACJ01021827 | IBS 1, 2, & 3 | Whole organism, Norway |
| | | Neocalanus flemingeri | GRUD01065088 | All 4 | Adult female |
| | Caligidae | Caligus rogercresseyi | GAZX01011857 | IBS 1, 2, & 3 | Mixed developmental stages and sexes |
| | Metridinidae | Pleuromamma xiphias | GFCI01346245 | All 4 | Whole adults, both sexes, BATS, Atlantic Ocean |
| | Pontellidae | Labidocera madurae | GFWO01190644 | All 4 | |
| | Temoridae | Eurytemora affinis | GBGO01037152 | All 4 | Whole females |
| Cyclopoida | Cyclopettidae | Paracyclopina nana | GCJT01014574 | All 4 | Whole animals, Gangneung, South Korea |
| | Cyclopidae | Eucyclops serrulatus | GARW01011119 | All 4 | |
| | Oithonidae | Oithona nana | FTRT01002078 | IBS 1 & 2 | |

| Harpacticoida | Harpacticidae | Tigriopus californicus | JW502496 | All 4 | San Diego population |
|---------------|---------------|----------------------------|--------------|-------|--|
| | Harpacticidae | Tigriopus japonicas | GCHA01002206 | All 4 | Whole animals, Gangneung, South Korea |
| | Harpacticidae | Tigriopus sp. 1 SL-2012 | GDFW01016749 | All 4 | Tidal pool, Maxwell Bay, Antarctica |

- ⁶⁰³ ¹ Trace sequence archive database at NCBI
- 604 ² Nucleotide database at NCBI

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