Cleavage of dimethylsulfoniopropionate and reduction of acrylate by Desulfovibrio acrylicus sp. nov


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Abstract  From anoxic intertidal sediment, a dimethylsulfoxoniopropionate (DMSP)-cleaving anaerobe (strain W218) was isolated that reduced the acrylate formed to propionate. The bacterium was vibrio- to rod-shaped and motile by means of multiple polar flagella. It reduced sulfate, thiosulfate, and acrylate, and used lactate, fumarate, succinate, malate, pyruvate, ethanol, propanol, glycerol, glycine, serine, alanine, cysteine, hydrogen, and formate as electron donors. Sulfate and acrylate were reduced simultaneously; growth with sulfate was faster than with acrylate. Extracts of cells grown in the presence of DMSP contained high DMSP lyase activities (9.8 U/mg protein). The DNA mol% G+C was 45.1. On the basis of its characteristics and the 16S rRNA gene sequence, strain W218 was assigned to a new Desulfovibrio species for which the name Desulfovibrio acrylicus is proposed. A variety of other sulfate-reducing bacteria (eight of them originating from a marine or saline environment and five from other environments) did not reduce acrylate.

Key words  Dimethylsulfoxoniopropionate · Dimethylsulfide · Acrylate · Anaerobic respiration · Sulfate-reducing bacterium · Desulfovibrio acrylicus sp. nov.

Abbreviations  DMSP Dimethylsulfoxoniopropionate · DMS Dimethylsulfide

Introduction

Dimethylsulfoxoniopropionate (DMSP), an important osmolyte of many marine algae and some plants, is the major precursor of dimethylsulfide (DMS) in the marine environment. DMS is thought to play a role in climate regulation through its atmospheric oxidation products methanesulfonic acid and sulfuric acid (Charlson et al. 1987) and to act as a carrier of sulfur from the marine to the terrestrial environment (Lovelock et al. 1972). Microbial breakdown of DMS and DMSP has received considerable attention in recent years. Several aerobic marine bacteria have been shown to degrade DMSP either by cleavage to DMS and acrylate or via an initial demethylation (Kiene 1990; Taylor and Gilchrist 1991; Diaz et al. 1992; Ledyard et al. 1993; Visscher and Taylor 1994; De Souza and Yoch 1995). Relatively high concentrations of DMSP are found in marine sediments (Kiene 1988; Visscher et al. 1994). In anoxic sediments, DMSP has been found to be either demethylated to 3-mercaptopropionate, with 3-methiolpropionate as a possible intermediate, or cleaved to DMS and acrylate (Kiene and Visscher 1987; Kiene and Taylor 1988). The combined activities of marine DMSP-demethylating Desulfovibacterium strains (Van der Maarel et al. 1993) and certain 3-methiolpropionate-demethylating Methanosarcina strains (Van der Maarel et al. 1995) might be responsible for the observed conversion of DMSP to 3-mercaptopropionate. Anaerobic marine DMSP-cleaving bacteria have not been described. The only known DMSP-cleaving anaerobe, a strain of Clostridium propionicum, was isolated from freshwater (Wagner and Stadtman 1962) and fermented acrylate to propionate and acetate. This report describes an anaerobic marine DMSP-cleaving bacterium that does not ferment acrylate, but uses it only as an electron acceptor.
Materials and methods

Organisms, media, and growth conditions

Strain W218 was isolated from anoxic intertidal sediment from the Wadden Sea near Westermeland, The Netherlands. The sediment sampling and preparation of diluted sediment slurry have been described previously (Van der Maarel et al. 1995). A bicarbonate-buffered, sulfide-reduced basal medium with a N₂/CO₂ (80/20%) headspace according to Heijthuijsen and Hansen (1989) and supplemented with 0.05–1 g/l yeast extract (Difco, Detroit, Mich., USA) was used. From an enrichment culture, a pure culture was obtained by repeated plating on a sulfide-reduced agar medium with 10 mM DMSP as substrate in the presence of 1 g/l yeast extract; plating was carried out in an anaerobic chamber. Sulfate, acrylate, and substrates were added from neutralized stock solutions (1 M). Incubations were in 120-ml bottles with 50 ml medium at 28°C (for the enrichment cultures and isolation) or 30°C.

The following strains were grown in the media as described in the Deutsche Sammlung von Mikroorganismen und Zellkulturen catalogue of strains (1993): Desulfovoccus multivorans DSM 2059 (5 mM benzoate), Desulfosarcina variabilis DSM 2060 (5 mM benzoate), Desulfofexi postgatesi DSM 2034 (10 mM acetate), Desulfovibrio salexigens DSM 2638, Desulfovibrio vulgaris DSM 1744, Desulfovibrio halophilus DSM 5663, Desulfovibrio sp. strain HDv DSM 6830, Desulfitomaculum kuznetsovii DSM 6115 (60°C), Desulfobulbus propionicus DSM 2032, Desulfovibrio desulfiticans strain BH (laboratory collection) and Desulfovibrio gigas NCIMB 9332 were grown in the freshwater medium of Kremer and Hansen (1987) and Desulfovibrio sp. strain 20028 (DSM 3100) were grown in their marine medium. Desulfuromonas acetoxidans DSM 864 and Desulfuromonas sp. strain PM1 (laboratory collection) were cultivated on fumarate as described by Heijthuijsen and Hansen (1989). The incubation temperature was 30°C and the growth substrate was 20 mM lactate unless otherwise indicated.

Growth measurements

Growth rates were determined by following the increase in optical density at 660 nm in screw-capped tubes (17 ml) with 10 ml medium using a Biotron BTR 101 colorimeter. Molar growth yields of two cultures grown in 250 ml mineral medium with 0.1 g/l yeast extract plus 20 mM lactate and 20 mM sulfate or 10 mM acrylate were determined. Cultures were harvested at the end of the exponential growth phase by centrifugation (10 min at 16,000 × g), and the pellet was washed twice with 10 mM potassium phosphate (pH 7.1) containing NaCl (25 g/l) and MgCl₂:6H₂O (3 g/l).

Preparation of cell-free extracts and DMSP lyase assay

Cells were grown in the basal medium with 20 mM sulfate, 40 mM lactate, 5 mM DMSP, and 1 g/l yeast extract. Twelve hours before harvesting of the cells, an additional 3 mM DMSP was added. The following steps were performed under air: After harvesting the cells by centrifugation (10 min at 16,000 × g) and washing in 25 mM potassium phosphate (pH 7.1), a cell-free extract was prepared by three additional passages of the cells through a French pressure cell operated at 106 MPa, followed by 30 min centrifugation at 39,000 × g. DMSP lyase was assayed by measuring the absorbance increase at 220 nm due to the formation of acrylate (ε220 1.14 mM⁻¹ cm⁻¹) with a Hitachi U-1100 spectrophotometer. The assay mixture (1.0 ml) contained 25 µmol potassium phosphate (pH 7.1) and 5 µmol DMSP; the reaction was started by adding an appropriate volume of cell extract. Protein in cell-free extracts was measured according to Bradford (1976).

DNA analysis

The G+C content of the DNA was determined by the Deutsche Sammlung von Mikroorganismen und Zellkulturen (Braunschweig, Germany) by HPLC based on the method of Mesbah et al. (1989). For 16S rRNA gene sequence analysis, cells were suspended in 2 ml 10 mM Tris-HCl (pH 7.5) containing 1 mM EDTA and incubated for 30 min at 37°C with lysozyme (2 mg/ml), followed by lysis of the cells with SDS (0.5% w/v), N-lauroylsarcosine (1% w/v), and RNase (50 µg/ml). Subsequently, the lysed cell suspension was incubated for 30 min at 55°C with proteinase K (100 µg/ml). DNA was extracted and precipitated according to Sambrook et al. (1989). The 16S rRNA gene was amplified via the polymerase chain reaction (PCR) using a set of oligonucleotide primers corresponding to positions 50–68 (5′-AACACATGCAA-GTCGAACG-3′) and 1378–1392 (5′-ACGGGCCGTTGTAGAC-3′) of Escherichia coli numbering. The PCR reaction mixture (100 µl) contained 20 ng DNA, 10 pmol of each primer, 10 µl Vent® polymerase buffer (New England Biolabs, Beverly, Mass., USA), 20 pmol of each nucleoside triphosphate and 2 units of Vent® DNA Polymerase (New England Biolabs). The reaction mixture was preincubated at 94°C for 3 min, and subsequently subjected to 26 cycles of 94°C for 60 s, 55°C for 120 s, and 72°C for 120 s. In the final cycle of the PCR reaction the incubation at 72°C was extended to 300 s. The amplified 16S rRNA gene was ligated into two oligonucleotides primers complementary to conserved sequences in the 16S rRNA gene corresponding to positions 342–356 (5′-TACGGGAGGCAGACG-3′), 500–517 (5′-CACAGCAGGC-GTATAC-3′), 900–919 (5′-AACTCTAAATAATGAGTCGCCG-3′), and 1115–1130 (5′-CAAGCGAGCGACCCT-3′) of E. coli numbering. To rule out possible artifacts introduced in the 16S rRNA gene by PCR, the nucleotide sequences of two independently amplified 16S rRNA genes were determined. The nucleotide sequence was aligned manually with other 16S rRNA gene sequences derived from the NCBI collection. The sequence phylogeny was analyzed using PHYLIP 3.5c (Felsenstein 1987) and PAUP (Swafford 1993). A distance matrix was calculated with DNADIST using the two parameter model of Kimura (1980). A phylogenetic tree was constructed via the neighbour-joining method of Saitou and Nei (1987) as implemented in the NEIGHBOR program.

Restriction fragment analysis was done by incubating the 16S rRNA gene cloned in the pBlueScribe vector or PCR-amplified 16S rRNA (positions 50–1392) from a culture with one of the following restriction enzymes (1 U for 60 min): TaqI, Sau3a, Rsul, and Alul, followed by separation of the fragments on a 2% agarose gel.

Chemicals and analytical procedures

DMSP was synthesized from DMS and acrylic acid according to Chambers et al. (1987). The purity was checked by analyzing the amount of DMS or acrylate before and after alkaline treatment and by determining the amount of organic carbon present.

Acetate, propionate, and acrylate were determined after removal of the cells by centrifugation and analyzed on a Packard 437 gas chromatograph (Packard, Delft, The Netherlands) equipped with a flame ionization detector. A 1.4-m glass column (inner diameter, 2 mm) was used with Chromosorb WAW 100/120 mesh coated with 10% SP-1000 + 1% H₃PO₄ (Chrompack Nederland, Bergen op Zoom, The Netherlands). Injector and detector temperature were 220 and 240°C, respectively; a temperature program (115–140°C, with an increase of 1.3°C/min) was used. Volatile fatty acids were extracted in 0.3 ml diethyl ether after the addition of 0.2 g NaCl, 1 ml internal standard (20 mM formic acid), 0.1 ml of the ether phase was injected directly on the column. DMS was analyzed as described by Van der Maarel et al. (1993). DMSP was determined as acrylate after overnight treatment.
with 1 M NaOH. Hydrogen was determined by the method described for measuring methane (Heijthuijsen and Hansen 1989). Sulfide was measured colorimetrically (Trüper and Schlegel 1964). The presence of desulfoviridin in cell extract was determined according to Postgate (1959). Cytochromes were detected by recording an air-oxidized/dithionite-reduced difference spectrum of cell-free extract. Protein was determined according to Lowry et al. (1951) after treatment of the cell pellet with 1 M NaOH at 100°C for 10 min; before protein analysis, the pellet obtained after centrifugation and washing with 10 mM phosphate buffer (pH 7.1) containing 25 g/l NaCl and 3 g/l MgCl₂·6H₂O was treated with 5 ml methanol to remove interfering sulfur compounds.

Results

Enrichment, isolation, and general characteristics of strain W218

An enrichment culture with anoxic intertidal sediment as inoculum in mineral medium with 1.5 mM DMSP and 500 mg/l yeast extract resulted within a few days in the rapid formation of DMS and the transient accumulation of acrylate. After transfer into fresh medium with 3.5 mM DMSP, an enrichment culture was obtained that converted DMSP to DMS and acrylate, which was slowly degraded further. Microscopically, mainly vibrio- and rod-shaped cells were present. A pure culture of a DMSP-cleaving bacterium, strain W218, was obtained by using anoxic agar plates with 10 mM DMSP and 1 g/l yeast extract; colonies of strain W218 were beige. The pure culture, which grew poorly on DMSP and 1 g/l yeast extract, grew much better in basal media with lactate and sulfate and was shown to be a facultatively acrylate-reducing strain of Desulfovibrio (see below).

Cells ranged from short rods to vibrios (Fig. 1) and were motile. Electron microscopic examination revealed the presence of multiple flagella at one of the poles (Fig. 2). Motility was lost in some cases when colonies were repeatedly transferred to anoxic agar plates containing 20 mM acrylate and 2 g/l yeast extract. Spores were never observed. Cells of strain W218 stained gram-negative, behaved negative in the KOH lysis test (Gregersen 1978), and showed a gram-negative cell wall structure in electron micrographs (data not shown). The strain was catalase-negative. Desulfoviridin was detected in cell extracts. Absorption spectra of dithionite-reduced minus air-oxidized cell extract showed peaks at 422, 524, and 553 nm, which are indicative for c-type cytochromes. The G+C content of the DNA of strain W218 was 45.1 mol%.

Growth characteristics

When grown in medium containing 0.5 g/l yeast extract and DMSP, strain W218 rapidly converted DMSP to DMS. After an increase in the optical density and the concentration of propionate and a decrease in the concentration of acrylate during the first 30 h, hardly any acrylate was further metabolized (Fig. 3). However, when extra yeast extract was added after 123 h, acrylate was metabolized fur-
ther, and the optical density and the concentration of propionate increased. At the end of the incubation (262 h), 5.2 mM acrylate had been converted, and 4.6 mM acetate had been formed. These results indicated that strain W218 did not ferment acrylate, but that it might have utilized acrylate as an electron acceptor. Indeed, strain W218 was able to grow on H₂ and acrylate; under these conditions, equimolar concentrations of propionate were formed. In a medium with 1 mM acetate without yeast extract under an atmosphere of H₂/CO₂ (80/20%), the strain grew poorly, but still reduced acrylate to propionate.

In the presence of acrylate as an electron acceptor and 1 g/l yeast extract, strain W218 was able to grow on several amino acids (glycine, l-serine, l-alanine, and l-cysteine), organic acids (l-lactate, fumarate, succinate, malate, pyruvate, and formate), alcohols (ethanol, propanol, and glycerol), and H₂; organic substrates were tested at 10 mM. Lactate was oxidized to acetate. Growth was also observed with yeast extract, peptone, or casamino acids (all 1 g/l) as substrates. No growth was observed with the following compounds (all at 10 mM unless otherwise mentioned): l-threonine, l-asparagine, l-methionine, l-glutamate, l-histidine, l-arginine, l-lysine, l-aspartate, adenine, thymine, acetate, propionate, butyrate, 3-methylpropionate, 3-mercaptopropionate, betaine, choline, sarcosine, nicotinate, glycolate, methanol, 2,3-butanediol, glucose, galactose, fructose, lactose, maltose, mannose, xylose, sorbitol, cellobiose, melibiose, cellulose (1 g/l), pectin (1 g/l), xylan (1 g/l), and inulin (1 g/l). Growth on lactate plus acrylate was fastest between 30 and 37°C, at a pH of 7.4, and with an NaCl concentration of 18 g/l (range: 6–40 g/l). Other electron acceptors were tested with H₂ as an electron donor in the presence of 1 g/l yeast extract; both sulfate and thiosulfate gave good growth and were reduced to sulfide. No growth was observed with nitrate, elemental sulfur, sulfite (5 mM), crotonate, vinylacetate, acrylamide, cinnamate, caffeate, 3-methylpropionate, and 3-mercaptopropionate as potential electron acceptors; unless otherwise indicated the soluble compounds were added to 10 mM. No growth (with 1 g/l yeast extract as a substrate) was observed in stationary cultures under air.

Growth with sulfate as electron acceptor resulted in rod- to vibrio-shaped cells (usually 2–4 µm long and 0.8 µm wide). However, cells grown with acrylate became longer and tended to become spiral-shaped, as is shown for an extreme example in Fig. 2. Acrylate was reduced simultaneously with sulfate when 5 mM acrylate was added to a culture growing on 25 mM lactate and 20 mM sulfate (Fig. 4). The maximum specific growth rate on lactate and sulfate was considerably higher than on lactate and acrylate; the molar growth yield on lactate plus acrylate was slightly higher than on lactate and sulfate (Table 1). A concentration of 40 mM acrylate as electron acceptor resulted in a longer lag phase and lower growth rate and molar growth yield. Since addition of propionate (up to 40 mM) did not affect the growth characteristics, the results indicate that higher concentrations of acrylate are inhibitory.

Extracts prepared from cells grown in a medium with 20 mM sulfate and 40 mM lactate and DMSP (8 mM total, see Materials and methods) contained high activities of DMSP lyase [9.8 µmol min⁻¹ (mg protein)⁻¹]. The apparent $K_m$ for DMSP in cell extracts was approximately 1 mM. From the DMS production rate of a cell suspension, the apparent $K_m$ for DMSP of whole cells was determined to be approximately 0.4 mM.

Phylogenetic position of strain W218

The morphological and physiological characteristics and the presence of desulfoviridin indicated that strain W218 is a member of the genus Desulfovibrio. This was confirmed by a phylogenetic analysis of the 16S rRNA gene of strain W218 (positions 48–1392). Strain W218 clearly groups within the genus Desulfovibrio (Fig. 5), but shows only approximately 85% similarity with other species of

<table>
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<th>Electron acceptor</th>
<th>Sulfate</th>
<th>Acrylate</th>
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<tr>
<td>Lactate used (mM)</td>
<td>17.2</td>
<td>5.2</td>
</tr>
<tr>
<td>Acetate formed (mM)</td>
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<td>6.1</td>
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<tr>
<td>Electron acceptor used (mM)</td>
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<td>11.4</td>
</tr>
<tr>
<td>$\mu_{max}$ (h⁻¹)</td>
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<td>0.06</td>
</tr>
<tr>
<td>$Y_{mol}$ (g protein/mol substrate)</td>
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<td>2.7</td>
</tr>
<tr>
<td>Carbon recovery (%)</td>
<td>97</td>
<td>107</td>
</tr>
<tr>
<td>Electron recovery (%)</td>
<td>95</td>
<td>107</td>
</tr>
</tbody>
</table>

Table 1  Lactate oxidation with sulfate or acrylate as electron acceptor by strain W218 in media with 0.1 g/l yeast extract. Values of the molar growth yields ($Y_{mol}$) and on the substrate conversions are the means of two cultures; values of the specific growth rate ($\mu_{max}$) are the means of three cultures. Carbon and electron recoveries were calculated assuming $C_4H_7O_2N_4O_3S_0$ as empirical formula for cell material (Stouthamer 1979)
this genus. A maximum parsimony analysis using PAUP resulted in one most parsimonious tree with the same topology as the distance tree shown in Fig. 5. The sequence has been deposited with GenBank (U32578).

In view of the pleomorphic character of the culture, the purity of the culture was checked by isolation of 46 strains from individual colonies from a 10^8 dilution in lactate/sulfate agar tubes. All strains consisted of rod- to vibrio-shaped cells and reduced sulfate to sulfide upon transfer to fresh medium containing 20 mM lactate and 20 mM sulfate. When transferred to medium containing 20 mM lactate and 20 mM acrylate, all strains consisted of rod-shaped to small spiral-shaped cells and reduced the acrylate to propionate. The restriction fragment pattern of PCR amplified 16S rDNA from a culture on lactate/sulfate was identical to the pattern of the cloned 16S rRNA gene; both patterns matched the expected pattern based on the sequence information completely.

Acrylate reduction by other sulfate-reducing bacteria?

A wide variety of sulfate-reducing bacteria from different habitats and two sulfur-reducing strains were tested for the ability to reduce acrylate (see Materials and methods for a list of strains). None of the 15 strains was able to grow with the combination of their original growth substrate and acrylate as an electron acceptor.

Discussion

Strain W218 is the first example of an anaerobic marine bacterium that is able to cleave DMSP. It has a higher DMSP lyase level than the aerobic Alcaligenes sp. strain M3A and Pseudomonas doudoroffii (de Souza and Yoch 1995). Unlike the DMSP-cleaving strain of Clostridium propionicum (Wagner and Stadtman 1962) and the acrylate-utilizing Clostridium propionicum strain 19acyr3 (Janssen 1991), strain W218 does not ferment acrylate, but reduces it to propionate with a suitable electron donor. Kiene and Taylor (1988) found that after the addition of DMS to low micromolar concentrations, DMS and acrylate did not accumulate. They suggested that fermenting organisms like C. propionicum could be responsible for the cleavage of DMS and the degradation of acrylate. Another indication for the involvement of fermenting organisms in the degradation of acrylate came from the observation of Kiene and Taylor (1989) that addition of micromolar levels of acrylate to sediment suspensions in the presence of molybdate as inhibitor of sulfate reduction resulted in the formation of acetate and propionate in a 1:2 ratio. The nature and the quantitative importance of fermenting bacteria in the degradation of DMSP remains to be examined.

Strain W218 is a sulfate-reducing bacterium that can also carry out the reduction of acrylate to propionate. This reduction can be considered as a novel type of anaerobic respiration. The ability to grow by the reduction of an unsaturated organic compound has been found for a variety of organisms. Acetobacterium woodii is able to reduce the acrylic side chain of caffeate (3,4-dihydroxyphenylacrylate) with H_2 and to couple this process to the synthesis of ATP (Tschech and Pfennig 1984; Hansen et al. 1988). The most common and best understood type of anaerobic respiration with an unsaturated organic compound is the reduction of fumarate (Kröger et al. 1992). Organisms such as Wolinella succinogenes, Escherichia coli, and Desulfovibrio gigas are able to grow by the reduction of fumarate. The fumarate reductase is localized at the cytoplasmic side of the cell membrane and converts fumarate to succinate without energy-rich intermediates (Kröger et al. 1992). It would be interesting to know whether the mechanism of acrylate reduction in strain W218 is similar to the mechanism of fumarate reduction or to the propionate-forming pathway in bacteria with a non-randomiz-

Fig. 5 Phylogenetic relationship of Desulfovibrio acrylicus W218 with other sulfate-reducing bacteria. The tree is based on a distance analysis of the 16S rRNA gene sequence (positions 48–1392) and was constructed using the neighbour-joining method as implemented in PHYLIP 3.5c. Bar 0.05 substitutions per sequence position.
ing propionic acid fermentation, where acrylyl-CoA is reduced (Brockman and Wood 1975).

Acrylate reduction is energetically more favorable than sulfate reduction: per mol of hydrogen consumed the ΔG°
values are −75 kJ/mol and −38 kJ/mol, respectively (calculated from Thauer et al. 1977). At limiting electron donor concentrations, this thermodynamic advantage may be translated into a competitive advantage for the acrylate-reducing strain. This may explain why a sulfate-reducing bacterium is able to reduce acrylate in an environment where sulfate is present in high concentrations (up to 28 mM). The molar growth yield with acrylate as an electron acceptor was slightly higher than with sulfate. This was expected because in the activation of 1 mol of sulfate, 1 mol of ATP is eventually converted to 1 mol of AMP.

On the basis of its morphology, physiology, and 16S rRNA gene sequence, strain W218 clearly is a member of the genus *Desulfovibrio*. Devereux et al. (1990) concluded that *Desulfovibrio* comprises a coherent, but phylogenetically diverse group of species that is distinct from other groups of sulfate-reducing bacteria and proposed to create the family Desulfovibrionaceae, which includes at least five different lineages. Strain W218 does not group closely within one of these lineages and could represent a new lineage within the family. We propose to place strain W218 in the genus *Desulfovibrio* as the type strain of the new species *Desulfovibrio acrylicus*.

Description of *Desulfovibrio acrylicus* sp. nov.

*Desulfovibrio acrylicus* (acr‘yl‘icus L. adj. derived from *acidum acrylicum* acrylic acid). Small rods to vibrios; 2–4 µm long and 0.8 µm wide when grown with sulfate, but longer when grown with acrylate. Motile by multiple polar flagella; no spores formed. Gram-negative and catalase-negative. Strictly anaerobic. H₂, formate, lactate, fumarate, succinate, malate, pyruvate, ethanol, propanol, glyceral, glycerine, serine, alanine, and cysteine serve as electron donor; sulfate, thiosulfate, or acrylate used as electron acceptor. Growth optimal at pH 7.4, a temperature of 30–37°C, and 18 g/l NaCl. Contains desulfoviridin and cytochromes of the c-type. Forms dimethylsulfide from dimethylsulfiniopropionate. G+C content of the DNA is 45.1 mol%. Habitat anoxic marine sediments. Type strain W218 deposited with the Deutsche Sammlung von Mikroorganismen und Zellkulturen (DSM 10141). The 16S rRNA gene sequence has been deposited with GenBank (U32578).

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References


