

1 **Title: Hagfish: Champions of CO₂ tolerance question the origins of vertebrate gill function**

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5 **Supplementary Information:**

6 **Methods:**

7 **Experimental Animals**

8 Pacific hagfish (*Eptatretus stoutii*; 100-400g) were captured using baited traps placed at a
9 depth of 100 meters in Barkley Sound, west of Vancouver Island, British Columbia (B.C.),
10 Canada. These traps were transported to the Bamfield Marine Science Centre, Bamfield, B.C.,
11 Canada, where animals were transferred to opaque, covered, flow-through seawater tanks (12
12 °C) for 1-2 weeks prior to experiments. Hagfish were fed freshly killed salmon weekly.

13 **Experimental Protocol**

14 The night prior to experiments, hagfish were carefully transferred under water (to
15 minimize stress and slime production) into 4 l clear plastic containers. Containers were
16 transferred to a wet table, plumbed with flow-through seawater (12°C), and animals were
17 allowed to acclimate for 24 h. Animals were not fed during this time. Following overnight
18 acclimation, the containers were supplied with re-circulating seawater pumped from a header
19 tank previously equilibrated with approximately 10, 30 and 50 mm Hg pCO₂ accomplished using
20 a Cameron, Model 250 gas mixer and verified using a thermostatted (12°C) pCO₂ electrode

21 (E5036) displayed on a Radiometer PHM 73 (Radiometer Copenhagen). Animals remained
22 coiled and motionless during these exposures. Animals were then sampled either a) immediately
23 after transfer, time 0, or b) after 3, 6, 12, 24, 48 (only 30 and 50 mm Hg pCO₂) or 96 (only 50
24 mm Hg pCO₂) h of exposure to elevated pCO₂. To sample hagfish, the box was disconnected
25 from the re-circulating system, and a concentrated solution of benzocaine (dissolved in 70%
26 ethanol) was mixed into the box (final concentration 0.5 g benzocaine/L H₂O). Animals were
27 considered to be anaesthetized when they no longer responded to external prodding (<10 min.),
28 and a 1ml blood sample was drawn from the caudal sinus into a heparinized syringe, and placed
29 on ice. The animal was then quickly transferred to a surgical table, decapitated and the entire
30 heart and sections of liver and white muscle were removed and frozen in liquid nitrogen for later
31 measurement of intracellular pH (pHi). Tissues for buffer value assessment were collected from
32 non-CO₂ exposed animals using an otherwise similar protocol.

33 Blood was removed for measurement of hematocrit, haemoglobin, and mean cell
34 haemoglobin concentration (MCHC) as previously described²⁹. Blood pH was measured using a
35 thermostated capillary electrode (BMS Mark 2, Radiometer Copenhagen) and digital pH meter
36 (PHM 73, Radiometer Copenhagen). Another blood sample (1.5 ml) was centrifuged for 3 min.
37 at 10000 rpm and plasma total CO₂ (TCO₂) was measured using a Corning CO₂ analyzer. The
38 remaining packed red blood cell pellet was used to measure red blood cell pHi using the freeze
39 thaw method³⁸ and the same capillary electrode as used for whole blood.

40 Frozen plasma and tissues were transported to the Department of Zoology, University of
41 British Columbia where plasma samples were analysed for osmolarity (Vapor Point Osmometer,
42 Westcor), [Cl⁻] (Digital Chloridometer, Radiometer), and [Na⁺], [Mg²⁺] and [Ca²⁺] (AA

43 spectrophotometer, Varion, FS240) and total protein (BSA method). Tissue pHi was measured
44 from frozen tissues using the metabolic inhibitor method³⁹ validated for tissues from fish
45 exposed to high pCO₂ tensions⁴⁰.

46 Non-bicarbonate whole blood buffer capacity was determined on a caudal blood sample
47 (4 ml) maintained in an Eschweiller thermostated (12°C) glass tonometer, equilibrated for 45
48 min to a pCO₂ of 3.5, 7.5, 11, 21, and 45 mmHg using Sierra Mass flow controllers that mixed
49 air and CO₂ (n=4 for each CO₂ level). At each CO₂ tension, blood was removed and TCO₂ and
50 pH were measured as described above. Tissue non-bicarbonate buffer capacity was determined
51 as described previously²⁹. Tissue homogenates (n=4) were equilibrated at 2, 7.5, 15 and 30 mm
52 Hg pCO₂ and sampled at each CO₂ level for pH and TCO₂. Tissue water fraction was determined
53 for calculation of intracellular fluid based on the difference between wet mass and dry mass
54 (following drying to constant weight), which were compared well with previously reported
55 values. CO₂ solubility constants and pK' were calculated using previously derived equations⁴¹. In
56 both cases, intrinsic non-bicarbonate buffer capacity (β_{NB}) was calculated from the slope of
57 $\Delta[\text{HCO}_3^-] \Delta\text{pH}^{-1}$, and then expressed in mmol $[\text{HCO}_3^-] \text{pH}^{-1} \text{ l}^{-1}$ of blood or kg^{-1} of intracellular
58 tissue water, over an *in vivo* relevant pH range.

59 An estimate of the increase in whole animal net acid excretion rates in hagfish was
60 calculated for comparison with other species (Supplementary Table 1) as the inverse of the net
61 increase in whole body $[\text{HCO}_3^-]$ following CO₂ exposure relative to pre-exposure (i.e., time 0)
62 values as has been done previously for other aquatic species^{29,31}. The hagfish was simplified to a
63 two compartment model consisting of 33% extracellular fluid⁴²; assumed to be similar in
64 composition to plasma) and 67% intracellular fluid (assumed to be similar in composition to

65 parietal muscle which makes up the largest volume of the animal). Plasma $[\text{HCO}_3^-]$ was
66 calculated as above and parietal $[\text{HCO}_3^-]$ was calculated based upon previous work^{29,31}. Briefly,
67 buffer capacity and pH and HCO_3^- at normocapnia (i.e., control values) of the parietal muscle
68 was used to calculate pH and $[\text{HCO}_3^-]$ values during passive equilibrium of the tissue with
69 treatment CO_2 levels. Then, pH change and HCO_3^- accumulation were calculated as the
70 difference between these estimates and the measured pH and calculated $[\text{HCO}_3^-]$ values at 3 and
71 24 h. The rate of net acid excretion during exposure to 30 and 50 mm Hg pCO_2 respectively is
72 1.2 and 1.4 $\text{mmol h}^{-1} \text{kg}^{-1}$ body water at 3 h (Supplementary Table 1). Most of the increase in
73 HCO_3^- at this time point is in the intracellular compartment. When calculated at 24 h, net acid
74 excretion rates remained high (0.8 and 0.61 $\text{mmol h}^{-1} \text{kg}^{-1}$ body water at 30 and 50 mm Hg pCO_2
75 respectively), but a much greater proportion of HCO_3^- was in the extracellular space at this time.
76 Overall, net acid excretion rates in hagfish were similar to those determined in other fish
77 (Supplementary Table 1).

78 It is important to note that this method likely underestimates net acid excretion at two
79 levels. First, using parietal muscle as a proxy for the whole animal results in a conservative
80 estimate of acid excretion because a) this tissue demonstrated the slowest rate of HCO_3^-
81 accumulation and the smallest degree of pH compensation of all tissues measured, partially due
82 to its high buffer capacity, and b) the method with which we obtained buffer capacity in muscle
83 is known to overestimate intracellular buffering⁴². Second, we assumed that active net acid
84 excretion began immediately after exposure to hypercarbia (i.e., time 0), which is unlikely to be
85 the case. Thus, the high rates of net acid excretion relative to other fishes reported in
86 Supplementary Table 1 are likely conservative and they are achieved in an animal with one of
87 the lowest resting metabolic rates ever measured.

88 **Statistical Analyses**

89 Data are presented as mean \pm SEM. All statistical differences were detected using a one-way
90 ANOVA and, when necessary, a post-hoc Dunnett's test. All statistical analyses were conducted
91 using SigmaStat for Windows 3.5.0.54 (Systat Software, Inc., 2006), and all analyses were
92 interpreted using $\alpha = 0.05$ to determine statistical significance.

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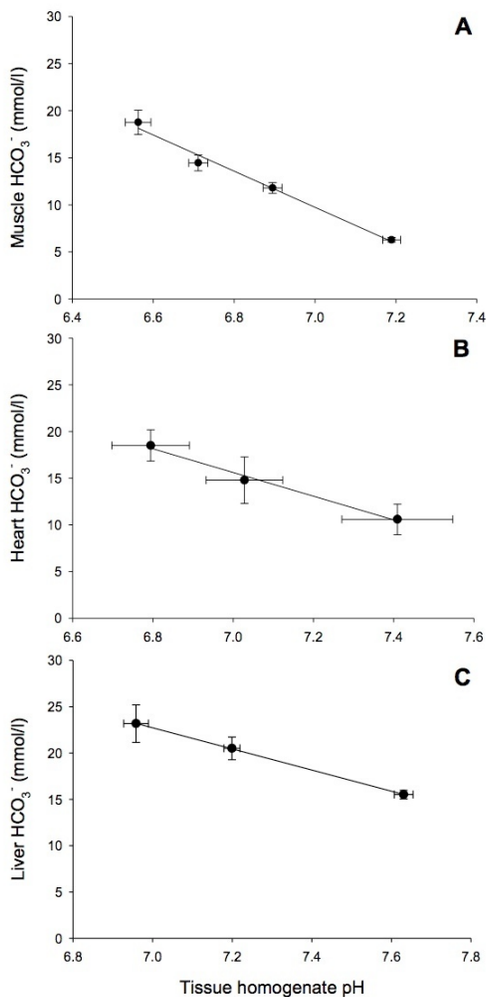
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Supplementary Table 1: Summary of calculated net acid excretion rate in aquatic fishes listed approximately from most basal to most derived.

Species	Net Acid Excretion Rate (mmol h ⁻¹ kg ⁻¹)	Origin of Acidosis	Medium
<i>Eptatretus stoutii</i> *†	0.9-1.4 ^a	1.5-6% CO ₂	SW
<i>Eptatretus stoutii</i> ¹⁶ †	~ 0.9 ^b	acid infusion	SW
<i>Myxine glutinosa</i> ¹⁴ †	~ 0.85 ^b	5% CO ₂	SW
<i>Myxine glutinosa</i> ¹⁵ †	~0.6 ^b	acid infusion	FW
<i>Acipenser transmontanus</i> ²⁹ ††	0.6-1.6 ^a	1.5-6%	FW
<i>Pterygoplichthys pardalis</i> ³¹ ††	0.7-1.4 ^a	2-4% CO ₂	FW
<i>Petromyzon marinus</i> ⁴³	~ 0.5 ^b	exhaustive exercise	FW
<i>Scyliorhinus stellaris</i> ⁴⁴ ‡	~ 0.6 ^b	1% CO ₂	SW
<i>Conger conger</i> ⁴⁵	~ 0.6 ^b	1% CO ₂	SW
<i>Opsanus beta</i> ⁴⁶	~ 0.5 ^b	5% CO ₂	SW
<i>Myoxocephalus octodecimspinosus</i> ⁴⁷	~0.5 ^b	acid infusion	SW
<i>Carpus carpio</i> ⁴⁸	~ 0.2 ^b	1% CO ₂	FW

100 *present study
 101 † *E. stoutii* and *Myxine glutinosa* are osmo- and ionoconforming vertebrates.
 102 ‡ *S. stellaris* is an ionoregulating and osmoconforming vertebrate.
 103 †† *Acipenser transmontanus* and *Pterygoplichthys pardalis* are preferential pHi regulators, able to protect
 104 tissue pHi while blood pH is severely depressed.
 105 a. Estimated from values of buffering capacity and net HCO₃⁻ accumulation (see supplemental text).
 106 b. Estimated from the difference between net ammonia and titratable acidity excretion rates in
 107 surrounding aquatic environment (net acid flux).
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111 **Supplementary Figure 1:** The relationship between pH and $[\text{HCO}_3^-]$ of tissue homogenates
 112 from the Pacific hagfish at different CO_2 tensions. Values are means \pm SEM. Intrinsic (i.e., non-
 113 bicarbonate) buffer capacities were calculated as the following: parietal muscle = 29.3, entire
 114 heart = 22.5, and liver = 16.7 mmol $\text{HCO}_3^- \text{ pH}^{-1} \text{ kg}^{-1}$ intracellular water. These values were
 115 calculated from best-fit linear regression and using estimates of extra- and intracellular tissue
 116 water for *E. stoutii* from previous work⁴⁵.