Metabolic flexibility of sulfate-reducing bacteria

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Dissimilatory sulfate-reducing prokaryotes (SRB) are a very diverse group of anaerobic bacteria that are omnipresent in nature and play an essential role in the global cycling of carbon and sulfur. In anoxic marine sediments sulfate reduction accounts for up to 50% of the entire organic mineralization in coastal and shelf ecosystems where sulfate diffuses several meters deep into the sediment. As a consequence, SRB would be expected in the sulfate-containing upper sediment layers, whereas methanogenic archaea would be expected to succeed in the deeper sulfate-depleted layers of the sediment. Where sediments are high in organic matter, sulfate is depleted at shallow sediment depths, and biogenic methane production will occur. In the absence of sulfate, many SRB ferment organic acids and alcohols, producing hydrogen, acetate, and carbon dioxide, and may even rely on hydrogen- and acetate-scavenging methanogens to convert organic compounds to methane. SRB can establish two different life styles, and these can be termed as sulfidogenic and acetogenic, hydrogenogenic metabolism. The advantage of having different metabolic capabilities is that it raises the chance of survival in environments when electron acceptors become depleted. In marine sediments, SRB and methanogens do not compete but rather complement each other in the degradation of organic matter. Also in freshwater ecosystems with sulfate concentrations of only 10–200 μM, sulfate is consumed efficiently within the top several cm of the sediments. Here, many of the δ-Proteobacteria present have the genetic machinery to perform dissimilatory sulfate reduction, yet they have an acetogenic, hydrogenogenic way of life. In this review we evaluate the physiology and metabolic mode of SRB in relation with their environment.

Keywords: sulfate-reducing bacteria, metabolic flexibility, syntrophy, metabolic interactions
enzymatic steps to acetate resulting in ATP synthesis (Heidelberg et al., 2004). Besides substrate-level ATP synthesis, additional ATP is generated from a proton gradient according to a chemiosmotic model (Peck, 1966), in which the protons and electrons produced during lactate oxidation react with cytoplasmic hydrogenases to form H₂, which then diffuses across the membrane where it is oxidized by periplasmic hydrogenases to form a proton gradient (Odom and Peck, 1981; Heidelberg et al., 2004). The electrons generated during lactate oxidation are channeled to sulfate through a vast network of hemes that is created by various interconnected c-type cytochromes and involving several transmembrane complexes (Aubert et al., 2000; Heidelberg et al., 2004). Nowadays, research efforts with SRB are significantly aided by the availability of over 20 genome sequences of which 12 representatives of the genus Desulfovibrio (http://img.jgi.doe.gov; Integrated Microbial Genomes). Numerous research groups have since then reported global transcriptomic and proteomic analyses of D. vulgaris under various growth or stress conditions (Chhabra et al., 2006; Clark et al., 2006; He et al., 2006; Mukhopadhyay et al., 2006, 2007; Zhang et al., 2006a,b,c; Bender et al., 2007; Tang et al., 2007; Pereira et al., 2008; Walker et al., 2009; Plugge et al., 2010). As a result, there has been a better understanding of the electron transfer and energy conservation mechanisms of D. vulgaris mainly associated with lactate oxidation during sulfidogenic growth. Yet, the energy metabolism of D. vulgaris is very complex and flexible and as such deserves further study (Pereira et al., 2008).

While the physiology of the symbiotic/syntrophic relationship has been studied for more than 40 years (Bryant et al., 1967, 1977; Stams, 1994; Schink, 1997; Stams and Plugge, 2009), relatively little is known about the metabolic and regulatory networks involved in syntrophic interactions. This may be due to the technical difficulties to establish stable mixed-culture systems and lack of analytical tools for direct large-scale measurement of various biological components (i.e., RNA, proteins or metabolites). However, availability of complete genome sequences and various functional genomics tools in recent years have provided the most needed methodologies to analyze mixed-culture systems. In a recent study, a syntrophic pair, D. vulgaris and a hydrogenotrophic methanogen Methanococcus maripaludis, was cultured syntrophically on lactate in the absence of sulfate (Stolyar et al., 2007). Syntrophic associations were initiated by mixing equal volumes of stationary phase D. vulgaris and M. maripaludis cultures. Experimental measurements with the co-culture were used to test predictions derived from the first multi-species stoichiometric metabolic model involving the two species (Stolyar et al., 2007). The D. vulgaris and M. maripaludis flux-balance models were combined to form one model describing growth and metabolite accumulation when the organisms were growing together. To model the interaction between the two species, a system of three “compartments” was proposed. The first two compartments each contained the metabolite fluxes for one of the single-species models analyzed above. These species’ compartments could each represent the action of single cells, or the combined flux of many cells of the same species. To model the interaction between the two species, a third compartment was added to the model, through which metabolites could be transferred between organisms. Exchange fluxes were added to the model in this compartment. With this modification, species could take up metabolites excreted by the other species. The results of these initial model simulations predicted that D. vulgaris, growing optimally, converts the majority of the carbon contained in the substrate lactate into acetate, and that only 4.8% of the carbon is directed to biomass. The remaining carbon is lost as CO₂, or formate. To maintain redox balance in the absence of an external electron acceptor, evolution of a reduced compound, either formate or hydrogen, is predicted. The model shows that when acetate is available, it is the preferred source of biomass carbon for M. maripaludis over the CO dehydrogenase pathway to fix carbon dioxide. This prediction is consistent with published data and the experimental results showed that M. maripaludis consumes acetate and presumably uses it as a carbon source (Stolyar et al., 2007).

Comparative transcriptomic analysis to investigate syntrophic systems was done recently. In one study, comparative transcriptional analysis of D. vulgaris in two culture conditions was performed: syntrophic co-cultures with M. maripaludis strain S2 (without sulfate) and sulfate-limited monocultures (Walker et al., 2009). During syntrophic growth on lactate with a hydrogenotrophic methanogen, numerous genes involved in electron transfer and energy generation were up-regulated in D. vulgaris compared with their expression in sulfate-limited pure cultures. In another study, whole-genome D. vulgaris microarrays were used to determine relative transcript levels when D. vulgaris shifted its lifestyle from syntrophic in a lactate-oxidizing co-culture with Methanosarcina barkeri to a sulfidogenic lifestyle (Plugge et al., 2010). In the study, syntrophic co-cultures were grown in two independent chemostats and perturbation was introduced after six volume changes with the addition of sulfate. Functional analyses revealed that genes involved in cell envelope and energy metabolism were the most regulated when comparing syntrophic and sulfidogenic metabolism. These two studies are similar in many ways, however, there are four major differences in experimental design between the two studies: (i) the methanogenic partner in the study of Walker et al. (2009) was M. maripaludis, whereas it was M. barkeri in the study of Plugge et al. (2010); (ii) the D. vulgaris and M. maripaludis co-culture and the D. vulgaris monoculture were cultivated in parallel in different chemostats by Walker et al. (2009), while a perturbation experiment by adding sulfate was performed to the chemostat co-culture to produce the D. vulgaris sulfidogenic metabolism in Plugge et al. (2010); (iii) the sulfidogenic monoculture of Walker et al. (2009) was sulfate-limited, whereas the D. vulgaris sulfidogenic metabolism was lactate-limited (Plugge et al., 2010); and (iv) the cell ratio (between D. vulgaris and M. maripaludis) during steady-state co-culture growth was higher (4:1) compared with the 1:1 in the D. vulgaris M. barkeri co-culture. Nevertheless, some similar results were obtained. These included the identification of a five-gene cluster encoding several lipo- and membrane-bound proteins which was down-regulated when cells were shifted to a sulfidogenic metabolism (Figure 1). Interestingly, this gene cluster has orthologs found only in the syntrophic bacterium Syntrophobacter fumaroxidans and four recently sequenced Desulfovibrio strains, suggesting that these genes are possible “syntrophic genes.” Both studies demonstrated that syntrophic growth and sulfate respiration use mostly independent energy generation pathways, implying that the molecular mechanism of microbial syntrophic processes cannot be fully interpreted by studying only
pure cultures (Walker et al., 2009; Plugge et al., 2010). In addition, the studies also identified the upregulation during the *D. vulgaris* syntrophic metabolism of the high-molecular-mass cytochrome complex (DVU0533, encoding Hmc protein 4), the DVU0145–0150 cellular membrane gene cluster of unknown function and heterodisulfide reductase (hdrAB), and the downregulation of genes involved in iron transport (feoB and feoA; Walker et al., 2009; Plugge et al., 2010).

Among the SRB members of the genus *Desulfovibrio* are easy to grow and they grow rapidly. Therefore, they have been the subject of the most intensive biochemical and molecular research (Postgate, 1984; Peck, 1993; Voordouw, 1993). However, multiple studies have described physiology and metabolic mode of SRB in relation with their environment. In the following paragraphs we will review a variety of these SRB-containing communities and their role in anaerobic biodegradation.

**METABOLIC FLEXIBILITY OF DESULFOTOMACULUM CLUSTER IH**

Members of the Gram-positive *Desulfotomaculum* cluster I are commonly considered as regular sulfate-reducing bacteria. However, in the last decade new representatives have been isolated that lack the ability of sulfate reduction, all phylogenetically grouping in *Desulfotomaculum* cluster Ih. These representatives are isolated from environments typically low in sulfate and producing methane: anaerobic bioreactors and rice paddy soils. Syntrophic propionate-oxidizing species of the genus *Pelotomaculum* are *Pelotomaculum schinkii*, *P. thermopropionicum*, and *P. propionicicum* (de Bok et al., 2005; Imachi et al., 2002, 2006, 2007). Additional members of *Desulfotomaculum* cluster Ih are *P. terephthalicum* sp. nov., and *P. isophthalicum* sp. nov. (Qiu et al., 2006) involved in syntrophic degradation of phthalate isomers and also lacking the ability of sulfate reduction. Imachi et al. (2006) demonstrated that none of these species was able to use sulfate, sulfite, or organosulfonates as electron acceptors. A PCR-based screening for *dsr*AB genes (key genes of the sulfate respiration pathway encoding the alpha and beta subunits of the dissimilatory sulfite reductase) of all cultures tested was negative with the exception of *P. propionicicum*. Based on these results it was proposed that subcluster Ih bacteria have adapted to anoxic, low-sulfate conditions and thus constitute a significant fraction of the *Desulfotomaculum* cluster I population in methanogenic microbial communities in a wide variety of methanogenic environments, highlighting its ecological impact in anoxic environments low in sulfate (Imachi et al., 2006; Figure 2). As a consequence of this evolutionary process, they have lost the capability of dissimilatory sulfate reduction and adopted the syntrophic life style, in close proximity to hydrogen- and formate-consuming methanogens. In such a way they can maintain an energetically favorable, low-hydrogen partial pressure that is necessary for the syntrophic oxidation of organic substrates. Given their recognized phenotypes and wide occurrence in low-sulfate, methanogenic environments, descendants of the *Desulfotomaculum* subcluster Ih branch most likely function as non-sulfate-reducing, syntrophic degraders of organic substrates in situ. This hypothesis received further support from a study that showed, by using rRNA-based stable-isotope probing, that *Pelotomaculum* species were not only dominant, but also actively involved in syntrophic propionate oxidation in a rice paddy soil (Lueders et al., 2004). Interestingly, the recently sequenced genome of *P. thermopropionicum* (Kosaka et al., 2008) points out the presence of *dsr* and *aps* genes (*aps* is coding for
benzene degradation in the studies by Kleinsteuber et al. (2008) and Herrmann et al. (2010). Benzene was completely mineralized in the presence of sulfate by a consortium consisting of syntrophs, hydrogenotrophic sulfate reducers and to a minor extent aceticlastic methanogens. Also Laban et al. (2009) showed that organisms phylogenetically related to the Gram-positive genus Pelotomaculum were responsible for benzene degradation coupled to sulfate reduction. However, the 16S rRNA gene-based sequence similarity to the next cultivated representative of Desulfotomaculum cluster Ih constituted only 95%. These sequences could be clustered between the genera Desulfotomaculum and Pelotomaculum. The similarity of the sequences of the Pelotomaculum-related phylotypes described by Kleinsteuber et al. (2008) to those identified by Laban et al. (2009) range from 88.8 to 95.7% indicating that these phylotypes are indeed different. Based on their results with a highly enriched culture, where only one dominant species was present, Laban et al. (2009) proposed that these bacteria with Pelotomaculum-related 16S rRNA gene sequences oxidize benzene directly coupled to sulfate reduction. Clearly we cannot exclude that members of the Desulfotomaculum cluster 1h are capable of sulfate reduction, but apparently they can not be easily adapted to do so. The information presented here is based on our present knowledge.

Desulfotomaculum cluster 1h related species are also involved in the anaerobic mineralization of benzene in the presence of sulfate (Kleinsteuber et al., 2008; Laban et al., 2009; Herrmann et al., 2010). Identification of the benzene-degrading, sulfate-reducing communities was in all three studies based on culture-independent methods. A Pelotomaculum/Cryptanaerobacter-like phylotype represented a syntrophic community responsible for the initiation of the anaerobic benzene degradation in the studies by Kleinsteuber et al. (2008) and Herrmann et al. (2010). Benzene was completely mineralized in the presence of sulfate by a consortium consisting of syntrophs, hydrogenotrophic sulfate reducers and to a minor extent aceticlastic methanogens. Also Laban et al. (2009) showed that organisms phylogenetically related to the Gram-positive genus Pelotomaculum were responsible for benzene degradation coupled to sulfate reduction. However, the 16S rRNA gene-based sequence similarity to the next cultivated representative of Desulfotomaculum cluster 1h constituted only 95%. These sequences could be clustered between the genera Desulfotomaculum and Pelotomaculum. The similarity of the sequences of the Pelotomaculum-related phylotypes described by Kleinsteuber et al. (2008) to those identified by Laban et al. (2009) range from 88.8 to 95.7% indicating that these phylotypes are indeed different. Based on their results with a highly enriched culture, where only one dominant species was present, Laban et al. (2009) proposed that these bacteria with Pelotomaculum-related 16S rRNA gene sequences oxidize benzene directly coupled to sulfate reduction. Clearly we cannot exclude that members of the Desulfotomaculum subcluster 1h are capable of sulfate reduction, but apparently they can not be easily adapted to do so. The information presented here is based on our present knowledge.

**METABOLIC FLEXIBILITY BY MEMBERS OF THE ORDER SYNTROPHOBACTERALES**

The Syntrophobacterales are an order of the δ-Proteobacteria, with three families, the Syntrophaceae, Syntrophobacteraeae, and Syntrophorhabdaceae (McInerney et al., 2008). Many of...
the members of these families are SRB, but also representatives of these families have been isolated that lack the ability for anaerobic sulfate respiration, or can grow as SRB or fermentative organism, depending on the environmental conditions (Wallrabenstein et al., 1994, 1995; Van Kuijk and Stams, 1995; Liu et al., 1999; Chen et al., 2005). With respect to 16S rRNA phylogeny, no predictions can be made on their metabolic flexibility.

Research has focused on propionate-degrading bacteria from the order Syntrophobacterales, specifically members that were enriched and isolated in the absence of sulfate. All species studied (Table 1) were capable of propionate degradation in syntrophic co-culture with a syntrophic partner but also as a pure culture coupled to dissimilatory sulfate reduction.

It remains unknown why some bacteria have actively expressed dsrAB genes, but cannot utilize sulfate, sulfite, and/or organosulfonates for anaerobic respiration. One might speculate that these microbes were formerly active sulfate reducers, but have lost this trait since they have to deal with low-sulfate and/or sulfite levels in methanogenic environments (Imachi et al., 2006). For this reason, the presence of dsrAB in these bacteria, which most often live in close association with hydrogen-consuming microorganisms for the syntrophic oxidation of substrates, would be a genetic remnant and thus indicative of an ancient sulfate/sulfite-respiring potential. The physiological data on syntrophs from the Syntrophobacterales order, which are abundantly present in methanogenic environments (Loy et al., 2004; Lueders et al., 2004; Stams and Plugge, 2009), indicate that all have retained their sulfate-reducing capability (Wallrabenstein et al., 1994; Van Kuijk and Stams, 1995). Syntrophic dsrAB-containing non-SRBs, syntrophic SRBs, and real sulfate reducers are phylogenetically fused, indicating an evolutionary connection between the sulfate-reducing and syntrophic lifestyle.

### Table 1 | Growth rates of selected propionate-degrading bacteria with and without sulfate.

|          | Propionate | Propionate + sulfate | References            |
|----------|------------|----------------------|-----------------------|
| Syntrophobacter fumaroxidans | 0.17       | 0.024                | Van Kuijk and Stams (1995), Hammsen et al. (1998) |
| Syntrophobacter pfeffnergii | 0.066      | 0.063                | Wallrabenstein et al. (1995) |
| Syntrophobacter sulfatireducens | n.d.*      | 0.12                 | Chen et al. (2005)     |
| Syntrophobacter wolinii | 6.7**      | 0.062                | Wallrabenstein et al. (1994) |
| Smithella propionica | n.d.*      | n.d.***              | Liu et al. (1999)      |

*Capable of syntrophic growth on propionate, but growth rate not determined.

**Only a doubling time (days) was determined.

***Capable of sulfate reduction, but growth rate not reported.

### NON-EXISTING METABOLIC FLEXIBILITY BY BUTYRATE OXIDIZERS

Bacteria that grow on butyrate syntrophically with methanogens can only be found within the Gram-positive genera *Syntrophomonas* and *Syntrophus*. None of the members of these genera has the ability to reduce sulfate. The recently sequenced genomes of *Syntrophomonas wolfei* and *Syntrophus aciditrophicus* clearly show the lack of dsr and *aps* genes which explains on a molecular basis the inability of these organisms to do so. On the other hand, Gram-positive and Gram-negative butyrate-degrading sulfate-reducing genera have been described (e.g., *Desulfovibacterium*, *Desulfoarcinina*, and *Desulfoarcinus*). None of the isolated sulfate-reducing species was ever shown to be able to grow in the absence of sulfate in syntrophy with methanogens (Rabus et al., 2006; Muyzer and Stams, 2008).

This suggests that butyrate-degrading communities have a different response to changes in sulfate availability than propionate-degrading communities. In granular sludge butyrate-degrading bacteria that grow in syntrophy with methanogens are not easily outcompeted by sulfate-reducing bacteria (Oude Elferink et al., 1994). Instead, hydrogen-consuming methanogens seem to be replaced by hydrogen-consuming sulfate reducers, while propionate-degrading communities are easily outcompeted by typical propionate-degrading sulfate-reducing bacteria. Recently, it was observed that in communities that degrade long-chain fatty acids *Syntrophomonas* sp. persisted when methanogenic sludge was exposed to increasing levels of sulfate, but the hydrogen-consuming archaea were outcompeted by hydrogen-consuming sulfate-reducing bacteria (Sousa et al., 2009).

### METABOLIC FLEXIBILITY OF SRB IN MARINE ENVIRONMENTS

In sulfate-rich marine sediments, sulfate-reducing bacteria typically use all the products of primary fermentations and oxidize them to CO₂ coupled to sulfate reduction (Muyzer and Stams, 2008). Therefore, fermentation by syntrophic communities is thought to be unimportant. However, when sulfate becomes limited the organic matter is no longer mineralized coupled to sulfate reduction, but through methanogenesis. Kendall et al. (2006) described marine syntrophic propionate- and butyrate-degrading cultures. This study implied that syntrophic communities contribute to the vast methane reservoirs in marine sediments. Microbial populations identified in shallow methanogenic sediments in the Gulf of Mexico revealed the presence of members of *Syntrophobacteriaceae* (Lloyd et al., 2006; McInerney et al., 2008). Cultured members of this family include syntrophic propionate oxidizers that can also reduce sulfate and as such have the capability to switch from sulfate-reducing life style to syntrophic life style as was discussed in paragraph “metabolic flexibility by members of the order Syntrophobacterales” (McInerney et al., 2008). Members of the *Syntrophobacteriaceae* were also found in the sulfate–methane transition zone in the Black Sea (Leloup et al., 2007, 2009). It was speculated that in these sediments, sulfate-reducing bacteria and methanogens do not compete but rather co-exist when mineralizing organic matter. In the absence of sulfate the reducing equivalents produced (hydrogen or formate) by SRB can be shuttled to a methanogen, serving as the syntrophic partner. More recently in Aarhus Bay it was found, that sulfate reducers were still very abundant in the methanogenic zones (Leloup et al., 2009). It is still...
unclear which type of metabolism sulfate reducers encompass in the methane zone and vice versa. Non-competitive substrates in marine environments (used by methanogens but not by sulfate reducers) methanol and methylamine were indeed converted by methanogens in sulfate-containing sediments from Aarhus Bay. However, sulfate reducers could scavenge the nanomolar hydrogen that leaked from the methanogenic cells (Finke et al., 2007), once more illustrating the complexity of metabolic interactions.

**EVOLUTION OF STABILITY AND PRODUCTIVITY AT THE ORIGIN OF A SYNTROPIC COMMUNITY OF AN SRB AND METHANOGEN**

A syntrophic community was established by combining the *D. vulgaris* Hildenborough with the methanogenic archaeon *M. maripaludis* S2 (Hillesland and Stahl, 2010). Both members in the community could be grown in pure culture with appropriate substrates, but under the conditions of the experiments they could grow only through syntrophic cooperation. In the absence of hydrogen and sulfate, the species feed together by cooperating to complete the following energy-yielding reaction:

$$2\text{Lactate}^- + H_2O \rightarrow 2\text{Acetate}^- + CH_4 + H^+ + HCO_3^-$$

Twenty-four independent pairings (co-cultures) of the bacterium *D. vulgaris* and the archaeon *M. maripaludis* were established and followed for 300 community doublings in two environments, one allowing for the development of a heterogeneous distribution of resources and the other not. Evolved co-cultures grew up to 80% faster and were up to 30% more productive (biomass yield per mole of substrate) than the ancestors. Unfortunately, the authors did not include quantitative data on substrate depletion, product formation, and biomass production in the evolved co-cultures to support their hypothesis.

Unlike eukaryotes that evolve principally through the modification of existing genetic information, bacteria can obtain a significant proportion of their genetic diversity through the acquisition of sequences from distantly related organisms (Scholten et al., 2007). Horizontal gene transfer produces extremely dynamic genomes in which substantial amounts of DNA are introduced into and deleted from the chromosome. These lateral transfers have effectively changed the ecological and pathogenic character of bacterial species (Ochman et al., 2000). Previous studies indicated that the scale of horizontal gene transfer may have previously been underestimated and that horizontal gene transfer may be a major force in the evolution of prokaryotic genomes (Boucher and Doolittle, 2000; Calteau et al., 2005). In a more recent study, a combination of transcriptomics and phylogenomics, codon, and amino acid usage analyses of the syntrophic interaction between *D. vulgaris* and *M. bacteri* has led to the discovery of a functionally unknown gene set in the genome of *D. vulgaris*. This may be the result of ancient horizontal gene transfer from an archaeal methanogen. The gene set consists of three closely located genes on the chromosome, DVU2103 and DVU2104 encoding iron–sulfur cluster-binding ATPase domain proteins and DVU2108 encoding an MTH1175-like domain family protein (Scholten et al., 2007). Similar lateral gene transfer was also reported between other syntrophic bacteria and methanogenic archaea (Kosaka et al., 2008; Fournier, 2009), suggesting that lateral gene transfer among microorganisms whose niches are closely associated is a common and important evolutionary process (Kato and Watanabe, 2010).

**FUTURE PERSPECTIVES**

The metabolic flexibility of syntrophs and SRBs deserves future attention. The analysis of available and future genome sequences of SRBs, syntrophs and their partners in more detail will provide more insight into horizontal gene transfer, loss of genes (such as *dsr* and *aps*), genotypic and phenotypic similarities and its consequences for their ecological specialization. Dedicated physiological studies will unravel the occurrence and significance of less obvious metabolic capabilities of microorganisms in general and of more specifically syntrophic and sulfate-reducing bacteria. Molecular ecological studies such as comparative metagenomics on environments with fluctuating sulfate levels may reveal the *in situ* metabolism of the microorganisms present.

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