Coversheet

Oxidoreductases and metal cofactors in the functioning of Earth

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Abstract

Life sustains itself using energy generated by thermodynamic disequilibria, commonly existing as redox disequilibria. Metals are significant players in controlling redox reactions, as they are essential components of the engine that life uses to tap into the thermodynamic disequilibria necessary for metabolism. The number of proteins that evolved to catalyze redox reactions is extraordinary, as is the diversification level of metal cofactors and catalytic domain structures involved. Notwithstanding the importance of the topic, the relationship between metals and the redox reactions they are involved in has been poorly explored. This work reviews the structure and function of different prokaryotic organometallic-protein complexes, highlighting their pivotal role in controlling biogeochemistry. We focus on a specific subset of metal-containing oxidoreductases (EC1 or EC7.1), which are directly involved in biogeochemical cycles, i.e., at least one substrate or product is a small inorganic molecule that is or can be exchanged with the environment. Based on these inclusion criteria, we select and report 59 metalloenzymes, describing the organometallic structure of their active sites, the redox reactions in which they are involved, and their biogeochemical roles.

Keywords: Redox reactions, metalloproteins, organometallic compounds, ligands, metabolism, biogeochemistry

Introduction

Life is fundamentally electric [1]. The thermodynamic disequilibria present in the environment as geochemical gradients are exploited by life to drive its metabolic reactions. The two energy sources used by life, chemical for chemotrophs and light for phototrophs, are always linked or converted to redox disequilibria. Thus life's need for thermodynamic disequilibrium is ultimately a requirement for redox chemistry imbalance. Thermodynamically favorable redox reactions (e.g., glucose oxidation coupled to oxygen respiration) are chopped by life into sub reactions decoupling the

flow of electrons and protons through the electron transport chain and the cell membrane to create a chemiosmotic gradient (Figure 1). This separation effectively converts a scalar (directionless) redox chemical reaction into a vectorial (gradient-forming) process, producing chemical and mechanical work. In a sense, life has solved the need for energy to drive biochemical reactions anticipating Alessandro Volta's battery by nearly four billion years [2].

Biology has evolved proteins that act stepwise to control redox reactions, transferring electrons across redox states between the opening donor and the ultimate acceptor. These proteins, called oxidoreductases (classified under the Enzyme Commission classes 1 and 7.1), are overwhelmingly metal-containing. To precisely and efficiently transfer electrons to and from a wide range of molecules, they finely tune their conjugated metals' midpoint electric potential by controlling the coordination sphere, geometry, and accessibility of the active site [3]. Elements incorporated in the oxidoreductases' catalytic centers include transition metals such as Fe, Mo, W, Zn, Cu, V, Mn, Ni, and Co and non-metals like S and Mg, coordinated either directly or through organometallic structures in the active center [4] (Figure 2). Despite the critical role of metalloproteins in biology, our understanding of the diversity of elements and structures they use is still limited.

The functioning of our planet: a focus on biogeochemistry

Redox couples are recycled on a planetary scale by coupled geological and biological processes happening at diverse spatial and temporal scales. Within biology, redox cycling of key macromolecule-building elements (e.g., carbon, hydrogen, nitrogen, oxygen, and sulfur, also known as CHNOS elements) is primarily carried out by microorganisms inhabiting diverse ecosystems [5]. Most key reactions that control biogeochemistry are carried out by a small set of microbial-encoded proteins containing a redox-sensitive transition metal as core catalytic center [6]. Life can exploit thermodynamic disequilibria present in natural systems using these enzymes whenever the kinetics of the abiotic reactions is slow enough or the activation energy required is big enough for life to outcompete it [7].

Here we discuss the diversity of metal-containing catalytic structures in essential biogeochemical redox proteins and their importance in our planet's functioning. While all enzymes participating in a given metabolic pathway are essential, and all are critical in biogeochemistry regardless of the metabolism itself, this review focused on a small subset of enzymes selected following these criteria:

- 1. They are exclusively oxidoreductases (EC1 or EC7.1), given the dependence of life on redox chemistry;
- 2. They are metal-containing proteins (metalloproteins). Metals often occur in multiple subunits participating in the redox reaction and passing electrons within the enzyme complex. Here we have considered only oxidoreductases in which the metal directly participates in the primary redox reactions;
- 3. They are biogeochemically relevant, i.e., they catalyze a reaction where the

substrate/product is a small inorganic molecule that is (or can be) directly exchanged with the environment. Enzymes interacting with molecules like CO₂, CO, H₂, NO₃⁻, NH₄⁺, SO₄²-, H₂S, and many other compounds fall in this category. Methane (CH₄), considered an organic molecule, is included in this work's list of valid biogeochemical compounds.

These criteria exclude all the enzymes that, while fundamental for the functioning of metabolism, interact with metabolic intermediates and all the key enzymes that do not deal with redox reactions—for example, the key enzyme for the Calvin-Benson-Bassam cycle, Rubisco (EC 4.1.1.39), and many essential genes involved in carbon fixation. In addition, metal-containing oxidoreductase complexes without a metal in the active site are excluded. An example in this category is the flavocytochrome c sulfide dehydrogenase (EC 1.8.5.4), responsible for the reversible conversion of sulfide to elemental sulfur in several sulfide oxidizers and anoxygenic phototrophs. While the heterodimer contains two heme cofactors (making it an iron-containing metalloprotein) and interacts with both H₂S and elemental sulfur, the active site of the catalytic subunit does not contain any metal. It uses instead two flavin-adenine dinucleotide (FAD) cofactors [8]. Under these criteria, the number of biogeochemically-relevant metalcontaining oxidoreductases involved in key steps of the CHNOS cycles is reduced to 59 (Table 1 and Supplementary Figure S1). These effectively control the biogeochemistry at the interface between the geosphere and biosphere and are more likely to be influenced by the environmental availability of their metal cofactor [9].

Carbon Cycle

At the most fundamental level, life is carbon-based. Hence, life plays a vital role in mediating the biogeochemical cycles of carbon on Earth's surface [10,11]. Inorganic carbon is a building block for assembling complex C molecules [10] through autotrophybased metabolic strategies. At the same time, CH₄ can be oxidized to yield energy for cellular growth and maintenance, ultimately releasing CO₂ [12,13]. While oxidoreductases are involved in the pathways responsible for the uptake and release of inorganic carbon compounds, few are metal-containing oxidoreductases. The KEGG database lists 66 enzymatic classes involved in carbon fixation pathways, of which 21 are classified as oxidoreductases. However, only three (5 % of all enzymatic classes involved in carbon fixation pathways) fall within our definition (Table 1). Other carbon-related metabolisms important at the biogeochemical level are methanogenesis, aerobic and anaerobic methane oxidation, and carbon monoxide utilization. KEGG lists 33 enzymatic classes involved in these pathways; 15 are oxidoreductases, and only three fall within our definition (9 %; Table 1).

Most oxidoreductases in the seven known carbon fixation pathways do not contain free metals in their active center. Instead, they interact with cofactors such as ferredoxin, FAD, and NAD (see supplementary online materials). A few exceptions exist. *Escherichia coli* formate dehydrogenase (*fdhF* [1FDO], Figure 2A and 3D) and *Methanothermobacter wolfeii* formylmethanofuran dehydrogenase (*fwdA* [5T5I], Figure

3E and Supplementary Figure S2) are homologous enzymes involved in CO₂ assimilation in the Wood-Ljungdahl (WL) pathway of carbon fixation. Their active site enzymes contain a molybdenum ion bound to two molybdopterin guanine dinucleotide (PDB accession MGD) and a selenocysteine or a tungsten ion bound to two MGD and a cysteine, respectively.

The Carbon monoxide dehydrogenase (coxL [1ZXI], Figure 2C) is also a metal-containing oxidoreductase of biogeochemical interest. It uses either a Cu-S-Mo cluster for the aerobic variant or a Cu-Ni or Ni-only cofactor for the anaerobic variant of the enzyme (codh [1MGJ], Figure 2D) [14,15]. The Cu-S-Mo cluster associated with the aerobic CODH interacts with a single molybdopterin cytosine dinucleotide (MCN) rather than two (as in the FDH). The two oxygens of the cluster replace the dithiolate group of the second MCN in defining the metal geometry (here constrained to be distorted pyramidal) [16]. For anaerobic CODH, Ni is integrated within a Fe-[NiFe₃S₄] cluster rather than being bridged to a cubane [Fe₄S₄] [17].

Within the methane cycle, two additional enzymes match our definition of biogeochemically relevant metal oxidoreductase: the membrane-bound particulate Methane monooxygenase (pMMO, pmoB1/B2 [3RGB]), which uses Cu as a catalytic cofactor, and the cytoplasmic, copper starvation-induced soluble Methane monooxygenase (sMMO, mmoX [1MHY], Figure 2B), which in turn uses Fe-Fe. Albeit both catalyze methane oxidation, they are entirely different from a structural standpoint.

Despite the low number of biogeochemical metal-containing oxidoreductases present, the carbon cycle is very diverse in its metal requirement, with Fe, Mo, W, Cu, and Ni involved in key steps of the cycle (Figure 4A).

Nitrogen Cycle

Nitrogen is abundant in Earth's atmosphere in the form of dinitrogen (N_2) gas and it is present in significant quantities also in the mantle. This element is vital in building nucleic acids, proteins, and enzymes. At the enzymatic level, nitrogen can be transformed between different compounds with different redox states, *e.g.*, NH_4^+ , NO, NO_2 , N_2O , NO_3^- , NO_2^- , hydroxylamine, and amino acids, moving from +5 in NO_3^- to -3 in ammonia [18] (Figure 4C). The 22 different EC numbers present in the energetic nitrogen cycle on KEGG are oxidoreductases, and 16 (73 %) of them are metalloenzymes relevant in our context. The most frequent metal is Fe, followed by Mo, Cu, and V (Table 1).

Nitrification is governed by Fe, except for the Cu-containing cofactor known as cupredoxin [19]. The most utilized metal cofactor for denitrification involves Mo, followed by Fe and Cu, with different geometry inside the enzymatic cofactors. The dissimilatory nitrite reduction and the assimilatory nitrate/nitrite reduction are controlled by Fe, except

for the nitrate reductases, in which the catalytic metal is Mo [20]. The anaerobic oxidation of ammonia is carried out by Fe-containing enzymes (Figure 4C). The nitrogen fixation pathway is carried out by the Nitrogenase enzyme (Figure 2F, 3G, 3H), which exists in three different isoforms partnering with a unique cofactor: FeMoco (*nifD* [3U7Q], Figure 3H), FeVco (*vnfD* [5N6Y], Figure 2I, 3G and 3I and Supplementary Figure S3), or FeFeco [21]. Fe is the leading metal in every step of the nitrogen cycle associated with the more reduced nitrogen molecules. In contrast, Mo and Cu are associated with the most oxidized forms of nitrogen or enzymes directly involving molecular oxygen (e.g., Ammonia monooxygenase) (Figure 4C).

Sulfur cycle

Sulfur is the 10^{th} most abundant element on Earth. Despite only a small fraction of it being bound to biomass, it is essential in all organisms. Life plays key roles in the global sulfur cycle through its assimilation into methionine and cysteine, enzyme cofactors (*i.e.*, iron-sulfur clusters), and through its use as electron donor/acceptor in dissimilatory energy-yielding reactions (mainly restricted to prokaryotes)[22]. The sulfur cycle involves reactions between eight valence states, from the most reduced H_2S (-2) to the most oxidized SO_4^{2-} (+6, Figure 4D). Among the 19 enzymes involved in the cycle, 14 are oxidoreductases, and 11 of these fall within our definition (79 % of all sulfur cycle enzymatic classes), relying on the presence of either Mo or Fe for their catalytic activity and having a direct biogeochemical impact through their function (Table 1).

The aerobic sulfur disproportionation, assimilatory sulfate reduction, and sulfate reduction pathways are catalyzed by Fe-containing enzymes (Table 1 and Figure 4D). DMSO reduction, sulfite oxidation, sulfur disproportionation, and thiosulfate reduction pathways are catalyzed by Mo-containing enzymes. Interestingly, both Dimethyl sulfide:cytochrome c2 reductase (DMSO reduction) and Sulfite dehydrogenase (sulfite oxidation) contain a molybdenum-bis (molybdopterin guanine dinucleotide) geometry. Additionally, some pathways of the sulfur cycle involve steps catalyzed by enzymes containing both Fe and Mo. For instance, sulfur reduction is catalyzed by Sulfhydrogenase (Fe-containing, part of a NiFe hydrogenase multienzyme complex) and sulfur reductase (Mo-containing). The same pattern is observed in the Thiosulfate oxidation pathway, with Sulfane dehydrogenase (Mo-containing) and Thiosulfate dehydrogenase (Fe-containing). This difference could be due to the different substrates these enzymes interact with, as Sulfhydrogenase interacts with hydrogen and sulfur reductase with oxygen, suggesting that the redox potential of these substrates could provide selective pressures for specific metal utilization.

Oxygen cycle

The great availability of oxygen in Earth's extant atmosphere results from the

emergence of oxygenic photosynthesis, which, coupled with a complex series of geological feedbacks, was responsible for the Great Oxidation Event (GOE, 2.5-2.3 billion years ago) [23,24]. Photosystem II (PS-II) is the main protein complex involved in oxygenic photosynthesis. The oxygen-evolving complex (OEC) represents the PS-II catalytic site where the manganese-dependent photo-oxidation of water occurs, with subsequent release of oxygen (Figure 4B)[25,26]. The presence of Mn ions in the OEC catalytic center is supposedly a consequence of its abundance in the Archean oceans and its hypothetical former use as a phototrophic electron donor [27–29]. Furthermore, enhanced oxygen availability prompted the evolution of both O_2 -respiratory and -detoxifying mechanisms [30].

Oxygen high electronegativity makes it a suitable terminal acceptor in oxidative phosphorylation, the hallmark of aerobic respiration, where oxygen reduction to water is carried out by cytochrome oxidases (classified as translocases, EC 7.1) (Figure 4B) [31–33]. These enzymes generally require Cu as a metal cofactor, directly located in the catalytic center and coordinated by a heme group. Cytochrome bd ubiquinol oxidases make an exception, as their only metal cofactor is Fe, complexed in a heme group (cydA, appC [6RKO, 7OY2] Figure 2L; Table 1) [26,33–37]. In $E.\ coli$, Cu-containing cytochrome bo is maximally synthesized under high oxygen availability. Conversely, iron-containing cytochromes bd predominate in microaerophilic conditions [38], showing a very low K_m for oxygen and a less efficient proton motive force [39,40]. This evidence suggests that the nature of the metal cofactor is crucial in determining cytochromes' performance and their affinity for oxygen.

On the other hand, aerobic respiration induces the formation of reactive species of oxygen (ROS) (Figure 4B), which are responsible for cell damage [41]. Superoxide radical anions can be detoxified by three Superoxide dismutase (SOD) families, which differ in the catalytic metal (e.g., Fe/Mn, Cu, and Ni) (sodA, sodC, sodN [1Y67/3KKY, 1BZ0, 1Q0G] Figure 3J, 3K, 3L; Table 1)[42–44]. Among them, the Fe/Mn family is highly flexible in cofactor utilization, representing a clear example of a cambialistic enzyme (sodA/B [1y67/3kky])[43,45]. Hydrogen peroxide produced by SODs is rapidly detoxified by the Catalase-peroxidase, whose metal cofactor is Fe in a heme conformation (Figure 4B) [46].

Hydrogen cycle

Hydrogen is a key reduced compound in the redox balance of the planet. It is produced by several abiotic processes, including water photolysis/radiolysis, hydrothermal reactions, magmatic degassing, and hydration of iron-rich ultramafic rocks [47]. Hydrogen is also produced and consumed by microorganisms and used as an electron donor — it is one of the main energetic currencies exchanged within microbial communities [48]. Microorganisms can interact with molecular hydrogen through a

group of diverse enzymes called hydrogenases, which catalyze the conversion of molecular hydrogen to protons and electrons and H₂ regeneration through the reverse reaction [49,50](Figure 4B). Their specialized metallic centers coordinate dihydrogen, polarizing the molecule to induce its heterolytic splitting into a proton and a hydride ion.

There are three main groups of hydrogenases, NiFe containing hydrogenases (hydA [6N59] Figure 2E, 3A), FeFe hydrogenases (hydB [5XLF] Figure 2G, 3B), and Fe-only hydrogenases (hmD [6HAV] Figure 3C). [NiFe]-hydrogenases are found in many bacteria and archaea, [FeFe]-hydrogenases in bacteria and some eukaryotes, and [Fe]hydrogenases only in archaea [48]. Of the 30+ classes of hydrogenases known, we report here an example of H₂-consuming and H₂-producing hydrogenases from the main [NiFe], [FeFe], and [Fe] hydrogenases (Table 1; Figure 4). [NiFe]-hydrogenases are mainly involved in H₂ oxidation but have many other functions such as H₂ evolution, sensing, CO respiration, electron bifurcation, and cofactors reduction [48,51,52]. In selenium-rich conditions, some bacteria, like Desulfovibrio vulgaris, downregulate the production of [NiFe]-hydrogenases in favor of protein variants with selenocysteine as one of Ni ligands, displaying lower inhibition by molecular hydrogen and lower O2 sensitivity [53]. [FeFe]-hydrogenases also serve diverse physiological functions such as H₂ uptake, sensing, evolution, electron bifurcation, and CO₂ fixation [54]. [Fe]hydrogenases, the least characterized type of hydrogenases, have only been detected in methanogenic archaea where they are crucial for the archaeal methanogenic pathway [55]

Cofactor cambialism at the core of biogeochemistry

Diverse factors constrain the choice of metals at the core of metabolisms: the environmental availability of the element of interest, its suitability for the specific redox reaction to be catalyzed, and the ability to control its binding to the target enzyme. Theoretically, metal-binding affinities of natural proteins are defined by the ligand field stabilization energy of metal ions and follow the Irving-Williams (IW) series (Mn²⁺ < Fe²⁺ $< Co^{2+} < Ni^{2+} < Cu^{2+} > Zn^{2+}$, [56]. In practice, cells tend to maintain the availability of metal ions inverse to the IW series [57] so that binding is more regulated by ion availability in the immediate environment of the metalloprotein metallochaperone), with very high spatial granularity—cells are not ideal solutions. Moreover, ions' concentration can change to the point that different metals can be acquired when folding in different places [58]. The environmental concentration is modulated by metal transport and the metal bioavailability in the outer environment computational studies have shown that if one removes metallochaperones, metal specificity becomes strongly correlated with metal abundance in the environment. The situation is further complicated because cations have overlapping characteristics that impede absolute specificity. For some metals, similarity in binding affinity and preference over coordination environments is associated with different redox chemistry (e.g., Mn²⁺/Mg²⁺/Fe²⁺ and Mo/W). In this context, excluding the wrong metals from proteins may be more challenging than acquiring the right ones [59], and having a metallochaperone or an additional metal center (as in binuclear Mg²⁺, [60] could reduce mismetallation.

At the environmental level, the (bio)availability of metals might control to a first order its utilization by biology [9]. However, the metal used also depends on the enzyme's evolutionary trajectory. The idea that ancient, promiscuous oxidoreductases were constrained to use bioavailable metals to catalyze redox reactions and that a contingency shaped evolution of more "focused" metalloenzymes differing in metal utilization is supported by comparison of proteomes across life domains [61]. It is worth remembering that many of the transition metals detailed in this review were readily available in ancient times due to the low oxygen/high sulfur environment, except for Cu, Mo, and Zn (that are sparingly soluble in those conditions) and that the Paleozoic oxidation event (GOE) reverted this trend [62]. At the same time, it is essential to consider that selection "locked in" some crucial enzymes (e.g., Fe-S proteins, [63] relying on once-plentiful metal species (after the GOE, iron is primarily available in the low-solubility ferric form).

Currently, biogeochemical cycles are dominated by Fe as a key catalytic metal (Figure 4; Supplementary Figure S4). However, its ability to interact with oxidized substrates is often limited to low-concentration conditions requiring high affinity (like in the Fecontaining cytochromes used under microaerophilic conditions). As a result, cells rely on Cu and Mo to attain the higher redox potential needed to interact with powerful oxidants—such as oxygen in full aerobic conditions, nitrate, and other oxidized nitrogen species. Determining the *in vivo* utilization of metal ions by biomolecules is challenging since complicated metal centers can remain poorly defined even after structure determination due to, e.g., experimental procedure-related substitutions.

Conclusion

The requirement of life for metals as cofactors in key biogeochemical reactions attests to the vital role that metals play in the functioning of Earth and the intricate relationship between the biosphere and the geosphere. Complex stellar processes, protoplanetary disk accretion, and planetary differentiation [64], changing redox conditions during planetary evolution [65,66], plate tectonics, supercontinent assembly [67], and changes in dominant volcanism [68], all contributed to the complex interactions between metal bioavailability and the evolution of biogeochemistry. Nevertheless, our understanding of the role of metals in controlling microbial metabolism and biogeochemistry is still in its infancy. Critical questions about selective pressures imposed by redox potentials of substrates and reaction products in selecting specific metals and the effect of metal environmental availability remain open. In addition, we still need a complete catalog of the elements life uses in protein structure; the diversity of organometallic structures has been poorly examined in environmental—and mostly unculturable [69]— microbes, making it problematic to investigate protein structures and cofactors using traditional approaches. For example, recent work has demonstrated that lanthanides, a group of

elements previously believed to be inert for life, are used by an enzyme catalyzing a key step in the aerobic respiration of methane [70,71]. Increasing our knowledge of organometallic cofactors from uncultured microbial groups can revolutionize our understanding of how redox chemistry mediates the interaction between life and our planet, offering promising possibilities in the green chemistry industry and opening our transition to a more sustainable economy [72–74].

Summary

- There is a universal need for redox chemistry by life to use thermodynamic disequilibrium.
- Biogeochemical cycles, and therefore the functioning of our planet, are controlled by a small number of biogeochemically relevant redox proteins, most of which use metal cofactors. The metal used is tuned together with the protein structure to the midpoint potential of the reaction catalyzed.
- Metal choice is dictated on the first order by availability and active transport and refined by protein structure. Besides, evolution contributed to "frozen accidents" that irreversibly paired some metals to specific cycles.
- The correlation between the diversity of metal cofactors and the biogeochemical redox reactions in which they are involved is still unclarified.
- Despite their importance for our planet's functioning, we have limited information regarding the organometallic structure found in oxidoreductases of uncultured lineages of microorganisms.

Figures and tables

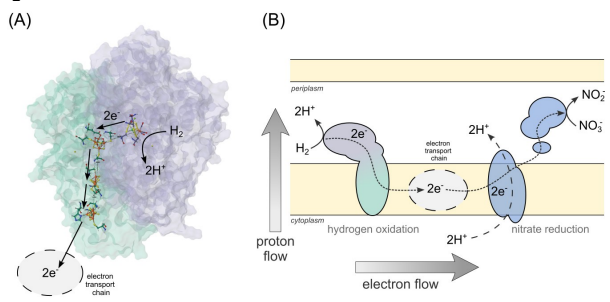


Figura 1. (A) The heterodimeric structure of the periplasmic [NiFe]-hydrogenase from *Desulfovibrio vulgaris* (hyd, 5XLF) showing the [NiFe] catalytic center and the iron-sulfur clusters responsible for the electron transport. (B) Cellular model of the coupling between the periplasmic membrane-bound [NiFe]-hydrogenase Hyd and the periplasmic molybdopterin-containing nitrate reductase Nap in *Thermovibrio ammonificans* showing the decoupling between electrons and protons across the membrane (adapted from [75]).

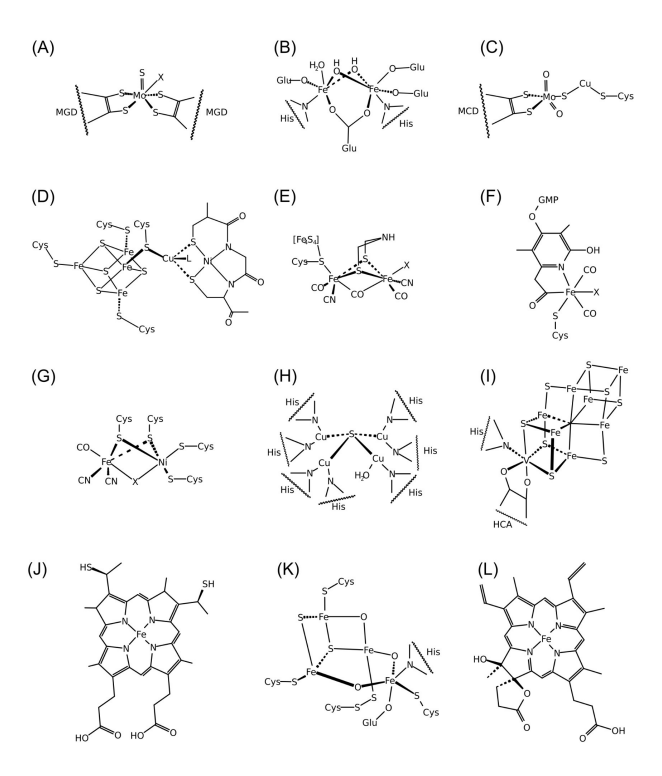


Figure 2. Chemical structures of biogeochemically relevant metal-containing cofactors from prokaryotic oxidoreductases: (A) the Mo-containing catalytic site of Formate dehydrogenase (*fdhF*, 1FDO; shown in Figure 3D); (B) the FeO cluster of the soluble methane monooxygenase hydroxylase (*mmoX*,1MHY); (C) the Mo-Cu-containing cluster in the active site of CO dehydrogenase (*coxL*, 1ZXI); (D) the Ni-Fe-Cu center of

the anaerobic carbon monoxide dehydrogenase (*codh*, 1MJG); (E) the catalytic centers of the [FeFe]-hydrogenase (*hydA*, 6N59; shown in Figure 3A), [Fe]-hydrogenase (F) (*hmd*, 6HAV; shown in Figure 3C) and [NiFe]-hydrogenase (G) (*hydB*, 5XLF; shown in Figure 3B); (H) the [Cu₄S] cluster of the nitrous oxide reductase (*nosZ*, 1FWX); (I) the FeVco cofactor of the V-containing nitrogenase (*vnfD*, 5N6Y; shown in Figure 3G); (J) the Heme C contained in several oxidoreductases (*hzsA*, *hdh*, *nrfA*, *nirS*, *hao*, *tsdA* [5C2V, 6HIF, 2J7A, 6TSI, 1FGJ, 4V2K]); (K) the hybrid cluster from the the Hybrid Cluster Protein from Desulfovibrio vulgaris (*hcp*, 1E1D); (L) the Cis-heme hydroxychlorin gamma-spirolactone (*cydA* and *appC*, 6RKO and 7OY2).

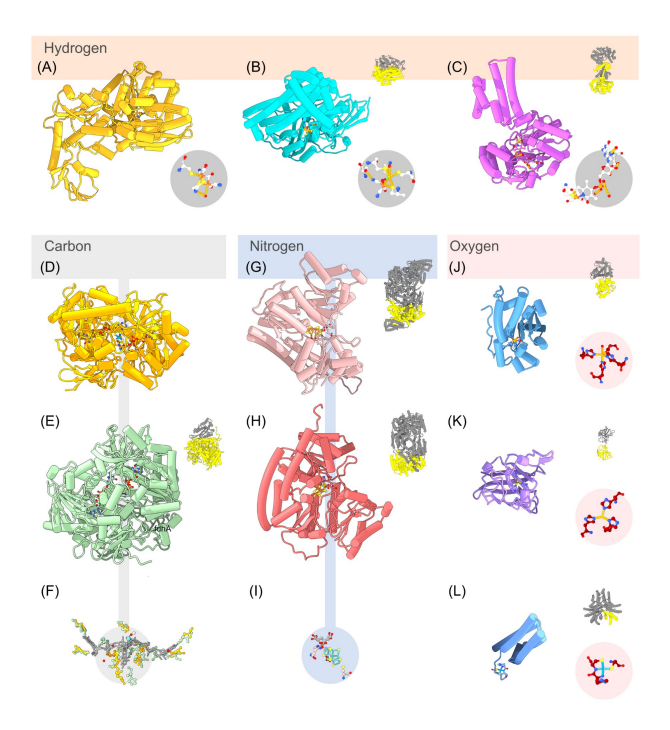


Figure 3. Structures of metal-containing subunits associated with prokaryotic metalloproteins relevant for biogeochemical cycles. The structures reported are relative to the hydrogen, carbon, nitrogen, and oxygen cycle and have been selected because they have isoforms containing different metals in their catalytic site or show some degree of cofactor cambialism. For each pdb, the coordinated metal/organometallic complex is displayed within a circle, together with a miniature of the assembly highlighting in yellow the catalytic subunit. **Hydrogen cycle-related structures**: (A)

FeFe-hydrogenase small subunit (hydA) from Clostridium pasteurianum (6N59); (B) NiFe-hydrogenase large subunit (hydB) from C. pasteurianum (5XLF; hydA-hydB heterodimeric assembly [AB]); (C) activated Fe-hydrogenase (hmd) Methanococcus aeolicus Nankai-3 (6HAV). Carbon cycle-related structures: (D) formate dehydrogenase H α -chain (fdhF) from Escherichia coli (1FDO); (E) Tungsten formylmethanofuran dehydrogenase chain α (fwdA) from Methanothermobacter wolfeii (5t5i, dodecameric assembly 2x[ABCDFG]); (F) the Mo/W-bis(molybdopterin guanine dinucleotide) cofactor common to both enzymes. Nitrogen cycle-related structures: (G) V containing nitrogenase α-chain (vnfD) from Azotobacter vinelandii (5N6Y; hexameric assembly 2x[ABC]); (H) Mo containing nitrogenase α -chain (*nifD*) from A. vinelandii (53U7Q; tetrameric assembly 2x[AC]); (I) the FeMoco/FeVco cofactor. Oxygen cycle-related structures: (J) Superoxide dismutase (sodA) from Deinococcus radiodurans (1Y67, 3KKY; homodimeric assembly); (K) Superoxide dismutase (sodC) from Photobacterium leiognathi (1BZ0; homodimeric assembly); (L) Ni-containing superoxide dismutase (sodN) from Streptomyces selenosis (1Q0G; hexameric assembly).

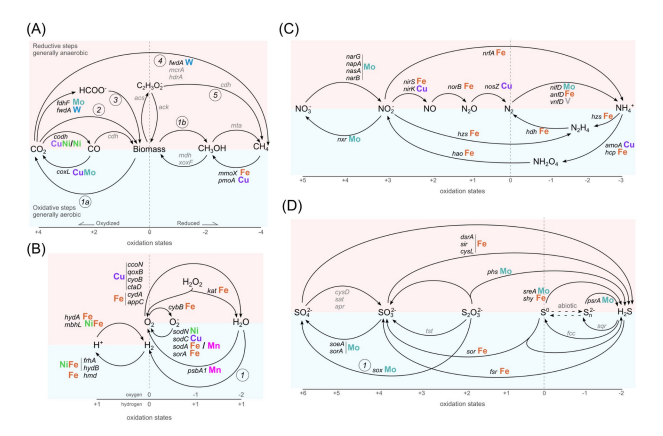


Figure 4. Biogeochemical cycles of the major CHNOS elements. The reductive side of each cycle is reported on the upper side of each cycle and highlighted in light red, while the oxidative side is on the bottom and highlighted in light blue. Molecules in each element are ordered left to right based on the oxidation state starting with the most oxidized form. Key enzymes for each step of each cycle are reported: in black the names of enzymes that do not meet our criteria of biogeochemically relevant metal containing oxidoreductases, while in light gray other enzymes. Enzyme names are based on the KEGG names and reported in Supplementary Table 1. The catalytic metal is reported for each biogeochemical oxidoreductase colored according to Jmol color scheme. The numbers in a circle represent complex pathways/processes. (A) Carbon cycle. 1a and 1b, heterotrophy and fermentation: no enzyme meets our criteria in these pathways: 2, carbon fixation; 3, formate assimilation; 4, methanogenesis pathway: the reported enzyme catalyze key steps in this multi-enzyme pathway; 5, acetoclastic methanogenesis. (B) Hydrogen and Oxygen cycle: 1, abiotic and biotic hydrogen formation; note that the oxidation scale for oxygen and hydrogen are distinct, and hydrogen is reported with the same oxidation state on both sides. (C) Nitrogen Cycle. (D) Sulfur cycle: 1, sulfur/thiosulfate oxidation is accomplished by a complex group of enzymes (sox) of which soxCD meets our criteria (Table 1).

Table 1. List of biogeochemically relevant metals containing oxidoreductases controlling the major CHNOS cycles. PDB and UniProt accessions are reported for each gene, together with Ligand ID; AlphafoldDB codes are reported when a crystallographic structure is unavailable. For enzymes known to be cambialistic (*i.e.*, accept alternative metals in the active site) in experimental setups, the alternative metals are reported separated by a "/". An extended version of the table reporting all the other cofactors present in the catalytic subunit of the enzyme is available as supplementary online material and published on a permanent archive with doi: 10.5281/zenodo.7934782. *a* - the structure, organometallic structure and Uniprot accession number for the Fe nitrogenase is on hold at the time of writing and awaiting release.

Cycle	Pathway	Step	Uniprot	Protein Name	Gene	EC	ко	PFAM	Metal in the active center	Organometallic structure [PDB ligand accession]	PDB	Representative organism
	Aerobic methane oxidation	Oxygenation of	G1UBD1	Particulate methane monooxygenase alpha subunit	pmoB1 , pmoB2	1.14.18.3	K10944	PF04744	Cu	Cu (II)-Cu (II) [CUA]	3RGB	Methylococcus capsulatus
		methane to methanol	P27354	(soluble) Methane monooxygenase component A beta chain	mmoX	1.14.13.2 5	K16157	PF02332	Fe	2 Fe (III) [FE]	1MHY	Methylosinus trichosporium
	Carbon fixation	CO ₂ reduction to formate	P07658	Formate dehydrogenase H	fdhF	1.17.98.4	K22015	PF04879	Мо	Mo (VI) [MO] molybdenum- bis(molybdopterin guanine dinucleotide) [MGD]	1FDO	Escherichia coli
Carbon		CO ₂ reduction to formylmethanofuran	O74030	Tungsten formylmethanofuran dehydrogenase subunit fwdA	fwdA	1.2.7.12	K00200	PF00384	W	W (VI) [W], molybdenum- bis(molybdopterin guanine dinucleotide) [MGD]	5T5I	Methanothermobacter wolfeii
		Oxidation of CO to CO ₂	P19920	(aerobic) Carbon monoxide dehydrogenase medium chain	coxL	1.2.5.3	K03520	PF02738 , PF20256	CuMo	Cu (I) -S- Mo (VI) (=O) OH Cluster [CUM]	1ZXI	Oligotropha carboxidovorans
		Reduction of CO ₂ to CO	P27988	(anaerobic) Carbon monoxide dehydrogenase/acetyl-CoA synthase subunit alpha	codh	1.2.7.4	K00192	PF03063	CuNi / Ni	Fe(4)-Ni(1)-S(4) Cluster [XCC] + Cu Ion [CU1]	1MJG	Moorella thermoacetica
Hydrogen		F420 Reduction	D9PYF9	(NiFe) F420-reducing hydrogenase, subunit alpha	frhA	1.12.98.1	K00440	PF00374	NiFe	Formyl [bis(hydrocyanato- 1kappaC)] Fe-Ni [NFU]	4OMF	Methanthermobacter marburgensis str. Marburg
	Hydrogen oxidation	H₂-respiration	Q58194	5,10-methenyl tetrahydromethanopterin hydrogenase	hmd	1.12.98.2	K13942	PF03201	Fe	Fe (II) [FE2] coordinated by 5'-O- [(S)-hydroxy {[2- hydroxy-3,5-dimethyl- 6-(2-oxoethyl) pyridin- 4-yl]oxy} phosphoryl] guanosine [[2C]	3F47	Methancaldococcus jannaschii
			P21852	Periplasmic [NiFe] hydrogenase large subunit	hydB	1.12.2.1	K00437	PF00374	NiFe / NiFeSe	Ni-Fe oxidized active center [NFV] or NiFeSe	5XLF	Desulfovibrio vulgaris
	Hydrogen production	H₂-production	P29166	(FeFe) Iron hydrogenase 1	hydA	1.12.7.2	K00533	PF02906	Fe	dicarbonyl [bis(cyanide- kappaC)]-mu-(imin dimethanethiolatato- 1kappaS:2kappaS)- mu-(oxomethylidene) diiron (II) [402]	6N59	Clostridium pasteurianum
		H ₂ -production/Mrp Antiporter	Q8U0Z6	(NiFe) Membrane-bound hydrogenase subunit alpha	mbhL	1.12.7.2	K18016	PF1434	NiFe	Formyl [bis(hydrocyanato-	6CFW	Pyrococcus furiosus

										1kappaC)] Fe-Ni [NFU]		
Nitrogen	Anomaly	From Ammonia to Hydrazine	Q1Q0T3	Hydrazine synthase subunit gamma	hzsA	1.7.2.7	K20932	PF18582	Fe	HEME C [HEC]	5C2V	Candidatus Kuenenia stuttgartiensis
	Anammox	From Hydrazine to Nitrogen	Q1PW30	Hydrazine dehydrogenase	hdh	1.7.2.8	K20935	PF13447	Fe	HEME C [HEC]	6HIF	Candidatus Kuenenia stuttgartiensis
	Assimilatory nitrate reduction	From Nitrate to Nitrite	P73448	Nitrate reductase	narB	1.7.7.2	K00367	PF00355	Мо	Mo (VI [MO] + 2 molybdenum- bis(molybdopterin guanine dinucleotide) [MGD]	AF-P73448-F1	Synechocystis sp. PCC 6803
		From Ammonia to Nitrite	P9WJ03	Ferredoxin-nitrite reductase	nirA	1.7.7.1	K00366	PF01077 , PF03460	Fe	Siroheme [SRM]	1ZJ8	Mycobacterium tuberculosis H37Rv
	Dissimilatory Nitrite Reduction	From Ammonia to nitrite	Q72EF3	Cytochrome c nitrite reductase subunit NrfA	nrfA	1.7.2.2	K03385	PF02335	Fe	HEME C [HEC]	2J7A	Desulfovibrio vulgaris str. Hildenborough
	Denitrification	From Nitrate to Nitrite	P09152	Respiratory nitrate reductase 1 alpha chain	narG	1.7.5.1	K00370	PF00384 PF01568	Мо	Mo (VI) [MO] + 2 PO4-(2-amino-4-oxo- 3,4,5,6,-tetrahydro- pteridic-6-YL)-2- hydroxy-3,4- dimercapto-butenyl ester guamylate [MD1]	1Y4Z	Escherichia coli
			P81186	Periplasmic nitrate reductase	napA	1.9.6.1	K02567	PF04879 PF00384 PF01568	Мо	Mo (VI) [MO] + 2 molybdenum- bis(molybdopterin guanine dinucleotide) [MGD]	2JIM	Desulfovibrio desulfuricans
		From Nitric oxide to	E8PLV7	Copper-containing nitrite reductase	nirK	1.7.2.1	K00368	PF00394 , PF00732	Cu	Cu (II) [CU]	6HBE	Thermus scotoductus
		Nitrite	P24474	Nitrite reductase	nirS	1.7.2.1	K15864	PF02239 , PF13442	Fe	HEME C [HEC]	6TSI	Pseudomonas aeruginsa
		From Nitrite to Nitrate	P49050	Nitrate reductase [NADPH]	nasA	1.7.7.2	K00372	PF04879 , PF00384 , PF01568	Мо	(Molybdopterin-S,S)- dioxo-thio-Mo (IV) [MTV]	2BIH	Ogataea angusta
		From Nitrogen to Nitrous oxide	Q51705	Nitrous-oxide reductase	nosZ	1.7.2.4	K00376	PF00116 PF18764 PF18793	Cu	Cu4S [CUZ]	1FWX	Paracoccus denitrificans
		From Nitrous oxide to Nitric Oxide	B3Y963	Nitric oxide reductase	norB	1.7.2.5	K04561	PF00115	Fe	Protoporphyrin IX containing Fe [HEM]	3AYF	Geobacillus stearothermophilus
	Nitrification	From Ammonia to	Q04508	(Cupredoxin) Ammonia monooxygenase beta subunit	amoB1 ; amoB2	1.14.99.3 9	K10944	PF02461	Cu	Cu (II) [CU]	AF-Q04508-F1	Nitrosomonas europaea
		Hydroxylamine	P31101	Hydroxylamine reductase	hcp	1.7.99.1	K05601	PF03063	Fe	Iron/Sulfur/Oxygen Hybrid Cluster [FSO]	1E1D	Desulfovibrio vulgaris
		From Ammonia to Nitrite	P08201	Nitrite reductase (NADH) large subunit	nirB	1.7.1.15	K00363	PF04324 PF01077 PF03460 PF07992 PF18267	Fe	-	AF-P08201-F1	Escherichia coli (strain K12)
		From Hydroxilamine to Nitrite	Q50925	Hydroxylamine oxidoreductase	hao	1.7.2.6	K10535	PF13447	Fe	HEME C [HEC]	1FGJ	Nitrosomonas europaea
		From Nitrite to Nitrate	Q1PZD8	nitrite oxidoreductase subunit A	nrxA	1.7.99	K00370	PF09459	Мо	Mo (VI) [MO] + 2 PO4-(2-amino-4-oxo- 3,4,5,6,-tetrahydro-	7B04	Candidatus Kuenenia stuttgartiensis

										pteridic-6-YL)-2- hydroxy-3,4- dimercapto-butenyl ester guamylate [MD1]		
			_a	Nitrogenase iron-iron protein Nitrogenase molybdenum-iron protein	anfD	1.18.6.1	K00531	PF00148	Fe	FeFeco ^a	8OIE ^a	Azotobacter Vinelandi
	Nitrogen fixation	From Nitrogen to Ammonia	P07328	alpha chain	nifD	1.18.6.1	K02586	PF00148	Мо	FeMoco [ICS]	3U7Q	Azotobacter Vinelandi
			P16855	Nitrogenase vanadium-iron protein alpha chain	vnfD	1.18.6.1	K22896	PF00148	V	FeVco [8P8]	5N6Y	Azotobacter Vinelandi
		Hydrogen peroxide detoxification	Q3JNW6	Catalase-peroxidase	katG	1.11.1.21	K03782	PF00141	Fe	Protoporphyrin IX Containing FE [HEM]	5SW4	Burkholderia pseudomallei
		Oxygen detoxification	P0ABE5	Superoxide oxidase CybB	cybB	1.10.3.17	K12262	PF01292	Fe	Protoporphyrin IX Containing FE [HEM]	5OC0	Escherichia coli
			P80734	Superoxide dismutase [Ni]	sodN	1.15.1.1	K00518	PF09055	Ni	Ni (II) [NI]	1Q0G	Streptomyces seoulensis
	Oxygen radicals detoxification		P00446	Superoxide dismutase [Cu-Zn]	sodC	1.15.1.1	K04565	PF00080	Cu	Cu (II) [CU]	1BZO	Photobacterium leiognathi
		Superoxide detoxification	Q9RUV2	Superoxide dismutase [Mn]	sodA	1.15.1.1	K04564	PF02777 , PF00082	Fe / Mn	Fe (III) [FE] / Mn (II) [MN]	1Y67, 3KKY	Deincoccus radiodurans
			P82385	Superoxide reductase	sorA	1.15.1.2	K05919	PF06397 , PF01880	Fe	Fe (III) [FE]	1DQI	Desulfovibrio desulfuricans
Oxygen	Oxygen respiration	Oxidative phosphorylation	D9IA44	Cbb3-type cytochrome c oxidase (subunit II)	ccoN	7.1.1.9	K00404	PF00115	Cu	Cu (II) [CU] + Protoporphyrin IX [HEM]	5DJQ	Stutzerimonas stutzeri
			P34956	Cytochrome ba quinol oxidase subunit 1	qoxB	7.1.1.5	K02827	PF00115	Cu	Cu (II) [CU] + Heme- A [HEA]	6КОВ	Bacillus subtilis
			P0ABJ9	Cytochrome bd-l ubiquinol oxidase subunit 1	cydA	7.1.1.7	K00425	PF01654	Fe	Cis-heme D hydroxychlorin gamma-spirolactone [HDD]	6RKO	Escherichia coli
			P24244	Putative cytochrome bd-II ubiquinol oxidase subunit AppX	аррС	7.1.1.7	K00425	PF01654	Fe	Cis-heme D hydroxychlorin gamma-spirolactone [HDD]	7OY2	Escherichia coli
			P0ABJ6	Cytochrome bo(3) ubiquinol oxidase subunit 4	суоВ	7.1.1.3	K02298	PF00115	Cu	Cu (II) [CU] + HEME- O [HEO]	7N9Z	Escherichia coli
			P98005	Cytochrome-c oxidase polypeptide I + III	ctaD	7.1.1.9	K02274	PF00115	Cu	Cu (II) [CU] + Heme- AS [HAS]	2YEV	Thermus thermophilus
	Oxygenic photosynthesis	Water oxidation to oxygen	P0A444	Photosystem II protein D1 1	psbA1	1.10.3.9	K02703	PF00124	Mn	Oxygen evolving system [OEC]	3KZI	Thermosynechococcus elongatus
Sulfur		From S- sulfanylglutathione to glutathione + sulfite	A5VWI3	Sulfur dioxygenases	sdoA	1.13.11.1 8	-	PF00753	Fe	Fe (III) [FE]	4YSK	Pseudomonas putida
	Aerobic sulfur disproportionation	Catalyzes the simultaneous oxidation and reduction of elemental sulfur in the presence of oxygen	P29082	Sulfur oxygenase/reductase	sor	1.13.11.5 5	K16952	PF07682	Fe	Fe (III) [FE]	2CB2	Acidianus ambivalens
	Assimilatory sulfate	Reduction of sulfite to sulfide	A0A920E3E6	Assimilatory sulfite reductase (ferredoxin)	sir	1.8.7.1	K00392	PF03460 , PF01077	Fe	SIROHEME [SRM]	-	Synechococcus sp. PCC7942
	reduction		P17846	Sulfite reductase [NADPH] hemoprotein beta-component	cysL	1.8.1.2	K00381	PF01077 , PF03460	Fe	SIROHEME [SRM]	1AOP	Escherichia coli
	DMSO reduction	Catalyzes the conversion of DMSO to dimethyl sulfide	Q57366	Dimethyl sulfoxide/trimethylamine N- oxide reductase	dmsA	1.8.5.3	K07306	PF04879 PF01568 PF00384	Мо	Mo (VI [MO] + 2 molybdenum- bis(molybdopterin guanine dinucleotide) [MGD]	1EU1	Rhodobacter sphaeroides
		DMSO reduction	Q8GPG4	Dimethylsulfide dehydrogenase subunit alpha	ddhA	1.8.2.4	K16964	PF00384 PF01568	Мо	Mo (VI [MO] + 2 molybdenum- bis(molybdopterin guanine dinucleotide)	AF-Q8GPG4- F1	Rhodovulum sulfidophilum

									[MGD]		
Sulfate reduction	Catalyzes the reduction of sulfite to sulfide	Q59109	Sulfite reductase, dissimilatory-type subunit alpha	dsrA	1.8.99.5	K11180	PF03460 , PF01077	Fe	Siroheme [SRM]	3MM5	Archeoglobus fulg
Sulfite oxidation	Sulfite oxidation to sulfate	D3RNN8	Sulfite dehydrogenase subunit A	soeA	1.8.5.6	K21307	PF04879 , PF00384 , PF01568	Мо	Mo (VI [MO] + 2 molybdenum- bis(molybdopterin guanine dinucleotide) [MGD]	AF-D3RNN8- F1	Allochromatium vi
Sulfur disproportionation	From sulfite to sulfate	Q9LA16	Sulfite:cytochrome c oxidoreductase subunit A	sorA	1.8.2.1	K05301	PF00174 , PF03404	Мо	(molybdopterin-S,S)- oxo-Mo [MSS]	2BPB	Starkeya novella
Sulfur reduction	Catalyzes the cytoplasmic production of hydrogen sulfide in the presence of elemental sulfur	-	Sulfhydrogenase	shyB	1.12.98.4	K17995 , K17996	PF17179 PF00175 PF10418	Fe	-	-	Pyrococcus furios
Sullul Teduction	Sulfur reduction	Q8NKK1	Sulfur reductase molybdopterin subunit	sreA	1.97.1.3 / 1.12.98.4	K17219	PF04879 , PF01568 , PF00384	Мо	Mo (VI) [MO] + 2 molybdenum- bis(molybdopterin guanine dinucleotide) [MGD]	AF-Q8NKK1- F1	Acidianus ambiva
Thiosulfate	From thiosulfate to sulfate	O07819	Sulfur-oxidation complex	soxCD	1.8.2.6	K17225	PF00174 , PF03404	МоСо	Mo (IV) oxide [2MO] + Co (II) [CO]	2XTS	Paracoccus pantotrophus
oxidation	From thiosulfate to tetrathionate	D3RVD4	Thiosulfate dehydrogenase	tsdA	1.8.2.2	K19713	PF13442	Fe	HEME C [HEC]	4V2K	Allochromatium v
Thiosulfate reduction	From thiosulfate to hydrogen sulfide	Q72LA6	Polysulfide reductase chain A	phsA/ psrA	1.8.5.5	K08352	PF04879 , PF00384 , PF01568	Мо	Mo (VI) [MO] + 2 molybdenum- bis(molybdopterin guanine dinucleotide) [MGD]	2VPX	Thermus thermop
Sulfite reduction	Reduces sulfite to sulfide	Q58280	Coenzyme F420-dependent sulfite reductase	fsr	1.8.98.3	K21816	PF00037 PF04432 PF04422 PF01077 PF03460	Fe	SIROHEME [SRM]	7NP8	Methanocaldococ jannaschii

Competing Interests

The authors declare that there are no competing interests associated with the manuscript.

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Supplementary Online Materials Supplementary Methods

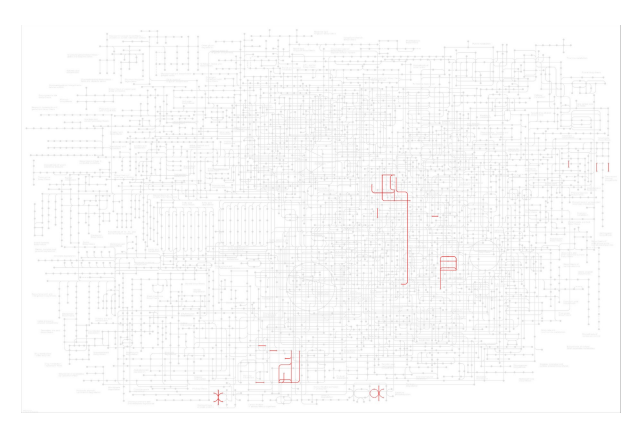
The enzymatic classes involved in the energetic pathways of the different biogeochemical cycles were counted from KEGG, querying the KEGG REST API service for cross-references between pathway (map entries) or modules (Module entries) and enzyme classes. The service deduplicates repeated enzymatic classes. Enzymatic classes not cross-referenced by the service were manually added. These are:

- EC:1.7.1.10, EC:1.7.2.7, EC:1.7.2.8 for nitrogen cycle, as they are involved in the anammox pathway;
- EC:1.2.5.3 for carbon cycle, as it is essential for aerobic oxidation of CO to CO₂;
- EC:1.13.11.18, EC:1.13.11.55, EC:1.8.2.1, EC:1.8.5.5, EC:1.8.5.6, EC:1.97.1.3, EC:1.12.98.4, as they are crucial for energetic sulfur metabolism.

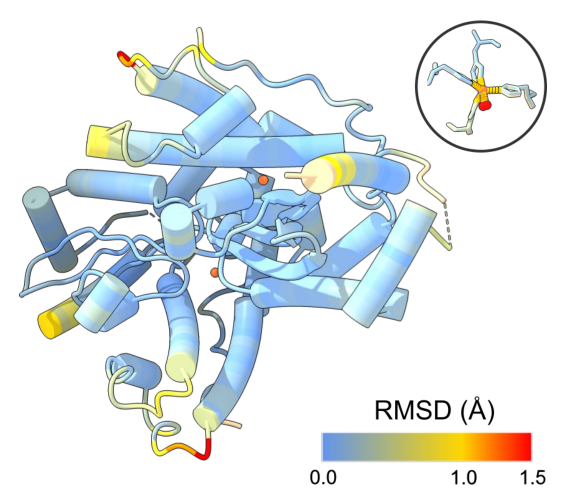
Supplementary Table 1. Enzyme names used in Figure 4.

Enzyme abbreviation	Enzyme name
fdhF	Formate dehydrogenase
fwdA	Tungsten formylmethanofuran dehydrogenase
codh	Anaerobic carbon monoxide dehydrogenase
coxL	Aerobic carbon monoxide dehydrogenase
cdh	acetyl-CoA decarbonylase/synthase
acs	Acetate synthase
ack	Acetate kinase
mcrA	Methyl-coenzyme M reductase
hdrA	Heterodisulfide reductase
mta	Methyl-Co(III) coenzyme M methyltransferase
mmoX	Soluble methane monooxygenase
ртоА	Particulate methane monooxygenase
mdh	Malate dehydrogenase
xoxF	Lanthanide-dependent methanol dehydrogenase
hydA	(FeFe) Iron hydrogenase 1
mbhL	(NiFe) Membrane-bound hydrogenase
frhA	(NiFe) F420-reducing hydrogenase
hydB	Periplasmic [NiFe] hydrogenase
hmd	5,10-methenyltetrahydromethanopterin hydrogenase
ccoN	Cbb3-type cytochrome c oxidase
qoxB	Cytochrome ba quinol oxidase
суоВ	Cytochrome bo(3) ubiquinol oxidase
ctaD	Cytochrome-c oxidase polypeptide I + III
cydA	Cytochrome bd-I ubiquinol oxidase
аррС	Putative cytochrome bd-II ubiquinol oxidase
kat	Catalase-peroxidase
cybB	Superoxide oxidase
sodN	Superoxide dismutase [Ni]
sodC	Superoxide dismutase [Cu-Zn]
sodA	Superoxide dismutase [Mn]
sorA	Superoxide reductase
psbA1	Photosystem II protein D1 1
narG	Respiratory nitrate reductase 1
napA	Periplasmic nitrate reductase
nasA	Nitrate reductase [NADPH]
narB	Nitrate reductase
nxr	Nitrite oxidoreductase

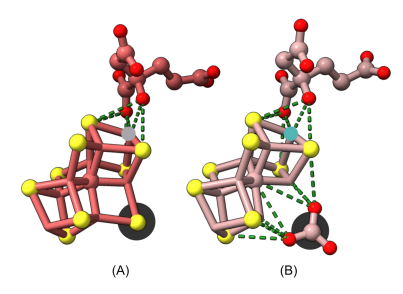
nrfA	Cytochrome c nitrite reductase
nirS	Nitrite reductase
nirK	Copper-containing nitrite reductase
norB	Nitric oxide reductase
nosZ	Nitrous-oxide reductase
nifD	Nitrogenase molybdenum-iron protein
anfD	Nitrogenase iron-iron protein
vnfD	Nitrogenase vanadium-iron protein
hsz	Hydrazine synthase
hdh	Hydrazine dehydrogenase
amoA	(Cupredoxin) ammonia monooxygenase
hcp	Hydroxylamine reductase
hao	Hydroxylamine oxidoreductase
cysD	Sulfite reductase [NADPH] hemoprotein
sat	Sulfate adenylyltransferase
apr	Assimilatory adenylylsulfate reductase
soeA	Sulfite dehydrogenase
sorA	Superoxide reductase
sox	Sulfur-oxidation complex
tst	Thiosulfate sulfurtransferase
dsrA	Sulfite reductase, dissimilatory-type
sir	Assimilatory sulfite reductase (ferredoxin)
cysL	Sulfite reductase [NADPH] hemoprotein
phs/psrA	Polysulfide reductase
sreA	Sulfur reductase
shy	Sulfhydrogenase
sqr	Sulfide:quinone oxidoreductase
fcc	Sulfide dehydrogenase
fsr	Coenzyme F420-dependent sulfite reductase
sor	Sulfite:cytochrome c oxidoreductase



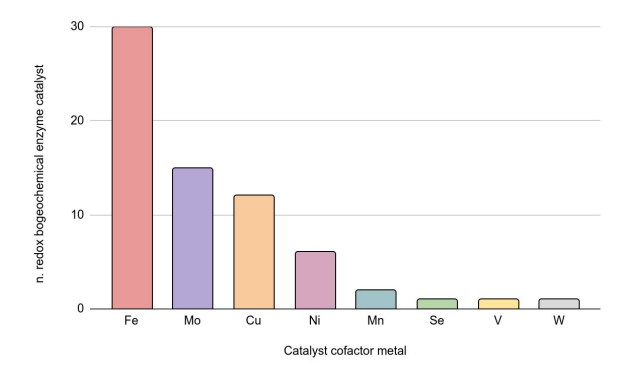
Supplementary Figure S1. Biogeochemically-relevant oxidoreductases as defined in this work highlighted in the KEGG metabolic map (PATH:MAP0100).



Supplementary Figure S2. Structural alignment of cambialistic [Fe-Mg] Superoxide dismutase gene (sodA) from *Deinococcus radiodurans* (1Y67, 3KKY; homodimeric assembly). <RMSD> = 0.277, RMSD_{residue} as per color key, using 1Y67 as reference. Inset: detail of the inner coordination sphere for the two metals.



Supplementary Figure S3. Details of *vnfD*-associated FeVco (A) and *nifD*-associated FeMoco (B). Vanadium and molybdenum are filled disks (JMol color code); positional equivalents (Fe in FeVco and CO3 in FeMoco) are highlighted by dark gray disk.



Supplementary Figure S4. Number of metals found in biogeochemically-relevant oxidoreductases as defined in this work.