



Diversity and evolution of nitric oxide reduction in bacteria and archaea

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Edited by José J. Moura, Universidade Nova de Lisboa, Caparica, Portugal; received September 27, 2023; accepted April 24, 2024 by Editorial Board Member Marcetta Y. Darensbourg

Nitrous oxide is a potent greenhouse gas whose production is catalyzed by nitric oxide reductase (NOR) members of the heme-copper oxidoreductase (HCO) enzyme superfamily. We identified several previously uncharacterized HCO families, four of which (eNOR, sNOR, gNOR, and nNOR) appear to perform NO reduction. These families have novel active-site structures and several have conserved proton channels, suggesting that they might be able to couple NO reduction to energy conservation. We isolated and biochemically characterized a member of the eNOR family from the bacterium *Rhodothermus marinus* and found that it performs NO reduction. These recently identified NORs exhibited broad phylogenetic and environmental distributions, greatly expanding the diversity of microbes in nature capable of NO reduction. Phylogenetic analyses further demonstrated that NORs evolved multiple times independently from oxygen reductases, supporting the view that complete denitrification evolved after aerobic respiration.

denitrification | heme-copper oxygen reductase | nitric oxide reductase | *Rhodothermus marinus* | aerobic denitrification

The heme-copper oxidoreductase (HCO) superfamily is extremely diverse, with members playing crucial biogeochemical roles in both aerobic (oxygen reductases) and anaerobic [nitric oxide reductases (NORs)] respiration (1–3). While NO reduction can also be performed by fungal NORs (4) and flavodiiron proteins (5), in this paper we focus on NORs from the HCO superfamily. Fungal NO reduction is performed by cytochrome P450 (6), and flavodiiron proteins are primarily used for detoxification of NO. Respiratory denitrification in both Bacteria and Archaea involves NORs from the HCO superfamily. The HCO superfamily consists of three well-characterized oxygen reductase families (A, B, and C) and three NOR families (cNOR, qNOR, and qCu_ANOR) (1–3). The oxygen reductases catalyze the reduction of O₂ to water (O₂ + 4e_{out}⁻ + 4H_{in}⁺ + nH_{in}⁺ → 2H₂O + nH_{out}⁺) and share a conserved reaction mechanism (3, 7), wherein three of the electrons required to reduce O₂ are provided by the active-site metals, heme-Fe and Cu_B, while the fourth electron is derived from a unique redox-active cross-linked histidine-tyrosine cofactor (8) (Fig. 1). The free energy available from this reaction is converted into a transmembrane proton electrochemical gradient, allowing microbes to harness energy from aerobic respiration. The generation of electrochemical gradient occurs via two different mechanisms: charge separation across the membrane and proton pumping (9, 10). Both the protons used for chemistry (i.e., O₂ reduction to water) and those separately pumped protons are taken up from the electrochemically negative side of the membrane (bacterial cytoplasm) by conserved proton-conducting channels that are composed of conserved polar residues and internal water molecules. The different oxygen reductase families exhibit differential proton pumping stoichiometries (*n* = 4 for the A-family, and *n* = 2 for the B and C-families) (10–12) and thus conserve energy differentially depending on their proton channels—though there is some dispute regarding the proton pumping stoichiometry of the C-family, with some studies reporting *n* = 4 (13). The oxygen reductases also vary in their secondary subunits that function as redox relays from electron donors in the electron transport chain (e.g., cytochrome *c*) to the protein complex active site, with the A and B-families utilizing a Cu_A-containing subunit (14–16) and the C-family containing one or more cytochrome *c* subunits (17) (Fig. 1).

NORs catalyze the reduction of NO to nitrous oxide (2NO + 2H_{out}⁺ + 2e_{out}⁻ + nH_{in}⁺ → N₂O + H₂O + nH_{out}⁺). NO reduction requires 2 molecules of NO to form nitrous oxide. With each N atom decreasing in oxidation state by 1, it is only a 2-electron reaction and does not require the cross-linked histidine-tyrosine cofactor for catalysis

Significance

With the advent of culture-independent techniques for studying environmental microbes, our knowledge of their diversity has exploded, uncovering unique organisms, pathways, and proteins carrying out important processes in the biosphere. Novel biochemical reactions are often proposed based on sequence data, but experimental validation is difficult and rare. In this work, we used environmental sequence data to find enzymes that produce the greenhouse gas N₂O from NO and validated our hypothesis with experiments. These new enzymes likely contribute to global N₂O fluxes and expand the breadth of nitrogen cycling. We also demonstrated that these enzymes evolved multiple times from oxygen reductases, indicating that the evolutionary histories of aerobic respiration and denitrification—and more broadly the oxygen and nitrogen cycles—are tightly connected.

The authors declare no competing interest.

This article is a PNAS Direct Submission. J.J.M. is a guest editor invited by the Editorial Board.

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This article contains supporting information online at <https://www.pnas.org/lookup/suppl/doi:10.1073/pnas.2316422121/-/DCSupplemental>.

Published June 20, 2024.

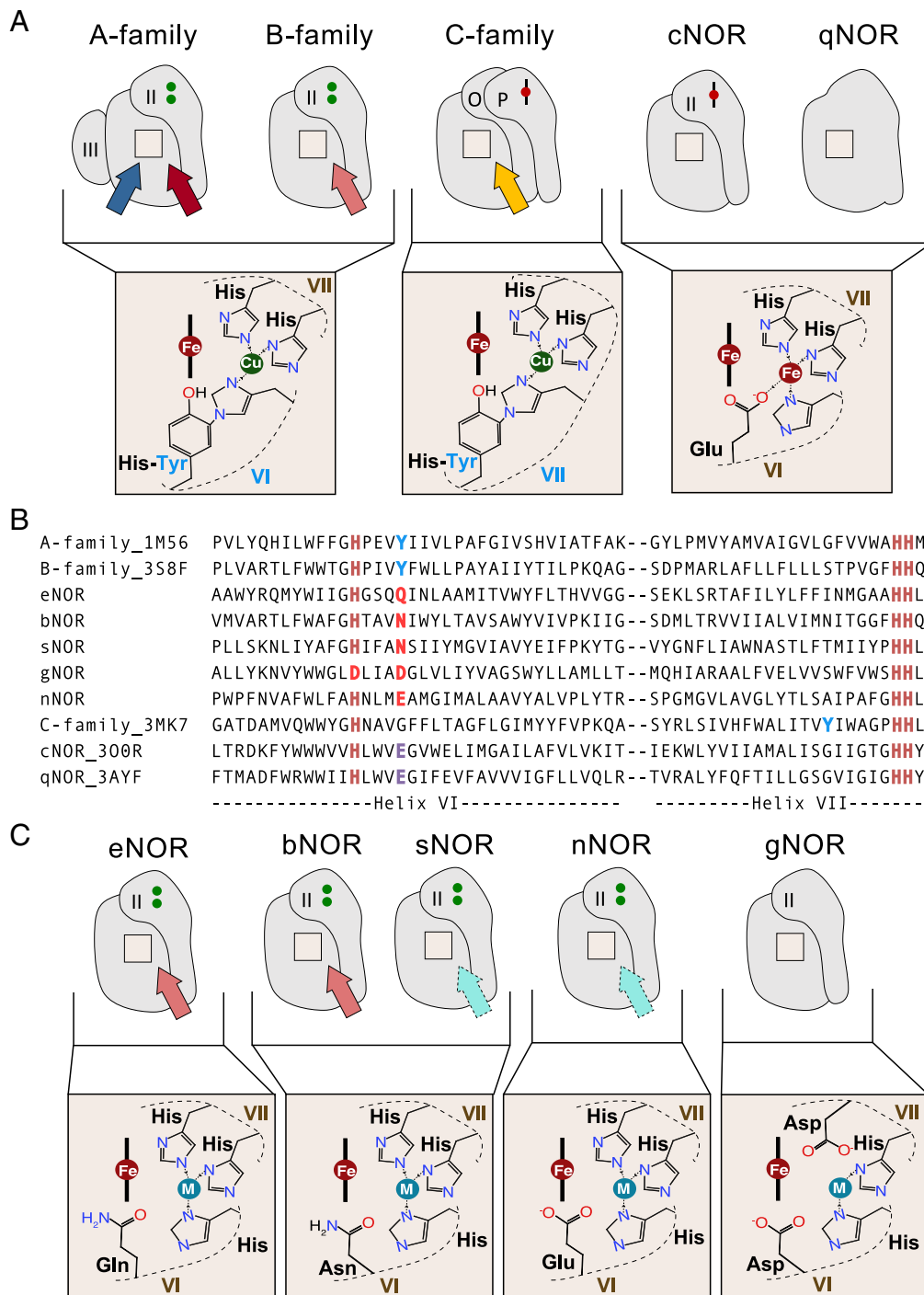


Fig. 1. Comparison of HCO active sites. (A) Active-site and proton channel properties of the five characterized HCO families (A-family, B-family, C-family, cNOR, and qNOR). The oxygen reductases all have an active site composed of a high-spin heme, a redox-active cross-linked tyrosine cofactor, and a copper (Cu_B) ligated by three histidines. The A-family has two conserved proton channels, whereas the B- and C-families only have one. The active sites of the NORs are composed of a high-spin heme and an iron (Fe_B) that is ligated by three histidines and a glutamate. Notably, they are missing the tyrosine cofactor. The cNOR and qNOR are also missing conserved proton channels. (B) Sequence alignment of the active sites of the recently found HCO families that are related to the B-family. (C) Predicted active sites and proton channels for the recently identified HCO families. The eNOR, bNOR, sNOR, and nNOR families contain completely conserved proton channels shown here as arrows. The putative proton channel in the bNOR and eNOR families are highly similar to the K-channel from the B-family oxygen reductase and are colored in red. The K-channel in the B-family is also similar to the K-channel in the A-family oxygen reductase which is colored in dark red. The proton channel in the C-family is different from these channels and is marked in yellow. The putative proton channels in sNOR and nNOR are marked in cyan and differentiated from the other channels with a dashed black outline.

(18)—providing one metric for identifying putative NORs from environmental sequence data. There are currently three biochemically characterized NORs within the HCO superfamily, the cNOR, qNOR, and qCu_A NOR. The cNOR and qNOR families have a four amino acid coordinated Fe_B ion in their active sites,

in contrast to the three amino acid coordinated Cu_B found in the HCO oxygen reductases (18–20). The cNOR and qNOR families are closely related to the C-family oxygen reductases (21). Like the C-family O_2 reductase, cNOR has a secondary cytochrome *c* subunit, while qNOR appears to be the result of

a gene fusion of the primary and secondary subunits forming a single polypeptide that lacks the heme *c* binding motif (22–24). cNOR does not conserve energy, with the enzyme taking both electrons and protons for NO reduction from the periplasmic side (25). Although qNOR is proposed to take up protons from the cytoplasm for NO reduction (20, 26), it does not have conserved residues that could form a proton channel from the cytoplasm, and it is not clear whether this enzyme conserves energy via either charge separation or proton pumping. The qCu_ANOR from *Bacillus azotoformans* (2, 27) is not closely related to cNOR and qNOR and is instead derived from within the B-family O₂ reductases, leading it to be reclassified as the bNOR family (23). bNOR is fundamentally different from cNOR and qNOR, containing a Cu_A cofactor in the secondary subunit and a conserved proton channel for proton uptake from the cytoplasm. bNOR was shown to be electrogenic (27) and thus capable of generating more energy than previously characterized NORs. In earlier work, Hemp and Gennis demonstrated that the HCO superfamily was more diverse than previously thought, working with data from archaeal genomes (1). With recent work in a larger dataset including Bacteria and Archaea, we expanded that diversity to 12 families and demonstrated that quinol oxidation evolved within the HCO superfamily multiple times (23). In this work, we used phylogenomics of both isolates and environmental sequence data to study the diversity and evolution of multiple putative NOR families (eNOR, gNOR, nNOR, and sNOR) within the HCO superfamily and verified the biochemical NO reduction activity of eNOR from the bacterium, *Rhodothermus marinus*. We also identified a new family of putative N₂O reductases. Our findings expanded the number of

denitrification pathways in Bacteria and Archaea, increased the breadth of modern N₂O production and further constrained the evolutionary history of one of the key protein scaffolds involved in aerobic and anaerobic respiration.

Results and Discussion

Expansion of the HCO Superfamily. Phylogenomic analyses of genomic and metagenomic data identified at least six new families belonging to the HCO superfamily (Figs. 1 and 2) that are missing the active-site tyrosine, indicating that they do not catalyze O₂ reduction. Analysis of structural models and sequences (*SI Appendix*) for each of these families showed no evidence for the sequence migration of a conserved tyrosine that could form an active-site cross-linked cofactor, as was observed in the C-family O₂ reductases (8). Furthermore, their active sites exhibited structural features never before seen within the superfamily (Fig. 1). One of these families is closely related to qNOR and has been proposed to be a NO dismutase (NOD) contributing to intracellular O₂ production in “*Candidatus Methyloirabidilis oxyfera*” (28, 29). Another family is closely related to cNOR and might serve as a unique sulfide and acetylene-insensitive nitrous oxide reductase (N₂O red) (30, 31). The remaining four families (eNOR, sNOR, nNOR, and gNOR) are closely related to the B-family of O₂ reductases (Fig. 2) and encode homologs of the Cu_A-containing secondary subunits. This is consistent with the presence of Cu_A-containing subunits in the B-family of oxygen reductases (Fig. 1 and *Dataset S1*). Based on modeled active-site structures and genomic context, we proposed that these novel families perform NO reduction (Fig. 1).

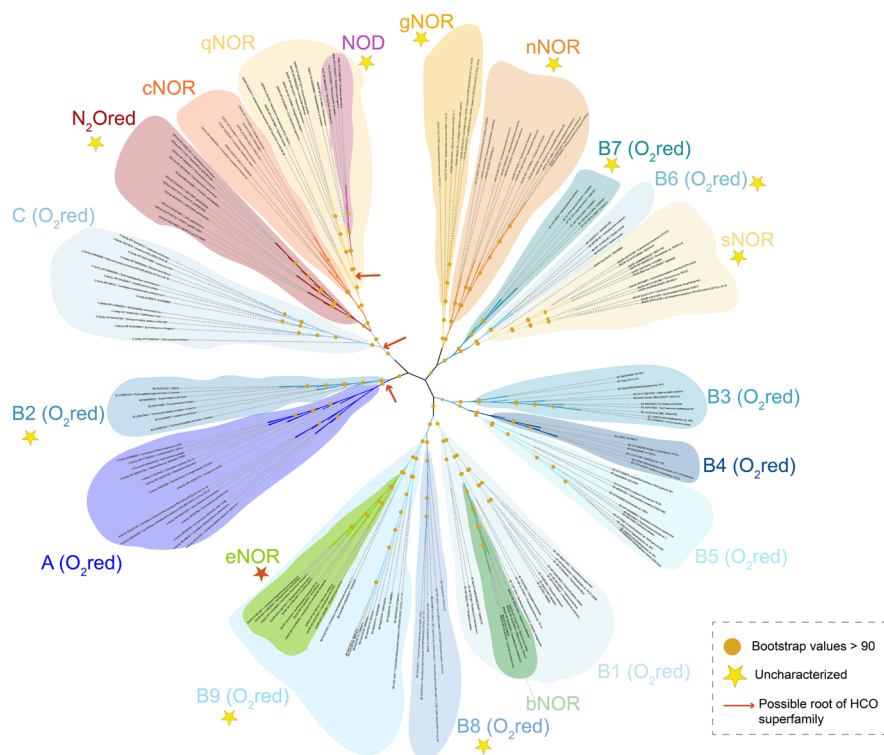


Fig. 2. Evolution of NORs. An unrooted phylogenetic tree of HCO sequences was inferred from a multiple sequence alignment (*Dataset S2*—Multiple sequence alignment) of a representative set of HCO sequences using IQ-Tree as described in *Materials and Methods*. Each of the HCO families is shaded in a different color. Oxygen reductases are in shades of blue, whereas NORs are in shades of yellow, green, and red. The putative N₂O red family is depicted in a light shade of red. The eNOR, bNOR, nNOR, gNOR and sNOR families are derived from oxygen reductase ancestors. Putative root positions (within the A-family, in the qNOR or between the A- and C-families) for the HCO superfamily are noted with a red arrow based on previous literature (32, 33). Uncharacterized enzymes are indicated with a yellow star while the eNOR is indicated with a red star. The Newick tree file is available as *Dataset S3* and a list of the leaf labels is available as *Dataset S4*.

Biochemical Characterization of eNOR. To validate these predictions, we isolated and biochemically characterized a member of the eNOR family from *R. marinus* DSM 4252, a thermophilic member of the Bacteroidetes phylum. *R. marinus* was originally classified as a strict aerobe (34), but its genome encoded a periplasmic nitrate reductase (NapA), two nitrite reductases (NirS and NirK), and a N₂O reductase (NosZ), suggesting that it may also be capable of denitrification (SI Appendix, Fig. S1). Denitrification was not observed under strictly anaerobic conditions, however, under microoxic conditions, we observed that isotopically labeled ¹⁵NO₃⁻ was converted to ³⁰N₂ (SI Appendix, Fig. S2), demonstrating that *R. marinus* DSM 4252 was capable of complete aerobic denitrification (NO₃⁻ → N₂). Blockage of the N₂O red (NosZ) with acetylene led to the accumulation of N₂O (Fig. 3), implying that a NOR was also present in *R. marinus* DSM 4252. No known NORs (cNOR, qNOR, qCu_ANOR/bNOR, or flavodiiron proteins) were found in the genome. However, *R. marinus* DSM 4252 encoded a member of the eNOR family (SI Appendix, Fig. S1).

Isolation and biochemical characterization of the *R. marinus* DSM 4252 eNOR protein verified that it catalyzed NO reduction [at 25 °C, k_{cat} = 0.68 ± 0.21 NO s⁻¹ (n = 4)] (Fig. 3). This turnover number is lower than catalytic turnover rates reported for NORs purified from mesophilic bacteria such as *Pseudomonas stutzeri* [16 NO s⁻¹] (35) or *Neisseria meningitidis* [30 NO s⁻¹] (20) but is higher than activities reported for cNOR purified from other thermophilic microorganisms such as *Thermus thermophilus* [0.09 NO s⁻¹] (25). eNOR was unable to catalyze O₂ reduction using a range of electron donors (SI Appendix, Fig. S3), showing that it

functioned solely as a NOR. UV-Vis spectroscopy and heme characterization via mass spectrometry demonstrated that the *R. marinus* DSM 4252 eNOR contained a unique modified heme *a* that is used in both heme sites (Fig. 3 and SI Appendix, Figs. S3 and S4). Another member of the eNOR family was previously isolated from the aerobic denitrifier *Magnetospirillum magnetotacticum* MS-1 (36, 37); however, its function was never determined. The UV-Vis spectra of the *M. magnetotacticum* eNOR (36) were identical to the *R. marinus* eNOR, implying that the modified heme *a* is a general feature of the family. Mass spectroscopic analysis of the hemes extracted from eNOR revealed that this heme was A₅—a previously isolated heme *a* with a hydroxyethylgeranylgeranyl side chain first identified in the B-family oxygen reductase from *Sulfolobus acidocaldarius* (38). Many eNOR operons contain a CtaA homolog, an O₂-dependent enzyme that converts heme *o* to heme *a* (39). This is consistent with the observation that eNOR required microoxic conditions to be expressed. Organisms performing denitrification with eNOR appear to be obligate aerobic denitrifiers, and future work will establish the extent of their role in environmental aerobic denitrification (40).

Unique Active-Site Features of recently identified NORs. In addition to the experimental evidence that both eNOR and bNOR enzymes are NO reductases, there are several reasons to predict that the other recently identified families also perform NO reduction. The sNOR family has the same active-site structure as the bNOR family, strongly suggesting that it also performs NO reduction. However, the sNOR and bNOR families are not closely

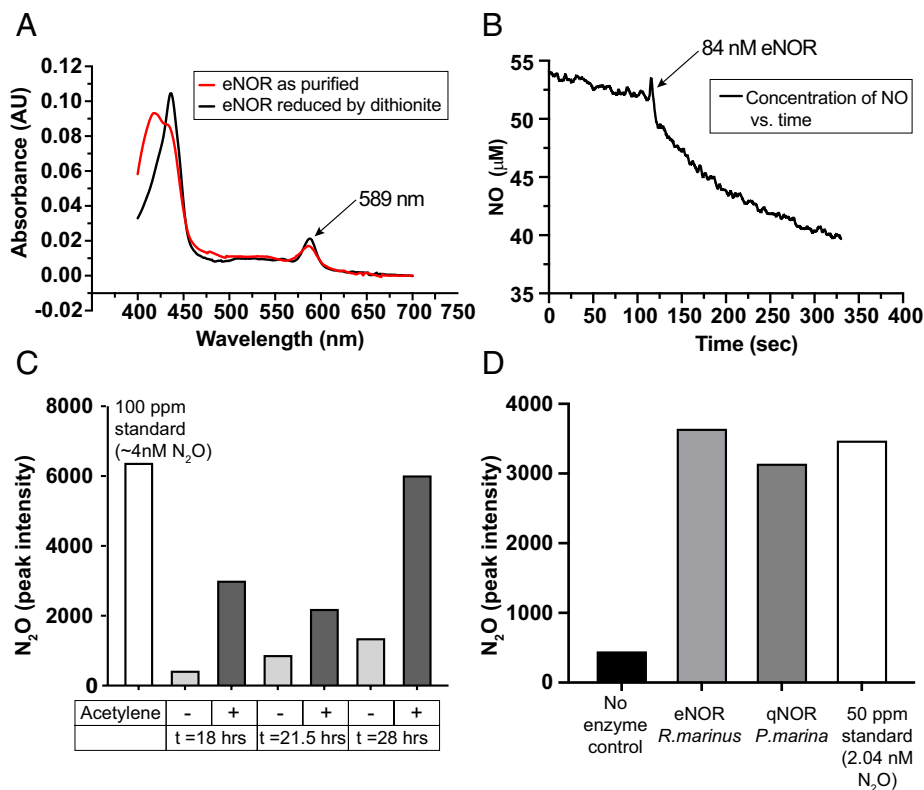


Fig. 3. Biochemical Characterization of the eNOR from *R. marinus*. (A) UV-Visible spectrum of isolated eNOR indicated the presence of an unusual heme *a* signature at 589 nm. (B) NO reductase activity was measured with the use of a Clark electrode in the presence of N, N, N', N'-tetramethyl-p-phenylenediamine (TMPD) and ascorbate as electron donor. (C) N₂O accumulation was observed in an actively growing culture of *R. marinus*. Then, 5 mL was subsampled from a 1 L culture of *R. marinus* and incubated at 42 °C for 30 min in an anaerobic stoppered serum vial with or without acetylene, an inhibitor of the terminal enzyme in denitrification, N₂O red or NosZ. The headspace gas from this incubation was sampled and N₂O concentrations were measured with gas chromatography (GC) followed by analysis by an electron capture detector (ECD). (D) N₂O production by eNOR from *R. marinus* and qNOR from *Persephonella marina* were measured using GC-ECD following incubation of the pure enzyme with TMPD, Ascorbate, and NO under the same conditions as described for subsamples of the culture in C.

related and mark an example of convergent evolution of active-site structures within the HCO superfamily (Figs. 1 and 2). We identified another example of convergent evolution in the nNOR family. Members of this family have the same conserved active-site residues as the cNOR and qNOR families (Fig. 1) but are only very distantly related to them; nNOR is related to the B-family, whereas cNOR and qNOR are related to the C-family. Interestingly, the low-spin heme in nNOR is ligated by a histidine and methionine, which likely raises its redox potential by ~150 mV (41). This feature is similar to a modification found in some eNORs, wherein the low-spin heme is ligated by histidine and lysine. The gNOR is the first example of a HCO family member that has replaced one of the active-site histidines—residues completely conserved in all other families. The gNOR active site, with an aspartate in place of histidine, is likely capable of catalyzing NO reduction, since NO reduction was demonstrated in a bioinorganic mimic of the gNOR active-site (42). The gNOR has a secondary subunit with a cupredoxin fold that is missing the residues required to bind Cu_A, similar to the quinol-oxidizing oxygen reductase cytochrome *bo*₃ from *Escherichia coli*. Conserved residues that could bind quinol have been identified in gNOR, so it may be a quinol-oxidizing NOR similar to qNOR (23).

The biochemically characterized eNOR and bNOR and proposed sNOR and gNOR families within the HCO superfamily have active sites that differ significantly from those found in the well-characterized cNOR and qNOR enzymes (Fig. 1). Importantly, while oxygen reduction chemistry is constrained to require a redox-active tyrosine cofactor, multiple HCO active-site structures appear to be compatible with NO reduction chemistry. Oxygen reductases from the A-family, B-family, and C-family (18, 43, 44) appear to catalyze NO reduction, albeit less efficiently and with a different mechanism than the NORs. This difference in biochemical constraints between NO and oxygen reduction chemistry suggests that the evolutionary transition of oxygen reduction to NO reduction is relatively simple within the HCO superfamily. Another useful chemical constraint that appears to differentiate the catalysis of O₂ reduction and NO reduction is the active site metal: in the currently characterized HCOs, Cu_B is utilized for O₂ reduction chemistry, whereas Fe_B is used for NO reduction chemistry. If this pattern is verified for the other predicted NOR families, it would indicate that the chemistry performed by HCOs is determined to a certain degree by the electronic properties of the active-site metal. It is important to note that the above biochemical constraints for NO and O₂ reduction chemistry are only applicable within the HCO superfamily, since other enzymes such as the flavodiiron proteins (5) or cytochrome *bd* oxygen reductases (45) are capable of NO and O₂ reduction with entirely different active site characteristics. Interestingly, tryptophan/tyrosine chains that are predicted to prevent oxidative damage in redox-active proteins (46) are conserved in both the A-family and B-family O₂ reductases and several of the NO reductases that have evolved from the B-family (bNOR, eNOR, and gNOR) (Dataset S5). In these chains, radicals generated during substrate turnover move by hole hopping through a series of tryptophan and tyrosine residues to the surface of the protein where they are safely quenched by redox buffers within the cell (e.g., glutathione). Despite the difference in catalytic mechanisms between O₂ reductases, 2 out of 3 residues implicated in oxidative protection are found in bNOR, eNOR, and gNOR; they are missing in sNOR and nNOR. Future investigation of the catalytic differences between these NORs will provide insight into the role played by these residues.

Bioenergetics of Denitrification Pathways with Recently Identified NORs. Although both denitrification and aerobic respiration are highly exergonic processes, most of the enzymes in the denitrification pathway are not directly coupled to energy

conservation in cells, making denitrification less energetically efficient than aerobic respiration (47). In the HCO oxygen reductases, conserved proton channels deliver protons from the cytoplasm to the active site for chemistry. These same channels are used to pump protons to the periplasmic side (9, 11, 12, 48). In contrast, previously characterized NORs do not appear to pump protons or conserve as much energy as the oxygen reductases. cNOR does not have conserved proton channels from the cytoplasm, which makes this enzyme incapable of conserving energy (25). The evidence regarding qNOR is currently ambiguous: although qNOR does not have conserved proton channels either, there is some evidence suggesting that it can conserve energy (20, 26).

We found that eNOR family has conserved hydrophilic residues, similar to the electrogenic bNOR, that closely resemble those found in the proton-conducting K-channel within the B-family of oxygen reductases (12, 27) (Dataset S1 and SI Appendix, Fig. S5). The sNOR family also has conserved residues in the K-channel region. However, this putative proton channel is slightly different from those found in the B1-subfamily of O₂ reductases (that contains the *T. thermophilus ba*₃) and the eNOR and bNOR families (Dataset S1). The conserved serine (S309, B1-subfamily *T. thermophilus ba*₃ numbering) found in those families is missing in the sNOR, and instead, this enzyme has a conserved glutamate residue in a structurally different location from other HCO enzymes. Interestingly, the nNOR family, which has the same active site as cNOR and qNOR, also has a conserved proton channel (Dataset S1 and SI Appendix, Fig. S5). This implies that these recently identified NORs may be capable of energy conservation and the lack of a proton channel in the cNOR and qNOR may not be due to energetic constraints universal to NO reduction (49). The conserved proton channels in the eNOR, bNOR, sNOR, and nNOR families would allow them to conserve energy via charge separation and potentially by proton pumping. Detailed characterization of these new NOR families will be helpful for understanding the mechanism of proton pumping in the HCO superfamily—one of the long-standing questions in bioenergetics (50).

Environmental Distribution of NORs. The recently identified HCO NOR families have broad phylogenetic and environmental distributions that substantially expand the scope of denitrification occurring in nature (Table 1 and Datasets S6 and S7). The eNOR, sNOR, gNOR, and nNOR families are all found in both Bacteria and Archaea, whereas the bNOR family was only found in the Bacillales order of Firmicutes (Dataset S6). Phylogenetic analysis of metagenomic data shows that the majority of eNOR, sNOR, gNOR, and nNOR enzymes appear in uncharacterized taxa, hinting at many more organisms capable of NO reduction than previously suspected. Furthermore, the new HCO NOR families were found in a wide variety of environments (Table 1 and Dataset S7). sNORs are broadly distributed in many environments, however, they are rarely found in Archaea. sNORs are found in most ammonia-oxidizing bacteria (AOB) sequenced to date, suggesting that the capability of NO reduction is an important difference in ammonia oxidation pathways between Bacteria and Archaea. Given the importance of AOB, it is likely that sNOR plays a role in this key biogeochemical process in nature (Dataset S7). The gNORs were predominantly found in microbes inhabiting sulfidic environments and as mentioned above may reflect an adaptation that allows for denitrification in the presence of free sulfide, which inhibits other NOR families (Dataset S7). Our analyses revealed that the eNOR family—the new HCO enzyme characterized in detail here—is extremely common in nature and has a broad distribution, similar to the cNOR and qNOR families (Datasets S6 and S7).

Table 1. Distribution of families from the HCO superfamily in various public databases

	NCBI-Genomes	IMG-metagenomes	GTDB-genomes
A-family	20,290	102,368	45,135
B-family	1,238	4,683	2,021
C-family	13,976	23,015	14,981
qNOR	4,388	7,680	3,458
cNOR	2,801	4,824	2,594
eNOR	68	2,709	547
sNOR	95	872	344
bNOR	51	12	200
nNOR	6	289	32
gNOR	10	913	156
NOD	8	539	108
N₂O red	25	597	293

Distribution of NOR families in sequenced genomes vs. environmental datasets. The recently found NOR families account for approximately 2/3 of currently known diversity and 1/2 of the abundance of NORs in nature.

eNORs were found in many strains of *Candidatus Accumulibacter phosphatis*, a critical microbe utilized in wastewater treatment plants for enhanced biological phosphorus removal. The eNOR is highly expressed in transcriptomic datasets from these facilities, demonstrating that *Ca. Accumulibacter phosphatis* is capable of complete denitrification in situ (51). eNOR has also been found in microbes capable of performing autotrophic nitrate reduction coupled to Fe(II) oxidation (NRFO). *Gallionellaceae* KS and related strains express an eNOR under denitrifying conditions, suggesting that an individual organism is capable of complete NRFO (52). eNOR is also common in hypersaline environments (Dataset S7), where it might play a role in the adaptation of denitrification to high salt conditions.

Many organisms encode NORs from multiple families (e.g., *Candidatus Methylophilus oxyfera* has qNOR, sNOR, and gNOR; *B. azotoformans* has qNOR, sNOR, and bNOR). The reasons underlying this apparent redundancy remain unclear, but it suggests that selection for different enzymatic properties (NO affinity, enzyme kinetics, energy conservation, or sensitivity to inhibitors) or the concentration of O₂ may be important factors in determining their distribution and use, similar to what was observed for the HCO oxygen reductase families (10). Analysis of the presence of denitrification genes (nitrite reductases, NORs, and the NosZ-type N₂O reds) within sequenced genomes revealed that many more organisms are capable of complete denitrification than previously realized (Dataset S8). Our current understanding of the diversity of organisms capable of performing denitrification in nature is far from complete but stands to grow with the recognition of these new families of NORs.

The Intertwined Evolutionary History of Aerobic Respiration and Denitrification. Combining our biochemical results and insights with the phylogenetic relationships among different groups in the HCO superfamily, which contains both oxygen reductases and NORs, allowed us to better ordinate the evolutionary histories of aerobic respiration and denitrification. Previous work had demonstrated close evolutionary relationships between the A and B-family oxygen reductases (3), as well as close ancestry between the C-family oxygen reductases, cNOR, and qNOR (21). Yet, the question of which arose first—denitrification or aerobic respiration—has been harder to resolve.

Our analysis of the distribution of oxygen reductases and NORs across the wide diversity of microbial life revealed that oxygen reductases are far more widely distributed; over 30,238 of the 47,894 species in the genome taxonomy database (GTDB) encoded oxygen reductases, whereas NORs were only found in 6,626 species (Datasets S6 and S8). This distribution illustrated the massive impact that oxygen has had on the energetics of the biosphere. The A-family is by far the most widely distributed of the HCO enzymes. It is found in all three domains of life and in more phyla than any other enzyme of the HCO superfamily. This is consistent with the view that the A-family oxygen reductase holds greater antiquity (1, 10, 32, 53), wherein the B-family and C-family O₂ reductases each evolved independently from within the A-family to facilitate specific metabolic and ecological challenges associated with exotic flavors of aerobic biology, like in hyperthermophiles (15) or in chemoautotrophic iron oxidation (54). An evolutionary transition from the A-family to B-family due to selection for higher oxygen affinity, which led to the loss of the D-proton channel to facilitate greater access of oxygen to the active site has been inferred from structural and phylogenetic data with a putative intermediate enzyme suggested in *Nitrosopumilis maritimus* (10). With that in mind, a clear transition from A to the C-family oxygen reductases has not been demonstrated.

The C-family branch of the HCO superfamily consists of the closely related C-family oxygen reductase, N₂O red, cNOR, qNOR, and NOD (Fig. 2) (3, 21, 29, 33, 55). The transition from cNOR to qNOR as the result of a gene fusion of subunits I and II has been reasonably inferred based on the sequence similarity between the N-terminal domain of qNOR and subunit II of cNOR (24). That NOD was derived from qNOR is supported by the high level of sequence similarity between qNOR and NOD, as well as its branching topology within the qNOR clade (Fig. 2). These relative constraints support a simple interpretation of the evolutionary history within this branch—the C-family is the oldest, followed by the evolution of the N₂O red and cNOR. The cNOR then is the ancestor of qNOR, followed by NOD. The sparse distribution of the C-family oxygen reductases, cNOR, and NOD in Archaea supports the hypothesis that these families evolved after the A-family oxygen reductases. While qNOR is widely distributed, it is rarely associated with energetically efficient denitrification and has been proposed to function in nature as a detoxification enzyme (21). Therefore, the presence of qNOR cannot be used as a robust constraint for the antiquity of denitrification. The wide distribution of the A-family, the indications of an evolutionary transition from the A- to B-family, and the relatively sparse distribution of the C-family branch members in Archaea, all suggest that the A-family likely hosts the root of the HCO superfamily (Fig. 2). In the future, the use of different comparative biological approaches—particularly those that might better capture the evolution of paralogs—to root the phylogenetic tree of HCOs could be used to test this idea.

What is clear from our new observations—regardless of the placement of the root of the HCO superfamily—is that NORs have evolved independently multiple times from the B-family and C-family oxygen reductases (Fig. 2). There are key underlying factors that enabled this, both chemical and environmental. It is biochemically straightforward to adapt an oxygen reductase (4 e⁻ chemistry) for NO reduction (2 e⁻ chemistry). B and C-family oxygen reductases can reduce NO at high concentrations in vitro (43, 56). It is thus unsurprising that small evolutionary modifications would lead to a cascade of enzyme descendants each capable of NO reduction at lower concentrations to enable more effective denitrification. It is also clear that in many environments, denitrification

and aerobic respiration often co-occur (57, 58), and that many microorganisms display the respiratory flexibility to shift from aerobic to anaerobic respiration, especially at lower O₂ concentrations (59). This respiratory flexibility is reflected in the fact that denitrification and oxygen respiration share much of the same bioenergetic logic, conserving energy via complex III or alternative complex III (60). Thus, the biochemical promiscuity of O₂ reductases toward NO, the ecological proximity of NO₃⁻ and O₂, and the close similarity between their respiratory pathways help explain why the evolutionary transition of O₂ reduction to NO reduction is both favorable and readily achievable.

Nitrate is derived from biogeochemical processes involving oxygen (61) and consequently opportunities for denitrification prior to Earth's great oxygenation event (GOE) were muted compared to those after the GOE with the rise of nitrate in seawater. Finally, both aerobic respiration and denitrification are constrained by the presence of copper. Copper is an essential bioinorganic component of the active site in the O₂ reductases, nitrite reductase NirK, and for N₂O red (NosZ) and is therefore essential for their biochemical activity. The environmental abundance of copper increased significantly after the GOE (62–64), suggesting that both of these metabolic pathways evolved and expanded after the accumulation of O₂ in Earth's surface environments.

Materials and Methods

This is an abbreviated version of the materials and methods used in this work. A detailed version is available as part of *SI Appendix*.

Purification of eNOR from *R. marinus* Grown under Denitrification Conditions. *R. marinus* DSMZ 4252 was grown in DSM Medium 630 with 30 mM NO₃⁻ added and shaken at 75 rpm to induce denitrification. The microoxic conditions that result from slow shaking were essential for denitrification in *R. marinus*. We used labeled nitrate (¹⁵NO₃⁻) to verify that *R. marinus* DSM 4252 was capable of complete denitrification (NO₃⁻ to N₂). The experiments detailed below established that eNOR was expressed under these conditions and functions as a NOR.

Cultures of *R. marinus* were grown in 1L of this medium in 24 × 2L Erlenmeyer flasks for 36 h or to stationary phase, to generate sufficient biomass for protein purification. The cell pellet recovered from this culture was subject to lysis as described in *SI Appendix, Materials and Methods* and the membrane fraction was recovered by ultracentrifugation. eNOR was purified in a protocol similar to that described for *caa*₃ from *R. marinus* (65). Purification of eNOR was improved when the membranes were first solubilized in 1% 3-[(3-Cholamidopropyl)dimethylammonio]-1-propanesulfonate (CHAPS), apparently recovering peripheral membrane proteins. The membranes not solubilized in this step were then pelleted with ultracentrifugation and solubilized in 1% N-dodecyl-β-D-maltoside (DDM). These solubilized membrane proteins were subject to a protein purification protocol detailed in *SI Appendix, Materials and Methods*. Purified protein was confirmed to be eNOR using electrophoretic analysis and mass spectrometric identification.

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Biochemical Characterization of eNOR. Purified eNOR was verified to perform NO reduction by measuring NO consumption using a Clark Electrode (World Precision Instruments) in a protocol previously described (25) and by measurement of the product, N₂O using GC.

The heme cofactors of eNOR were first analyzed using a pyridine heme-chrome assay (66) and then analyzed by Liquid Chromatography-Mass Spectrometry (LC-MS) after solvent extraction. Further details for both assessment of activity and cofactor identification are available in *SI Appendix, Materials and Methods*.

A detailed description of the phylogenomic analysis of NORs by taxonomy and environment is provided in *SI Appendix, Materials and Methods*.

Data, Materials, and Software Availability. All the protein accession numbers used for generation of trees in this study, as well as associated phylogenetic trees and multiple sequence alignments are included in the [supporting information](#). The Hidden Markov Models (HMMs) used for identification of HCO sequences can be found at <https://github.com/ranjani-m/HCO> (67).

ACKNOWLEDGMENTS. We thank the NIH for funding support (grant# U12AB123456 to PI: R.B.G.). This research was also supported by funding from the Agouron Institute (W.W.F. and J.H.) and by the Community Science Project 507064 (PI: R.H.) under the Joint Genome Institute (<https://ror.org/04xm1d337>), which is a Department of Energy (DOE) Office of Science User Facility. Resources were also used at Office of Biological and Environmental Research of the US Department of Energy Atmospheric System Research Program Interagency Agreement grant DE-AC02-05CH11231 (JGI). Resources were used at Office of Biological and Environmental Research of the United States Department of Energy Atmospheric System Research Program Interagency Agreement grant DE-AC05-76RL01830 Environmental Molecular Sciences Laboratory (EMSL). We thank Sylvia Choi for providing pure *ba*₃ oxygen reductase from *T. thermophilus* to use as a control for oxygen reductase assays and for heme extraction, Paige Sheridan for providing purified qNOR from *Persephonella marina*, Lici Schurig-Briccio for guidance in performing NOR assays with the Clark Electrode, and Peter Yau at the University of Illinois' Mass spectrometric facility for protein identification. We thank Alon Philosofo and Connor Skennerton for valuable discussions on bioinformatics analysis. Finally, we would like to thank our reviewers for their valuable comments that led to the improvement of this manuscript.

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