# **RESEARCH ARTICLE**



# **Biochemical Barriers on the Path to Ocean Anoxia?**

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**ABSTRACT** The kinetics of microbial respiration suggests that, if excess organic matter is present, oxygen should fall to nanomolar levels in the range of the Michaelis-Menten constants ( $K_m$ ). Yet even in many biologically productive coastal regions, lowest observed  $O_2$  concentrations often remain several orders of magnitude higher than respiratory  $K_m$  values. We propose the hypoxic barrier hypothesis (HBH) to explain this apparent discrepancy. The HBH postulates that oxidative enzymes involved in organic matter catabolism are kinetically limited by  $O_2$  at concentrations far higher than the thresholds for respiration. We found support for the HBH in a meta-analysis of 1,137  $O_2$  $K_m$  values reported in the literature: the median value for terminal respiratory oxidases was 350 nM, but for other oxidase types, the median value was 67  $\mu$ M. The HBH directs our attention to the kinetic properties of an important class of oxygen-dependent reactions that could help explain the trajectories of ocean ecosystems experiencing  $O_2$ stress.

**IMPORTANCE** Declining ocean oxygen associated with global warming and climate change is impacting marine ecosystems across scales from microscopic planktonic communities to global fisheries. We report a fundamental dichotomy in the affinities of enzymes for oxygen—the terminal proteins catalyzing respiration are active at much lower oxygen concentrations than oxygenase enzymes involved in organic matter catabolism. We hypothesize that this dichotomy in oxygen affinities will cause some types of organic matter to accumulate in hypoxic ecosystems and will slow rates of oxygen decline. This proposed biochemical barrier may explain why many ocean ecosystems rarely reach anoxia. Competition between intracellular enzymes for oxygen may also have impacted microbial strategies of adaptation to low oxygen, requiring cells to regulate oxygen respiration so that it does not compete with other cellular processes that also require oxygen.

**KEYWORDS** dissolved organic matter, ocean respiration, oxygen minimum zones, oxygenase  $K_m$ , ocean respiration, dissolved organic matter

Marine suboxic and anoxic zones are hot spots of microbially mediated biogeochemical transformations that regulate the nitrogen budget and air-sea fluxes of greenhouse gases of the global ocean (1). Because dissolved oxygen (DO) also organizes the structure and dynamics of ocean food webs, understanding the processes that regulate expansion of suboxic and anoxic zones in response to past and current climate changes is a pressing challenge (2). Suboxic and anoxic zones are embedded within broader oxygen minimum zones (OMZ) that comprise some 8% of the surface area of the ocean. While recent advances in nanomolar-scale DO measurement technologies have enabled precise delineation of the presence of suboxia and anoxia (3), we contend that **Citation** Giovannoni S, Chan F, Davis E, II, Deutsch C, Wolf S. 2021. Biochemical barriers on the path to ocean anoxia? mBio 12:e01332-21. https://doi.org/10.1128/mBio.01332-21.

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**FIG 1**  $K_m$  values are significantly smaller in respiratory oxidases compared to other oxygenases.  $K_m$  DO (dissolved oxygen) values for respiratory oxidases (yellow; n = 109) and other oxygenases (blue; n = 890) are depicted on a  $\log_{10}$  scale. The relatively high (e.g.,  $10^1 \mu$ M)  $K_m$  values reported for oxidase enzymes indicate a potential bottleneck in the supply of electrons from organic matter to respiration. The reported *P* value is from a *t* test using the Satterthwaite approximations to degrees of freedom of a linear mixed model fit by maximum likelihood. Data and citations can be found in Table S1 in the supplemental material.

a perplexing yet fundamental question has been overlooked. Given our canonical understanding of microbial respiration kinetics, why are suboxia and anoxia not a much more pervasive feature of the ocean's low-oxygen zones?

Of biological reactions that consume  $O_2$ , by far the most important, in terms of mass, is carbon respiration. Michaelis-Menten half-saturation ( $K_m$ ) constants for respiration are typically very low, on the order of a few nanomolar, although higher values have been reported (4) (Fig. 1). Thus, if labile organic carbon, i.e., compounds that readily can be used as a source of electrons for respiration, is delivered in excess to a microbial ecosystem, DO declines at a rate determined by the respiratory capacity of the microorganisms present and the supply of organic matter. Importantly, the minimum DO attainable should reflect the well-described high-affinity, nanomolar-scale  $K_m$  of microbial respiratory oxidases (5). Other factors that can influence DO in aquatic ecosystems include photosynthesis when light is present; oxygen transport by ocean currents and mixing; diffusion, which can limit respiration, particularly in aggregates of cells; impacts of low oxygen on grazing metazoa (6), which require higher oxygen concentrations than bacteria; nonrespiratory biochemical reactions that consume oxygen; and abiotic reactions that consume oxygen (7). Nonetheless, DOM formation and oxidation are the mechanistic centerpiece in our fundamental understanding of microbial-scale processes leading to low-oxygen states and predictions of global ocean oxygen dynamics.

Eastern boundary upwelling systems (EBUS) represent one of the ocean's most productive biomes. In these coastal ecosystems, oxygen-poor subsurface waters uplifted from the vertical periphery or core of open-ocean OMZs receive elevated organic carbon inputs from surface phytoplankton blooms. Figure 2 shows relative water volumes for DO concentrations across these systems. Of the ocean's four major EBUS, only the Peru-Chile current system in the Eastern Tropical Pacific Ocean persistently exhibits DO-deficient states in continental shelf waters. For both Pacific EBUS, there is an accumulation of water volumes below  $100 \,\mu$ M but a sharp drop-off in volume of waters that reach suboxic ( $<5 \,\mu$ M) or anoxic ( $\sim 0 \,\mu$ M) states in the upper ocean (0 to 400 m), including continental shelf waters where remineralization and oxygen loss are most active (Fig. 2b). The pattern is striking; despite the nanomolar-scale of respiratory  $K_m$ values, respiration in productive EBUS is able to draw down DO to hypoxic levels but rarely is able to consume the last 10 to  $60 \,\mu$ M DO. While the depth of OMZs extends below 400 m, the failure of suboxic and anoxic volumes to accumulate despite the presence of large volumes of hypoxic water persists when we expand our sampling to 1,000 m (Fig. 2d).



**FIG 2** Climatological distribution and volumetric census of DO. Observation-based maps of DO averaged over the upper (a) 0-400 m and (c) 0-1000 m of the global ocean reflect the combination of temperature-dependent gas saturation in surface water and the integrated consumption of DO during respiration of organic matter over decadal to centennial time scales of ocean circulation at subsurface depths. For each depth range, the volume of water at each DO level (panel b and d) is summed globally and for major EBUS regions: the California Current System (CCS), Eastern Tropical Pacific (EastTropPac), and Eastern Tropical Atlantic (EastTropAtl), which all exhibit a peak at low DO, albeit at different concentrations.

To explain the observations in Fig. 2, we propose the hypoxic barrier hypothesis (HBH), which states that dissolved  $O_2$  kinetically limits the activity of oxygenase enzymes involved in the breakdown of organic matter in the range of oxygenase  $K_m$  values (median value,  $67 \mu$ M) causing a decline in DOM oxidation rates in ecosystems experiencing oxygen stress and an accumulation of DOM that is catabolized by pathways that require oxygenases. The HBH ascribes the decline in  $O_2$  frequency distributions of suboxic and anoxic waters (Fig. 2 and Fig. S2 in the supplemental material) to fundamental biochemical properties of cells, particularly the mechanisms by which oxidative enzymes cleave semilabile organic matter, making it accessible to further oxidation.

Oxygen depletion by respiration in aquatic systems. How far can respiring marine bacteria lower oxygen concentrations when they are provided with an ample supply of reductant for respiration, as would be expected for plankton in the presence of an excess of labile organic carbon? The Pasteur point is an influential concept based on the observation that facultative anaerobes switch to fermentation at ca. 2.2  $\mu$ M O<sub>2</sub>, approximately an order of magnitude below the average  $K_m$  for oxygenases (Fig. 1), steep declines in suboxic and anoxic water volumes in EBUS (Fig. 2), and the ca.  $25-\mu$ M inflection in cumulative frequency distribution of DO observations recorded in the California current system (CCS) (Fig. S2). Newer information suggests that the limits of bacterial respiration are in the nanomolar range. This is consistent with the observation of high-affinity cytochromes that exhibit  $O_2 K_m$  between 3 and 200 nM (8) (Fig. 1). Higher O<sub>2</sub> Michaelis constants have sometimes been reported for marine bacteria, but it has been suggested that higher values obtained with whole cells reflect diffusion limitation, which can be expected to inflate apparent  $O_2$  Michaelis constants in proportion to cell sizes and respiration rates. Stolper et al. showed that Escherichia coli cells could grow at less than 3 nM O<sub>2</sub>, a sufficiently low concentration to limit growth by

diffusion but high enough to sustain growth through O<sub>2</sub> respiration (8). Our meta-analysis indicates that cells grown on highly labile carbon compounds, such as glucose, display whole-cell O<sub>2</sub>  $K_m$  values that extend to nanomolar O<sub>2</sub> concentrations. We conclude that a substantial background of observations and theory supports the conclusion that the respiration rate of chemoheterotrophic cells should not be limited by O<sub>2</sub> at concentrations found in ocean hypoxic zones or the Pasteur point at ca. 2.2  $\mu$ M.

The accumulation of hypoxic and scarcity of suboxic or anoxic volumes in the ocean nonetheless suggest that negative feedbacks between oxygen decline and respiration may be at play. Direct measurements of microbial  $O_2 K_m$  in natural systems are rare, but available evidence points to  $K_m$  values far higher than the nanomolar values reported from laboratory cultures with labile carbon sources. Working in the Arabian Sea OMZ, Keil et al. (9) observed an apparent  $K_m$  of 20  $\mu$ M O<sub>2</sub> for microbial community respiration. In the Nambian and Peruvian OMZ, Kalvelage et al. (10) reported a linear decline in respiration rate between 20 and  $0 \,\mu$ M O<sub>2</sub>. In the Chesapeake Bay, a hypoxiaprone system, microbial respiration rates saturate at O<sub>2</sub> above 25  $\mu$ M (11), a pattern that we have similarly found for the CCS OMZ (Fig. S3). Holtappels et al. (12) further reported linear declines in respiration rates between 14 and  $1 \mu M O_2$  in waters collected from a fjord in Denmark. These results are surprising because researchers using the same methods have also found many instances of nM K<sub>m</sub> values for microbial respiration. This suggests a bimodal distribution of  $O_2 K_m$  values that differ by upwards of 3 orders of magnitude. Telescoping out further, global models of ocean O2 and carbon export converge on  $K_m$  values of between 4 and 20  $\mu$ M O<sub>2</sub> in order to optimize fit between model and observations (13, 14). What accounts for the disparity between accumulation of hypoxic water volumes, the micromolar-scale  $K_m$ s reported from natural systems and used to fit models and nanomolar-scale K<sub>m</sub>s predicted by respiratory oxidases?

Biochemistry offers a mechanistic explanation for this apparent disparity. Evidence in the scientific literature suggests that microbial respiration of some types of organic matter slows when oxygen concentrations fall low enough to inhibit catabolic oxygenase enzymes: Kroonman et al. (15), studying 3-chlorobenzoate degradation by the bacterium *Alcaligenes*, reported two  $K_m$  values for O<sub>2</sub> uptake. They attributed the lower value (65 nM) to respiration and the higher value (7 to 17  $\mu$ M) to the activity of dioxygenases. Leahy and Olsen (16), studying toluene degradation by *Pseudomonads*, also reported biphasic kinetics for toluene catabolism as a function of oxygen concentration. The slope of the oxygen response declined with an inflection at 20 to 30  $\mu$ M O<sub>2</sub>. In both of these cases, the behavior of the cultured cells oxidizing recalcitrant compounds is remarkably similar to the generalized behavior of ocean ecosystems approaching hypoxia.

To further explore the distribution of  $O_2 K_m$  values among biological reactions, we conducted a meta-analysis of published data, shown in Fig. 1, Fig. S1A, and Table S1. For  $O_2 K_m$  values reported in the literature, the median value for terminal respiratory oxidases was 350 nM, but for other oxidase types, the median value was  $67 \mu$ M. The difference of ~100-fold in median values was supported by a *P* value of <2e-16 in a *t* test of the linear mixed-effect model coefficient comparing the log-transformed  $O_2 K_m$  values (17). The bimodal distribution of  $O_2 K_m$  observed at the enzyme scale is also repeated in whole-cell studies. Cells that are grown on more complex organic carbon sources have a median respiratory  $K_m$  value of 20  $\mu$ M, while cells grown on highly labile organic carbon such as glucose have median  $K_m$  of 690 nM (Fig. S1B).

Many enzymes that catalyze the biological breakdown of organic matter use oxygen as a substrate, yielding partially oxidized products that are metabolized further through catabolic pathways. These enzymes are often classified as either monooxygenases (mixed-function oxidases) or dioxygenases. Enzymes in both families evolved to use  $O_2$  as a substrate, but monooxygenases incorporate a single oxygen atom into the substrate, reducing the second atom to water, whereas dioxygenases typically add both atoms of the reacting  $O_2$  to the product. A geochemically important example of a

monooxygenase is the heme-dependent Mn peroxidase that catalyzes oxidation of lignin, a phenolic oligomer. Fungal ligninases belong in the heme-dependent peroxidase superfamily (18). Ligninases evolved in the Paleozoic era, and it has been postulated that their origin resulted in widespread biodegradation of wood, the end of the carboniferous period, and rises in global atmospheric  $CO_2$  (19). Another superfamily of oxidases, the flavin-dependent monooxygenases, are among the most diverse and prevalent proteins known. They catalyze a wide range of reactions, for example, hydroxylation, Baeyer-Villiger oxidation, oxidative decarboxylation, epoxidation, desulfurization, sulfoxidation, and oxidative denitration (20). Many of the reactions catalyzed by flavin-dependent monooxygenases initiate the catabolism of compounds that are otherwise recalcitrant to oxidation. These enzymes share a mechanism in which reduced flavin reacts with oxygen to produce a flavin C4a-(hydro)peroxide that then reacts with electrophilic or nucleophilic substrates, typically resulting in the consumption of one diatomic oxygen molecule, the addition of an oxygen atom to the substrate, and the release of water. Also important are the dioxygenases, which belong to a different protein family and also feature prominently in DOM degradation, particularly for aromatic compounds. These protein families came to the attention of oceanographers recently when it was discovered that cells of one of the important oceanic bacterial clades, SAR202, harbor expanded clusters of paralogous genes from both of these protein types (21). It has been proposed that these enzymes participate in the oxidation of semilabile organic matter, initiating its breakdown.

The meta-analysis of  $O_2 K_m$  values presented in Fig. 1 suggests that micromolar DO sensitivity is ubiquitous across metabolic processes. Among the "other oxidase" types, we observed no clear trends that associated  $O_2 K_m$  values with protein families sorted by the Clusters of Orthologous Groups (COG) database or other precise functional groups, such as enzyme commission classifications. In contrast, cytochrome respiratory proteins with heme cofactors consistently displayed a much higher affinity for oxygen than other protein types. Phylogenetically, the distribution of  $O_2 K_m$  values included diverse bacteria, including *Proteobacteria*, *Actinobacteria*, *Firmicutes*, and *Cyanobacteria*, as well as eukaryotic organisms, including fungi, humans, and other chordates.  $O_2 K_m$  values showed a 100-fold difference between respiratory and nonrespiratory oxidases regardless of taxonomic group. We conclude that this pattern is robust to phylogenetic bias in  $O_2 K_m$  value sampling. While our focus in the current work is marine systems, the HBH in principle applies to all ecosystems.

The role of oxidases in organic matter degradation. Oceanographers classify organic matter by its half-life, frequently using the category "labile dissolved organic matter" (LDOM) to describe dissolved organic matter that is oxidized in minutes to hours, or at most, a few days, while the term "semilabile" (SLDOM), and sometimes "recalcitrant," is used to refer to dissolved organic matter that persists longer but is eventually oxidized. Here, we introduce a new term, "oxygen-dependent DOM" (ODDOM), to describe DOM that is catabolized via reactions that require the activity of oxygenases and thus are susceptible to inhibition when  $O_2$  concentrations reach values in the range of ca. 10 to  $100 \,\mu$ M. We'll confine the discussion to dissolved forms of organic matter, although most organic matter enters ecosystems as particulate organic matter (POM) and is subsequently converted to DOM before being used by microorganisms. Implicit in the above categories is the idea that different kinds of organic matter are accessible to biological oxidation through different mechanisms and at different rates.

The HBH is consistent with the distribution of ocean anoxic zones if one assumes organic matter supplies are uneven. If LDOM is oversupplied relative to oxygen, for example, by high rates of export production in systems with restricted circulation, then the activities of respiratory terminal cytochrome complexes would be expected to readily draw down DO to nanomolar concentrations in accordance with their nanomolar  $K_m$  values. In natural systems, LDOM is rapidly depleted. As hypothesized, the activities of nonrespiratory oxidases limit the supply of reductant to respiratory oxidases. This acts as a bottleneck that slows the rate of respiration as DO declines. With sufficient time, DO should reach minimum values as expected from nanomolar  $K_m$  values of respiratory oxidases. Such conditions can be met in the core of OMZs that have been isolated from the atmosphere over decadal to century time scales, and evidence of this can be seen in Fig. 2.

The large disparity in  $K_m$ s we report between respiratory oxygenases and other oxygenase types has implications for microbial cell evolution and metabolic regulation at the cellular level. Inside of cells' respiratory oxygenases could outcompete other oxygenases, exacerbating the slowing of some oxygen-dependent cellular processes at low oxygen. To avoid this, cells may have evolved metabolic regulation that avoids such competitive interactions, for example, by shifting to alternate electron acceptors before  $O_2$  is depleted (22). This topic, which needs exploration, could help us understand how microbial cells have adapted to suboxic environments, which are far more common in the ocean than anoxic environments.

**Testing the HBH.** The HBH sets forth a number of central predictions that are testable by experimentation, observation, and modeling. The impact of biphasic oxygen dependence predicted by the HBH should be manifested as a broad potential for oxygen to limit microbial respiration across hypoxic systems in the range of oxygenase  $K_m$  values (median value,  $67 \mu$ M) when LDOM is depleted, but not if excess LDOM is present. To test that prediction, we measured rates of respiration (oxygen uptake) in water samples from the Northern California Current System OMZ, where DO minimum reach only  $\sim 5 \mu$ M, well above canonical nanomolar  $K_m$  for cytochrome oxidases. DO was increased by the simple expedient of allowing air to be momentarily entrained during filling (Fig. S3). In our experiments and other similar experiments, we found, among published work, the addition of DO caused respiration rates to rise relative to controls. This observation could be attributed to the limitation of respiration by diffusion (20), but alternatively, it could result from mechanisms described in the HBH model we propose.

There are many other experimental avenues to testing the HBH that have not been explored. Figure S2 scratches the surface of what could done with field experiments and mesocosms to verify predictions of the HBH. For example, experiments that test the biological availability of DOM at high (e.g., 200  $\mu$ M) and moderate (e.g., 20  $\mu$ M) DO could challenge these ideas. Mechanisms invoked by the HBH would lead to changes in the chemical composition of DOM as DO declines: the ratio of LDOM to ODDOM should decrease as DO approaches the  $K_m$  values of catabolic oxygenase enzymes for O<sub>2</sub>. Measurements of DOM chemistry could determine whether these changes occur as predicted. ODDOM, a term coined herein to segregate DOM into categories by chemical composition and oxygenase involvement in catabolism, is at present a theoretical concept, albeit grounded in the fundamentals of biochemistry. Although chemical oceanographers do not at present measure ODDOM, in principle methods such as high-resolution nuclear magnetic resonance (NMR), high-performance liquid chromatography (HPLC), and liquid chromatography-tandem mass spectrometry (LC-MS/MS) could be applied for this purpose and could be used to test predictions of the HBH. Omics approaches, including functional genomics, provide an avenue that could be applied in marine systems to measure the expression and activity of oxygenase enzymes involved in ODDOM metabolism and to characterize the responses of plankton cells and communities to suboxia.

The HBH has broader implications that could be explored with global data. It posits that rates of oxygen loss and DOM oxidation slow as DO approaches hypoxia, setting the upper bounds for the size of oceanic anoxic zones and organic carbon pools within. This can be evaluated in detail by modeling studies that test the sensitivity of model data comparisons to changes in assumptions about microbial kinetic constants for oxygen.

**Alternatives to the HBH.** While we propose HBH to explain declines in oxygen frequency distributions at unexpected high values (Fig. 2 and Fig. S2) and the rarity of suboxia and anoxia across productive, low-oxygen EBUS, alternate scenarios could explain this phenomenon. For example, consistent barriers to oxygen diffusion to the terminal oxidases of respiratory systems, feedback mechanisms involving the production of sulfides and/or depletion of DO in microhabitats, or oxygen limitation of metazoan grazing could play a role in suppressing respiration at low oxygen concentrations. Alternatively, there may be constraints on supply of organic carbon or positive feedbacks on the resupply of DO by advection or diffusion as DO approaches hypoxia.

Public interest and policy. The relevance of this issue to public interests in ecosystem management could not be more profound. Ocean deoxygenation, the decline in ocean oxygen inventories, has emerged as a leading pathway for climate change impacts in the sea. This decline has been linked with expansion of hypoxic and anoxic zones. Oxygen-deficient zones are hot spots of biogeochemical transformations whose growth can have profound impacts on marine biodiversity, vertical organic carbon flux, the sustainability of fisheries, and feedbacks that govern ocean nitrogen budgets and flux of radiatively active N<sub>2</sub>O. The ability to accurately forecast such ecosystem changes is central for informing responsive climate change mitigation and adaptation policies. However, the disagreement between observations and the textbook understanding of microbial respiration raises fundamental questions about the mechanisms that underlie our conceptual and numerical models of the ocean dynamics as climate change intensifies. The HBH offers a testable framework for examining a potentially flawed fundamental principle that governs our thinking about OMZ formation. If this hypothesis is correct, it will open previously overlooked avenues of research at the intersection of oxygenase enzyme evolution, oxygenase-dependent metabolism in microbial communities, and OMZ dynamics.

**Conclusion.** If these ideas have the power to even partially explain the kinetics of ocean oxygen depletion, they could contribute to a better understanding of climate change impacts on ocean deoxygenation and DOM chemistry. The data in Fig. 1 show us that the HBH is founded on sound basic principles, but the impact of oxygenase "barrier" we describe is relative to many other processes, mentioned above, that can also slow respiration, most notably diffusion. Sorting out the magnitude of catabolic oxygenase enzyme contributions to DOM oxidation, whether that number be large or small, will help us assess how the trajectories of aquatic systems experiencing oxygen declines are shaped by the fundamental biochemistry described in the HBH.

## **MATERIALS AND METHODS**

**Data collection.** Scientific literature was mined for characterized oxygenase enzymes with published  $K_m$  values for dissolved oxygen for both individual enzyme assays as well as whole-cell assays (Table S1 in the supplemental material). Metadata, including enzyme name and host scientific organism name, were extracted from each article. We used a combination of the BRENDA enzyme database, UniProt protein database, and KEGG database searches to determine putative protein accessions, KEGG ortholog IDs, and EC numbers associated with the published enzyme data. Repeated entries for the same organism-protein pairs were included due to the various testing conditions per study.

Basin and global inventories of DO volumes were compiled from the World Ocean Atlas 2018 (https:// www.ncei.noaa.gov/products/world-ocean-atlas). Dissolved oxygen observations (5 to 400 m) from CTD profiles were compiled from Chan et al. 2008, https://www3.mbari.org/bog/, and https://www.calcofi.org/ for the northern (n = 107,032; 1950 to 2006), central (n = 4,372; 1997 to 2013), and southern (n = 4,372; 1997 to 2013) CCS, respectively (23).

**Respiration rate experiment.** Water samples were drawn from above and within the CCS OMZ (46° 47.56'N, 125°11.83'W, 1,000 m station depth) and filled into 300-ml borosilicate glass biochemical oxygen demand (BOD) bottles that each contained an oxygen optode dot (PreSens Precision Sensing GmbH). On filling, DO was increased in a subset of samples by allowing air to be entrained momentarily in the Niskin outflow tubing. Bottles were incubated in an ~6°C water bath in the dark. DO change over 48 h was measured through the glass via detection of phase shift luminescence.

**Boxplot generation.**  $K_m$  DO values were split into two primary groups dependent upon general protein function, either respiratory oxidases or nonrespiratory oxidases.  $K_m$  DO values were also gathered for whole cells, whereby labile and semilabile carbon sources were compared, mirroring the respiratory and nonrespiratory individual enzyme assays. Plots were generated with the grouped  $K_m$  DO values using R v4.0.2 (24) and ggplot2 (25). A linear mixed-effects model was used to control for repeated  $O_2$   $K_m$  measurements from the same organism, with the formula  $\log(K_m) \approx \text{oxidase type } + (1 | \text{organism})$ . The model was fit using maximum likelihood, and the *t* test to confirm significant difference of the coefficient for oxidase type was done using Satterthwaite's method for degrees of freedom (17). Log (natural)-transformed values were used to approximate normality in the data. R code for this analysis can be found at github repo (https://github.com/davised/HBH-2021).

#### SUPPLEMENTAL MATERIAL

Supplemental material is available online only. **FIG S1**, TIF file, 0.5 MB. **FIG S2**, TIF file, 0.5 MB. **FIG S3**, TIF file, 0.4 MB. **TABLE S1**, XLSX file, 0.1 MB.

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