

1 **Elucidating the acid-base mechanisms underlying otolith overgrowth in fish**  
2 **exposed to ocean acidification**

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9

10 **Abstract**

11 Over a decade ago, ocean acidification (OA) exposure was reported to induce  
12 otolith overgrowth in teleost fish. This phenomenon was subsequently confirmed in  
13 multiple species; however, the underlying physiological causes remain unknown. Here,  
14 we report that splitnose rockfish (*Sebastes diploproa*) exposed to ~1,600  $\mu\text{atm}$   $p\text{CO}_2$   
15 (pH ~7.5) were able to fully regulate the pH of both blood and endolymph (the fluid that  
16 surrounds the otolith within the inner ear). However, while blood was regulated around  
17 pH 7.80, the endolymph was regulated around pH ~8.30. These different pH setpoints  
18 result in increased  $p\text{CO}_2$  diffusion into the endolymph, which in turn leads to  
19 proportional increases in endolymph  $[\text{HCO}_3^-]$  and  $[\text{CO}_3^{2-}]$ . Endolymph pH regulation  
20 despite the increased  $p\text{CO}_2$  suggests enhanced  $\text{H}^+$  removal. However, a lack of  
21 differences in inner ear bulk and cell-specific  $\text{Na}^+/\text{K}^+$ -ATPase and vacuolar type  $\text{H}^+$ -  
22 ATPase protein abundance localization pointed out to activation of preexisting  
23 ATPases, non-bicarbonate pH buffering, or both, as the mechanism for endolymph pH-

24 regulation. These results provide the first direct evidence showcasing the acid-base  
25 chemistry of the endolymph of OA-exposed fish favors otolith overgrowth, and suggests  
26 that this phenomenon will be more pronounced in species that count with more robust  
27 blood and endolymph pH regulatory mechanisms.

28

29 **Keywords**

30 Endolymph, climate change, calcification, biomineralization, rockfish, carbon dioxide

31

32 **Introduction**

33 The inner ear of teleost fishes contains three pairs of otoliths that contribute to  
34 hearing and maintaining balance. Otoliths are comprised of calcium carbonate ( $\text{CaCO}_3$ )  
35 embedded within a protein matrix, and are biomineralized within an acellular fluid called  
36 the endolymph (Payan et al., 2004a). Otoliths are biomineralized in a successive ring  
37 pattern correlated with the fish growth rate (Campana and Neilson, 1985; Kalish, 1989;  
38 Pannella, 1971), which are used by scientists and fishery managers to estimate fish age  
39 and length (Campana, 2001; Campana and Thorrold, 2001), estimate recruitment, and  
40 set fishery-specific catch limits (Methot, 2015; Vitale et al., 2019).

41 Originally, it was predicted that  $\text{CO}_2$ -induced ocean acidification (OA) would  
42 impair otolith biomineralization because the associated decreases in seawater pH and  
43  $[\text{CO}_3^{2-}]$  hamper  $\text{CaCO}_3$  precipitation (Ishimatsu et al., 2008). However, subsequent  
44 studies reported that fish exposed to OA developed enlarged otoliths (S. Bignami et al.,  
45 2013; Sean Bignami et al., 2013; Checkley et al., 2009; Faria et al., 2017; Hurst et al.,  
46 2012; Réveillac et al., 2015; Shen et al., 2016). These findings led to a broader

47 awareness that otolith biomineralization is strongly linked to endolymph and blood  
48 chemistries, and to the hypothesis that biological regulation of endolymph pH could lead  
49 to increased  $[CO_3^{2-}]$  resulting in otolith overgrowth (Checkley et al., 2009). In addition,  
50 fish exposed to hypercapnia typically accumulate  $[HCO_3^-]$  in their plasma to compensate  
51 the respiratory acidosis; this could result in enhanced  $HCO_3^-$  flux into the endolymph  
52 and further contribute to otolith overgrowth (Heuer and Grosell, 2014). However,  
53 experimental support for these hypotheses is lacking, as there are no reports of  
54 endolymph acid-base parameters under OA-relevant conditions, and only a few studies  
55 have measured blood acid-base parameters in fish exposed to OA-relevant  $CO_2$  levels  
56 (Esbaugh et al., 2016, 2012; Montgomery et al., 2019). This knowledge gap is in large  
57 part due to the disrupting effects of blood sampling by caudal puncture on the acid-base  
58 status of fish internal fluids and the challenge of blood vessel cannulation in small fish  
59 and species with convoluted dorsal aorta anatomy, coupled with the difficulty of  
60 collecting sufficient endolymph for analyses. Moreover, the cellular heterogeneity of the  
61 inner ear complicates the quantification of ionocyte-specific responses using standard  
62 molecular and biochemical assays on bulk tissue. As a result, the underlying acid-base  
63 and physiological causes of OA-induced otolith overgrowth remain unknown.

64 The chemistry of the endolymph is actively controlled by the inner ear epithelium  
65 to maintain acid-base conditions that promote biomineralization, namely, higher pH,  
66  $[HCO_3^-]$ ,  $[CO_3^{2-}]$ , and total  $CO_2$  than the blood (Payan et al., 1999, 1997; Takagi, 2002;  
67 Takagi et al., 2005). This gradient is actively maintained by two types of ion-transporting  
68 cells (“ionocytes”): the Type-I ionocyte, which transports  $K^+$  and  $Cl^-$  into the endolymph  
69 and removes  $H^+$  powered by  $Na^+/K^+$ -ATPase (NKA) (Kwan et al., 2020; Mayer-Gostan

70 et al., 1997; Payan et al., 1997; Takagi, 1997) and the Type-II ionocyte, which secretes  
71  $\text{HCO}_3^-$  into the endolymph driven by V-type  $\text{H}^+$ -ATPase (VHA) (Kwan et al., 2020;  
72 Mayer-Gostan et al., 1997; Payan et al., 1997; Takagi, 1997; Tohse et al., 2006, 2004).  
73 However numerous other cells within the inner ear organ also express NKA and VHA,  
74 including the sensory hair cells and the endothelial cells that make up the blood vessels  
75 (Kwan et al., 2020; Mayer-Gostan et al., 1997; Shiao et al., 2005).

76 In the current study, splitnose rockfish (*Sebastes diploproa*) were exposed to  
77  $\sim 1,600 \mu\text{atm CO}_2$  ( $\text{pH} \sim 7.5$ ), a condition readily experienced in their natural habitat  
78 (Culberson and Pytkowicz, 1970; Love et al., 2002) and predicted for the surface ocean  
79 by the year 2300 (Goodwin et al., 2018). The OA exposure spanned three days, a  
80 duration previously documented to result in otolith overgrowth in fish larvae (Faria et al.,  
81 2017). Blood acid-base chemistry was measured after taken samples using a  
82 benzocaine-based anesthetic protocol that yields measurements comparable to those  
83 achieved using cannulation (Montgomery et al., 2019). Additionally, we took advantage  
84 of the large rockfish inner ear organ to collect sufficient endolymph for acid-base  
85 chemistry analysis, and inner ear tissue for quantification of NKA and VHA protein  
86 abundances. Finally, we performed immunohistochemical analyses on six inner ear cell  
87 types to explore potential cell-specific changes in protein expression patterns. This  
88 multidimensional approach allowed us to explore the mechanistic acid-base causes that  
89 underlie otolith overgrowth in fish exposed to OA.

90

## 91 **Methods**

### 92 *Specimens*

93 Juvenile splitnose rockfish (*S. diploproa*) were caught from drifting kelp paddies  
94 off the shores of La Jolla and raised in the Hubbs Experimental Aquarium (La Jolla,  
95 USA) in accordance to the permit (#SCP13227) issued by the California Department of  
96 Fish and Wildlife. Rockfish were raised for >2 years in a flow-through system with  
97 seawater continuously pumped from the Scripps Coastal Reserve, and were fed frozen  
98 market squids and food pellets (EWOS, Cargill Incorporated, Minneapolis, MN, USA).  
99 Average rockfish total length ( $12.33 \pm 0.16$  cm) and weight ( $49.11 \pm 2.18$  g) (N=9) were  
100 not significantly different between treatments. All experiments were approved under the  
101 Institutional Animal Care and Use Committee protocol (#S10320) by the Scripps  
102 Institution of Oceanography, University of California San Diego animal care committee.

103

104 *Experimental Aquarium Setup*

105 Two header tanks were supplied with ambient seawater from the Scripps Coastal  
106 Reserve, one was not manipulated and was considered as the control condition. The  
107 other header tank was bubbled with CO<sub>2</sub> using a pH-stat system (IKS Aquastar,  
108 Karlsbad, Germany) to maintain a seawater pH ~7.5 and generate the OA condition.  
109 Temperature and pH were continuously monitored and recorded every 2 minutes using  
110 the IKS Aquastar system (figure S1). Discrete seawater samples were collected from  
111 header tanks at the beginning and end of each experiment, and analyzed for alkalinity  
112 (via titration with LabView software Version 2.9j; National Instruments, Austin, Texas,  
113 United States), pH (using the indicator dye purified m-cresol purple (Liu et al., 2011) in  
114 an Agilent 8453 spectrophotometer (Agilent, Santa Clara, CA, USA)), and salinity (by  
115 converting density measurements using Mettler Toledo DE-45 (Mettler-Toledo,

116 Columbus, Ohio, United States)) by the Dickson Lab (Scripps Institution of  
117 Oceanography). The pH values from the discrete seawater samples were used to  
118 validate and back-correct the IKS pH measurements. Subsequently, the pH, alkalinity,  
119 and salinity values were used to calculate  $p\text{CO}_2$  using CO2SYS (Lewis and Wallace,  
120 1998). These analyses indicated control pH and  $p\text{CO}_2$  levels of  $7.89 \pm 0.012$  and  $571.90$   
121  $\pm 4.88 \mu\text{atm}$ , respectively, which are typical for La Jolla, USA (Frieder et al., 2012;  
122 Hofmann et al., 2011; Takeshita et al., 2015). In contrast, pH and  $p\text{CO}_2$  in the OA  
123 treatment were  $7.49 \pm 0.01$  and  $1,591.56 \pm 18.58 \mu\text{atm}$ , respectively (table S1).

124 Each header tank supplied water to three opaque 3-L experimental tanks at a  
125 flow rate of  $0.3\text{-L min}^{-1}$ . Individual rockfish were acclimated within an experimental tank  
126 for 12 hours, followed by a 72-hour exposure to control or OA conditions. To ensure  
127 similar metabolic state among individuals, rockfish were not fed during the 48 hours  
128 prior to the acclimations or during the experiment. Three separate experiments were  
129 conducted during March 2020, each time with three control and three OA-exposed fish.  
130 No mortality was observed.

131

132 *Blood, endolymph, and inner ear sampling*

133 Sampling and acid-base determinations were performed in a temperature-  
134 controlled room at  $18^\circ\text{C}$  (i.e. same as that of seawater). Fish were anesthetized by  
135 stopping the seawater flow into the individual experimental tank and slowly adding  
136 benzocaine through to achieve a final concentration of  $0.15 \text{ g/L}$ . After fish lost  
137 equilibrium ( $\sim 5$  minutes), they were moved to a surgery table where the gills were  
138 irrigated with aerated seawater from their respective treatment (control or OA)

139 containing benzocaine (0.05 g/L) using a pump (Harter et al., 2021; Montgomery et al.,  
140 2019, 2022). Blood was drawn from the caudal vein using a heparinized syringe and pH  
141 was immediately measured using a microelectrode (Orion<sup>TM</sup> PerpHecT<sup>TM</sup> Ross<sup>TM</sup>,  
142 ThermoFisher Scientific, Waltham, MA, USA). Next, blood was centrifuged for 1 minute  
143 at 6,000xg using a microcentrifuge (VWR Kinetic Energy 26 Joules, Radnor, PA, USA),  
144 and the resulting plasma was measured for total CO<sub>2</sub> (TCO<sub>2</sub>) using a carbon dioxide  
145 analyzer (Corning 965, Ciba Corning Diagnostic, Halstead, Essex, United Kingdom).  
146 After blood sampling (N=8-9), the fish was euthanized by spinal pithing, and the gills  
147 were quickly removed. Endolymph (N=7-8) was drawn using a heparinized syringe from  
148 the ventral side of the skull, and pH and TCO<sub>2</sub> were measured as described above.  
149 CO<sub>2</sub> loss was minimized by measuring endolymph within 3 minutes after spinal pithing.  
150 Inner ear tissue was either flash frozen in liquid nitrogen and stored at -80°C, or fixed in  
151 4% paraformaldehyde (8 hours at 4°C), incubated in 50% ethanol (8 hours at 4°C), then  
152 stored in 70% ethanol until processing.

153

154 *HCO<sub>3</sub><sup>-</sup>, CO<sub>3</sub><sup>2-</sup>, and pCO<sub>2</sub> calculation*

155 Blood and endolymph pH and TCO<sub>2</sub> values were used to calculate [HCO<sub>3</sub><sup>-</sup>],  
156 [CO<sub>3</sub><sup>2-</sup>], and pCO<sub>2</sub> using the Henderson-Hasselbalch equation. The solubility coefficient  
157 of CO<sub>2</sub> (plasma: 0.0578 mmol L<sup>-1</sup> Torr<sup>-1</sup>; endolymph: 0.0853 mmol L<sup>-1</sup> Torr<sup>-1</sup>), ionic  
158 strength (plasma: 0.15 mol L<sup>-1</sup>, endolymph: 0.18 mol L<sup>-1</sup>), pK<sub>1</sub>' (plasma: ~6.20,  
159 endolymph: ~6.16), and pK<sub>2</sub>' (plasma: ~9.76, endolymph: ~9.71) were based upon  
160 (Boutilier et al., 1984) and (Takagi, 2002) for blood and endolymph, respectively. The

161  $[\text{Na}^+]$  (plasma: 170 mmol L<sup>-1</sup>, endolymph: 100 mmol L<sup>-1</sup>) used for calculating pK<sub>1'</sub> was  
162 based upon (Payan et al., 1997).

163

164 *Antibodies*

165 NKA was immunodetected using a monoclonal  $\alpha$ 5 mouse antibody raised against  
166 the  $\alpha$ -subunit of chicken NKA (a5, Developmental Studies Hybridoma Bank, Iowa City,  
167 IA, USA; (Lebovitz et al., 1989)), whereas the  $\beta$ -subunit of VHA was immunodetected  
168 using a custom-made polyclonal rabbit antibody (epitope: AREEVPGRRGFPGYC;  
169 GenScript, Piscataway, USA). These antibodies have been previously used in the inner  
170 ear of the Pacific chub mackerel (*Scomber japonicus*; (Kwan et al., 2020)), and were  
171 validated here for splitnose rockfish (figure S2). Secondary antibodies goat anti-mouse  
172 HRP-linked secondary antibodies (Bio-Rad, Hercules, CA, USA) and goat anti-rabbit  
173 HRP-linked secondary antibodies (Bio-Rad) were used for immunoblotting.

174

175 *Western Blotting and Relative Protein Abundance Analysis*

176 Frozen inner ear samples were immersed in liquid nitrogen, pulverized using a  
177 handheld motorized homogenizer (Kimble®/Kontes, Dusseldorf, Germany), and  
178 suspended in ice-cold homogenization buffer containing protease inhibitors (250 mmol  
179 L<sup>-1</sup> sucrose, 1 mmol L<sup>-1</sup> EDTA, 30 mmol L<sup>-1</sup> Tris, 10 mmol L<sup>-1</sup> benzamidine hydrochloride  
180 hydrate, 1 mmol L<sup>-1</sup> phenylmethanesulfonyl fluoride, 1 mmol L<sup>-1</sup> dithiothreitol, pH 7.5).  
181 Samples were centrifuged at low speed (3,000xg, 10 minutes, 4°C) to remove debris,  
182 and the resulting supernatant was considered the crude homogenate. Total protein  
183 concentration in all fractions was determined by the Bradford assay (Bradford, 1976).

184 Prior to SDS-electrophoresis, samples were mixed with an equal volume of 90% 2x  
185 Laemmli buffer and 10%  $\beta$ -mercaptoethanol, and heated at 70°C for 5 minutes. Proteins  
186 (crude homogenate: 10  $\mu$ g per lane; membrane-enriched fraction: 5  $\mu$ g per lane) were  
187 loaded onto a 7.5% polyacrylamide mini gel (Bio-Rad, Hercules, CA, USA) – alternating  
188 between control and high CO<sub>2</sub> treatments to avoid possible gel lane effects. The gel ran  
189 at 200 volts for 40 minutes, and the separated proteins were then transferred to a  
190 polyvinylidene difluoride (PVDF) membrane using a wet transfer cell (Bio-Rad) at 100  
191 mAmps at 4°C overnight. PVDF membranes were incubated in tris-buffered saline with  
192 1% tween (TBS-T) with milk powder (0.1 g/mL) at RT for 1 hour, then incubated with  
193 primary antibody (NKA: 10.5 ng/ml; VHA: 3  $\mu$ g/ml) in blocking buffer at 4°C overnight.  
194 On the following day, PVDF membranes were washed with TBS-T (three times; 10  
195 minutes each), incubated in blocking buffer with their respective secondary antibodies  
196 (1:10,000) at RT for 1 hour, and washed again with TBS-T (three times; 10 minutes  
197 each). Bands were made visible through addition of ECL Prime Western Blotting  
198 Detection Reagent (GE Healthcare, Waukesha, WI) and imaged and analyzed in a  
199 BioRad Universal III Hood using Image Lab software (version 6.0.1; BioRad). Following  
200 imaging, the PVDF membrane was incubated in Ponceau stain (10 minutes, room  
201 temperature) to estimate protein loading. Relative protein abundance (N=6-8) were  
202 quantified using the Image Lab software (version 6.0.1; BioRad) and normalized by the  
203 protein content in each lane.

204

205 *Whole-mount immunohistochemistry and confocal microscopy*

206 Immunolabeling was performed based on the protocol described in Kwan *et al.*,  
207 (2020) for tissue sections and optimized for whole tissues as follows. Fixed inner ear  
208 tissue was rehydrated in phosphate buffer saline + 0.1% tween (PBS-T) for 10 min.  
209 Autofluorescence was quenched by rinsing in ice-cold PBS-T with sodium borohydride  
210 (1.5 mg/mL; six times; 10 minutes each), followed by incubation in blocking buffer (PBS-  
211 T, 0.02% normal goat serum, 0.0002% keyhole limpet hemocyanin) at room  
212 temperature for one hour. Samples were incubated with blocking buffer containing  
213 primary antibodies (NKA: 40 ng/mL; VHA: 6 µg/mL) at 4°C overnight. On the following  
214 day, samples were washed in PBS-T (three times at room temperature; 10 minutes  
215 each), and incubated with the fluorescent secondary antibodies (1:500) counterstained  
216 with DAPI (1 µg/mL) at room temperature for 1 hour. Samples were washed again in  
217 PBS-T as before and stored at 4°C until imaging.

218 Immunostained inner ear samples were immersed in PBS-T, mounted onto a  
219 depressed glass slide fitted with a glass cover slip (No. 1.5, 0.17 mm) and imaged using  
220 a Zeiss LSM800 inverted confocal microscope equipped with a Zeiss LD LCI Plan-  
221 Apochromat 40x/1.2 Imm Korr DIC M27 objective and Zeiss ZEN 2.6 blue edition  
222 software (Cambridge, United Kingdom). The following channels were used for imaging:  
223 VHA (excitation 493 nm with 1% laser power, emission 517 nm, detection 510– 575  
224 nm), NKA (excitation 577 nm at 1% laser power, emission 603 nm, detection 571–617  
225 nm), and DAPI (excitation 353 nm at 0.7% laser power, emission 465 nm, detection  
226 410–470 nm). Z-stacks (range: ~70–400 optical sections; thickness: ~0.27 µm per  
227 section) of the various inner ear cell types were visualized as maximum intensity  
228 projection, and through orthogonal cuts to capture fluorescent signal across the X-Z and

229 Y-Z planes. Inner ear organs from four control and four OA-exposed rockfish were  
230 imaged.

231

232 **Statistical Analysis**

233 Normality was tested using the Shapiro-Wilk normality test, and homogeneity  
234 was tested using the F-test. Datasets that failed to meet the assumptions of normality  
235 were log- (i.e.  $[\text{CO}_3^{2-}]$ , pH) or inverse-transformed (i.e.  $[\text{H}^+]$ ). Acid-base parameters were  
236 analyzed using two-way analysis of variance (2-way ANOVA), with “CO<sub>2</sub> level” (control  
237 or OA) and “internal fluid” (blood or endolymph) as factors. If significant interaction  
238 effect was detected, subsequent Tukey honest significant difference (HSD) tests were  
239 used. NKA and VHA protein abundances were analyzed using two-tailed Student’s t-  
240 tests. Values are reported as mean  $\pm$  s.e.m., and an alpha of 0.05 was employed for all  
241 analyses. Statistical tests were performed using Prism (version 7.0a) and R (version  
242 4.0.3; R Development Core Team, 2013).

243

244 **Results and discussion**

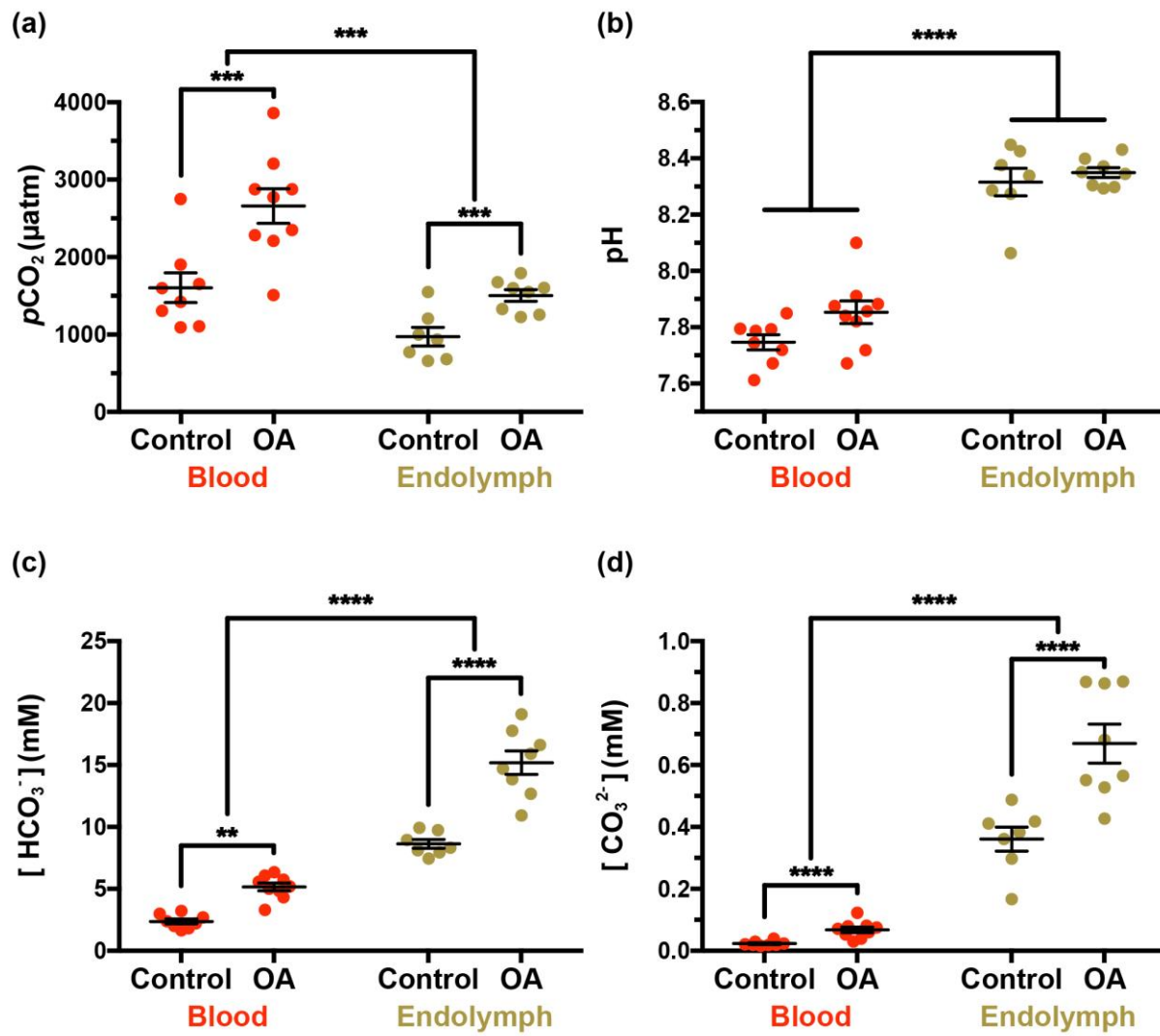
245 The difference in seawater  $p\text{CO}_2$  between the control and OA-condition was  
246  $\sim 1,000 \mu\text{atm}$ , which induced an equivalent elevation in blood  $p\text{CO}_2$  from  $1,603.25 \pm$   
247  $190.69 \mu\text{atm}$  in control fish to  $2,659.20 \pm 223.87 \mu\text{atm}$  in OA-exposed fish (figure 1a).  
248 However, blood pH was fully regulated (control:  $7.75 \pm 0.03$ ; OA:  $7.85 \pm 0.04$ ) (figure  
249 1b). As is typical for regulation of blood acidosis (Tresguerres and Hamilton, 2017), OA-  
250 exposed fish demonstrated a significant accumulation of  $\text{HCO}_3^-$  in blood plasma, from  
251  $2.37 \pm 0.20 \text{ mM}$  in control fish up to  $5.16 \pm 0.31 \text{ mM}$  in OA-exposed fish (figure 1c). This

252 response matches the magnitude of the hypercapnic stress according to classic  
253 Davenport acid-base physiology, as well as the three previous studies on blood acid-  
254 base chemistry in fish exposed to OA-relevant CO<sub>2</sub> levels (Esbaugh et al., 2016, 2012;  
255 Montgomery et al., 2019). In addition, the increased plasma TCO<sub>2</sub> at unchanged pH led  
256 to the tripling of plasma [CO<sub>3</sub><sup>2-</sup>] from ~0.02 to ~0.07 mM (figure 1d). These increases in  
257 blood [HCO<sub>3</sub><sup>-</sup>] and [CO<sub>3</sub><sup>2-</sup>] may contribute to the skeletal hypercalcification (Di Santo,  
258 2019) and deformities (Pimentel et al., 2014) reported in some OA-exposed fishes.

259 The endolymph of control rockfish had higher TCO<sub>2</sub> compared to the blood (9.06  
260  $\pm$  0.38 vs 2.46  $\pm$  0.20 mM; figure S3) and also a higher pH (8.32  $\pm$  0.05 vs. 7.75  $\pm$  0.03),  
261 resulting in lower pCO<sub>2</sub> (971.38  $\pm$  120.70 vs. 1,603.25  $\pm$  190.69  $\mu$ atm), higher [HCO<sub>3</sub><sup>-</sup>]  
262 (8.63  $\pm$  0.35 vs. 2.37  $\pm$  0.20 mM), and much higher [CO<sub>3</sub><sup>2-</sup>] (0.36  $\pm$  0.04 vs 0.02  $\pm$  0.01  
263 mM) (figure 1a-d). Importantly, these measurements revealed higher pH and lower  
264 pCO<sub>2</sub>, TCO<sub>2</sub>, [HCO<sub>3</sub><sup>-</sup>] and [CO<sub>3</sub><sup>2-</sup>] compared to previous studies that collected  
265 endolymph without previously anesthetizing the fish (Payan et al., 1999, 1997), and to  
266 others that used 2-phenoxyethanol as anesthetic but did not irrigate the gills during  
267 endolymph collection (Takagi, 2002; Takagi et al., 2005) (table S2). This finding  
268 highlights the crucial importance of sampling procedures for accurate acid-base  
269 measurements in fish physiological fluids. Indeed, fish struggling during handling and  
270 hypoxia due to gill collapse during emersion are known to greatly affect blood acid-base  
271 measurements, and our results indicate that these disturbances extend to the  
272 endolymph.

273

274 In rockfish exposed to OA, endolymph  $p\text{CO}_2$  increased from  $971.38 \pm 120.70$  to  
275  $1503.21 \pm 73.72$   $\mu\text{atm}$  (figure 1a). Crucially, this  $\sim 500$   $\mu\text{atm}$  increase was half of that  
276 observed in the blood and therefore the  $p\text{CO}_2$  difference between blood and endolymph  
277 increased from  $\sim 600$  to  $\sim 1,100$   $\mu\text{atm}$ , which is predicted to induce a proportional  
278 increase in  $\text{CO}_2$  flux into the endolymph following Fick's law of diffusion. Endolymph  
279  $\text{TCO}_2$  in OA-exposed rockfish also nearly doubled (control:  $9.06 \pm 0.38$  mM; OA=  $15.96$   
280  $\pm 1.02$  mM; figure S3) and, since pH remained unchanged at  $\sim 8.30$  pH (figure 1b), it  
281 was reflected as increased  $[\text{HCO}_3^-]$  (control:  $8.63 \pm 0.35$  mM, OA:  $15.19 \pm 0.95$  mM vs)  
282 (figure 1c) and  $[\text{CO}_3^{2-}]$  (control:  $0.36 \pm 0.04$  mM; OA  $0.67 \pm 0.06$  mM) (figure 1d). Since  
283 aragonite saturation state ( $\Omega_{\text{aragonite}}$ ) is directly proportional to  $[\text{CO}_3^{2-}]$ , it implies that  
284 biomineralization in the endolymph of OA-exposed fish is nearly twice more favorable  
285 than in that of control fish. To our knowledge, this is the first direct evidence that the  
286 acid-base chemistry in the endolymph of OA-exposed fish favors otolith overgrowth.  
287



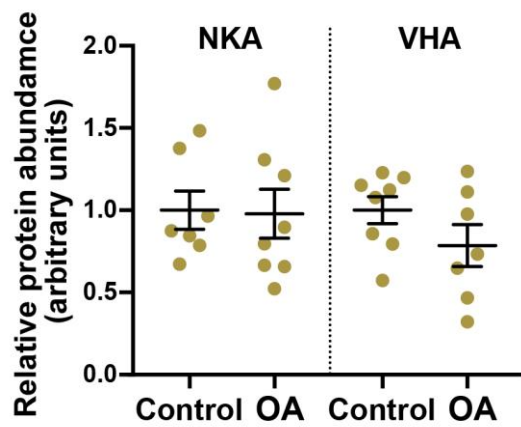
288

289 **Figure 1:** Blood and endolymph acid-base parameters in control and OA-exposed  
 290 rockfish. **A)**  $p\text{CO}_2$ , **B)** pH, **C)**  $[\text{HCO}_3^-]$ , and **D)**  $[\text{CO}_3^{2-}]$ . Data is presented as mean and  
 291 s.e.m. for each group and the individual measurements are shown as red (blood) or  
 292 beige (endolymph) points ( $N= 7-9$ ). Statistical significance between fluids, and between  
 293 treatments for a given fluid are indicated by the connecting lines and asterisks (2-way  
 294 ANOVA, \* $p<0.05$ , \*\* $p<0.005$ , \*\*\* $p<0.001$ , \*\*\*\* $p<0.0001$ ). Statistical details are reported  
 295 in tables S3-S5, and  $\text{TCO}_2$ ,  $[\text{CO}_2]$  and  $[\text{H}^+]$  are shown in figure S3.

296

297 The increased  $p\text{CO}_2$  diffusive rate into the endolymph and subsequent  
298 generation of  $\text{H}^+$  as a result of  $\text{CO}_2$  hydration and  $\text{CaCO}_3$  biomineralization would be  
299 expected to induce a decrease in pH. Thus, the lack of change in endolymph pH in OA-  
300 exposed rockfish indicates robust pH regulation. Hence, we hypothesized that OA-  
301 exposed fish may have increased abundance of NKA and VHA, as these ATPases are  
302 proposed to provide the driving force for transepithelial  $\text{H}^+$  and  $\text{HCO}_3^-$  transport across  
303 the inner ear epithelium (Kwan et al., 2020; Mayer-Gostan et al., 1997; Payan et al.,  
304 1997; Shiao et al., 2005). However, Western blotting on bulk inner ear tissue revealed  
305 no significant differences between control and OA-exposed fish (figure 2, table S6).

306



307  
308 **Figure 2:**  $\text{Na}^+/\text{K}^+$ -ATPase (NKA) and V-type  $\text{H}^+$ -ATPase (VHA) protein abundance in  
309 the inner ear organ of control and OA-exposed rockfish. Data is presented as mean and  
310 s.e.m. and the individual measurements are shown as beige points ( $N= 7-8$ ). Relative  
311 protein abundance was calculated for each ATPase; NKA and VHA abundances are not  
312 comparable to each other. There were no significant differences for NKA ( $p=0.9104$ ) or  
313 VHA ( $p=0.1695$ ). Statistical details are reported in tables S6.

314

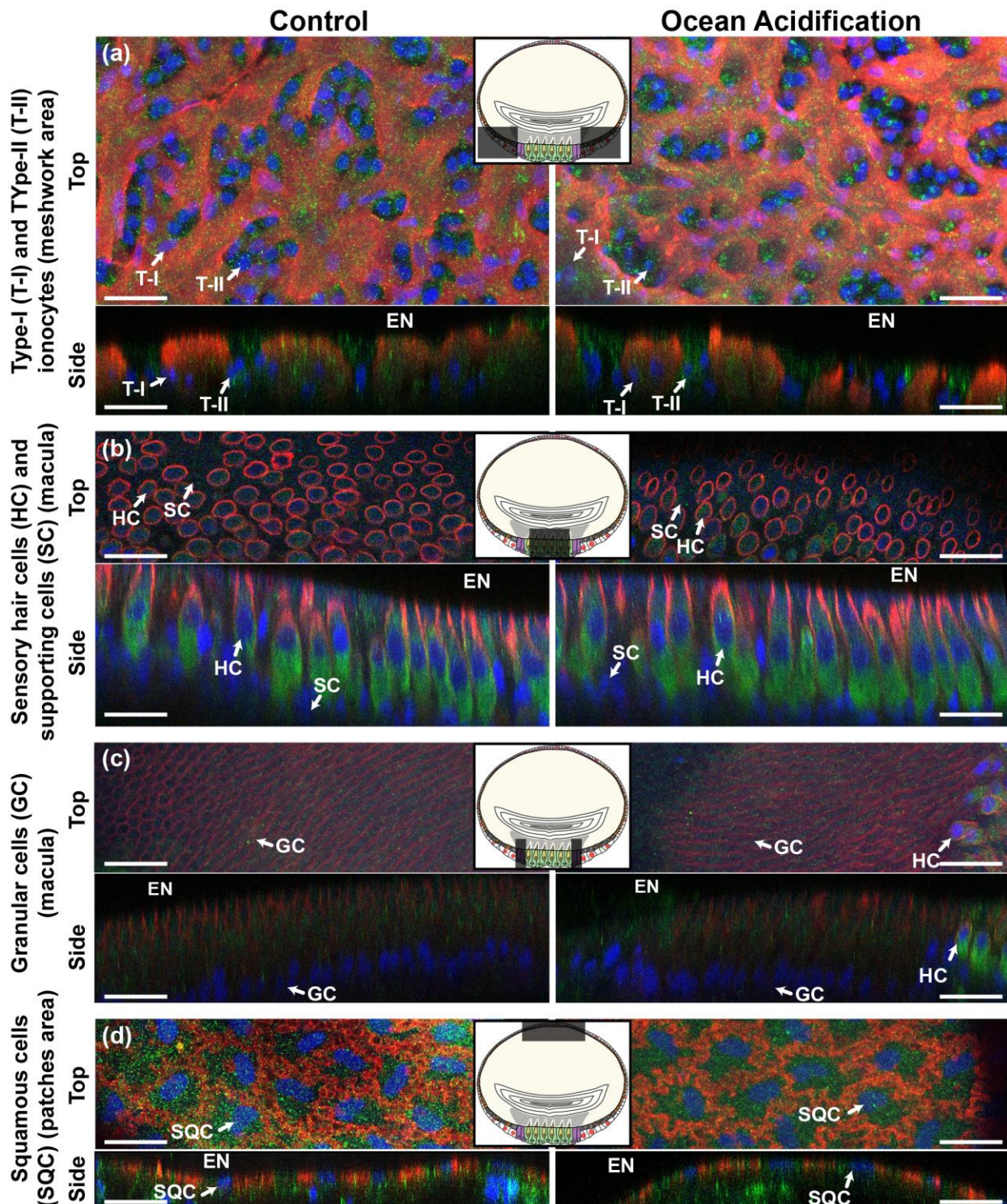
315 Next, we used immunocytochemistry and confocal microscopy to examine  
316 potential changes in NKA and VHA abundance or sub-cellular localization in specific  
317 inner ear epithelial cell types. The NKA and VHA immunostaining in rockfish inner ear  
318 epithelial cells generally matched reports from other fish species (Kwan et al., 2020;  
319 Pisam et al., 1998; Shiao et al., 2005; Takagi, 1997) (figure 3; figure S4), and there  
320 were no apparent differences between control and OA-exposed fish in any cell type in  
321 terms of signal intensity of subcellular localization. The Type-I ionocytes are  
322 characterized by intense NKA signal in their highly infolded basolateral membrane and  
323 by a much fainter cytoplasmic VHA signal (figure 3a). These ionocytes are most  
324 abundant in the meshwork area, where they contact each other by their pseudopods  
325 giving the appearance of an interconnected matrix. The Type-II ionocytes are  
326 interspersed between the Type-I ionocytes in the meshwork area and have cytoplasmic  
327 VHA signal of comparable intensity to that in the Type-I ionocytes; however, they lack  
328 NKA signal (figure 3a). The sensory hair cells are in the macula; they express intense  
329 NKA signal in their basolateral membrane and very intense cytoplasmic VHA signal,  
330 which was especially concentrated towards their basal area consistent with synaptic  
331 vesicles (figure 3b). The supporting cells surround each sensory hair cell; they display  
332 faint cytoplasmic VHA signal and no detectable NKA signal (figure 3b). The granular  
333 cells flank the macula and have a characteristic columnar shape. These cells have faint  
334 NKA signal along their lateral plasma membrane and faint cytoplasmic VHA signal  
335 (figure 3c). Finally, the squamous cells are found in the patches area in the distal side of  
336 the epithelium; these cells are very thin and have NKA signal on their ribbon-like lateral

337 membrane as well as faint cytoplasmic VHA signal (figure 3d). A summary of the NKA  
338 and VHA relative signal intensities in each cell type is reported in table S7.

339 The lack of apparent differences in NKA and VHA abundance and localization  
340 cellular patterns between control and OA-exposed fish indicates that preexisting levels  
341 of NKA and VHA were sufficient to mediate the endolymph pH regulation observed in  
342 our study. Overall, these findings are consistent with models suggesting that  $H^+$   
343 extrusion from the endolymph into the blood passively follows the transepithelial  
344 potential that is established by active  $K^+$  excretion into the endolymph (Payan et al.,  
345 2004). And since the function of the sensory hair cells requires a high  $[K^+]$  in the  
346 endolymph, modulation of inner ear transepithelial potential for the sole purpose of  
347 decreasing  $H^+$  extrusion seems unlikely.

348 In our recent paper (Kwan et al., 2020), we proposed that  $HCO_3^-$  transport into  
349 the endolymph and  $H^+$  removal could be upregulated by insertion of VHA into the  
350 basolateral membrane of Type-II ionocytes; however, we found no evidence for such  
351 mechanism in OA-exposed rockfish (figure 3a, *right panels*). Instead, upregulation of  
352 ATPase activity could have occurred *via* other post-translational modifications or by  
353 increased substrate availability (c.f. Kwan et al., 2021). The expression of carbonic  
354 anhydrases, ion exchangers, and other acid-base relevant proteins must be examined  
355 in future studies, ideally through an approach that includes cell-specific analyses. Lastly,  
356 a contribution of non-bicarbonate buffering to endolymph pH regulation cannot be ruled  
357 out; unfortunately, performing the required titrations are not trivial due to the small  
358 volume of this fluid.

359



361 **Figure 3:** Immunocytochemistry of the inner ear epithelium of control and OA-exposed  
362 rockfish.  $\text{Na}^+/\text{K}^+$ -ATPase is in red, V-type  $\text{H}^+$ -ATPase is in green, and nuclei are in blue.  
363 There were no apparent differences in NKA or VHA signal intensities or localization

364 patterns between control and OA-exposed fish. **(a)** Type-I (T-I) and Type-II ionocytes  
365 (T-II), **(b)** sensory hair cells (HC) and supporting cells (SC), **(c)** granular cells (GC), and  
366 **(d)** squamous cells (SQC). The top view shows the X-Y plane in maximum projection,  
367 whereas the side view shows the X-Z or Y-Z plane using orthogonal cuts. EN =  
368 endolymph. Scale bar = 20  $\mu$ m. Images are representative of inner ear from four control  
369 and four OA-exposed rockfish. The shaded boxes in the diagrams indicate the location  
370 of each cell type within the otolith sac. A larger diagram showcasing the heterogeneous  
371 cellular anatomy of the inner ear epithelium is provided in figure S4.

372

### 373 **Conclusions**

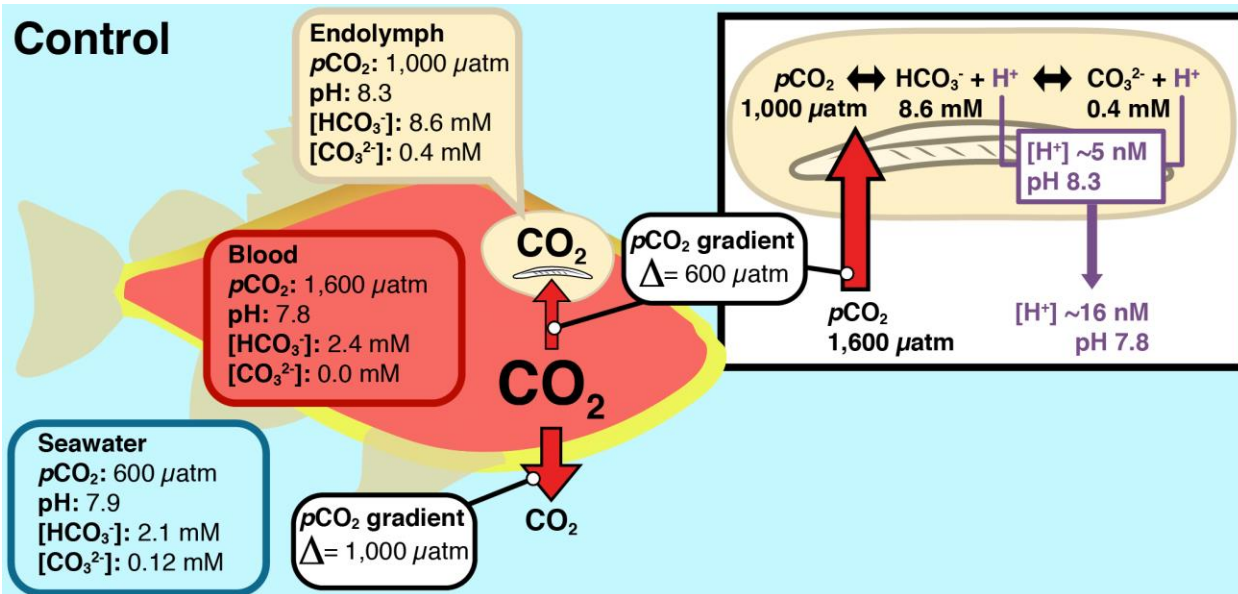
374 Increased endolymph  $[HCO_3^-]$  and  $[CO_3^{2-}]$  provides a mechanistic explanation for  
375 otolith overgrowth in OA-exposed fish, a phenomenon that was first described over a  
376 decade ago (Checkley et al., 2009). The ultimate cause is an interplay between blood  
377 and endolymph acid-base regulation, which results in increased  $CO_2$  flux into the  
378 endolymph coupled with endolymph pH regulation. As a result, the carbonate equilibria  
379 reactions shift to the right, promoting  $[HCO_3^-]$  and  $[CO_3^{2-}]$  accumulation bound to  
380 increase  $\Omega_{\text{aragonite}}$ , and thus promote biomineralization (figure 4). This implies that otolith  
381 overgrowth in response to OA will be more pronounced in fish species with more robust  
382 acid-base regulatory mechanisms; however, this hypothesis must be experimentally  
383 tested. Future studies should also investigate whether the fish inner ear epithelium can  
384 curb otolith overgrowth during prolonged OA exposure, if the long-term response  
385 requires changes in NKA and VHA protein abundance, and if species-specific  
386 differences exist. Potential mechanisms include a change in the endolymph pH setpoint,

387 modulation of glycoprotein or  $\text{Ca}^{2+}$  secretion, and engagement of other compensatory  
388 mechanisms. Coupled with functional studies (e.g. Radford et al., 2021; Shen et al.,  
389 2016), this information will help predict whether the inner ear vestibular and auditory  
390 sensory systems of fish will be affected by OA. Furthermore, understanding the  
391 mechanisms responsible for otolith biomineralization and overgrowth during OA  
392 exposure can help improve the accuracy of otolith-reliant aging techniques in the future  
393 ocean.

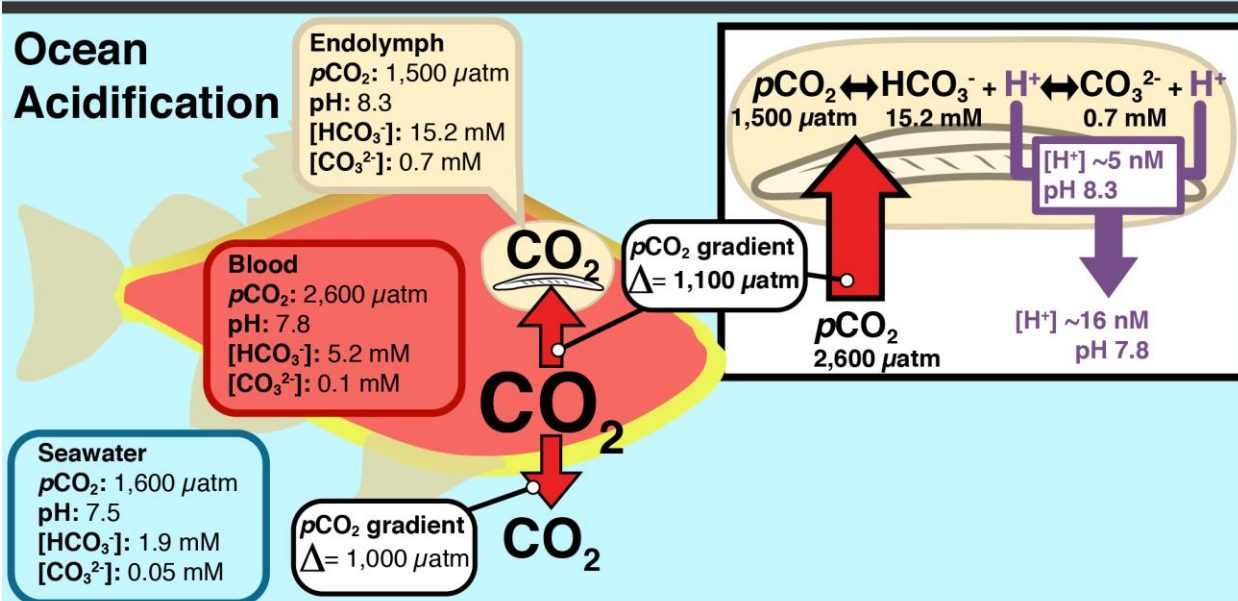
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395

## Control



## Ocean Acidification



396

397 **Figure 4:** Effect of blood and endolymph acid-base regulation on otolith overgrowth  
398 during exposure to ocean acidification. Under control conditions, metabolically produced  
399 CO<sub>2</sub> results in higher levels within the fish blood (~1,600  $\mu\text{atm}$ ) than those in seawater  
400 (~600  $\mu\text{atm}$ ) and endolymph (~1,000  $\mu\text{atm}$ ). As a result, blood CO<sub>2</sub> diffuses into  
401 seawater ( $\Delta = \sim 1,000 \mu\text{atm}$ ) as it passes through the gills, and into the endolymph ( $\Delta =$   
402  $\sim 600 \mu\text{atm}$ ) as it passes through the inner ear. Under ocean acidification, the 1,000

403  $\mu\text{atm}$  increase in seawater  $p\text{CO}_2$  (to  $\sim 1,600 \mu\text{atm}$ ) induces an equivalent increase in the  
404 blood (to  $\sim 2,600 \mu\text{atm}$ ), but a lesser increase in the endolymph (to  $\sim 1,500 \mu\text{atm}$ ). Thus,  
405 the  $p\text{CO}_2$  diffusion gradient from the blood into seawater remain constant, but the  $p\text{CO}_2$   
406 diffusion gradient from the blood into the endolymph increases ( $\Delta = \sim 1,100 \mu\text{atm}$ ). This  
407 process is driven by pH regulation from the endolymph by the inner ear epithelium,  
408 presumably by increased  $\text{H}^+$  removal into the blood (although non-bicarbonate buffering  
409 cannot be ruled out). The increased  $\text{CO}_2$  diffusion rate into the endolymph coupled with  
410 endolymph pH regulation results in the accumulation of  $[\text{HCO}_3^-]$  and  $[\text{CO}_3^{2-}]$ , thereby  
411 increasing  $\Omega_{\text{aragonite}}$  and promoting otolith calcification. The size of the arrows is  
412 proportional to the fluxes of  $\text{CO}_2$  or  $\text{H}^+$ .

413

414

415

416

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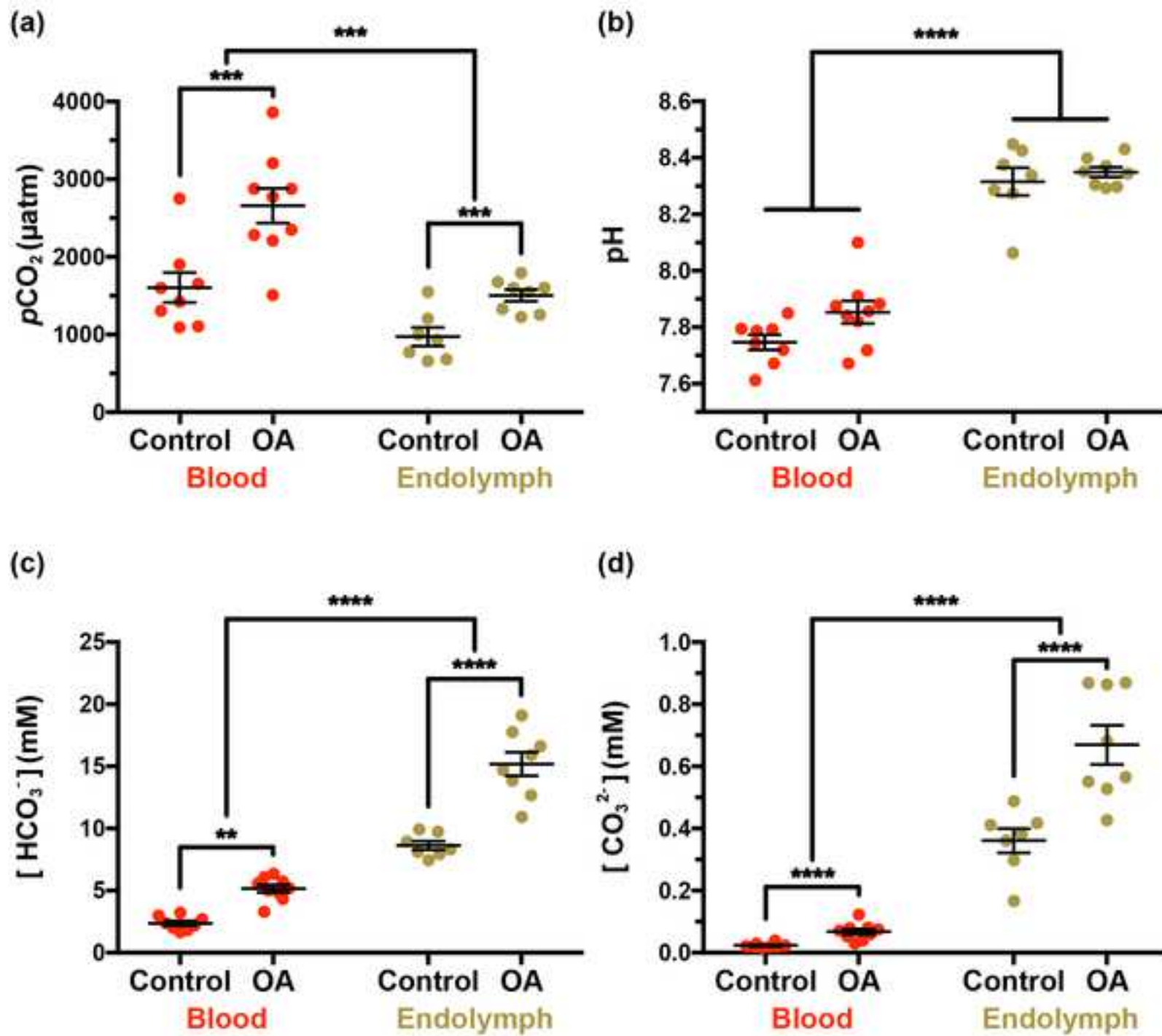
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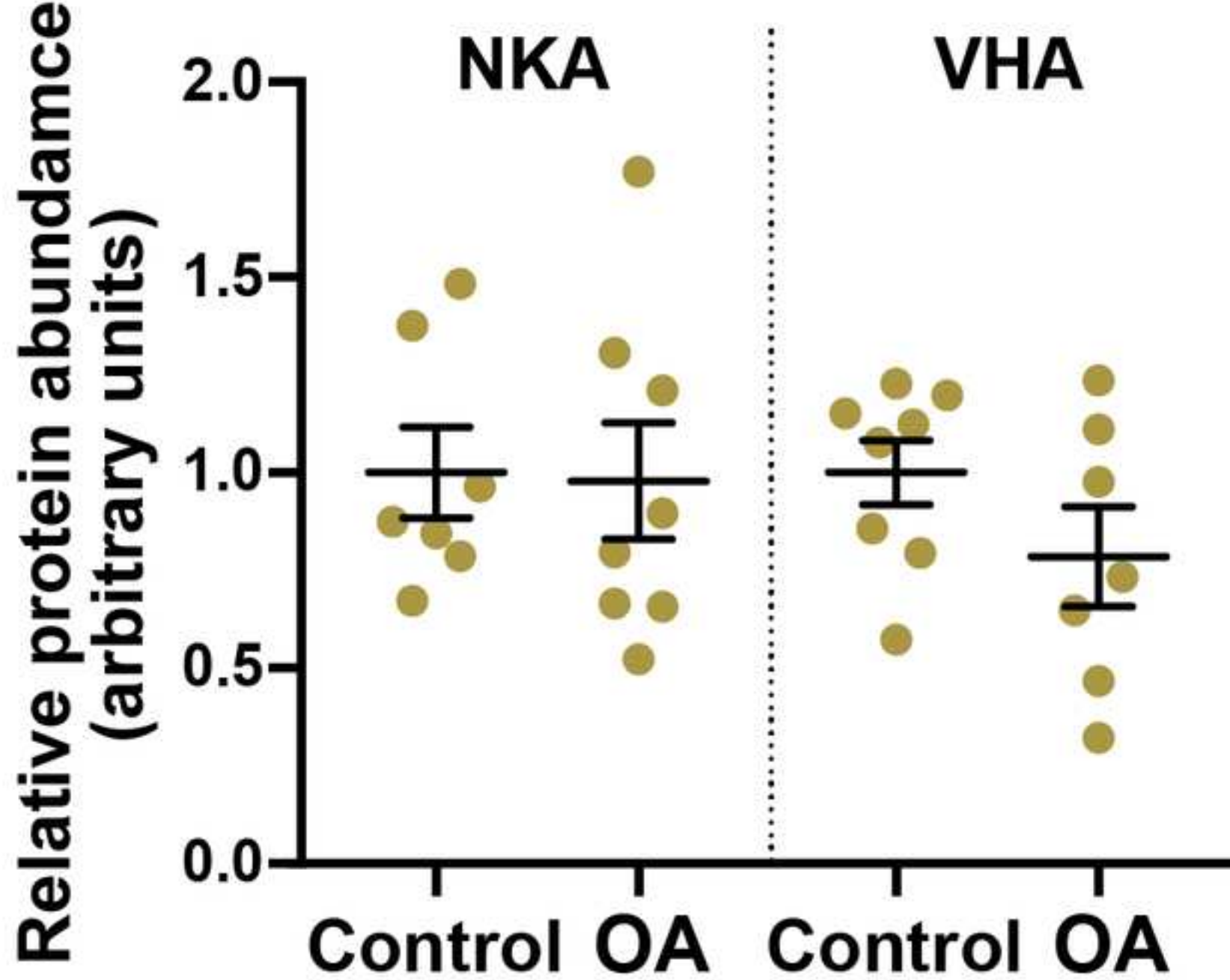
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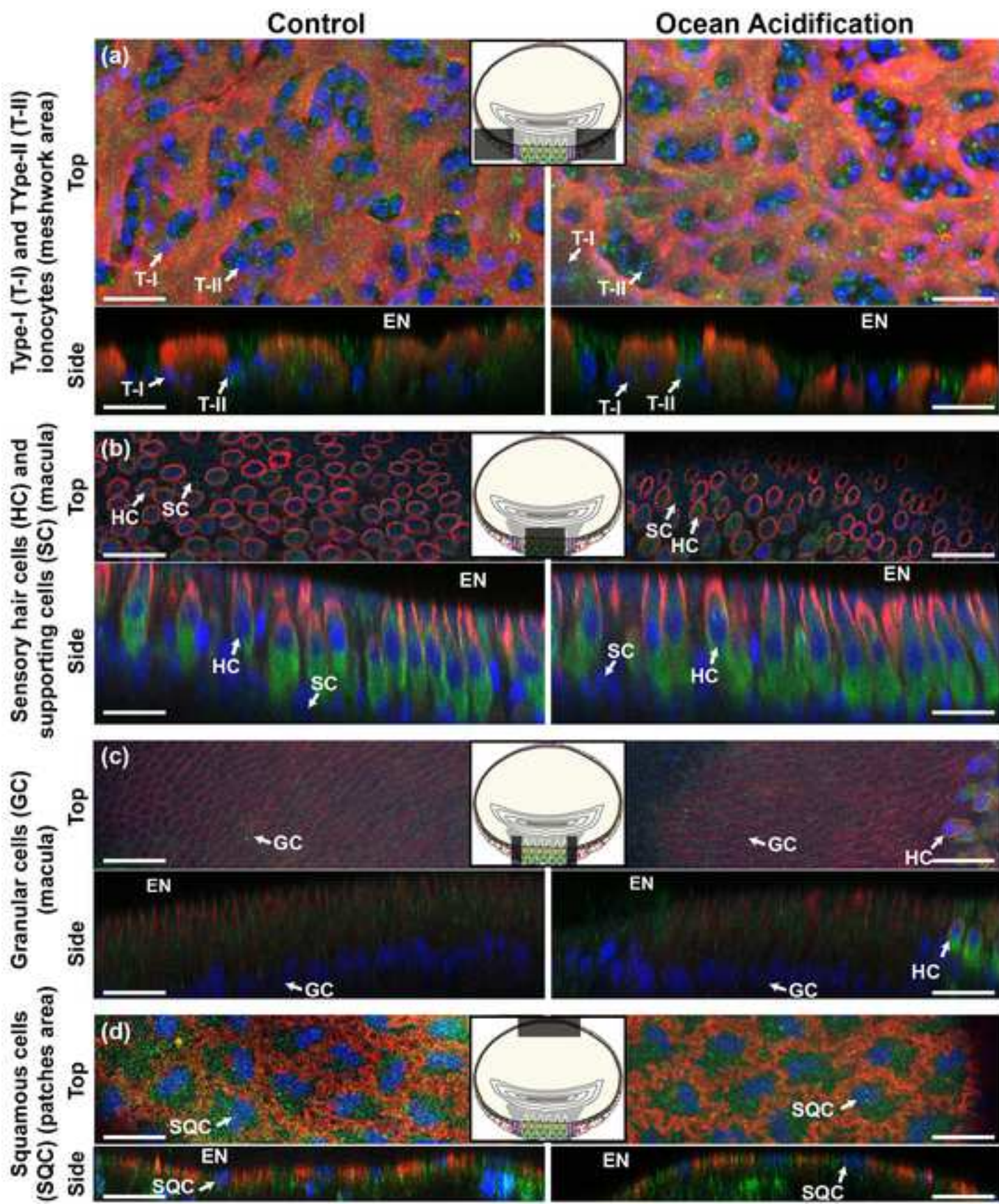
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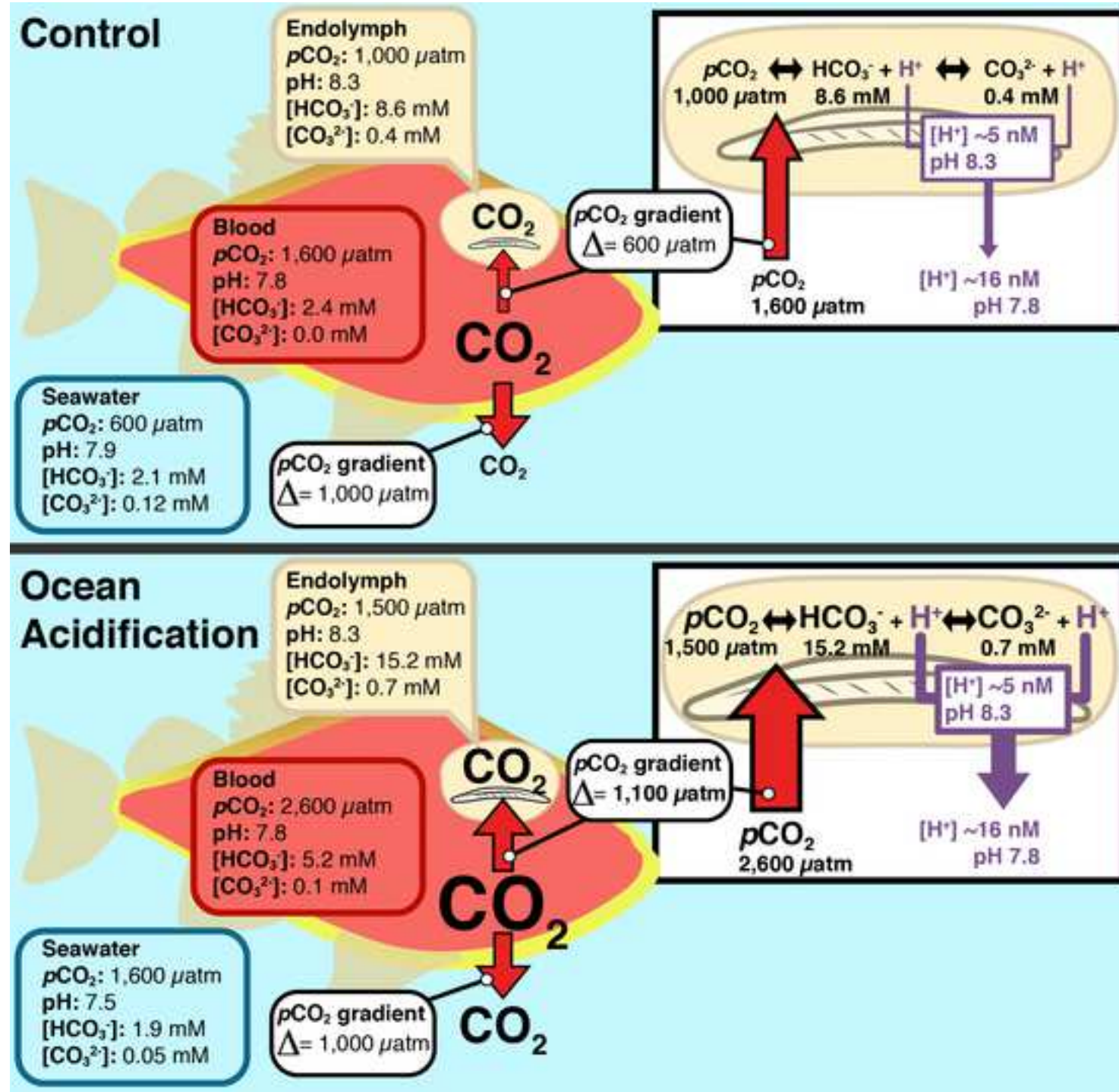
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**Declaration of Interest Statement**

No competing interests declared.

### **Author contributions**

GTK and MT conceived and designed the experiments, analysed the data, and wrote the manuscript. GTK executed the experiments. All authors gave final approval for publication.