Haloplanus natans gen. nov., sp. nov., an extremely halophilic, gas-vacuolate archaean isolated from Dead Sea–Red Sea water mixtures in experimental outdoor ponds

Rahel Elevi Bardavid, Lily Mana and Aharon Oren

The Institute of Life Sciences and The Moshe Shilo Minerva Center for Marine Biogeochemistry, The Hebrew University of Jerusalem, 91904 Jerusalem, Israel

To study biological phenomena in the Dead Sea and to simulate the effects of mixing Dead Sea water with Red Sea water, experimental mesocosms were operated at the Dead Sea Works at Sedom, Israel. Dense communities of red halophilic archaeeae developed in mesocosms filled with 80% Dead Sea water and 20% Red Sea water after enrichment with phosphate. The most common type of colonies isolated from these brines belonged to the genus Halorubrum. A few white–pinkish opaque colonies contained pleomorphic flat cells with gas vesicles. Three strains isolated from the latter colonies were characterized in depth. Their 16S rRNA gene sequences showed only 91% similarity to the closest cultured relative (Halofex mediterranei), indicating that the new strains represent a novel species of a new genus. The name Haloplanus natans gen. nov., sp. nov. is proposed for this novel organism. The type strain of Haloplanus natans is RE-101\(^T\) (= DSM 17983\(^T\) = JCM 14081\(^T\)).

The Dead Sea, located at the lowest point of the Syrian–African Rift Valley, is a unique, athalassohaline, salt-saturated lake with extremely high divalent cation concentrations. Its waters contain about 1.9 M Mg\(^{2+}\), 1.6 M Na\(^+\), 0.44 M Ca\(^{2+}\), 0.20 M K\(^+\), 6.35 M Cl\(^-\), 0.07 M Br\(^-\) and 0.005 M SO\(_4^{2-}\). The green alga Dunaliella sp. is the sole primary producer in the lake. Currently, the Dead Sea environment is too extreme to support extensive microbial life, but whenever the upper water layers become diluted by massive rain floods in winter, the algae multiply rapidly, followed by development of dense blooms of halophilic archaeeae: up to 1.9 x 10\(^7\) cells ml\(^{-1}\) and 3.5 x 10\(^7\) cells ml\(^{-1}\) were observed in 1980 and 1992, respectively. A number of novel species of the Halobacteriaeaceae have been isolated in the past from the lake: Haloferax volcanii, Haloarcuca marismortui, Halorubrum sodomense and Halobaculum gomorrense (Oren, 1999, 2000).

In the framework of plans to construct a water conduit between the Red Sea and the Dead Sea (the ‘Peace Conduit’), simulation experiments were performed on the grounds of the Dead Sea Works at Sedom, Israel, to study the effect of mixing Dead Sea water and Red Sea water. These experiments are intended to provide information on the microbiological properties of the Dead Sea when the ‘Peace Conduit’ plans are implemented and massive quantities of Red Sea water will enter the Dead Sea and lower the salinity of the upper water layers. Previous experiments have shown that phosphate is the limiting inorganic nutrient in the Dead Sea (Oren & Shilo, 1985), and therefore the effect of phosphate addition was investigated as well. Experimental mesocosms (0.9 m\(^3\)) filled with 80% Dead Sea water, 20% Red Sea water and enriched with low concentrations of phosphate became strongly red within 1–2 months as a result of dense communities of halophilic archaeeae (Oren et al., 2004).

We plated 0.1 ml samples of brine from these mesocosms and of dilutions in sterile liquid medium on the following media (contents per litre): (I) 175 g NaCl, 20 g MgCl\(_2\).6H\(_2\)O, 5 g K\(_2\)SO\(_4\), 0.1 g CaCl\(_2\).2H\(_2\)O and 5 g yeast extract, pH 7; and (II) 206 g NaCl, 36 g MgSO\(_4\).7H\(_2\)O, 0.373 g KCl, 0.5 g CaCl\(_2\).2H\(_2\)O, 0.013 mg MnCl\(_2\).4H\(_2\)O and 5 g yeast extract, pH 7. Media were solidified with 2% agar. Most colonies (99% at least) that appeared after 3 weeks incubation at 37 °C were translucent and red; preliminary 16S rRNA gene sequencing indicated that these organisms were affiliated with the genus Halorubrum. In addition, a few white–pinkish opaque colonies were obtained, and these comprised pleomorphic flat cells with gas vesicles. Three

The GenBank/EMBL/DDBJ accession numbers for the 16S rRNA gene sequences of strains RE-101\(^T\), RE-102 and RE-103 are DQ417339–DQ417341, respectively.

A phylogenetic tree based on 16S rRNA gene sequences that includes environmental sequences showing a high degree of similarity to strains RE-101\(^T\), RE-102 and RE-103 is available as supplementary material in IJSEM Online.

Correspondence
Aharon Oren
oren@cc.huji.ac.il

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representative strains from these latter colonies were isolated, designated RE-101\textsuperscript{T}, RE-102 and RE-103. Strains RE-101\textsuperscript{T} and RE-102 were grown routinely in medium (I), whereas strain RE-103 grew somewhat better in medium (II), i.e. at a slightly higher salt concentration. In view of their extremely halophilic nature, the isolates may be assumed to have been derived from the Dead Sea (340 g total dissolved salts l\textsuperscript{-1}) and not from the Red Sea (41 g total dissolved salts l\textsuperscript{-1}).

Cell morphology was examined via a Zeiss Axiovert microscope equipped with phase-contrast optics. Tests for phenotypic properties were carried out in accordance with the proposed minimal standards for the description of new taxa in the order \textit{Halobacteriales} (Oren \textit{et al.}, 1997). Most tests were performed as outlined by Holding & Collee (1971). Appropriate positive and negative controls were included in all tests; test organisms included, among others, the type strains of \textit{Haloferax volcanii}, \textit{Haloferax mediterranei}, \textit{Haloarcula marismortui}, \textit{Haloarcula vallismortis}, \textit{Halogeometricum borinquense} and \textit{Natrialba asiatica}. Growth and gas formation with nitrate as electron acceptor were tested in 15-ml stoppered bottles, completely filled with growth medium to which NaNO\textsubscript{3} (5 g l\textsuperscript{-1}) had been added, and containing an inverted test tube. The formation of nitrite was monitored colorimetrically. Anaerobic growth in the presence of L-arginine hydrochloride (5 g l\textsuperscript{-1}) was tested in completely filled 15-ml stoppered tubes. Controls without arginine were included, and incubations were performed in the dark; \textit{Halobacterium salinarum} R1 (=DSM 671) served as the positive control. Hydrolysis of starch was examined on agar plates supplemented with 2 g soluble starch l\textsuperscript{-1}. Starch hydrolysis was detected by flooding the plates with iodine solution. Gelatin hydrolysis was determined by growing colonies on agar plates amended with 0.1 % gelatin and flooding the plates with a solution of 15 % (w/v) HgCl\textsubscript{2} in 20 % (w/v) HCl after growth was established. Hydrolysis of Tween 20 was tested as outlined by Gutiérrez & González (1972). Starch, gelatin and Tween 20 were each added at a concentration of 1 ml l\textsuperscript{-1} to autoclaved medium supplemented with 1 g CaCl\textsubscript{2}.2H\textsubscript{2}O l\textsuperscript{-1}. Indole production was detected with Kovacs’ reagent after having grown the cells in media supplemented with 0.1 g L-tryptophan l\textsuperscript{-1}. To test for growth on single carbon sources, yeast extract was omitted from the medium and the compound to be tested was added at a concentration of 5 g l\textsuperscript{-1}, together with 1 g NH\textsubscript{4}Cl l\textsuperscript{-1} and 1.36 g KH\textsubscript{2}PO\textsubscript{4} l\textsuperscript{-1}.

Cells grown in liquid culture (2 weeks with shaking at 37 \textdegree C) were flat, extremely pleomorphic, 2.5–8 \textmu m in size and contained numerous gas vesicles (Fig. 1). Colonies were light pink in colour due to the presence of bacterioruberin pigments. The three new strains were extremely halophilic; optimum growth was observed at 3 M NaCl (range for growth 2.6–4.3 M), at pH 7.0 (range pH 6.5–8.0) and at about 40 \textdegree C (range 37–52 \textdegree C). All three strains were catalase- and oxidase-positive. Gelatin, starch and Tween 20 were not hydrolysed. They were able to reduce nitrate to nitrite. Anaerobic growth on nitrate and L-arginine was not observed. Acid was not produced from glucose, fructose, sucrose or maltose. The strains differed in indole production and in the range of single carbon sources supporting growth. The phenotypic properties of the three strains are given in the species description below, and properties for which the three isolates differed are shown in Table 1.

### Table 1. Differential characteristics of strains RE-101\textsuperscript{T}, RE-102 and RE-103

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>RE-101\textsuperscript{T}</th>
<th>RE-102</th>
<th>RE-103</th>
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<tr>
<td>Formation of indole</td>
<td>+</td>
<td>+</td>
<td>–</td>
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<td>Growth on single carbon sources:</td>
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<td>Sucrose</td>
<td>–</td>
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<td>Succinate</td>
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<td>Glycerol</td>
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<td>Glutamate</td>
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<td>Maltose</td>
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<td>Galactose</td>
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Polar lipids were extracted and analysed according to Oren et al. (1996). One- and two-dimensional TLC of the polar lipid fraction revealed that all three strains contained diphytanyl derivatives of phosphatidylglycerol, phosphatidylglycerol phosphate methyl ester, phosphatidyl glycerol sulfate and one major glycolipid chromatographically identical to S-DGD-1 previously reported for members of the genus *Halofex*.

For determination of the DNA base composition and DNA–DNA hybridization experiments, genomic DNA was extracted according to Cashion et al. (1977). The DNA G + C content was determined by HPLC of deoxribonucleosides (Mesbah et al., 1989; Tamaoka & Komagata, 1984). The DNA G+C content of strains RE-101T, RE-102 and RE-103 was 66.2, 66.4 and 66.1 mol%, respectively. DNA–DNA hybridization analyses were performed according to the renaturation method of De Ley et al. (1970) as modified by Huß et al. (1983). Mean levels of DNA–DNA relatedness, based on duplicate assays in 2 × SSC plus 10% formamide at 69 °C, were as follows: between strain RE-101T and RE-102, 78.1 ± 0.5%; RE-101T and RE-103, 81.1 ± 2.5%; and RE-102 and RE-103, 93.4 ± 2.3%. For amplification of the 16S rRNA gene, genomic DNA was extracted by using EZ-DNA-Genomic DNA isolation reagent (Biological Industries). The 16S rRNA gene was amplified via PCR by using primers 21F and 1492R (Martinez-Murcia et al., 2003). Sequencing was performed in the PCR by using the Big Dye Terminator reagent and the purified PCR products were electrophoresed on an ABI 373A DNA sequencer.

The almost-complete 16S rRNA gene sequences (1398–1400 bp) of the three new strains were determined and compared with sequences of members of the family *Halobacteriaceae*. A neighbour-joining phylogenetic tree (Saitou & Nei, 1987) was constructed with the MEGA package version 3.1 (Kumar et al., 2004) according to the Jukes–Cantor algorithm, and the robustness of the phylogeny was tested by bootstrap analysis (Jukes & Cantor, 1969) after multiple alignment of data by using CLUSTAL W version 1.8 (Thompson et al., 1994). The three strains showed high levels of 16S rRNA gene sequence similarity to each other: 98% between strain RE-101T and RE-102, 97% between strain RE-101T and RE-103 and 99% between strain RE-102 and RE-103. 16S rRNA gene sequence similarity to other members of the family *Halobacteriaceae* was low. The closest related recognized species were representatives of the genera *Halofex* and *Halorubrum* (about 91% similarity) and *Halogeometricum* (about 89%) (Fig. 2). The GenBank database contained two sequences of more closely related (about 96% similarity), yet uncultured halophilic archaea: the first (GenBank accession no. DQ432015) was recovered from an oilfield at an undisclosed location and the second (GenBank accession no. DQ103676) originated from a gypsum crust in a saltern (200 g l−1 total salt concentration) in Eilat, Israel, near the coast of the Gulf of Aqaba, Red Sea (Sørensen et al., 2005). A phylogenetic tree including these environmental sequences is available as Supplementary Fig. S1 in IJSEM Online. The finding of a related environmental sequence near the Red Sea coast is intriguing in view of the fact that the experimental mesocosms from which isolates RE-101T, RE-102 and RE-103 were obtained contained 20% Red Sea water sampled in Eilat. Given the high salt requirement of these organisms, it is not probable that they can live and multiply in seawater containing only 40 g total salts l−1. However, it is interesting to note that closely related environmental 16S rRNA gene sequences belonging to halophilic archaea have been recovered from geographically close locations.

In view of the unusual morphological properties of the new isolates and the low levels of 16S rRNA gene sequence similarity with other genera within the family *Halobacteriaceae*, we suggest that these three isolates represent a novel species of a new genus, for which the name *Haloplanus natan* gen. nov., sp. nov. is proposed.

**Description of Haloplanus gen. nov.**

*Haloplanus* (Ha.lo.pla’nus. Gr. n. hals, halos salt; L. adj. planus flat; N.L. masc. n. *Haloplanus* flat salt-life form).

Cells are pleomorphic, flat and contain gas vesicles. In static liquid culture the cells float to the surface. Gram-negative. Strictly aerobic. Extremely halophilic. Oxidase- and catalase-positive. The genomic DNA G+C content is...
66.1–66.4 mol% (as determined by HPLC). The type species is *Haloplanus natans*. Recommended three-letter abbreviation: Hpn.

Description of *Haloplanus natans* sp. nov.

*Haloplanus natans* (na’tans. L. part. adj. natans swimming, floating).

Cells are 2.5–8 μm in size. Colonies are about 2 mm in diameter after 3 weeks incubation at 37 °C on 2% agar plates containing 17.5–20% salt; they are translucent, entire, smooth, opaque and pinkish. Growth occurs at 2.6–4.3 M NaCl (optimum at 3 M NaCl, at pH 6.5–8.0 (optimum at pH 7.0) and at 37–52 °C (optimum at 40 °C). Chemoorganotrophic. Gelatin, starch and Tween 20 are not hydrolysed. Reduces nitrate to nitrite. No anaerobic growth on nitrate. Does not grow anaerobically in the presence of L-arginine. Some strains produce indole from tryptophan. Glucose, ribose and acetate support growth as single carbon and energy sources. Galactose, sucrose, succinate, glycerol, glutamate and maltose are not used. No acid is produced from sugars. The polar lipids are diphytanyl diether derivatives of phosphatidylglycerol, phosphatidylglycerol phosphate methyl ester, phosphatidyl glycerol sulfate and one major glycolipid that is chromatographically identical to S-DGD-1. Poly-β-hydroxybutyrate is not produced. Resistant to ampicillin, penicillin G, chloramphenicol, rifampicin and neomycin; sensitive to bacitracin, anisomycin and novobiocin.

The type strain, RE-101T (=DSM 17983T =JCM 14081T), was isolated from an experimental mesocosm filled with a mixture of water from the Dead Sea and the Red Sea, Israel. Strains RE-102 (=DSM 17984) and RE-103 (=DSM 17985) are additional strains of this species.

Acknowledgements

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References


