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Review

Unifying concepts in anaerobic respiration: Insights from dissimilatory sulfur metabolism

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ABSTRACT

Behind the versatile nature of prokaryotic energy metabolism is a set of redox proteins having a highly modular character. It has become increasingly recognized that a limited number of redox modules or building blocks appear grouped in different arrangements, giving rise to different proteins and functionalities. This modularity most likely reveals a common and ancient origin for these redox modules, and is obviously reflected in similar energy conservation mechanisms. The dissimilation of sulfur compounds was probably one of the earliest biological strategies used by primitive organisms to obtain energy. Here, we review some of the redox proteins involved in dissimilatory sulfur metabolism, focusing on sulfate reducing organisms, and highlight links between these proteins and others involved in different processes of anaerobic respiration. Noteworthy are links to the complex iron-sulfur molybdoenzyme family, and heterodisulfide reductases of methanogenic archaea. We discuss how chemiosmotic and electron bifurcation/confurcation may be involved in energy conservation during sulfate reduction, and how introduction of an additional module, multiheme cytochromes c, opens an alternative bioenergetic strategy that seems to increase metabolic versatility. Finally, we highlight new families of heterodisulfide reductase-related proteins from non-methanogenic organisms, which indicate a widespread distribution for these protein modules and may indicate a more general involvement of thiol/disulfide conversions in energy metabolism. This article is part of a Special Issue entitled: The evolutionary aspects of bioenergetic systems.

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1. Introduction

The dissimilatory metabolism of sulfur compounds is likely to have been among the earliest energy-yielding processes to sustain life [1,2]. In the early anoxic Earth H_2S and SO_2 were emitted by volcanic and hydrothermal sources, and photolysis of these compounds would also generate elemental sulfur and sulfate [3,4]. Both H_2S and S^0 could sustain anoxygenic photosynthesis that would produce sulfate, or other oxidized sulfur species, and organic matter. Sulfate and S^0 could serve as electron acceptors for H_2 oxidation, and disproportionation of S^0 and sulfur compounds of intermediate oxidation state (thiosulfate,

Abbreviations: SRO, Sulfate reducing organisms; SOB, Sulfur oxidizing bacteria; LUCA, Last universal common ancestor; CISM, Complex iron–sulfur molybdoenzymes; $Tplc_3$, $Type\ I$ cytochrome c_3 ; $Tpllc_3$, $Type\ I$ cytochrome c_3

sulfite) was another possible biological strategy. There is evidence that photosynthetic processes were established at least 3.5 billion years ago [5,6], and dissimilatory sulfur metabolism was also already present at this time, either as sulfate reduction or sulfur disproportionation, as indicated by sulfur isotope fractionation studies [7-9] and microfossil records [10]. However, this biological activity had little impact on the biogeochemical cycling of sulfur until ~2.45 billion years ago [11], when a rise in atmospheric oxygen levels (Great Oxidation Event) promoted the increase of the oceanic sulfate concentration from weathering of sulfide minerals on land [12-15]. The increased oxygenation of the atmosphere was likely due to the activity of oxygen-producing cyanobacteria, which seem to have emerged at approximately the same time when O₂ started to increase, and much later than once believed [16–18]. The rising O₂ promoted weathering of continental pyrite and an increase in oceanic sulfate concentration to low mM levels [12,13,15]. However, for most of the Proterozoic the deep ocean waters remained anoxic and sulfidic or ferruginous, overlaid by an oxygenated surface layer [12,19-21], a state that may have been perpetuated until as recently as ~600 million years ago by anoxygenic photosynthesis with sulfide as electron donor [22]. After a second major oxidation event in the Neoproterozoic, the deep ocean waters became oxygenated and the sulfate levels rose to present day levels (28 mM), marking the start of the modern sulfur cycle, where biological sulfate reduction plays a major role, particularly in marine sediments

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where it is responsible for about 50% of carbon remineralization [23]. Overall, it is clear that there was an intimate connection between the history of Earth's atmosphere and the biogeochemical cycle of sulfur (reviewed in [24,25]).

The start of widespread biological sulfate reduction between 2.45 and 2.35 billion years ago is derived from the large increase in mass-dependent sulfur isotope fractionations observed during this period (reviewed in [24,26]). A limited incidence of biological sulfate reduction in the very early Earth is also reflected in the fact that this metabolic trait is not dispersed among prokaryotic organisms, and might have initially been restricted to some early branching thermophilic sulfate reducers. The emergence of mesophilic sulfate reducing organisms (SRO) apparently coincided, or shortly followed the increase in oceanic sulfate levels [27,28]. This radiation of mesophilic SRO seems to have taken place after the rapid diversification of bacterial lineages observed during the Archaean eon, where a significant expansion of energy metabolism genes apparently occurred [29].

A striking feature of energy metabolism/respiratory proteins is their modular character, which has been described as being based on a "redox protein construction kit" [30], from which different combinations of a limited number of protein modules originate different protein complexes with diverse physiological functions. This modular character, which is observed in many protein families, denotes a conservative approach from Nature in using a limited number of original parts to derive new metabolic features. However, it probably also reflects the

high level of gene exchange that was present in the pool of LUCA organisms [31], as well as the high incidence of lateral gene transfer in later prokaryotes [32]. In sulfur-metabolizing organisms we find interesting and unique variations of respiratory proteins that reflect their ancient origin and their close environmental association with other anaerobic organisms, in particular with methanogens. Here, we present a short review of respiratory proteins involved in dissimilatory sulfur metabolism, focusing on SRO, and discuss new "parts" of the "redox protein construction kit" that are strongly associated with sulfur metabolism but show also links to other respiratory proteins (Fig. 1) [33]. We will not discuss several respiratory membrane proteins that are present in SRO, but also in many other classes of prokaryotes, and thus are not specifically related to sulfur metabolism. A discussion of these can be found in [33].

2. The AprBA and DsrAB terminal reductases and their evolution

There are two biological pathways of sulfate reduction. In the assimilatory pathway, which is widespread in the three domains of life, sulfate is reduced to sulfide in small amounts and this is transformed into cysteine, from which other biological sulfur-containing molecules are derived [34]. In the dissimilatory pathway, which is restricted to five bacterial and two archaeal lineages, sulfate is the terminal electron acceptor of the respiratory pathway producing large quantities of sulfide [35–37]. The two pathways (Fig. 2) start with activation of sulfate by reaction with ATP to form adenosine-5′-phosphosulfate (APS), a step catalyzed

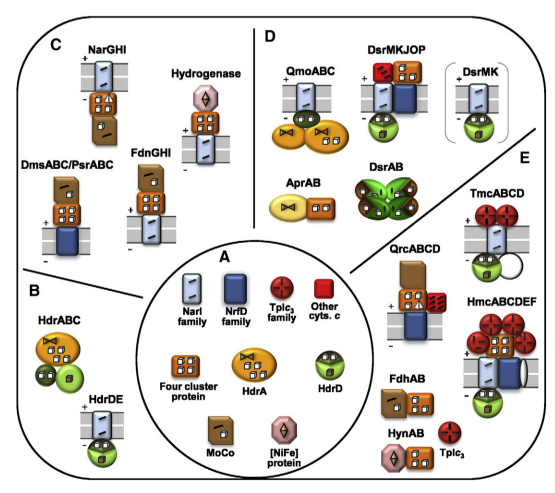


Fig. 1. The modular nature of sulfate respiration and related proteins. A) Redox modules i.e. building blocks from the "redox construction kit" [30] that pertain to SRO. B) Heterodisulfide reductases of methanogens. C) Trimeric respiratory enzymes including the CISM family and others (Hyn hydrogenase). D) Conserved respiratory proteins of SRO (for exceptions see text; only the "minimum" unit DsrMK is present in a few organisms). E) Periplasmic and membrane complexes of cytochrome-rich SRO (mainly *Deltaproteobacteria*). The proteins and respective cofactors are represented schematically (see text for descriptions).

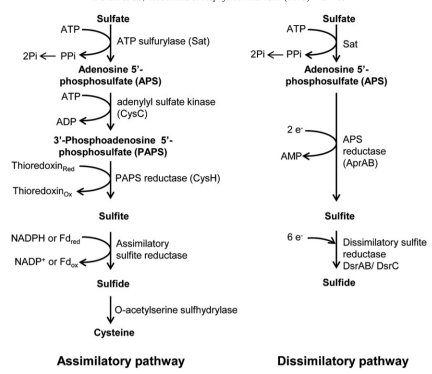


Fig. 2. The prokaryotic assimilatory and dissimilatory pathways of sulfate reduction.

by the trimeric sulfate adenylyl transferase (Sat), also known as ATP sulfurylase [38,39]. The formation of APS is endergonic and is driven by hydrolysis of the pyrophosphate formed by a pyrophosphatase (soluble or membrane-bound). So, the activation of sulfate to APS is considered to consume two ATP equivalents. In the prokaryotic assimilatory pathway APS is converted to 3'-phosphoadenosine-5'-phosphosulfate (PAPS) by the adenylyl sulfate kinase (CysC), PAPS is reduced to sulfite by a thioredoxin-dependent PAPS reductase (CysH), and finally sulfite is reduced to sulfide by an assimilatory sulfite reductase that is either multimeric and NADPH-dependent (CysII) or a monomeric ferredoxindependent enzyme [40]. In the dissimilatory pathway APS is reduced to sulfite by the APS reductase (AprBA), a heterodimeric iron-sulfur flavoenzyme [41-44]. Sulfite is reduced by the dissimilatory sulfite reductase DsrAB, a siroheme containing protein [45,46], with the involvement of the small protein DsrC (see below) [47-51]. Another small protein DsrD, which is often encoded downstream of dsrAB, might also be involved in sulfite reduction, possibly in a regulatory role, but its exact function is still unknown [52]. Interestingly, the dsrD gene is strongly downregulated in the presence of high sulfide concentrations [53]. In many anoxygenic phototrophic and chemolithotrophic sulfur oxidizing bacteria (SOB), the Sat, AprBA, DsrAB and DsrC proteins are also present, and thought to be involved in reverse oxidative reactions (reviewed in [54]). DsrAB and DsrC (and the associated DsrMKJOP complex, see Section 4.2) are also present in organisms that reduce sulfite, thiosulfate or organosulfonate compounds.

The evolution of the dissimilatory sulfate reduction pathway has been investigated by phylogenetic analysis of the *sat* [32,55], *aprBA* [32,56–59] and mostly of the *dsrAB* genes [32,57,60–65]. These studies indicate a mostly vertical inheritance for these genes, but also several episodes of lateral gene transfer (LGT).

The APS reductase is an $\alpha\beta$ heterodimer containing a FAD group in the AprA subunit and two $[4\text{Fe}-4\text{S}]^{2+/1}+$ clusters in the AprB subunit. AprBA is an example of a modular redox protein, as the AprA subunit shows strong structural similarity (although low sequence identity) to the module/family of flavoproteins containing fumarate reductase and aspartate oxidase, and AprB includes a domain similar to the bacterial ferredoxin module [42]. The *aprA* and *aprB* genes share a similar

evolutionary profile resulting from vertical inheritance and concurrent LGT. Several aprBA genes of SRO were acquired by LGT, namely among members of the Syntrophobacterales, Thermodesulfobacterium, Thermodesulfovibrio, Archaeoglobus and some deltaproteobacterial lineages [56,59]. The aprBA of SOB diverge into two phylogenetic lineages in which one, represented by AprBA from Allochromatium vinosum, is the authentic SOB group (lineage I, congruent with the monophyletic DsrAB phylogeny), and the other, represented by AprBA from Chlorobium tepidum (lineage II, discordant with DsrAB phylogeny) was acquired by LGT from SRO [58,59]. These two lineages correlate with different gene organizations (Fig. 3) and different physiological partners of AprBA, which are the integral membrane protein AprM in the case of SOB lineage I, and the OmoABC membrane complex [66] in the case of SRO and SOB lineage II [58,59]. This is further supported by homology modeling of AprBA from the two groups, which suggests different interacting partners for AprB [67].

The DsrAB sulfite reductase forms an $\alpha_2\beta_2$ unit, containing two siroheme cofactors, per $\alpha\beta$ unit, coupled to a [4Fe-4S] iron-sulfur cluster through the cysteine heme axial ligand. However, only one of the cofactors is catalytically active [50,68]. This protein is part of a large family, all sharing the same coupled cofactor, that includes also the assimilatory sulfite and nitrite reductases, and other proteins [40,69]. This family constitutes another module of the "redox construction kit", and probably diverged from a very ancient and primitive organism. The DsrA/DsrB proteins have also a modular character since they include a ferredoxin domain, which was probably the electron donor to a precursor enzyme that was later incorporated into the reductase gene sequence [45,69]. The dsrA and dsrB genes are paralogous, and seem to have derived from a gene duplication event preceding the divergence of the Archaea and Bacteria domains [45,57,64,65], in agreement with a very early onset of biological sulfite reduction. Furthermore, the assimilatory sulfite/nitrite reductases also display an internal two-fold symmetry of a module that is similar to DsrA/DsrB, suggesting they also resulted from a gene duplication event [40,68-70]. In fact, the core domains of DsrAB form a unit that is superimposable with the structures of the assimilatory enzymes (Fig. 4) [50], further stressing the common origin of the assimilatory

Sulfate Reducing Organisms

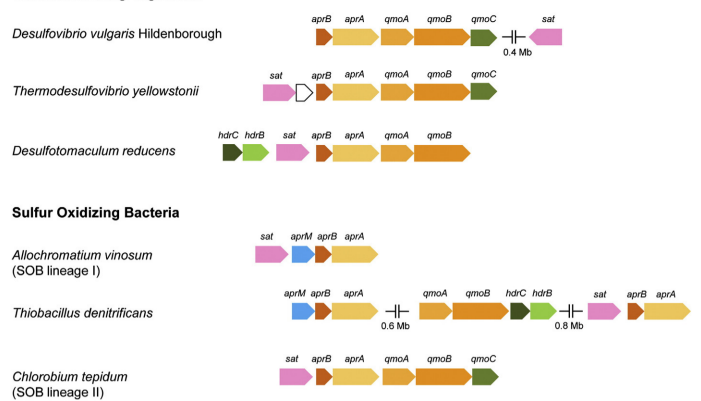


Fig. 3. Genomic organization of the *sat*, *apr* and *qmo* genes in selected SRO and SOB. *sat*, ATP sulfurylase; *aprBA*, APS reductase; *aprM*, transmembrane protein; *qmoABC*, subunits of the Qmo complex; *hdrBC*, subunits of heterodisulfide reductase. Adapted from [58,59].

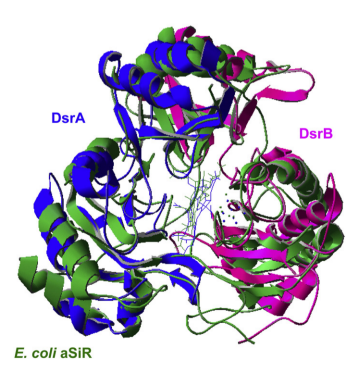


Fig. 4. Superposition of the core structures of assimilatory and dissimilatory sulfite reductases. The A1A2 and B1B2 domains of DsrA (blue) and DsrB (pink) subunits of the *D. vulgaris* Hildenborough dissimilatory sulfite reductase (PDB ID: 2v4j) [50] are superimposed on the structure of the *E. coli* assimilatory sulfite reductase (PDB ID: 1aop, green) [70]. The ferredoxin domains and the N- and C-terminal regions of the *D. vulgaris* DsrAB are omitted for clarity.

and dissimilatory enzymes from an ancestral gene that was present in one of the earliest life forms on Earth [64,69,70]. In the assimilatory enzymes the second cofactor was lost during evolution, indicating that in both families the process of gene duplication was associated with loss of function from one of the catalytic sites. A key difference between the assimilatory and dissimilatory sulfite reductases is that the former reduce sulfite directly to sulfide, whereas the latter form, *in vitro*, a mixture of products including also trithionate and thiosulfate, in relative proportions that depend on reaction conditions [71]. The physiological significance of these products is doubtful, as they may result from the absence of an essential component in the system, DsrC [50] (see Section 4.2), and be produced by further reaction of sulfite with semi-reduced intermediates present at the active site.

The phylogeny of DsrAB has been thoroughly investigated [32,57,60-65], and indicates a main pathway of vertical transmission, with a few episodes of LGT involving members of Thermodesulfobacterium and some low-GC Gram-positive bacteria of the phylum Firmicutes (Desulfotomaculum subclusters Ib, Ic, Id and Ie, Moorella thermoacetica and Ammonifex degensii) that acquired dsrAB from a deltaproteobacterial donor. The archaeal Archaeoglobi also have dsrAB genes of bacterial origin, indicating a cross-domain LGT. The DsrAB from SOB form a group clearly separated from SRO, while the DsrAB from the crenarchaeotal genus Pyrobaculum form a third group that represents the deepest branch in the dsrAB tree [62,63]. In the purple SOB A. vinosum [48,72], and the green SOB C. tepidum [73] it has been shown that the dsrAB and other dsr genes are essential for oxidation of sulfur globules stored in the periplasm, which are intermediates in the oxidation of sulfide and thiosulfate. In fact, most DsrAB-containing SOB are sulfur-storing members of the Chlorobi and Proteobacteria phyla [62]. If DsrAB from Pyrobaculum is of true archaeal origin, then the duplication of the dsr genes preceded the divergence of Bacteria and Archaea, and the ancestral DsrAB functioned in the reductive direction [63].

The spread of the sulfate reduction genes through a mobilizable metabolic island has been suggested [60], and gained support from the identification of genomic fragments from unidentified marine organisms containing a complete set of sulfate reduction genes [74]. However, this is unlikely to be a general mechanism, given the patchy distribution of *apr*, *dsr* and related genes in SRO, and the fact that the DsrAB tree topology is not congruent with that of AprBA, indicating independent LGT events. Nevertheless, the *Thermodesulfobacteriacae* and *Archaeoglobi* have similar branching positions in both the AprBA and DsrAB trees pointing to a concomitant acquisition of these genes conferring the capacity to reduce sulfate to sulfide [59]. In contrast, the ancestors of *Thermodesulfovibrio* may have been sulfite reducers (congruent DsrAB and 16S rRNA phylogenies, but not AprBA) that acquired the ability to respire sulfate later.

3. Modularity of simple respiratory membrane complexes

The modular nature of redox proteins is particularly evident in membrane-associated respiratory complexes. The simplest family of such complexes is the complex iron-sulfur molybdoenzyme family (CISM) that operates on a variety of reducing or oxidizing substrates, including formate, nitrate and several sulfur compounds (thiosulfate, DMSO, polysulfide and tetrathionate) [75,76]. This family is widespread in bacteria and greatly contributes to the flexibility of their respiratory chains [76,77]. Phylogenetic analysis indicates that most members of this family are very ancient and were likely present in LUCA [78]. CISM proteins include three subunits, or redox modules: i) a catalytic subunit that binds a pterin-guanine dinucleotide cofactor (that includes either Mo or W), and an [Fe-S] cluster; ii) a four-cluster subunit that binds four $[4Fe-4S]^{2+/1+}$ centers and is responsible for electron transfer between the membrane and catalytic subunits; and iii) a membrane subunit that has a quinone-binding site and is responsible for anchoring the other subunits to the membrane and for quinol oxidation/quinone reduction [76] (Fig. 1B). The quinone-interacting membrane subunit is the one showing more variation, and it can be broadly divided in two families: the first one comprises smaller proteins with 4 or 5 transmembrane helices (TMH), as in the case of FdnI of formate dehydrogenase and NarI of nitrate reductase, respectively. This family, which is usually referred to as the NarI-family, binds two hemes b on opposite sides of the membrane [79–81]. The hemes are coordinated by histidines present in two TMH in the case of NarI and three in the case of FdnI. The second family, which is usually referred to as the NrfD/PsrC family, includes between 8 and 10 TMH [82,83]. Sequence alignments indicate no conserved histidines to serve as heme ligands, and the structurally characterized member of this family (PsrC) does not contain hemes [82]. However, heterologous production of another protein from this family (DsrP from A. vinosum; see Section 4.2) unexpectedly resulted in a heme b-containing protein [84]. Therefore, it cannot be excluded that some members of this family may bind hemes.

The archetypal trimeric organization including a membrane anchor protein, an electron transfer subunit and a catalytic subunit is also found in a variety of other respiratory enzymes such as membranebound uptake hydrogenases, succinate dehydrogenases/fumarate reductases and others [30,85,86]. In many cases these membrane complexes are involved in energy conservation through charge separation and redox loops [87,88]. However, many succinate dehydrogenases/ fumarate reductases are tetrameric, containing two membrane-bound subunits, which suggests that single membrane anchor subunits may have resulted from gene fusions (or vice-versa). In SRO several respiratory membrane complexes are variations of this archetypal organization and have a specific role in dissimilatory sulfur metabolism. The first two examples, QmoABC and DsrMKJOP (see Section 4), are strictly conserved in SRO and are physiological partners of the two terminal reductases AprBA and DsrAB. These complexes are also present in other organisms that dissimilate sulfur compounds such as SOB and, in the case of the Dsr complex, in sulfite/thiosulfate/organosulfonate reducers, so they seem to have a dedicated role in sulfur metabolism. An interesting feature of QmoABC and DsrMKJOP is that they both contain subunits that are closely related to subunits of heterodisulfide reductases (Hdr). These enzymes, present in methanogenic archaea, are responsible for reducing the heterodisulfide of coenzymes B and M (CoB-S-S-CoM), which is formed in the last step of methanogenesis and is the terminal electron acceptor of the respiratory chain [89–91]. The second group of SRO membrane complexes, which includes Qrc and the Hmc/Tmc/Nhc family (see Section 6), is specific for those SRO that are rich in multiheme cytochromes *c* (mainly of the *Deltaproteobacteria* class).

4. The strictly conserved Qmo and Dsr membrane complexes

Energy conservation in SRO remains to be fully elucidated. One of the main questions that persisted for many years was the nature of the physiological electron donors to the APS and sulfite reductases. The demonstration that SRO could grow with H₂ as sole energy source [92] was a landmark achievement, since it demonstrated that sulfate reduction was associated with energy conservation through oxidative phosphorylation, and thus a membrane-associated electron transfer chain had to be present to generate a proton-motive force [93]. The role of quinones in sulfate respiration was disregarded for a long time (despite menaguinones being widespread in SRO [94]), because the redox potential of menaquinol ($E^{\circ\prime} = -75 \text{ mV}$) was not thought to be low enough to allow reduction of APS to sulfite (E° ' = -60 mV) or sulfite to sulfide ($E^{\circ\prime} = -116$ mV). The recent study of membrane complexes in SRO, together with genetic and transcriptomic studies, and the explosion in genomic information, have contributed to a better understanding of how membrane complexes may be involved in sulfate reduction and contribute to energy conservation [33,95–97]. The nature and mechanisms of these complexes are not straightforward, as discussed below, which has hampered our understanding of how they contribute to energy conservation. We have put forward some proposals, but these still need experimental validation. Elucidating the complete pathway and mechanism of sulfate respiration is important to understand this major biogeochemical process, and a key requirement for models used to track sulfur-isotope fractionations in ancient geological samples [98].

4.1. QmoABC

The Omo complex was first described through its isolation and characterization from Desulfovibrio desulfuricans ATCC 27774 [66]. It is composed of three subunits, one membrane-bound (QmoC) and two cytoplasmic (QmoA and QmoB), which are all related to subunits of Hdrs. The modular character of QmoABC is quite unique and interesting, relative to the trimeric arrangement of membrane/electron transfer/catalytic proteins discussed above (Figs. 1 and 5). In fact, the QmoC subunit is unprecedented among respiratory complexes as it is constituted by the fusion of two modules: a cytochrome b transmembrane domain (of the Narl family) homologous to HdrE, and a hydrophilic cytoplasmic domain containing two [4Fe-4S] cluster binding sites, homologous to the electron transfer subunit HdrC. The QmoA and QmoB subunits are both flavoproteins homologous to HdrA, the soluble Hdr subunit that has been proposed to perform flavin-based electron bifurcation in methanogens [99]. Curiously, HdrE is part of the membrane-associated HdrDE enzyme present in methylotrophic methanogens, whereas HdrC is part of the soluble HdrABC present in hydrogenotrophic methanogens [91] (see Section 7). QmoA is smaller than HdrA and includes only the flavin-binding site, whereas QmoB contains additionally two [4Fe-4S] cluster binding sites and a further domain, not present in HdrA, that is homologous to MvhD, an electron transferring subunit of the F₄₂₀-non-reducing hydrogenase that forms a complex with the soluble HdrABC [99,100]. Cofactor analysis confirmed

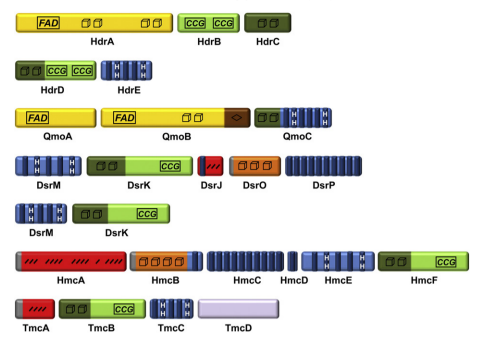


Fig. 5. Schematic representation of Hdr proteins and related complexes from SRO. Similar colors denote sequence identity. Cubes–[4Fe–4S] clusters, CCG–CX_nCCGX_mCX₂C sequence motif, transmembrane helices are in dark blue, signal peptide in grey, H–conserved histidines, and /–hemes c.

that the Qmo complex binds two hemes b, two FAD groups and several [Fe-S] clusters [66]. The redox potentials of the two QmoC hemes are +75 and -20 mV. Since the two hemes are reduced by quinols and the qmo genes were found next to aprBA, it was proposed that QmoABC was involved in electron transfer from the quinone pool to AprBA [66]. Recently, a deletion mutant of the qmoABC genes in Desulfovibrio vulgaris Hildenborough confirmed that the Qmo complex is essential for sulfate, but not sulfite, reduction [101].

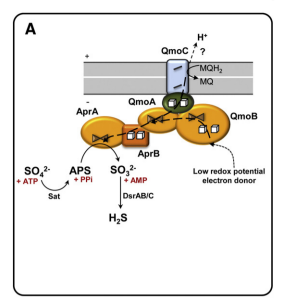
The *qmo* genes are conserved in all SRO genomes sequenced to date (except *Caldirvirga maquilingensis*), usually as part of a *sat–aprBA-qmoABC* gene cluster [33,59] (Fig. 3), and they have a phylogenetic profile congruent with *aprBA* from SRO and SOB lineage II [59,67]. However, in Gram-positive SRO the *qmoC* gene is absent, and is possibly replaced by the soluble *hdrBC* genes [33,59,102]. The QmoABC complex is also present in SOB of lineage II (see Section 2) [54,58], and in *C. tepidum* it was shown to be essential in oxidation of sulfite as an intermediary in the sulfur oxidation pathway [103,104]. In some SOB, as in Gram-positive SRO, the *qmoC* gene is replaced by two *hdrBC* genes (Fig. 3) [58,59].

Recently, we demonstrated that there is a direct interaction between the Qmo complex and AprBA, involving QmoA [105]. However, electron transfer between quinol analogues to AprBA through QmoABC could not be observed. We suggested that the reduction of APS by menaquinol has to be energy-driven due to the small difference in redox potential between menaquinol ($E^{\circ\prime} = -75 \text{ mV}$) and APS ($E^{\circ\prime}$ $APS/SO_3^2 = -60 \text{ mV}$), and to the fact that the membrane potential (~150 mV) has to be overcome when transferring electrons from the quinone binding site in QmoC (likely situated towards the periplasmic side of the membrane) to AprBA in the cytoplasm. This reaction cannot be driven by the membrane potential due to the topology of the Qmo subunits, as the electron flow goes against this potential. Instead, we proposed that a third partner is required to couple the reduction of APS by menaquinol to a second more favorable reaction. Based on the similarity of QmoB to HdrA, which is responsible for electron bifurcation [99,106], we proposed that a process of reverse electron bifurcation, i.e. electron confurcation, operates during APS reduction [105]. In such a process a low-redox potential electron donor is required to allow oxidation of menaquinol by APS. Menaquinol and a cytoplasmic reductant of low redox potential (from ferredoxin, H2, formate or NADH oxidation) could both serve as electron donors to the Qmo complex, which would confurcate electrons to the APS reductase (Fig. 6A). The favorable reduction of APS by this low potential electron donor would drive the unfavorable reduction of APS by menaquinol. The FAD group of QmoA or QmoB could serve as the confurcating center, where a high redox potential "hot" flavosemiquinone [107] would be generated by the first electron coming from the low potential donor, and would then be a favorable electron acceptor for a second electron coming from menaquinol, and in practice "pulling" this electron from the quinone. Overall, the advantage of this process is that it allows for the coupling of APS reduction with chemiosmotic energy conservation. This idea of electron confurcation during APS reduction expands the growing concept that electron bifurcation/confurcation may be an ancestral form of energy coupling involving two-electron centers such as flavins, quinones, or the Mo and W metals [91,99,106–112].

4.2. DsrMKJOP

The *dsrMKJOP* genes were first identified in the purple SOB *A. vinosum* as part of a large gene cluster involved in the oxidation of intracellular sulfur, containing also the *dsrAB* and *dsrC* genes (Fig. 7) [72]. These genes encode a multimeric transmembrane complex that was first isolated and characterized from *Archaeoglobus fulgidus* [113], where it was named Hme (for Hdr-like menaquinol-oxidizing enzyme complex), and later from *D. desulfuricans* ATCC 27774 [51], where the *dsr* nomenclature was adopted since it was already clear that in several genomes of SRO, SOB and sulfite reducers, the *dsrMKJOP* genes were associated with *dsrAB* and *dsrC*. In the SOB *A. vinosum* each subunit of this complex was shown to be essential for sulfur oxidation [114], and a membrane fraction enriched in DsrKJO contained also DsrAB and DsrC, suggesting an interaction between these proteins [48]. The whole complex was also recently purified using an affinity tagged Dsr [84].

The DsrMKJOP complex is another interesting variation in the family of respiratory complexes (Figs. 1 and 5). It includes two periplasmic subunits: DsrJ, a periplasmic tri-heme cytochrome c that shows no sequence similarity to any proteins in the databases; and DsrO that belongs to the family of four cluster proteins present in CISM (although in some SRO one of the [4Fe-4S] binding sites is missing). Two integral



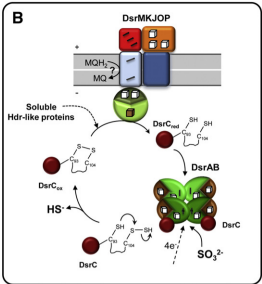


Fig. 6. Proposed mechanisms of APS and sulfite reduction. A) Electron confurcation hypothesis (grey dashed line): menaquinol (MQH₂) and a cytoplasmic low-redox potential partner both donate electrons to the Qmo complex, which transfers them to AprBA for APS reduction [105]. B) The four electron reduction of sulfite by DsrAB generates a persulfide in DsrC, which by displacement of sulfide forms an intramolecular disulfide bridge, DsrC_{ox}. This oxidized form of DsrC is reduced by the DsrK protein of the DsrMKJOP complex [50].

membrane subunits are present: DsrM that is homologous to HdrE, binds two hemes *b*, has five to six TMH, and belongs to the Narl family; and DsrP that has 10 TMH and belongs to the NrfD/PsrC family. Finally, DsrK is a cytoplasmic iron–sulfur protein homologous to the catalytic subunit HdrD, which is responsible for heterodisulfide reduction by the membrane-bound HdrDE complex [115,116]. DsrK has two typical

binding sites for $[4\text{Fe}-4\text{S}]^{2+/1+}$ clusters and one CCG domain (see Section 7) containing a five-cysteine motif that in Hdrs binds a catalytic $[4\text{Fe}-4\text{S}]^{3+}$ cluster responsible for heterodisulfide reduction [89]. The characteristic EPR signal of this $[4\text{Fe}-4\text{S}]^{3+}$ cluster is also detected in the *A. fulgidus* and *D. desulfuricans* complexes [51,113], suggesting the involvement of DsrK in thiol/disulfide chemistry. The Dsr complex

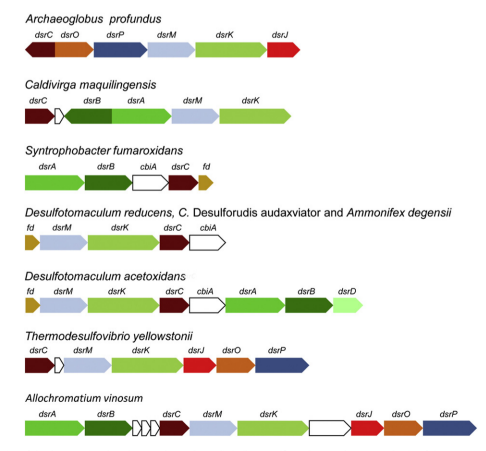


Fig. 7. Genomic organization of the dsr genes in selected SRO and SOB. dsrAB, dissimilatory sulfite reductase; dsrMKJOP, subunits of the Dsr complex; dsrC, DsrC protein; cbiA, cobyrinic acid a,c-diamide synthase; fd, ferredoxin. Adapted from [33].

isolated from *D. desulfuricans* contained the full complement of cofactors, including two hemes *b* and three hemes *c* per molecule [51].

The DsrMKJOP complex appears to be a combination of two sub-complexes: the DsrMK proteins are closely related to the HdrDE proteins of methanogens, and the DsrOP subunits correspond to two of the CISM units, interacting with a new module, DsrJ. The presence of two distinct quinone-interacting proteins is striking. The DsrJOP module is probably involved in electron exchange between the periplasm and the quinone pool, and DsrMK between the membrane pool and the cytoplasm, possibly involving some form of quinone cycling. However, the complex seems to function as a complete unit as all the subunits are detected upon isolation from A. fulgidus and D. desulfuricans, so direct transmembrane electron transfer cannot be discarded. An exception is found in Archaeoglobus profundus where only a DsrMK complex was isolated [117], even though a single complete dsrOPMKJ locus is present in the genome. In the archaeon C. maguilingensis and the Gram-positive SRO (with the exception of the recently sequenced species of Desulfosporosinus) only the dsrMK genes are present, suggesting this is a minimum functional unit of the complex [33]. Several SRO contain the dsrMKJOP genes, and one or more copies of dsrMK. Curiously, in the clostridial organisms (and in three Deltaproteobacteria) a ferredoxin gene is also present, which might be related to the absence of the dsrIOP genes [33]. The function of the DsrI cytochrome remains enigmatic. This is a quite unique cytochrome as the three hemes have distinct ligation: His/His, His/Met and the unusual His/Cys coordination [51,118]. There are few proteins carrying this type of heme c coordination, and one of them, SoxXA is involved in thiosulfate oxidation [119-121]. In A. vinosum the Cys coordinating this heme was shown to be essential for oxidation of sulfur globules, suggesting a catalytic role for DsrI [118]. However, in SRO no sulfur chemistry is thought to occur in the periplasm. The DsrJ cytochrome is poorly reduced by the periplasmic Type I cytochrome c_3 (see Section 5) [51,122].

The substrate of the DsrK subunit is proposed to be the small thiol protein DsrC, which has two conserved cysteines in a C-terminal swinging arm, and is strictly conserved in all organisms that contain a DsrAB [49-51,72,113,123]. DsrC is one of the most highly abundant energy metabolism proteins in SRO [124,125], and dsrC is also one of the most abundant genes in metatranscriptomic analysis of communities containing SRO and SOB [126,127], reflecting its key role in dissimilatory sulfur metabolism. DsrC belongs to a larger family (e.g. Escherichia coli TusE) where only the last Cys is conserved, that is involved in sulfur trafficking [128]. In A. vinosum it was recently shown that DsrC can accept sulfur from the DsrEFH proteins (present only in SOB) [129]. The two Cys of DsrC are redox active and can form a disulfide bond [49,123]. The crystal structure of the D. vulgaris DsrAB in complex with DsrC revealed that the C-terminal arm of this protein penetrates inside the DsrAB structure, such that its penultimate cysteine comes into close contact with the siroheme catalytic site [50]. This feature was confirmed in more recent structures from different organisms [130,131], and in the case of *Desulfovibrio gigas* a different conformation of the DsrC arm was also detected in which the arm is retracted and the two cysteines are in close contact [130]. We have proposed a new mechanism for sulfite reduction involving DsrC (Fig. 6B) [50], in which SO_3^{2-} is reduced by four electrons to an S^0 state ($Fe^{III} - S^0 - OH$ intermediate), which is attacked by the DsrC penultimate Cys forming a persulfide. This is displaced by the other DsrC Cys forming an intramolecular disulfide bridge (DsrCox), which is reduced by DsrK, forming sulfide and regenerating DsrC for another catalytic cycle. An analogous model was proposed for the sulfur oxidation pathway [84]. This mechanism is supported by the observations that DsrC interacts with DsrK in both A. vinosum [84] and D. vulgaris (S.S. Venceslau and I.A.C. Pereira, unpublished results). The reductant for DsrCox is presumably menaquinol, and the involvement of DsrMK in the process is analogous to the reduction of the CoM-S-S-CoB disulfide by methanophenazine catalyzed by HdrDE, which is coupled to energy conservation [132]. However, based on the presence of the DsrJOP module, it seems likely that the reduction of $DsrC_{ox}$ may involve a third partner and a more complex mechanism, which may also require electron bifurcation/confurcation, as discussed for the Qmo/Apr couple. The redox potential of $DsrC_{ox}$ has not been determined yet, but usually disulfides have $E^{\circ\prime}$ in the order of -150 to -200 mV, so menaquinol might also not work as sole reductant.

5. Multiheme cytochromes c as periplasmic redox modules in SRO

The most well studied genus of SRO, Desulfovibrio, belongs to the deltaproteobacterial sulfate reducers, which are characterized by an abundant pool of multiheme cytochromes c ([133,134] and references therein). The prototype of this family is the tetraheme cytochrome c_3 (more precisely called Type I cytochrome c_3 or TpI c_3), which was the first cytochrome c to have been described in an anaerobe [135,136], and is one of the most highly expressed proteins in Desulfovibrio spp. Although it was believed for a long time to be an essential protein for sulfate reduction [137], the $Tplc_3$ is in fact absent from many sulfate reducers, and some SRO such as C. maquilingensis, Desulfotomaculum acetoxidans and Candidatus Desulforudis audaxviator contain no cytochromes c at all [33]. The SRO can be divided in two physiologically distinct groups: the first group has a high content of cytochromes c and includes Thermodesulfovibrio yellowstonii and the deltaproteobacterial SRO (with the exception of the psychrophilic Desulfotalea psychrophila that has a small number [138]); the second group has few or no cytochromes c and includes the archaeal and clostridial SRO [33]. The two groups differ in the relevance of periplasmic electron transfer pathways to their energy metabolism, as discussed below. It should be pointed out that all genomes of deltaproteobacterial SRO sequenced so far belong to organisms that have the potential to grow by either or both, formate and hydrogen, which is likely to be a common trait among this group of organisms.

The $Tplc_3$ is the periplasmic electron acceptor of hydrogenases and formate dehydrogenases. The presence of the *cycA* gene coding for the $Tplc_3$ in the genomes of SRO (often in multiple copies) correlates with the presence of periplasmic hydrogenases and formate dehydrogenases that lack a membrane subunit for direct quinone reduction [33], in contrast to most bacteria [86,139]. In several cases these enzymes have a dedicated cytochrome c_3 subunit [133,140,141]. These soluble uptake hydrogenases and formate dehydrogenases are usually present in several copies in the deltaproteobacterial SRO [33,141]. The enzymes have distinct expression patterns that depend on factors such as substrate concentration or metal availability. For example in D. vulgaris Hildenborough, the expression of hydrogenases depends on hydrogen concentration [142] and whether Ni and Se are available [143]. Likewise, different formate dehydrogenases are expressed in this organism in the presence of either Mo or W [144].

The $Tplc_3$ is another redox module of the "construction kit", having a compact tetraheme arrangement that performs a proton-coupled two-electron transfer [145,146]. Other cytochromes of the same family are subunits of membrane-associated complexes [133,147], described below. These include the sixteen-heme high molecular mass cytochrome (HmcA) [148] and the nine-heme cytochrome c (NhcA) [149], both of which contain several $Tplc_3$ domains, but also the Type II cytochrome c_3 ($Tpllc_3$ or TmcA), which has small structural differences relative to $Tplc_3$, but lacks its characteristic positive surface region [150–152]. Further members of the $Tplc_3$ family are present in other Deltaproteobacteria, including the triheme cytochrome c_7 from $Desulfuromonas\ acetoxidans\ [153]\ and\ the\ PpcA\ cytochromes\ from <math>Geobacter\ spp.\ [154]$. Besides the $Tplc_3\ family$, the deltaproteobacterial SRO contain other types of multiheme cytochromes c, as described in detail in [33,134].

As a recipient of electrons from H_2 or formate oxidation, the $Tplc_3$ then functions as a hub for periplasmic redox networks, as it can deliver this reducing power to several membrane complexes (Qrc,

Tmc, Hmc, Nhc, and possibly others, but not the Dsr complex) or other cytochromes c [122,133,134]. We have argued that using soluble dehydrogenases and $TpIc_3$, rather than direct quinone reduction, confers to the deltaproteobacterial SRO a higher metabolic flexibility, as electrons can be shuttled through several alternative pathways [155]. Thus, the Deltaproteobacteria SRO may derive additional electrons from intracellular cycling of redox intermediates such as hydrogen and formate [95,137,156], relative to the other groups of SRO. A high content of multiheme cytochromes c seems to be characteristic of soil and sediment Proteobacteria, such as Geobacter, Shewanella, Anaeromyxobacter and Desulfovibrio, which are subjected to variable redox conditions. Thomas et al. have argued that having a high number of multiheme cytochromes c is a hallmark of metabolically versatile anaerobes that have to adapt to environments with fluctuating redox conditions [157]. It has also been suggested that the large pool of cytochromes c in Geobacter act as capacitors, sustaining viability and motility for short periods of time as cells move between heterogeneously dispersed metal oxides [158]. The versatile nature of SRO is reflected in the fact that they can even grow in the absence of sulfate, in syntrophy with other organisms that consume H₂ and/or formate, such as methanogens [159-161]. In fact, SRO were found to be still abundant in methanogenic zones of marine sediments [162].

6. Cytochrome c-associated membrane complexes of deltaproteobacterial SRO

The presence of $Tplc_3$ in SRO correlates also with the presence of several membrane redox complexes having a periplasmic cytochrome c subunit. These complexes have also a highly modular character, as discussed above, and they are either involved in quinone reduction (Qrc and Nhc) or transmembrane electron transfer (Tmc and Hmc). They accept electrons from the $Tplc_3$ and/or seem to be involved in syntrophic metabolism.

6.1. The QrcABCD complex

The membrane-associated Quinone Reductase Complex (Qrc) was first described as a molybdopterin oxidoreductase involved in H_2 oxidation, by screening a library of *Desulfovibrio alaskensis* G20 transposon mutants for strains deficient in syntrophic growth with a methanogen [163]. Three mutants were identified with mutations in the *cycA* gene (Tplc3), *hydB* ([FeFe] hydrogenase) and *mopB* (coding for a putative molybdopterin oxidoreductase). The *cycA* and *mopB* mutants were also impaired in their ability to grow with H_2 or formate (but not lactate) as electron donors for sulfate reduction, pointing to their involvement in the electron transfer chain from H_2 or formate to sulfate [163]. The Qrc complex was isolated from *D. vulgaris* Hildenborough, where it was shown to act as a Tplc3:menaquinone oxidoreductase, but not to be a molybdopterin oxidoreductase, as it lacks a molybdenum or tungsten cofactor [155].

The Orc complex is composed of four subunits, three periplasmic (OrcABC) and one integral membrane subunit (OrcD) (Figs. 1 and 8). OrcA is a membrane-anchored hexa- or pentaheme cytochrome c, QrcB is a membrane-anchored protein of the molybdopterin oxidoreductase family, but which does not contain a molybdopterin cofactor. QrcC is a four cluster protein and QrcD is an integral membrane protein of the NrfD/PsrC family. The three QrcBCD subunits are analogous to the three subunits of CISM complexes discussed above. Thus, Qrc is an interesting variation of the CISM family that includes additionally a cytochrome c subunit [155]. In addition, its subunits are also closely related to some subunits of the Alternative Complex III (Act), which performs the reverse reaction of oxidizing the quinone pool and reducing a periplasmic redox partner [164-166]. Like Qrc, the Act has a subunit related to molybdopterin oxidoreductases, which lacks a molybdopterin cofactor, as also observed for the Nqo3/NuoG subunit of Complex I [167]. The function of this protein in Orc is presently unknown, as it is also the case for its homologues in Act and Nuo complexes, and it may have only a structural role. The D. vulgaris OrcABCD complex contains six hemes c, one $[3Fe-4S]^{1+/0}$ cluster and three $[4Fe-4S]^{2+/1+}$ [155], whose redox potentials were determined by EPR [168].

The Orc complex is efficiently reduced by periplasmic hydrogenases and formate dehydrogenases only in the presence of TpIc₃, and can reduce menaguinone analogues, having activity as TpIc₃: menaquinone oxidoreductase [155]. Thus, Qrc constitutes the missing link between TpIc₃ and the quinone pool. The qrcABCD genes are present in Deltaproteobacteria SRO that have TpIc3 and hydrogenases or formate dehydrogenases lacking a membrane subunit for direct quinone reduction [33,155]. The fact that it is essential for growth of D. alaskensis G20 in H₂ or formate and sulfate [163], indicates that Orc is the physiological electron acceptor of the $TpIc_3$ in this organism, and cannot be replaced by other complexes such as Tmc and Hmc, which are also present in this organism. Furthermore, Qrc also seems to be implicated in syntrophic growth of this organism [163,169] and also D. vulgaris [170]. In D. vulgaris, Qrc forms a supramolecular complex with the TpIc3 and a periplasmic hydrogenase [168]. The quinone binding site in QrcD is located close to the [3Fe-4S]^{1+/0} cluster of QrcC [155]. Energy conservation by QrcABCD will depend on whether proton uptake for quinone reduction occurs on the periplasmic side of the membrane (electroneutral process), or from the cytoplasm (electrogenic process) as it has been proposed for PsrC [82]. We have suggested that the Qrc and Qmo complexes may be involved in a redox loop mechanism that sustains electron transport across the membrane to the cytoplasmic reduction of sulfate, coupled to proton motive force generation during sulfidogenic growth on H_2 or formate [155].

The evolutionary relationship between Qrc, CISM complexes and Act is an interesting issue that deserves further study. Qrc may have evolved from a CISM complex by association of a cytochrome *c* and loss of the molybdopterin cofactor. Yanyushin et al. have also proposed that the

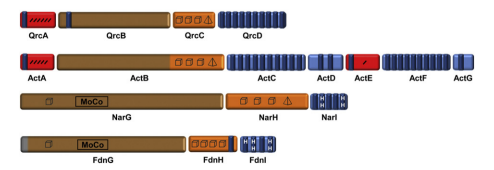


Fig. 8. Schematic representation of proteins from Qrc, Act and CISM complexes (NarGHI and FdnGHI). Similar colors denote sequence identity. Cubes—[4Fe-4S] clusters, pyramid—[3Fe-4S] cluster, transmembrane helices are in dark blue, signal peptide in grey, H—conserved histidines, /-hemes c, MoCo-molybdopterin cofactor.

Act complex arose from Qrc by acquisition of additional subunits [166], which would place Qrc as a stepping stone in evolution of bacterial complexes. Whatever the case, Qrc is an excellent example of how a different function can be achieved with a minimal modification of subunits, a strategy that forms the basis for the diversity and flexibility of bacterial energy metabolism.

6.2. The Hmc, Tmc and related complexes

The Hmc and Tmc are both transmembrane complexes with subunits in the periplasm, membrane and cytoplasm (Figs. 1 and 5). From a global perspective they share some features: a cytochrome c subunit of the Tpl c_3 family, one membrane cytochrome b and a cytoplasmic protein of the HdrD/DsrK family [96]. This suggests a similar function for both complexes in electron exchange between the periplasm and cytoplasm (or vice-versa), possibly involving thiol/disulfide exchanges. Curiously, most organisms that contain Hmc have also a Tmc complex [33].

The Hmc complex of *D. vulgaris* was the first transmembrane complex to be discovered in SRO [171]. This complex includes six subunits, HmcABCDEF, and has a composition that is strikingly similar to the Dsr complex (Fig. 1): one cytochrome c subunit (HmcA), one four-cluster protein (HmcB), two integral membrane proteins of the NrfD (HmcC) and NarI (HmcE) families and a cytoplasmic subunit homologous to HdrD (HmcF). However, the sequence identity between the Dsr and Hmc proteins is quite low, which suggests they may be paralogues that have diverged considerably. The cytochrome c subunit is actually completely different in the two complexes and points to a different function and/or physiological partner in the periplasm. HmcA is a large cytochrome with sixteen hemes organized in four TpIc₃-like domains (reviewed in [133]), and it can be reduced by this cytochrome [172]. Initial studies implicated Hmc in hydrogen uptake metabolism [173,174], but the hmc genes are downregulated during growth with H₂ [142,175]. These genes also have a low expression level in D. vulgaris Hildenborough grown in lactate/sulfate, relative to other membrane complexes [95]. A clear phenotype was observed for an hmc deletion mutant that was severely impaired in syntrophic growth on lactate with a methanogen [170]. Furthermore, comparative transcriptional analysis between syntrophic and sulphidogenic growth of D. vulgaris also indicated an upregulation of the Hmc complex in the former conditions [170]. A model was proposed in which reduced ferredoxin served as electron donor to Hmc, which then transferred electrons to TpIc₃ and this to periplasmic hydrogenases. However, there is no evidence to suggest that ferredoxin may interact with Hmc, and it seems more likely to be an electron donor to the Coo hydrogenase that is also upregulated, since energy-conserving hydrogenases are known to interact with ferredoxin [176]. This model also does not agree with previous observations that the hmc deletion mutant produced more H₂ than wild-type D. vulgaris from lactate, pyruvate or formate with limiting sulfate [156].

The *hmc* genes are present in all *Desulfovibrio* spp. sequenced to date, with the exceptions of *D. desulfuricans* ATCC 27774 and *D. piger* [33]. In these two species, instead of Hmc there is an Nhc complex, which is characterized by having a nine-heme cytochrome subunit (NhcA) that is very similar to the C-terminal domain of HmcA [149]. The Nhc complex is simpler than Hmc, lacking the cytochrome *b* and the cytoplasmic HdrD-like subunits. Thus, it should transfer electrons from the periplasm to the quinone pool [133].

The Tmc complex has four structural proteins, TmcABCD (in a $\alpha_2\beta\gamma\delta$ arrangement), and was isolated from *D. vulgaris* Hildenborough [177]. TmcA is a tetraheme cytochrome very similar to Tplc₃, also known as Type II cytochrome c_3 (Tpllc₃, previously also called acidic cytochrome c_3) [150–152]. TmcB is a cytoplasmic protein of the HdrD and DsrK family with a CCG domain, and which is very similar to HmcF. TmcC is an integral membrane cytochrome *b*, homologous to HmcE and of the Narl/HdrE/DsrM family. Finally, TmcD is a tryptophan-rich subunit with no

homology to any protein in the database [177]. TmcA is effectively reduced by the hydrogenase/TpIc₃ couple [150,151,178] and all the redox centers of Tmc are reduced with H₂ [177]. The tmcA gene is also upregulated during growth of *D. vulgaris* with H₂ [175], suggesting that Tmc is involved in transmembrane electron transfer from periplasmic H₂ oxidation. In *D. vulgaris* grown in lactate/sulfate the *tmc* genes are expressed at about the same level as the dsrMKJOP genes, indicating a functional role also in this growth condition [95]. However, a mutant deleted in the tmc genes had no apparent phenotype [95], which is not surprising as other proteins can most likely substitute for its function (Hmc, Qrc coupled with Qmo/Dsr, Ohc or others). In fact, a certain degree of interchangeability is presumably the reason for the presence of multiple cytochrome c-associated complexes in the Deltaproteobacteria SRO [33]. Another example of these is the Ohc complex (for Octaheme cytochrome complex) [141], whose function is unknown, and is expressed at low levels in D. vulgaris [95].

The Hmc and Tmc complexes have in common with the Dsr complex the presence of an HdrD-related subunit. The presence of the typical catalytic cofactor for thiol-disulfide catalysis in the Tmc complex was confirmed through the characteristic $\left[4Fe-4S\right]^{3+}$ cluster EPR signal [177]. Thus, Hmc and Tmc may also act as disulfide reductases, possibly on the $DsrC_{ox}$ protein, and thus link the periplasm with sulfite reduction. Given the considerable number of Hdr-related proteins in SRO and other organisms, we dedicate the following section to an analysis of these proteins, which have not been much investigated up to date outside of methanogens.

7. Hdr-related proteins as widespread redox modules in anaerobic respiration

In a recent genomic analysis of energy metabolism genes in 25 species of SRO we described the very high number of genes related to heterodisulfide reductases [33], which has also been pointed out by other authors in the context of individual genomes [102,179-182], or in other classes of organisms such as the acetogenic Moorella thermoaceticum [183]. The abundance of Hdr-like proteins in SRO [33,51,66,102,113,117,141,180] may indicate they were present in ancestral organisms, and/or that there was substantial exchange of genetic material between methanogens and SRO (and other classes of organisms), which could be due to their sharing common habitats. Hdrs are representative enzymes of a group of quite widespread proteins responsible for reduction of disulfides or oxidation of thiols [90], but they belong to a larger family that includes proteins that seem to have other functions (see below). In methanogenic archaea, the heterodisulfide is not an external substrate, but is produced in the final step of methanogenesis. By analogy, it is possible that thiol/ disulfides may be generated in other anaerobes and be involved in the respiratory chain, which would suggest that a sulfur-based energy metabolism, of obvious ancient origin, could be more widespread than presently considered [90,109].

There are two types of Hdr enzymes [91]: in methanogens without cytochromes a soluble HdrABC is present [89], which forms a complex with the MvhADG hydrogenase. This complex couples the endergonic reduction of ferredoxin by H_2 with the exergonic reduction of the heterodisulfide by H₂, through an electron bifurcation process [99]. In methanogens with cytochromes, a membrane-bound enzyme is present, HdrDE, which uses the quinone-like cofactor methanophenazine as electron donor in a process coupled to energy conservation [116,132,184]. The key subunits in Hdrs are the catalytic subunits (HdrB in the soluble enzyme and HdrD in the membrane-bound enzyme; actually HdrD resembles a fusion of the HdrBC proteins), and the HdrA subunit that contains a FAD group presumed to be responsible for bifurcation of electrons coming from the Mvh hydrogenase. There are several proteins related to both HdrA and HdrD in the genomes of SRO [33]. Several of these are multidomain proteins, and Strittmatter et al. proposed two new types of Hdr subunits, HdrF and HdrL based on proteins encoded in the

Desulfobacterium autotrophicum HRM2 genome (see below) [180]. Overall, both HdrA and HdrD (or more precisely the CCG domain) can be considered as additional modules of the "redox construction kit" that we discuss further below.

7.1. HdrA-related proteins

A complete set of hdrABC genes is found in many SRO, either next to a set of mvhDGA genes for an Mvh [NiFe] Hase, or next to a set of floxABCD genes (for flavin oxidoreductase) [33]. In many cases the hdrBC genes are absent (hdrA-mvhDGA or hdrA-floxABCD sets). We proposed that these proteins are part of electron-transfer pathways from oxidation of H₂ or ethanol involving energy coupling through electron bifurcation. A group of multidomain HdrA-like proteins was defined by Strittmater et al. as HdrL. These are large proteins containing an NADH binding site and, in some cases, a fumarate reductase domain [180] (Fig. 9). With a few exceptions, they are restricted to the sulfate/sulfite reducing Deltaproteobacteria and Firmicutes. It is noteworthy that some of the HdrL (and HdrA) proteins contain selenocysteine, and that there is a conserved CxxCxxCxxCxxCxxxC motif of unknown function present in all available HdrL sequences. The hdrL genes are usually found in loci together with hdrA and genes coding for a formate dehydrogenase or a pyruvate:ferredoxin oxidoreductase [33], indicating that pyruvate and formate may serve as electron donors for reduction of HdrA/L. Often an mvhD gene is found next to hdrA or fused to it (as also observed in QmoA).

Hdr proteins are also noteworthy in the genome of the acetogenic bacterium *M. thermoacetica* [183], where they are present in four gene loci including three HdrL proteins. One is an *hdrABC* locus, the other includes the *hdrLBC* genes next to the acetyl-CoA synthase (*acs*) genes, the third is an *hdrLBC-floxABCD* cluster and finally there is an *hdrDL* locus, where *hdrD* shows high similarity to *hmcF*.

HdrA and HdrD modules are also present in the benzoyl-coenzyme A reductase complex BamBCDEFGHI, present in several anaerobes capable of degrading aromatic compounds [185–187]. This large complex includes the active site subunit BamB, which contains a tungstopterin cofactor, and the iron–sulfur protein BamC that shows similarity to the electron transfer subunit of hydrogenases. BamD and BamE are

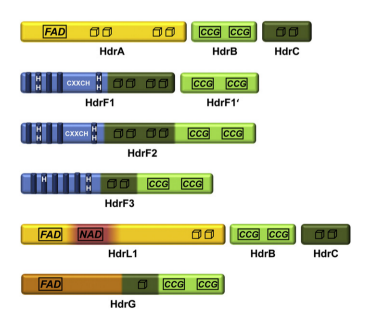


Fig. 9. Schematic representation of Hdr-related proteins. Similar colors denote sequence identity. Cubes—[4Fe-4S] clusters, CCG—CX $_n$ CCGX $_m$ CX $_2$ C sequence motif, transmembrane helices are in dark blue, H—conserved histidines, and CXXCH—possible heme c binding sequence.

HdrD- and HdrA-like proteins, while BamF shows similarity to MvhD and contains selenocysteine. BamGHI are similar to the soluble components of Complex I [187]. The BamBCDEFGHI complex is another striking example of the highly modular character of redox proteins, in this case with a quite intricate arrangement that suggests a complex mechanism.

7.2. The CCG protein family

The catalytic subunits of Hdrs (HdrB and HdrD) are characterized by the presence of the so called CCG domain (Pfam database accession number PF02754). In this domain up to five cysteines with the sequence CX_nCCGX_mCX₂C are usually found, where two tandem cysteines are followed by a glycine, which led to the designation of the CCG domain. To date more than 5000 protein sequences are present in the databases that contain either one or two CCG domains.

The family of CCG proteins can be divided into three main groups: I—proteins lacking TMH containing one or two CCG domains, and optionally additional [Fe-S] clusters; II—proteins predicted to be membrane bound; and III—proteins that contain a FAD binding site. The proteins in groups II and III are large proteins with a highly modular character as they include several distinct domains.

Group I includes the HdrB and HdrD proteins, which contain two CCG domains (Fig. 9). The C-terminal domain binds the catalytic [4Fe-4S] cluster, while the N-terminal domain provides ligands to a zinc site [188]. Many proteins in this group are membrane associated, although they lack TMH, suggesting a monotopic membrane anchoring [84,189–191]. Several of the proteins belonging to group I are part of membrane complexes in organisms capable of dissimilatory sulfur metabolism, as described above: DsrK, TmcB and HmcF. The [4Fe-4S]³⁺ characteristic EPR signal has been detected in the case of DsrK and TmcB [51,177]. The substrate of these proteins has been proposed to be DsrC.

Many group I proteins are subunits of oxidoreductases, including succinate:quinone oxidoreductase, thiol:fumarate reductase, glycolate oxidase, anaerobic glycerol-3-phosphate dehydrogenase and lactate dehydrogenase. The SdhE subunit of Sulfolobus solfataricus P2 succinate dehydrogenase has been studied in detail [190]. SdhE has two CCG domains that bind a $[4Fe-4S]^{3+/2+}$ cluster and a zinc, and serves as the monotopic membrane anchor of the enzyme. The function of the $[4Fe-4S]^{3+/2+}$ cluster is not known, but it has been speculated that it either mediates electron transfer to the quinone pool or that it has a structural role [190]. The thiol:fumarate reductase (Tfr) from Methanobacterium autotrophicum uses the thiols CoM and CoB as electron donors for the reduction of fumarate, producing succinate and CoB-S-S-CoM. TfrA is a flavoprotein carrying the catalytic site for the fumarate reduction, while TfrB is an iron-sulfur protein containing two CCG domains, that probably oxidizes the thiol substrates in analogy to Hdr [192]. CCG proteins present in E. coli are subunits of the glycolate oxidase (GlcF) and anaerobic glycerol-3-phosphate dehydrogenase subunit (GlpC). While Glc catalyzes the oxidation of glycolate to glyoxylate, Glp converts glycerol-3-phosphate into dihydroxyacetone phosphate. However, neither of these subunits has been biochemically characterized with respect to the CCG domain. A CCG protein is also a subunit of a recently described family of lactate dehydrogenases named LldEFG or LutABC [193-195] that is also widespread in SRO [33]. The BamB protein, which is part of the large benzoyl-CoA reductase complex BamBCDEFGHI discussed above [187], is also a member of this group.

Another member of the group I proteins is a subunit of the Isp [NiFe] uptake hydrogenases present in both archaea and bacteria [139,196,197]. Isp hydrogenases include two subunits (Isp1 and Isp2) similar to the HdrDE or DsrMK modules, besides the typical large and small hydrogenase subunits. The presence of an HdrD-like subunit suggests a link to sulfur metabolism, and in fact the best characterized

members of this family are from the SOB *Thiocapsa roseopersina* [197] and *A. vinosum* [196].

The CCG proteins belonging to group II are characterized by the presence of predicted TMH. This group includes the new family of HdrF proteins recently proposed by Strittmater et al. [180], based on proteins from *Db. autotrophicum* (Fig. 9). These multidomain proteins typically contain 4 to 6 predicted TMH at the N-terminus. In HdrF1 this transmembrane domain is followed by a GlpC-like and an HdrC-like domain binding four [4Fe – 4S] clusters. The *hdrF1* gene is followed by a gene similar to *hdrB* that was named *hdrF1'*. This arrangement is only found in 10 gene loci, and is restricted to deltaproteobacterial SRO. *Db. autotrophicum* contains also a fusion protein of HdrF1 and HdrF1' designated as HdrF2.

In a large number of HdrF proteins the transmembrane region is annotated as a Narl domain. In Narl, the hemes are ligated by conserved histidines located in TMH 2 and 5, respectively. These residues are conserved in the HdrF1 proteins and in each case they are predicted to be located in transmembrane helices. The distribution of the TMH along Narl is rather even. In contrast, a rather large soluble loop is predicted to be present in HdrF1 proteins. Interestingly, a conserved CXXCH motif can be found in all available sequences. Since this loop is predicted to be located in the periplasm this would allow covalently binding of heme c to this motif. If this is true, HdrF1 and HdrF2 are unique proteins containing hemes b, c and [Fe-S] clusters. However, only ten sequences are available to date limiting statistical validation of the alignments, and further experimental evidence will be needed to corroborate this.

Far more distributed is the HdrF3 protein, which is widespread amongst the *Deltaproteobacteria*, but also in other bacteria and in two archaea. It lacks the GlpC-like superdomain and contains either one or two CCG domains. Only 3 of the 4 histidines that ligate the hemes *b* in Narl are conserved. Thus, HdrF3 may bind only one heme or the second heme has an alternative coordination. However, there is another conserved histidine in helix 5, which may be another candidate ligand. The CXXCH motif present in HdrF1 is missing. In some SRO and other organisms, the gene for HdrF3 is located next to *etfAB* genes encoding an electron-transfer flavoprotein indicating that these proteins transport electrons to or from HdrF3.

The FAD-binding CCG proteins of group III are a new class of Hdr-like proteins that have not been described before and we propose to name HdrG. They are widespread among the Bacteria (in particular within the Proteobacteria and the Firmicutes), and are also present in the Euryarchaeota. These proteins include an N-terminal GlcD multidomain region fused to another GlpC multidomain region that includes either one or two CCG sites. The GlpC multidomain consists of a FAD binding region (PF01565) (not related to HdrA) that is followed by one or two FAD oxidase domains (PF02913). The glycolate oxidase D (GlcD) domain is related to several D-lactate dehydrogenases. In the Betaproteobacteria and in the Chlorobi there are HdrG proteins containing an additional N-terminal domain of unknown function (DUF3683). It appears that HdrG are most likely FAD linked oxidoreductases. In some cases the hdrG genes are located next to lactate transporters and in some cases also next to the three-subunit L-lactate dehydrogenase mentioned above. However, in the majority of cases the gene localization does not allow speculations about the protein function. As for the CCG proteins of group II there is yet no biochemically characterized HdrG protein.

In summary, it appears that the CCG domain represents a quite widespread module in redox proteins, where it may have developed different functions. These include the ligation of an [Fe-S] cluster with a catalytic or structural role, the ligation of a Zn site with a so far unknown function, and possibly others. Further research on this family is certainly needed, since only a few proteins are biochemically characterized, despite the large number of CCG proteins in the database, and so far no structural information is available. A phylogenetic analysis of this family is also warranted.

8. Conclusions

The modular nature of respiratory proteins is well apparent in proteins from SRO. In particular, several membrane-associated redox complexes from these organisms present new and interesting variations of the typical trimeric arrangement of simple respiratory proteins (catalytic, electron transfer and membrane anchor/quinone binding subunits). These variations may reflect the fact that the SRO complexes do not act directly on organic/inorganic substrates, as observed in the CISM family, but rather interact with other redox proteins, which considerably complicates in vitro studies and elucidation of their bioenergetic mechanisms. The SRO membrane complexes have a dedicated role in sulfur metabolism as they are also found in many SOB and organisms dissimilating other sulfur compounds, such as sulfite, thiosulfate and organosulfonates. Several of the proteins involved are related to subunits of heterodisulfide reductases of methanogenic archaea, which probably reflects a common ancient origin of sulfur-metabolizing organisms and methanogens, and their close environmental association. The mechanisms of energy conservation of these membrane complexes of SRO have not been clearly established, but may involve both chemiosmotic and electron bifurcation/confurcation processes that seem to be ancestral forms of energy coupling.

A subset of SRO, mainly of the *Deltaproteobacteria*, relies on multiheme cytochromes c as additional redox modules to diversify their respiratory metabolism. The prototype protein is the ${\rm Tpl}c_3$ that functions as a hub in periplasmic electron transfer pathways, with links to several membrane complexes having also a cytochrome c subunit. One of these complexes, ${\rm QrcABCD}$, is closely related to the CISM family and seems to be a cross-point in the evolution of bacterial complexes. It is an excellent example of how a different function can be achieved with a minimal modification of subunits.

Finally, the Hdr proteins, namely HdrD and HdrA, seem to be model proteins for a larger family with a wide distribution. In particular, a large group of proteins include the CX_nCCGX_mCX₂C sequence motif (CCG domain), characteristic of HdrD/HdrB. Novel proteins of the Hdr family have been proposed, namely HdrL (related to HdrA), and HdrF and HdrG (related to HdrD), which are constituted by multiple domains. The function of many of these proteins is still unknown, but their similarity to Hdrs may suggest that sulfur-based metabolic pathways may be more widespread than presently considered.

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