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# Rising CO<sub>2</sub> enhances hypoxia tolerance in a marine fish

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Global environmental change is increasing hypoxia in aquatic ecosystems. During hypoxic events, bacterial respiration causes an increase in carbon dioxide (CO<sub>2</sub>) while oxygen (O<sub>2</sub>) declines. This is rarely accounted for when assessing hypoxia tolerances of aquatic organisms. We investigated the impact of environmentally realistic increases in CO<sub>2</sub> on responses to hypoxia in European sea bass (*Dicentrarchus labrax*). We conducted a critical oxygen (O<sub>2crit</sub>) test, a common measure of hypoxia tolerance, using two treatments in which O<sub>2</sub> levels were reduced with constant ambient CO<sub>2</sub> levels (~530 µatm), or with reciprocal increases in CO<sub>2</sub> (rising to ~2,500 µatm). We also assessed blood acid-base chemistry and haemoglobin-O<sub>2</sub> binding affinity of sea bass in hypoxic conditions with ambient (~650 µatm) or raised CO<sub>2</sub> (~1770 µatm) levels. Sea bass exhibited greater hypoxia tolerance (~20% reduced O<sub>2crit</sub>), associated with increased haemoglobin-O<sub>2</sub> affinity (~32% fall in P<sub>50</sub>) of red blood cells, when exposed to reciprocal changes in O<sub>2</sub> uptake by the blood in low O<sub>2</sub> conditions, enhancing hypoxia tolerance. We recommend that when impacts of hypoxia on aquatic organisms are assessed, due consideration is given to associated environmental increases in CO<sub>2</sub>.

A lack of  $O_2$  is one of the greatest challenges that most life can face. In terrestrial ecosystems conditions of low  $O_2$  are rare. In contrast low  $O_2$ , referred to as hypoxia, is much more common in freshwater and marine ecosystems<sup>1-4</sup>. Hypoxia occurs because high biological demand for  $O_2$  can exceed the rate of  $O_2$  supply to the ecosystem, leading to a reduction in environmental  $O_2$  levels<sup>5,6</sup>. However, the challenges of hypoxia are not solely a result of reduced  $O_2$ . Organisms must also contend with simultaneous but reciprocal changes in the other respiratory gas,  $CO_2$ .

When  $O_2$  decreases in aquatic systems there is a corresponding increase in  $CO_2^{7,8}$ . This is a by-product of respiration, the same process that causes depletion of  $O_2$ . As such high  $CO_2$  during hypoxia is ubiquitous and unavoidable. This coupling of  $O_2$  and  $CO_2$  has been highlighted numerous times in oceanographic sciences, most recently by Robinson<sup>5</sup>. Yet unaccountably, despite the known link between decreasing  $O_2$  and increasing  $CO_2$  during hypoxia<sup>8,9</sup>, the issue of increased environmental  $CO_2$  during periods of low  $O_2$  has been relatively overlooked by biologists.

Implications of rising  $CO_2$  during hypoxia on aquatic organisms are particularly important to address in the face of human driven climate change. Hypoxic areas are predicted to become more common and more severe, particularly in marine systems, with the de-oxygenation of the world's oceans recently highlighted as a major component of climate change<sup>2,3,10–12</sup>. In addition, there will be an increase in ambient  $CO_2$  as rising atmospheric  $CO_2$  is absorbed by the world's oceans<sup>13</sup>. Non-linear interactive effects between higher atmospheric  $CO_2$  and  $CO_2$  accumulation during hypoxia will lead to increased  $CO_2$  levels during hypoxia in future oceans<sup>14</sup>. This means that effects of rising  $CO_2$  during hypoxia in marine systems will be amplified by climate change.

Typically, experiments which test responses to hypoxia or impacts from hypoxia on aquatic organisms create hypoxic conditions by off-gassing oxygen from water by gassing with pure nitrogen or a mix of nitrogen (N<sub>2</sub>) and O<sub>2</sub> (for examples see<sup>15-18</sup>). This creates low O<sub>2</sub> conditions without the concurrent CO<sub>2</sub> increase that would be expected in the environment. The lack of studies in which an environmentally realistic simultaneous decrease in O<sub>2</sub> and increase of CO<sub>2</sub> have been conducted may lead to mismeasurement of responses to hypoxia. Recently, several studies on marine fish and invertebrates have demonstrated interactive effects of low oxygen and increased CO<sub>2</sub><sup>9,19,20</sup>, with some species exhibiting loss of equilibrium (LoE) and death at higher O<sub>2</sub> concentrations when

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**Figure 1.** Calculated critical oxygen level ( $O_{2crit}$ ) of European sea bass, *Dicentrarchus labrax*, when  $O_{2crit}$  tests include a constant ambient CO<sub>2</sub> level (~500 µatm, N = 8) or an ecologically realistic rise in CO<sub>2</sub> (~500–2500 µatm, N = 7) during the test. \*Indicates significant difference between CO<sub>2</sub> regimes (p < 0.05). Boxes represent median value and inter-quartile range, whiskers represent minimum and maximum values.

 $CO_2$  is simultaneously elevated<sup>19,21</sup>. However, these experiments do not give insight into the physiological mechanisms underlying the influence of  $CO_2$  on hypoxic responses of fish.

Previously observed impacts of hypoxia-associated rises in CO<sub>2</sub> on hypoxia tolerance of fish could be a result of changes in  $O_2$  uptake, as  $CO_2$  has been shown to impact upon several aspects of organismal biology that are involved in  $O_2$  uptake and transport<sup>22,23</sup>. We aimed to assess whether concurrent increases in  $CO_2$  during decreases in  $O_2$  affect  $O_2$  uptake in a marine fish, the European sea bass (*Dicentrarchus labrax*) by conducting a standard critical O2 level test (O2crit). Under normal O2 (normoxic) conditions fish maintain a minimum level of  $O_2$  consumption rate ( $\dot{M}O_2$ ), referred to as the standard metabolic rate (SMR), in order to meet maintenance energetic demands of essential processes through aerobic respiration<sup>24</sup>. As the level of  $O_2$  in water drops fish deploy a number of responses (i.e. increased ventilatory water flow and cardiac output, increased haematocrit, functional changes in gill morphology, changes in Hb-O2 affinity) in order to maintain and regulate this minimum level of  $\dot{MO}_2^{25-27}$ . If environmental  $O_2$  continues to drop there comes a point at which fish are unable to regulate MO<sub>2</sub> to meet minimal energy demands, referred to as the critical O<sub>2</sub> level (O<sub>2crit</sub>). At O<sub>2</sub> levels below O<sub>2crit</sub> fish become oxy-conformers (where MO2 is directly proportional to environmental O2 availability) and fish become increasingly reliant on anaerobic metabolism which is unsustainable in the medium to long term. In the past the measure of O<sub>2crit</sub> has been used as a proxy for overall hypoxia tolerance but recently this approach has been questioned<sup>28,29</sup>. Nevertheless O<sub>2crit</sub> does provide information related to the ability of fish to maintain O<sub>2</sub> uptake and supply during hypoxia, and its prevalence in the literature allows comparison of responses between species<sup>30</sup>. Furthermore, we investigated whether any changes in  $O_{2crit}$  could be linked to changes in blood acid-base chemistry and blood gas transport via alteration of Hb-O<sub>2</sub> binding caused by rising environmental CO<sub>2</sub>. Our hypothesis was that the simultaneous increase in CO<sub>2</sub> during a progressive decrease in O<sub>2</sub> would decrease hypoxia tolerance (increase O<sub>2crit</sub>) and that this response may be a result of blood acid-base disturbance decreasing Hb-O<sub>2</sub> affinity and O<sub>2</sub> transport.

#### Results

**O**<sub>2crit</sub> **tests.** There was evidence of enhanced tolerance to hypoxia in sea bass exposed to rising compared to constant CO<sub>2</sub> conditions. This was indicated by measurements of O<sub>2crit</sub> in European sea bass being significantly different between fish exposed to either constant, or rising CO<sub>2</sub> levels during O<sub>2crit</sub> tests when accounting for variation in SMR (Fig. 1; ANCOVA, F<sub>1,12</sub>=7.525, p=0.0178). A CO<sub>2</sub> increase during O<sub>2crit</sub> tests resulted in a 20% reduction of O<sub>2crit</sub> (3.88 ± 0.19 kPa O<sub>2</sub>, 18.7 ± 0.9% air saturation, mean ± S.E.) when compared to tests in which CO<sub>2</sub> levels were maintained at ambient levels ( $4.87 \pm 0.22$  kPa O<sub>2</sub>,  $23.4 \pm 1.1\%$  air saturation, mean ± S.E.).

**Blood chemistry analysis.** A comparison of blood chemistry parameters between the two treatment groups indicated that sea bass fully compensated for the rise in CO<sub>2</sub> during hypoxia within 5 hours (the period of exposure prior to blood sampling). Blood pH (pH<sub>e</sub>) was not different between fish exposed to constant ambient CO<sub>2</sub> (7.87 ± 0.03) and fish exposed to progressively rising CO<sub>2</sub> (7.88 ± 0.02) (Fig. 2A, GLM, F<sub>1,14</sub> = 0.23, p = 0.64). The acidifying effect of the ~79% rise in blood pCO<sub>2</sub> levels in the rising CO<sub>2</sub> regime (0.272 ± 0.019 kPa CO<sub>2</sub>) compared to the constant ambient CO<sub>2</sub> regime (0.152 ± 0.015 kPa CO<sub>2</sub>) (Fig. 2C, GLM, F<sub>1,13</sub> = 23.9, p < 0.001) was fully compensated by elevating blood HCO<sub>3</sub><sup>-</sup> (Fig. 2D, GLM, F<sub>1,13</sub> = 40, p < 0.001). Plasma HCO<sub>3</sub><sup>-</sup> was 88% higher under the rising CO<sub>2</sub> regime (6.76 ± 0.29 mM) when compared to the constant ambient CO<sub>2</sub> regime (6.76 ± 0.29 mM) when compared to the constant ambient CO<sub>2</sub> regime (6.76 ± 0.29 mM) when compared to the constant ambient CO<sub>2</sub> regime (6.76 ± 0.29 mM) when compared to the constant ambient CO<sub>2</sub> regime (a.676 ± 0.29 mM) when compared to the constant ambient CO<sub>2</sub> regime (a.676 ± 0.29 mM) when compared to the constant ambient CO<sub>2</sub> regime ( $3.60 \pm 0.43$  mM). There were no differences in haematocrit (Fig. 2B, general linear model, F<sub>1,13</sub> = 0.69, p = 0.42) or plasma lactate (Fig. 2E, general linear model, F<sub>1,14</sub> = 1.48, p = 0.24) between fish sampled under a constant ambient CO<sub>2</sub> regime (haematocrit = 39.4 ± 0.8%, lactate = 0.88 ± 0.26 mM) or rising CO<sub>2</sub> regime (haematocrit = 38.2 ± 1.1%, lactate = 0.53 ± 0.12 mM). Blood glucose levels were ~26% lower in fish exposed to a progressively rising CO<sub>2</sub> regime (4.50 ± 0.53 mM) when compared to a constant ambient CO<sub>2</sub> regime (6.09 ± 0.31 mM) (Fig. 2F, GLM, F<sub>1,14</sub> = 6.74, p = 0.021).



**Figure 2.** Blood chemistry characteristics of European sea bass sampled at ~8.4 kPa O<sub>2</sub> (~40% air saturation) following a progressive O<sub>2</sub> decline accompanied by either constant CO<sub>2</sub> (~650 µatm CO<sub>2</sub>, N = 8) or a progressive increase in CO<sub>2</sub> (sampled at ~1770 µatm CO<sub>2</sub>, N = 8). Blood pH (**A**), haematocrit (**B**), plasma lactate (**E**) and plasma glucose (**F**) were directly measured whilst blood  $pCO_2$  (**C**) and plasma HCO<sub>3</sub><sup>-</sup> (**D**) were calculated (see Methods for details). \*Indicates statistical significance with p < 0.05 and \*\*\*indicates statistical significance with p < 0.001. Boxes represent median value and inter-quartile range, whiskers represent minimum and maximum values.

**Haemoglobin affinity for oxygen.** Oxygen affinity of haemoglobin was increased in fish sampled under the progressively rising CO<sub>2</sub> regime (Fig. 3A, GLM,  $F_{1,12} = 10.42$ , p = 0.0073). Haemoglobin  $P_{50}$  was decreased by ~32.5% in fish under the progressively rising CO<sub>2</sub> regime ( $1.64 \pm 0.15$  kPa O<sub>2</sub>,  $7.9 \pm 0.7\%$  air saturation) compared to fish sampled with a constant ambient CO<sub>2</sub> ( $2.43 \pm 0.20$  kPa O<sub>2</sub>,  $11.7 \pm 1\%$  air saturation). There was no

significant change in Hills number between treatments (Fig. 3B, GLM,  $F_{1,12} = 0.50$ , p = 0.494).

#### Discussion

Our results highlight the biological importance of simultaneously rising  $CO_2$  under conditions where  $O_2$  levels in water are depleted – a scenario that reflects the natural conditions during hypoxia which will be exacerbated by climate change – by demonstrating that ecologically relevant changes of  $CO_2$  impact physiological performance of a marine fish at both the molecular and whole organism level. We hypothesised that rising  $CO_2$  during progressive  $O_2$  decreases would lead to an increase in  $O_{2crit}$  as a result of increased blood  $CO_2$ , decreased blood pH and the associated Bohr/Root effect of fish haemoglobin (Hb), in which Hb-O<sub>2</sub> affinity (Bohr effect) and the total capacity of Hb for  $O_2$  (Root effect) are reduced when pH falls. In contrast, we show that increasing  $CO_2$  as  $O_2$  declined led to enhanced hypoxia tolerance of sea bass with a 20% lower critical oxygen level (Fig. 1). This change in whole organism hypoxic response was accompanied by an increase in Hb-O<sub>2</sub> affinity of blood cells in fish exposed to concurrent  $CO_2$  rises (Fig. 3). The change in Hb-O<sub>2</sub> prior to blood sampling may have contributed



**Figure 3.** Haemoglobin P<sub>50</sub> (**A**) and Hills number (**B**) for fish sampled at ~8.4 kPa O<sub>2</sub> (~40% air saturation) following a progressive O<sub>2</sub> decline accompanied by either constant CO<sub>2</sub> (~650 µatm CO<sub>2</sub>, N=7) or a progressive increase in CO<sub>2</sub> (sampled at ~1770 µatm CO<sub>2</sub>, N=7). Measurements were made using a gas mix which matched the calculated blood *p*CO<sub>2</sub> of each individual fish blood sample. \*\*Indicates statistical significance with p < 0.01. Boxes represent median value and inter-quartile range, whiskers represent minimum and maximum values.

to this result). This provides a potential mechanistic basis to explain improved  $O_{2crit}$ , enabling sea bass to enhance  $O_2$  uptake during hypoxia and thus maintain normal aerobic metabolism to lower environmental  $O_2$ .

The driver of increased Hb-O<sub>2</sub> affinity in sea bass exposed to concurrent  $O_2$  decline and  $CO_2$  rise is not clear from our results. Several allosteric factors that can modulate the affinity of haemoglobin for O<sub>2</sub> could be involved, including pH, organic phosphates and inorganic ions. Fish haemoglobin is highly sensitive to pH, which modulates Hb-O<sub>2</sub> affinity and carrying capacity via the Bohr and Root effects<sup>31</sup>, but we found no differences in blood pH of sea bass between treatment groups. In addition, the *in-vivo* increase in  $pCO_2$  in fish exposed to concurrent CO<sub>2</sub> rises during hypoxia led to an opposite response of Hb-O<sub>2</sub> affinity than would be expected by an in-vitro rise in pCO2 which would result in pH induced Bohr/Root effects. Increased Hb-O2 affinity could result from increased intracellular pH of erythrocytes<sup>32</sup>, as acute hypoxic exposure has been shown to stimulate a  $\beta$ -adrenergic stimulated increase in intracellular erythrocyte pH in rainbow trout<sup>33</sup>. Alternatively, increased Hb-O<sub>2</sub> affinity could be due to decreased red cell nucleoside triphosphates (NTPs)<sup>31,34</sup>, a known hypoxia adaptation, but this can take more than 6 days to complete<sup>35</sup>. Sea bass may also have a particularly strong  $\beta$ -adrenergic response and/or fast NTP response, although there is little evidence to suggest this because  $P_{50}$  measurements from fish exposed to hypoxia at ambient  $CO_2$  levels do not differ from  $P_{50}$  measurements in normoxic fish from the same population (Montgomery et al., Unpublished data). It is possible that rising CO<sub>2</sub> during hypoxia may modulate the  $\beta$ -adrenergic response and/or the red cell NTP response within the time frame (~4–6 hours) of our treatments. However, direct measurements of red cell pH<sub>i</sub> and NTP content of sea bass in each treatment group would be needed to confirm this.

A third possible driver of increased Hb-O<sub>2</sub> affinity in sea bass exposed to concurrent  $CO_2$  rise during hypoxia could be decreased erythrocyte chloride  $(Cl^{-})^{36}$ . Although erythrocyte  $Cl^{-}$  was not directly measured in our study, plasma  $HCO_3^{-}$  was approximately 3 mM higher in fish exposed to concurrent  $CO_2$  rises during hypoxia than fish which experienced constant ambient  $CO_2$  during hypoxia (Fig. 2D). The higher  $HCO_3^{-}$  in fish exposed to rising ambient  $CO_2$  during hypoxia is likely a result of rapid compensation for a respiratory acidosis due to rising blood p $CO_2$ . This change in plasma  $HCO_3^{-}$  is typically mirrored by a reciprocal change in plasma  $Cl^{-37,38}$  which is likely to be followed by a similar decline in erythrocyte  $Cl^{-}$ .

Analysis of  $O_{2crit}$  is a common measure of hypoxia tolerance in fish but concurrent  $CO_2$  increases during hypoxia have been generally unaccounted for. A recent meta-analysis by Rogers *et al.*<sup>39</sup> constructed a database of  $O_{2crit}$  research of fish (both freshwater and marine). This analysis identified two broad methods employed in  $O_{2crit}$  measurements:

- 1. Closed respirometry where O<sub>2</sub> is usually reduced by the O<sub>2</sub> consumption of the fish (52 identified studies) or;
- 2. Intermittent or flow-through respirometry in which  $O_2$  is usually reduced via gassing with pure  $N_2$  or combined  $N_2$  and  $O_2$  mixes (32 identified studies).

The use of closed respirometry in the majority of studies would result in concurrent  $CO_2$  rises as  $O_2$  is depleted by fish  $O_2$  consumption. The increase in ambient  $CO_2$  during closed respirometry is well known and often used as a criticism of this respirometry technique<sup>40</sup>. In contrast, use of intermittent-flow respirometry in  $O_{2crit}$  trials normally necessitates the reduction of  $O_2$  in the water by aeration with  $N_2$  or a mix of  $N_2 & O_2$ . As a result  $CO_2$  would likely decrease during the time course of hypoxia induction (as the gas mixture would contain zero  $CO_2$ , rather than ~400 µatm present in atmospheric air). Such a change in  $CO_2$  during the  $O_{2crit}$  trial would be the opposite of that seen in nature. Therefore it may be considered that closed respirometry provides conditions which give a more environmentally relevant measure of  $O_{2crit}^{28}$ .

As our results indicate that rising  $CO_2$  during hypoxia directly affects the ability of sea bass to maintain  $O_2$  uptake, it could be expected that the use of closed respirometry methods would result in lower measurements

of O<sub>2crit</sub> than intermittent-flow methods for the same species. This effect has not been documented for species in which a direct comparison has been made - with either there being no effect of respirometry method on O<sub>2crit</sub><sup>39,41</sup> or higher O<sub>2crit</sub> measurements when closed respirometry is used<sup>41</sup>. However, such comparisons are complicated by differences in the rate of hypoxia induction (RHI) by different studies, which in turn will influence how much time fish have to regulate blood pH when ambient CO<sub>2</sub> is rising. For example, Regan and Richards<sup>41</sup> have demonstrated that the faster rates of hypoxia induction (RHI) typical of closed respirometry O<sub>2crit</sub> trials lead to higher values of O<sub>2crit</sub> (i.e. lower hypoxia tolerance) when compared with longer trials using slower RHI's typical of the intermittent-flow method. The effect of RHI on O<sub>2crit</sub> was proposed by Regan and Richards<sup>40</sup> as a potential explanation of the results of Snyder et al. (i.e. higher O<sub>2crit</sub> in closed respirometry compared to intermittent-flow respirometry)<sup>42</sup>. The speed of RHI during closed respirometry will also effect the speed of CO<sub>2</sub> rise. Almost all studies using closed respirometry to measure O2crit do not report changes in CO2 over the course of measurement period. When accumulation of CO<sub>2</sub> during a closed respirometry O<sub>2crit</sub> trial was measured by Regan and Richards, CO<sub>2</sub> levels were ~8,000  $\mu$ atm after ~90 minutes<sup>41</sup>. However, we should note that this pCO<sub>2</sub> level was measured after the  $O_{2crit}$  point, when anaerobic metabolism continues to produce  $CO_2$  in the absence of  $O_2$  consumption, but also metabolic acid production and excretion further drives up water  $pCO_2$  in the respirometer due to excess  $H^+$  ions titrating ambient  $HCO_3^-$  to  $CO_2$ . Regardless, at  $O_2$  levels above  $O_{2crit}$  the rate of  $CO_2$  onset will be faster than those used in our current study as a result of the faster RHI. Increased speed of CO<sub>2</sub> onset in closed respirometry trials may 'outstrip' the ability of fish to acid-base regulate, causing an uncompensated respiratory acidosis during the time of the trial, which in turn would decrease Hb-O2 affinity via the Bohr & Root effects and potentially increase O<sub>2crit</sub>. Similarly, fish species which have reduced ability to acid-base regulate may have an increased O<sub>2crit</sub> when rising CO<sub>2</sub> is included in trials.

Our results indicate improved hypoxia tolerance during rising  $CO_2$  in European sea bass. This contrasts with previous research investigating interactive effects of  $CO_2$  and hypoxia on  $O_{2crit}$  of fish. Woolly sculpin, *Clinocottus analis* (an intertidal species that can breathe air), exposed to ~1100 µatm  $CO_2$  showed no impact on  $O_{2crit}$  after 7 days acclimation but after 28 days had  $O_{2crit}$  measurements ~34% higher than fish held in ambient (~400 µatm) conditions<sup>43</sup>. Higher  $O_{2crit}$  after 28 days corresponded with higher RMR and Na<sup>+</sup>, K<sup>+</sup>, ATPase activity. This contrast in results could indicate that the beneficial effect of acute rises in  $CO_2$  associated with natural hypoxia documented in our study are potentially reversed when fish are exposed to long term constantly high  $CO_2$  associated with anthropogenic climate change. Moreover, acute changes in  $CO_2$  had no effect on  $O_{2crit}$  of the estuarine fish species mummichog, *Fundulus heteroclitus*, and Norfolk spot, *Leiostomus xanthurus* when they were exposed to ~8,000–10,000 µatm  $CO_2$  immediately prior to an  $O_{2crit}$  trial<sup>44</sup>. As such the effect of  $CO_2$  on  $O_{2crit}$  will likely depend on differences in physiological responses to  $CO_2$  and  $O_2$  between species.

Simultaneously rising CO<sub>2</sub> also shows variable impacts on non-metabolic responses to hypoxia of several species. Cycling  $CO_2$  had no effect on aquatic surface respiration (ASR), the use of the thin surface layer of water for aquatic respiration<sup>45</sup>, or survival in juvenile Menidia menidia, Fundulus majalis, Fundulus heteroclitus or Morone saxatalis exposed to short term cycles of  $O_2^{46}$ . In contrast, combined hypoxia and acidification resulted in an increase in the O<sub>2</sub> level at which Menidia menidia and Menidia beryllina first performed ASR, consistently performed ASR, exhibited LoE, and finally died<sup>21</sup>. Additionally, combined high CO<sub>2</sub> (~2,000 µatm) and hypoxia had no effect on survival of larval Cyprinodon variegatus, an additive negative effect on larval Menidia beryllina, and a synergistic negative effect on larval Menidia menidia<sup>47</sup>. This variation in effect of  $CO_2$  on hypoxia responses could be a result of methodological differences (e.g. constant high CO<sub>2</sub> in Dixon et al.<sup>46</sup>, cycling DO/pH in DePasquale et al.<sup>47</sup>, and concurrent CO<sub>2</sub> rise/O<sub>2</sub> decrease in Miller et  $al^{21}$ , the level of CO<sub>2</sub> used in studies (e.g. CO<sub>2</sub> levels used by Miller *et al.*<sup>21</sup> were ~23,000  $\mu$ atm which is much higher than levels likely to be commonly found in the environment during hypoxia and may have contributed to the negative effects of rising CO<sub>2</sub> noted in the study), differences in species and life stages used (changes in physiological tolerance across life stages have been noted for thermal tolerance by Komoroske et al.48), or possibly variability in response as a result of differences in previous environmental experience<sup>49</sup>. The role of environmental variability in species sensitivities to  $CO_2$  has recently been outlined in the proposed Ocean Variability Hypothesis (OVH)<sup>50</sup> and warrants testing on various species in the future.

Overall our results indicate that the environmentally realistic, simultaneous rises in  $CO_2$  during a hypoxic event increased the hypoxia tolerance (i.e. reduces  $O_{2crit}$ ) of European sea bass which is at least partly explained by an enhanced ability of fish to uptake  $O_2$  via increased Hb- $O_2$  affinity. Miller *et al.*<sup>21</sup> also demonstrated impacts of concurrent  $CO_2$  rise on measurements of hypoxia tolerance, although in an opposite direction to that noted in our study. As concurrent  $CO_2$  rises during hypoxia are the norm in nature, evidence that this affects physiology of organisms exposed to hypoxia highlights an important shortcoming of research to predict tolerances to hypoxia of fish. More research on this issue is needed to clarify how common this modifying effect of  $CO_2$  on the response to hypoxia is and whether such measurements in the lab are ecologically relevant. A greater understanding of this issue may allow more accurate assessments of the impacts of hypoxic events on marine fish in nature, aiding management and conservation of fish species. With specific regard to measurements of  $O_{2crit}$  we believe future studies should include concurrent rising  $CO_2$  in the following ways:

- 1. Intermittent-flow respirometer studies should include increases of CO<sub>2</sub> relevant to hypoxic events that organisms may experience,
- 2. Closed respirometry studies should report the start and end  $CO_2$  levels in the respirometer.

In addition, both methods should aim to create an environmentally relevant rate of hypoxia induction/ $CO_2$  increase for the species studied, and consistently report  $CO_2$  levels measured. By incorporating these recommendations we believe that future studies of  $O_{2crit}$  will give more representative estimations of species hypoxia tolerance.

Time held in system (days)	Temperature (°C)	pН	Salinity	Total Alkalinity (mM/kgSW)	pCO <sub>2</sub> (µatm)
318	$18.01 \pm 0.03$	$8.04\pm0.04$	$33.25 \pm 0.70$	$2064.6 \pm 136.0$	$516.9 \pm 41.1$

**Table 1.** Water chemistry parameters of the recirculating aquaculture system in which sea bass were held prior to experimental work (means  $\pm$  S.E. shown).

### Materials and Methods

**Fish collection and husbandry.** We collected juvenile sea bass from estuaries and coastal lagoons on the south Dorset coast and Isle of Wight in June 2017 (Marine Management Organisation permit #030/17 & Natural England permit #OLD1002654). Prior to experimentation, these fish were held in the University of Exeter's Aquatic Resource Centre in an aerated recirculating aquaculture system and fed a commercial pellet at a ration of ~1–2% body weight per day three times a week (for system water chemistry see Table 1). All fish were starved for a minimum of 72 hours prior to the start of all measurements to ensure their metabolism was not affected by digestion (i.e. specific dynamic  $action^{24}$ ). All experimental procedures were carried out under home office licence P88687E07 and approved by the University of Exeter's Animal Welfare and Ethical Review Board.

**Measuring hypoxia tolerance.** We determined oxygen consumption rates ( $\dot{MO}_2$ ) of sea bass using an intermittent-flow respirometer system. The respirometer system set up followed recommendations set out by Svendsen *et al.*<sup>51</sup>. Briefly, the system comprised of a sealed 4.515 L respirometer chamber connected to a recirculating loop, including an in-line recirculating pump (Eheim universal 600, Deizisau, Germany), and a measurement chamber into which a temperature-compensated fibre optic oxygen optode (Firesting O<sub>2</sub> oxygen meter, Pyroscience GmBH, Germany) was placed. Oxygen optodes were calibrated in water at the start of experiments at 100% air saturation and 0% air saturation according to manual instructions. Respirometry was conducted in a semi-closed system consisting of three 100 L experimental tanks fed by a 100 L sump, with overflowing water from the experimental tanks recirculating back to the sump. A second pump was used to periodically flush the respirometer system with water from the surrounding tank. This pump was controlled by an automated computer program (AquaResp 3, AquaResp<sup>®</sup>) to intermittently flush the respirometer. Five respirometer chambers were distributed between the three experimental tanks (maximum of 2 chambers per experimental 100 L tank). The sump was temperature controlled (18.27 ± 0.02 °C, mean ± S.E.) using a heater/chiller unit (Grant TX150 R2, Grant Instruments, Cambridge, UK) attached to a temperature exchange coil. Together these tanks formed a 400 L system with the same temperature, oxygen and water chemistry parameters for all respirometers.

Individual sea bass (average mass =  $131.2 \pm 7.5$  g), chosen at random, were placed inside the respirometers and allowed an overnight recovery period, for a minimum of 13 hours, before O<sub>2crit</sub> tests began. While sea bass were in the respirometers measurements of  $\dot{MO}_2$  were conducted every 10 minutes, including a flush period of 300 s, a wait period of 60 s and a measurement period of 240 s. During the wait and measurement period the chamber was sealed by switching off the flush pump and the decline in dissolved O<sub>2</sub> within the chamber was continuously measured by the fibre optic O<sub>2</sub> electrode.

Following the overnight recovery period a standard  $O_{2crit}$  test was conducted. Oxygen levels in the respirometer system were reduced from ~100% air saturation to ~15% air saturation over the course of 6 hours (decline in  $O_2$  was ~20% air saturation per hour between ~100% air saturation and ~40% air saturation and ~10% air saturation per hour from ~40% air saturation until the end of trials). Oxygen was regulated by gassing the sump and experimental tanks with a mix of  $N_2$  and  $O_2$  (G400 Gas mixing system, Qubit Biology Inc.) at a rate of 10 L min<sup>-1</sup> following a pre-set automated protocol (Flowvision, Alicat software). Levels of  $CO_2$  within the system were controlled under one of two treatments (with 8 fish exposed to Treatment 1 and 8 separate fish exposed Treatment 2):

- Treatment 1 (Constant CO<sub>2</sub>) ambient levels of CO<sub>2</sub> were maintained by including 0.04% CO<sub>2</sub> as part of the gas mix delivered to the respirometer system.
- Treatment 2 (Rising CO<sub>2</sub>) the proportion of CO<sub>2</sub> in the gas mix was gradually increased as O<sub>2</sub> was decreased. This increase in CO<sub>2</sub> was designed to reflect environmentally realistic increases in CO<sub>2</sub> predicted as a result of depletion of O<sub>2</sub> by bacterial respiration (assumed respiratory quotient of 1)<sup>52</sup>, using the seawater carbonate chemistry calculator CO2sys (see supplementary material for predictions of increased CO<sub>2</sub> during hypoxia).

Water chemistry of treatment 2 was monitored once per hour by measuring pH<sub>NBS</sub>, temperature and salinity as well as taking a 12 mL water sample to measure total CO<sub>2</sub> (TCO<sub>2</sub>)/Dissolved Inorganic Carbon (DIC). Water chemistry of treatment 1 was monitored at the start and end of the treatment to ensure no change in water  $pCO_2$  occurred over the time course of the O<sub>2crit</sub> trial. Seawater DIC analysis was conducted using a custom built system described in detail by Lewis *et al.*<sup>53</sup>. These four parameters were then input into the seawater carbon calculator programme, CO2SYS to calculate  $pCO_2$  based on the NBS pH scale, equilibration constants from Mehrbach *et al.* refitted by Dickson and Millero, and KSO<sub>4</sub> dissociation constants from Dickson. The reciprocal changes in O<sub>2</sub> and CO<sub>2</sub> during O<sub>2crit</sub> tests for each treatment are illustrated in Fig. 4. O<sub>2crit</sub> tests were stopped once a minimum of 3  $\dot{M}O_2$  measurements showed a transition from an oxy-regulating to oxy-conforming state for each fish or fish showed a large drop in  $\dot{M}O_2$  and signs of distress in the respirometer. No fish exhibited LoE during trials. Following completion of O<sub>2crit</sub> trials experimental tanks were aerated with ambient air to swiftly restore O<sub>2</sub> and CO<sub>2</sub> levels.

Sea bass were left to recover in respirometers, for a minimum of 1 hour post-trial, until  $O_2$  levels reached ~21 kPa  $O_2$  (~100% air saturation). Fish were then removed from respirometers and background respiration was measured for a minimum of 1 hour (6 measurement cycles) for all respirometers immediately post trial.



**Figure 4.** Changes in partial pressure of  $O_2$  (expressed as kPa  $O_2$ ) and  $CO_2$  (µatm) during (**A**) two  $O_{2crit}$  trials representing treatment 1 in which  $O_2$  was reduced with no change in  $CO_2$ ; and (**B**) two trials representing treatment 2 where  $O_2$  was reduced with a corresponding rise in  $CO_2$ . Data presented are means  $\pm$  S.D.

*Oxygen consumption rate*  $(\dot{M}O_2)$  *analysis.* Following each 240 s measurement period  $\dot{M}O_2$  was automatically calculated by the AquaResp3 software. A linear regression was fitted to the  $O_2$  versus time data for each measurement period. The slope of this regression (s, kPa  $O_2$  h<sup>-1</sup>) was then used to calculate  $\dot{M}O_2$  (mg  $O_2$  kg<sup>-1</sup> h<sup>-1</sup>) using the equation outlined by Svendsen *et al.*<sup>51</sup>:

$$\dot{M}O_2 = sV_{resp}\alpha m^{-1}$$

where  $V_{resp}$  is the respirometer volume minus the volume of the fish (L),  $\alpha$  is the solubility of  $O_2$  in water (mgO\_2  $L^{-1} k Pa^{-1}$ ) for the relevant salinity and temperature, and m is the mass of the fish (kg). Calculations of  $\dot{M}O_2$  where s had a  $R^2$  of <0.98 were removed from subsequent analysis. For the purpose of establishing  $O_{2crit}$  values from a plot of  $\dot{M}O_2$  versus ambient  $O_2$  level, the oxygen saturation of each measurement period was defined as the average of the dissolved  $O_2$  measurement over the measurement period. The average background respiration, over the 1 hour post-trial measuring period, for each respirometer (average background respiration was <2% of fish  $\dot{M}O_2$ ) was subtracted from  $\dot{M}O_2$  measurements to correct fish  $\dot{M}O_2$  for background respiration.

We calculated the standard metabolic rate (SMR) in R v3.5.3<sup>54</sup> using function calcSMR in package fish  $\dot{M}O_2^{24}$ . All  $\dot{M}O_2$  values from the overnight recovery period and beginning of the  $O_{2crit}$  trial in which average dissolved  $O_2$  saturation was >80% air saturation were included for SMR calculations. This resulted in approximately 14–16 hours of  $\dot{M}O_2$  data to calculate SMR from for each fish. We estimated SMR for all fish using the mean of the lowest 10  $\dot{M}O_2$  measurements during the overnight period. Although Chabot *et al.*<sup>24</sup> recommend use of the mean of the lowest normal distribution (MLND) or quantile (q = 0.2) methods to calculate SMR we chose the mean of the 10 lowest values as it produced a value of SMR that most accurately matched the consistent low  $\dot{M}O_2$  measurements of oxy-regulating fish at values of  $pO_2$  above the  $O_{2crit}$  point. Additionally, for each fish the coefficient of variation in the mean lowest normal distribution (MLND) was assessed using a ROUT test (Q = 1%) to check whether variation in low  $\dot{M}O_2$  measurements was consistent between fish (to account for potential differences in activity in the respirometer prior to  $O_{2crit}$  trials). The ROUT test removed one fish from the rising CO<sub>2</sub> treatment which displayed a CoV of 34.6. Mean CoV of the remaining 15 fish was 11.85 ± 1.26 (±S.E.M).

We then used function calcO2crit from the package to calculate the  $O_{2crit}$  for each individual fish, using the estimated SMR of each individual, as detailed in the supplementary material of Claireaux & Chabot<sup>55</sup>. This function identifies the portion of the  $O_{2crit}$  test where metabolic rate data follows an oxygen conforming relationship and fits a linear regression line through this data,  $O_{2crit}$  is then calculated as the oxygen level at which this regression line crosses the calculated SMR of the individual fish. Plots of  $\dot{M}O_2$  against  $O_2$  during  $O_{2crit}$  trials showing calculated  $O_{2crit}$  for each individual can be seen in the supplementary material (Supplementary Material Figs 1–16). Calculations were conducted using a gap limit of 0.83 kPa  $O_2$  (4% air saturation) and a maximum number of 7  $\dot{M}O_2$  points to fit the regression line through the oxygen conforming component of the data used to estimate  $O_{2crit}$ .

**Measuring blood chemistry.** Following  $O_{2crit}$  trials sea bass were moved to individual 7 L chambers, which were aerated and supplied with seawater from the aquarium re-circulating system at a rate of about 4 L min<sup>-1</sup>.

After an overnight acclimation period we then exposed the fish to a decrease in  $O_2$  levels to ~6.4 kPa  $O_2$  (30% air saturation) over a period of 4 hours (equivalent to the rate of decrease in  $O_2$  used in the previous  $O_{2crit}$  tests). We chose to blood sample fish at an O<sub>2</sub> level above O<sub>2crit</sub> to ensure that anaerobic metabolism did not influence blood chemistry. This was combined with the same  $CO_2$  regime each fish experienced during the  $O_{2crit}$  test (i.e. either constant or reciprocally rising CO<sub>2</sub>). Once an O<sub>2</sub> level of ~6.4 kPa O<sub>2</sub> (30% air saturation) was achieved the fish were allowed to acclimate for 1 hour before being individually anaesthetised in situ using a dose of 100 mgL<sup>-1</sup> of benzocaine. Once fish were judged to be sufficiently anaesthetised (cessation of gill ventilation and lack of response to a pinch of the anal fin) they were immediately transferred within 5 seconds to a gill irrigation table (with the same  $pO_2$  and  $pCO_2$  levels of the respective treatment), where anaesthesia was maintained with a dose of  $37.5 \text{ mgL}^{-1}$  of benzocaine. Gill ventilation was artificially maintained by a micro-pump, so that the operculum were just open and exhalant water flow could just be visualised. Once a stable gill water flow was established blood was sampled by caudal vessel puncture using a 1 ml heparinised syringe. This method has been demonstrated to obtain accurate measurements of blood chemistry parameters comparable to those achieved using cannulation (Davison & Wilson, University of Exeter, personal communication). At the time of blood sampling water  $pCO_2$ was  $656 \pm 44 \mu \text{atm}$  (mean  $\pm$  S.E.) for fish in the constant ambient CO<sub>2</sub> regime and 1763  $\pm$  43  $\mu \text{atm}$  (mean  $\pm$  S.E.) for fish in the progressively rising CO<sub>2</sub> regime. Water  $pO_2$  was  $8.1 \pm 0.2$  kPa (mean  $\pm$  S.E., ~38% air saturation) for fish in the constant ambient CO<sub>2</sub> regime and  $8.7 \pm 0.2$  kPa (mean  $\pm$  S.E., ~41% air saturation) for fish in the progressively rising CO<sub>2</sub> regime. Following blood sampling fish were transferred to seawater isolation tanks containing ~20.8 kPa O<sub>2</sub> (100% air-saturated) to recover from the anaesthetic. They were then monitored over a period of 24 hours before we returned them to their original holding tanks.

Immediately after sampling, whole blood pO2 was measured at 18 °C in a temperature-controlled system (Strathkelvin 1302 electrode and 781 meter; Strathkelvin Instruments Ltd, Glasgow, UK). We measured extracellular pH on 30 µL of whole blood using an Accumet Micro pH electrode and Hanna HI8314 pH meter at 18 °C calibrated to pH<sub>NBS</sub> 7.04 and 9.21 specific buffers. Three 75 µL micro capillary tubes were then filled with whole blood and sealed with Critoseal capillary tube sealant (Fisher) and paraffin oil and centrifuged for 2 minutes at 10,000 rpm. Haematocrit was measured using a Hawksley micro-haematocrit reader. Plasma was extracted from these tubes for analysis of TCO<sub>2</sub> using a Mettler Toledo 965D carbon dioxide analyser. Plasma  $pCO_2$  and  $HCO_3^-$  were then calculated from TCO<sub>2</sub>, temperature and blood pH using the Henderson-Hasselbalch equation with values for solubility and pK<sup>1</sup><sub>app</sub> based on Boutilier *et al.*<sup>56,57</sup>. Haemoglobin content of the blood was also assessed by the cyanmethemoglobin method (using Drabkin's reagent, Sigma). Half the remaining whole blood was then centrifuged at 10,000 rpm for 2 minutes at 4 °C. The resulting plasma was separated and snap frozen in liquid nitrogen and stored at -80 °C before later being used to measure plasma glucose and lactate using a YSI 2900D Biochemistry Analyzer (Xylem Analytics, UK). All measurements or storage of blood occurred within 10 minutes of blood sampling.

**Measuring Hb-oxygen binding.** We measured Hb-O<sub>2</sub> affinity using a Blood Oxygen Binding System (BOBS, Loligo systems), detailed in Oellermann *et al.*<sup>58</sup>. A sample of the same whole blood used for blood chemistry measurements was diluted at a ratio of 1:4 in its own plasma. 1  $\mu$ L of this blood was then used for measurements. The BOBS exposed this blood sample to gas mixes with a progressive increase in O<sub>2</sub> whilst measuring absorbance of light across a spectrum ranging from 200 to 800 nm. For each individual fish the gas mix that blood was exposed to matched the calculated *p*CO<sub>2</sub> of the blood sample. The change in absorption of light at a wavelength of 435 nm was used to assess changes in oxygenation of Hb, as previously used by Verhille & Farrell<sup>59</sup>. Background changes in absorption of the blood sample were corrected using the isosbestic wavelength of 390 nm<sup>59</sup>. Following measurements the BOBS calculated the oxygen equilibrium curve of the sample using Hill's formula before estimating P<sub>50</sub><sup>58</sup>.

**Statistical analysis.** We conducted all statistical analysis in R v3.5.3<sup>54</sup>. There was no significant difference in mass of fish between the treatment groups (One-Way ANOVA,  $F_{1,13} = 2.821$ , p = 0.117) or of estimated SMR of fish between treatment groups (Unpaired t-test, t = 0.8455, d.f. = 13, p = 0.413). There was, however, a significant correlation between estimated SMR and calculated  $O_{2crit}$  for all fish (Pearson's correlation, df = 13, t = 3.32, R = 0.68, p = 0.0056). As such the effect of CO<sub>2</sub> treatment on O2<sub>crit</sub> was assessed using an ANCOVA with SMR as a covariate. All other analyses were conducted using general linear modelling (GLM). All values in the text are reported as mean  $\pm$  standard error (S.E.).

#### Data availability

Data is available via the University of Exeter's online repository at: https://doi.org/10.24378/exe.1523.

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#### **Author contributions**

D.W.M. contributed to study concept and design, conducted all data collection, analysed all data and drafted the manuscript. S.D.S. contributed to study concept and design, and revisions of the manuscript. G.H.E. and S.N.R.B. contributed to study concept and design, data analysis and revisions of the manuscript. R.W.W. was responsible for study supervision, contributed to study concept and design, assisted with data collection, helped with interpretation of data, and contributed to revisions of the manuscript.

#### **Competing interests**

The authors declare no competing interests.

#### Additional information

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