Thialkalivibrio nitratireducens sp. nov., a nitrate-reducing member of an autotrophic denitrifying consortium from a soda lake

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Strain ALEN 2T was isolated from a mixed culture capable of complete autotrophic denitrification with thiosulfate as electron donor at pH 10; the mixed culture was enriched from sediment from Lake Fazda (Wadi Natrun, Egypt), a hypersaline alkaline lake. The isolate had large, non-motile, coccoid or barrel-shaped cells with intracellular sulfur globules. The bacterium was obligately chemolithoautotrophic. It grew with reduced sulfur compounds aerobically and anaerobically with nitrate as electron acceptor, nitrate being reduced to nitrite. It was moderately halophilic and obligately alkaliphilic. On the basis of genetic analysis and its unique phenotype, strain ALEN 2T (=DSM 14787T = UNIQEM 213T) is proposed as the type strain of a novel species of the genus Thialkalivibrio, Thialkalivibrio nitratireducens.

Only a few species of lithotrophic sulfur-oxidizing bacteria (SOB) are capable of anaerobic growth with sulfur compounds and nitrogen oxides as electron acceptors. In particular, obligately autotrophic Thiobacillus denitrificans, Thiomicrospira denitrificans and facultatively autotrophic Paracoccus species perform the complete denitrification of nitrate to nitrogen gas, whereas filamentous, vacuolated, SOB Beggiatoa and Thioptea species appear to conduct dissimilatory nitrate reduction to ammonia with sulfide as electron donor (Kuenen et al., 1992; McHatton et al., 1996; Otte et al., 1999). Such SOB play an important role in mineral cycling and waste removal by linking the sulfur and nitrogen cycles (Kuenen & Robertson, 1987; Robertson & Kuenen, 1992). Of the haloalkaliphilic SOB isolated from soda lakes, currently represented by nearly 100 isolates (Sorokin et al., 2000, 2001a, 2002a, 2002b), only one, described as Thialkalivibrio denitrificans ALJD7, has the potential for anaerobic growth. This strain differs from its neutrophilic analogues Thiobacillus denitrificans and Thiomicrospira denitrificans in that it does not possess a dissimilatory nitrate reductase. It grows with nitrite or N2O as electron acceptor and thiosulfate or polysulfide as electron donor (Sorokin et al., 2001b). Recently, an enrichment culture has been obtained from a sediment sample from an alkaline hypersaline lake in Wadi Natrun (Egypt) that completely reduced nitrate to nitrogen gas, with thiosulfate as electron donor, at pH 10. In this report, the nitrate-reducing member of this haloalkaliphilic autotrophic denitrifying association is described.

Sediment from Lake Fazda, a hypersaline soda lake (250 g total salts l−1; pH 10) in the Wadi Natrun (Egypt), was used as inoculum to enrich for denitrifying SOB. The basic hydrochemical and microbiological characteristics of the Wadi Natrun lakes have been described by Taher (1999) and Imhoff et al. (1979), respectively. The enrichment was performed in 100 ml serum bottles with butyl-rubber stoppers filled with 50 ml alkaline base containing 0·6 M total Na+; pH 10 (Sorokin et al., 2001a), with 20 mM thiosulfate and 30 mM nitrate. Anoxic conditions were achieved by five cycles of evacuation–flushing with argon with active degassing of the liquid. Solid medium was prepared by 1:1 mixing of the above-mentioned alkaline base containing a double concentration of substrates and 4% (w/v) agar at 50 °C. Anaerobic plate incubation was performed using closed jars filled with pure argon in the presence of anaerobic catalyst (Oxoid). Growth with H2 as electron donor was tested in 100 ml bottles closed with butyl-rubber stoppers containing 10 ml medium under an

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Abbreviation: SOB, sulfur-oxidizing bacteria.

The GenBank accession number for the 16S rDNA sequence of Thialkalivibrio nitratireducens ALEN 2T is AY079010.
atmosphere of 98% (v/v) H₂ and 2% (v/v) O₂. Methylotrophy was tested with methanol and formate (5 mM) under aerobic and denitrifying conditions. The pH range for growth was tested on media prepared using 0·1 M HEPES/Na₂HPO₄ (pH 7–8) supplemented with 0·6 M NaCl or sodium carbonate/sodium bicarbonate (pH 8–11) with 0–6 M total Na⁺. Na⁺ tolerance was tested on sodium carbonate/bicarbonate-based mineral medium containing 0–1–2·0 M total Na⁺ at pH 10. The same pH and salinity buffers were used in experiments with washed cells. Respiration activity, reduction of nitrogen oxides by washed cells, sulfur-metabolizing activity, denitrifying enzymes and cytochrome spectra were measured as described previously (Sorokin et al., 2002c). Inorganic sulfur compounds, nitrate, nitrite, N₂O and protein concentrations were determined by spectrophotometric and GC methods as described previously (Sorokin et al., 2001a, b). DNA extraction, DNA G + C content determination and DNA–DNA hybridization were performed according to standard protocols (Marmur, 1961; De Ley et al., 1970). For amplification and sequencing of the 16S rRNA gene, DNA was obtained by standard phenol/chloroform extraction. The 16S rRNA gene was selectively amplified using primers 5'-AGAGTTTGATCC-TGGCTCA-G-3' (forward) and 5'-TACGGTATCC-TG-ACGACTT-3' (reverse). PCR products were purified from low-melting-point agarose using the Wizard PCR-Prep kit (Promega) according to the manufacturer’s instructions. Almost-complete sequencing (1420 nt) was performed using the Promega Silver Sequencing kit according to the manufacturer’s instructions, but with minor modifications. Primary comparative analysis of the 16S rRNA gene sequence of strain ALEN 2T with database sequences was done using BLAST. On the basis of results of the BLAST search, the 16S rRNA gene sequences of strain ALEN 2T and its closest relatives were aligned using CLUSTAL_X (Thompson et al., 1997). Regions that were not sequenced in one or more reference organisms were omitted from subsequent analyses. An unrooted phylogenetic tree based on 16S rRNA gene sequences of the studied bacteria was constructed by the neighbour-joining method available in the TREECON package (Van de Peer & De Wachter, 1994). Bootstrap analysis (100 replications) was used to validate reproducibility of the branching pattern of the tree.

The anaerobic enrichment resulted in a stable binary culture consisting of large irregular cocccoid cells with intracellular sulfur globules and thin straight rods that were occasionally motile. The latter were responsible for nitrate reduction. On the basis of their physiology and genetic properties, the thin straight rods resembled the previously described haloalkaliphilic SOB Thialkalivibrio denitrificans (Sorokin et al., 2001b). Further work focused on isolation and characterization of the unusual cocccoid morphotype that was responsible for the reduction of nitrate to nitrite.

The cocccoid organism was present in relatively low numbers in the mixed denitrifying culture, which made it difficult to isolate. When plated onto nitrate/thiosulfate alkaline agar under an argon atmosphere, a surprisingly low proportion of the total cells present in liquid culture produced colonies. It appeared that the numerically dominant small rods depended on the cocccoid phenotype for growth, resulting in formation of mixed colonies. By picking up the colony material containing mostly the cocccoid morphotype and replating it, the colony number of the desired organism gradually increased. Eventually, pure colonies of the cocccoid phenotype were obtained, and the isolated bacterium was designated strain ALEN 2T. It formed colonies of variable size and shape, 1–3 mm, some flat, some dome-like. The young colonies were shining white and full of sulfur, turning reddish and transparent with time. Cells in the colonies were extremely pleomorphic, mostly cocccoid, with multiple intracellular sulfur globules. Cells grown aerobically on liquid medium with thiosulfate at pH 10 were cocccoid, 0·8–2·0 μm in diameter and aggregated in chains of different lengths (Fig. 1a, c). The cells grown anaerobically in liquid culture were barrel-shaped and less aggregated (Fig. 1b, d).

The bacterium was obligately autotrophic. It grew well under fully aerobic conditions, oxidizing 40 mM thiosulfate at pH 10 within 3 days with prominent formation of intracellular sulfur (8–10 mM), which was finally oxidized to sulfate within another 3–5 day period. Maximum specific growth rate with thiosulfate under aerobic conditions was 0·08 h⁻¹, with a yield of 5·5–6·0 mg protein mmol⁻¹. The bacterium belonged to the obligate alkaliphiles (pH range for growth of 8·0–10·5, with optimum at 9·5–10·0) and moderate halophiles (salt range for growth of 0·2–1·5 M total Na⁺, with an optimum at 0·4–0·5 M). Respiration activity of the washed cells with sulfate and thiosulfate was maximal at pH 10 and still substantial at pH 11 (20% from maximum; data not shown). Strain ALEN 2T actively oxidized sulfide and, at much lower rates, polysulfide and thiosulfate. The oxidation activity of elemental sulfur was an order of magnitude lower than that of thiosulfate (Table 1), which accounts for a heavy sulfur accumulation in cultures.

In static aerobic cultures with thiosulfate, strain ALEN 2T reduced up to 15 mM nitrate to nitrite. Full aeration inhibited nitrite production. Under anaerobic conditions, the bacterium was able to grow with nitrate as electron acceptor and thiosulfate, sulfide and polysulfide (2 mmol 1⁻¹ portions, fed-batch supply for the latter two) as substrates. With 20 mM thiosulfate/40 mM nitrate, rapid nitrite production and transient sulfur accumulation were observed. When all the nitrate was consumed, growth was arrested. Finally, 38 mM nitrite was produced per 11 mM thiosulfate oxidized to sulfate, which corresponds to a two-electron nitrate reduction to nitrite (assuming that approximately 10% electrons are used for CO₂ reduction). No anaerobic growth was observed with either nitrite (10 mM) or N₂O as electron acceptors. The fed-batch growth with sulfide/nitrate proceeded in two phases. At first, when nitrate was still present, each 2 mM addition of sulfide was followed by cell growth, transient sulfur formation with its further
oxidation to sulfate and a build-up in nitrite concentration, similar to growth with thiosulfate. When nitrate was consumed, biomass increase stopped, but sulfide was still oxidized to elemental sulfur, accompanied by slow nitrite consumption with a molar HS\(^{-}/\)NO\(_2\)^{2-} ratio of 2.0–2.5:1. The latter implies a one-electron reduction of nitrite to NO. Although NO was not measured, at this stage some N\(_2\)O in the headspace was registered. However, no anaerobic growth was observed with sulfide and nitrite or N\(_2\)O as electron acceptors. From these experiments, it was concluded that strain ALEN \(2^T\) is a high-capacity nitrate-to-nitrite reducer and, therefore, may serve as a nitrite provider in a denitrifying association with a nitrite reducer. Further experiments with washed cells of strain ALEN \(2^T\) confirmed its inability to reduce nitrite and N\(_2\)O with thiosulfate as electron donor. However, low nitrite-reducing activity was observed with polysulfide and H\(_2\) as electron donors (Table 2). In both cases, N\(_2\)O was produced in the gas phase. Moreover, H\(_2\) also stimulated the reduction of elemental sulfur by washed cells of strain ALEN \(2^T\). Despite these facts, the bacterium was incapable of anaerobic growth with H\(_2\) as electron donor and nitrite or elemental sulfur as electron acceptors, which implies that some important links between the dissimilatory enzymes and the energy-generating system are missing in this unusual alkaliphilic SOB species.

In the soluble fraction of cell-free extract of strain ALEN \(2^T\), rapid nitrate-dependent oxidation of reduced methyl viologen was observed, but nitrite, the usual product of nitrate reductase activity, was not detected. Replacement of nitrate by nitrite in the reaction mixture resulted in extremely high rates of methyl viologen oxidation and nitrite consumption [8.0–10.5 μmol NO\(_2\) (mg protein)\(^{-1}\) min\(^{-1}\)].

Table 1. Respiratory activity of washed cells of strain ALEN \(2^T\) grown aerobically at pH 10 and 0.6 M Na\(^+\) with thiosulfate

<table>
<thead>
<tr>
<th>Substrate (50 μM)</th>
<th>Respiration rate [nmol O(_2) (mg protein)(^{-1}) min(^{-1})]</th>
<th>Products</th>
</tr>
</thead>
<tbody>
<tr>
<td>HS(^{-})</td>
<td>2810</td>
<td>Sulfur</td>
</tr>
<tr>
<td>S(_2)(^{3-})</td>
<td>430</td>
<td>Sulfur</td>
</tr>
<tr>
<td>S(_8)</td>
<td>32</td>
<td>ND</td>
</tr>
<tr>
<td>S(_2)O(_4)(^{-})</td>
<td>175</td>
<td>Sulfur + sulfate</td>
</tr>
<tr>
<td>S(_4)O(_3)(^{-})</td>
<td>38</td>
<td>ND</td>
</tr>
<tr>
<td>SO(_3)(^{-})</td>
<td>8</td>
<td>Sulfate</td>
</tr>
</tbody>
</table>

ND, Not determined.
This implies that strain ALEN 2T possesses both nitrate and nitrite reductase activities despite the fact that it is unable to use nitrite as an electron acceptor \textit{in vivo}. Antipov et al. (2003) have demonstrated the presence of a nitrate/nitrite reductase in strain ALEN 2T with unusual properties. Activities of the sulfur-metabolizing enzymes thiosulfate reductase, sulfite dehydrogenase and sulfide dehydrogenase were found in the soluble fraction of cell-free extract of strain ALEN 2T at pH 10 \([690, 260 \text{ and } 60 \text{ nmol (mg protein)}^{-1} \text{ min}^{-1}]\). The cell membranes of strain ALEN 2T contained high amounts of cytochrome types \(c\) and \(b\) (absorption maxima in the alpha region at 554 and 558 nm, respectively). CO difference spectra of the membranes indicated the presence of a cytochrome oxidase of type \(bb\) (troughs in alpha region at 558 and 563 nm).

The G + C content of the DNA of strain ALEN 2T was 64.8 ± 0.5 mol%. Phylogenetic analysis placed this bacterium in the genus \textit{Thialkalivibrio} (Fig. 2), which accommodates the high-G + C-containing species of haloalkaliphilic SOB (Sorokin et al., 2001a). The isolate had highest similarity (98.2 %) to the thiocyanate-utilizing species \textit{Thialkalivibrio paradoxus} (Sorokin et al., 2002b). Indeed, strain ALEN 2T resembled this species in its specific cell morphology and accumulation of intracellular sulfur due to a low elemental sulfur oxidizing capacity. These two strains also had the highest level of DNA–DNA relatedness (54 %) compared to values obtained with other species of \textit{Thialkalivibrio} (<30 %). However, strain ALEN 2T was substantially different physiologically from \textit{Thialkalivibrio paradoxus}: strain ALEN 2T was incapable of thiocyanate and carbon disulfide oxidation, whereas \textit{Thialkalivibrio paradoxus} cannot utilize nitrate either as electron acceptor nor as nitrogen source. Overall, these differences suggest that strain ALEN 2T should be regarded as a novel species of the genus \textit{Thialkalivibrio}, for which the name \textit{Thialkalivibrio nitratireducens} is proposed.

### Description of \textit{Thialkalivibrio nitratireducens} sp. nov.

\textit{Thialkalivibrio nitratireducens} (ni.tra.ti.re.du’en.s. N.L. n. \textit{nitras} nitrate; L. part. adj. \textit{reducens} converting to a different state; N.L. adj. \textit{nitratireducens} reducing nitrate).

Cells are mostly coccoid or barrel-shaped, often in chains and aggregates, 0.8–2.0 \(\mu\)m in diameter, often with large inclusions of sulfur.
sulfur globules inside. Non-motile. Colonies are up to 3 mm in size, full of sulfur, turning reddish with age. Obligately chemolithoautotrophic. Oxidizes thiosulfate, sulfide, polysulfide and, much less actively, elemental sulfur and tetrationionate to sulfate. Facultatively anaerobic. Grows anaerobically with nitrate as electron acceptor and thiosulfate, sulfide or polysulfide as electron donor. The sole product of nitrate reduction is nitrite. Obligately alkaliphilic and moderately halophilic. Genetically most closely related to a thiocyanate-oxidizing species, *Thialkalivibrio paradoxon*.

The type strain is ALEN $2^\dagger$ ($\approx$ DSM 14787$^T$ $=\text{UNIQUEM}$ 213$^T$), isolated from sediments of Lake Fazda (Wadi Natrun, Egypt), a hypersaline soda lake. Its DNA G+C content is $64\pm0.5$ mol% ($T_m$ method). Other properties are as for the genus.

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**References**


