Halophilic methylotrophic methanogens are classified in two genera of slight halophiles and one genus each of moderate and extreme halophiles. Members of the genera Methanolumbus and Methanococcoides are slight halophiles of marine origin, growing fastest in the presence of less than 0.7 M NaCl (10, 29, 30). Members of the genus Methanohalophilus are moderately halophilic, growing fastest in the presence of 0.5 to 2.5 M NaCl (23), and members of the genus Methanohaloobium are extremely halophilic, growing fastest in the presence of NaCl concentrations greater than 2 M (36). No genus other than these four contains halophilic methylotrophic methanogens; although several halophiles were reported in the closely related genus Methanosarcina (2, 28), other studies indicated that these species are not halotolerant rather than halophilic (12, 13). The genus Halomethanococcus has also been proposed (33), but we do not include it in our taxonomy because we have been unable to obtain the type strain of the only species described in this genus (Halomethanococcus doii) and because the name Halomethanococcus may be a subjective synonym of Methanohalophilus.

In addition to being unified by their halophilic, which is recognized as an important taxonomic characteristic (4), the four recognized genera of halophilic methylotrophic methanogens share a coccoid morphology and several physiological characteristics. They are all methylotrophic, growing on media containing methanol and d-l, and trimethylamines as catabolic substrates, but not on media containing dimethyl sulfide, methane thiol, H2, formate, or acetate; when cells were provided with H2 in addition to methanol or trimethylamine, they grew on the medium containing a methyl substrate but did not catabolize H2. All of the strains were capable of growth in mineral medium to which trimethylamine was added as a catabolic substrate, although some strains were greatly stimulated by biotin or p-aminobenzoate. DNA reassociation and denaturing electrophoresis of whole-cell proteins indicated that strains FDF-1T, FDF-2, SF-2, Ret-1, and Cas-1, together with previously described strains SF-1, Z-7302, Z-7401, Z-7404, and Z-7405, belong to a new taxon named Methanohalophilus portucalensis sp. nov.; FDF-1 (= OCM 59) is the type strain. These strains grew fastest at temperatures near 40°C and, in medium containing 0.5 to 2.5 M NaCl, at pH values near 7.

The two new strains excluded from the species on the basis of the results of phylogenetic tests, strains Cas-1 and SD-1, also differed from M. portucalensis in some minor physiological characteristics. Strain Cas-1 was less halophilic (fastest growth occurred in the presence of 0.5 to 1 M NaCl), and strain SD-1 was slightly alkaliphilic (fastest growth occurred at pH 7.8). The DNA reassociation study also showed that Methanohalophilus mahii SLP7 exhibited 52% sequence similarity with Methanohalophilus halophilus Z-79822, supporting the classification of these organisms as separate but closely related species.
other unassigned strains, and propose a new species, *Methanohalophilus portucalensis*.

(Portions of the results have been presented previously, including the isolation and characterization of strain SD-1 [16, 18] and the isolation and characterization of strains FDF-1<sup>T</sup> [T = type strain], FDF-2, Ret-1, and Cas-1 [16, 19].)

### MATERIALS AND METHODS

**Sources of cultures and sediment samples.** All of the following cultures used in this study are available from the Oregon Collection of Methanogens at the Oregon Graduate Institute (Table 1): strains Z-7302, Cas-1, FDF-1<sup>T</sup>, FDF-2, Ret-1, and Cas-1 [16, 19].

<table>
<thead>
<tr>
<th>Strain</th>
<th>Other designations</th>
<th>DNA guanine-plus-cytosine content (mol%)</th>
<th>Source</th>
<th>Reference(s)</th>
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<tr>
<td><em>Methanohalophilus mahii</em> SLP&lt;sup&gt;T&lt;/sup&gt;</td>
<td></td>
<td>68&lt;sup&gt;T&lt;/sup&gt; 5219&lt;sup&gt;T&lt;/sup&gt;</td>
<td>Great Salt Lake</td>
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<td><em>Methanohalophilus halophilus</em> Z-7982&lt;sup&gt;T&lt;/sup&gt;</td>
<td></td>
<td>160&lt;sup&gt;T&lt;/sup&gt; 3094&lt;sup&gt;T&lt;/sup&gt;</td>
<td>Saline cyanobacterial mat</td>
<td>34, 35</td>
</tr>
<tr>
<td><em>Methanohalophilus portucalensis</em> FDF-1&lt;sup&gt;T&lt;/sup&gt;</td>
<td></td>
<td>59&lt;sup&gt;T&lt;/sup&gt; 44</td>
<td>Salinarian</td>
<td>This study</td>
</tr>
<tr>
<td><em>Methanohalophilus portucalensis</em> FDF-2</td>
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<td>66</td>
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<td>This study</td>
</tr>
<tr>
<td><em>Methanohalophilus portucalensis</em> Ret-1</td>
<td></td>
<td>57</td>
<td>Retba Lake</td>
<td>This study</td>
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<td>130</td>
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<td>7302</td>
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<tr>
<td><em>Methanohalophilus sp.</em> strain TR-7</td>
<td></td>
<td>161&lt;sup&gt;T&lt;/sup&gt; 3721&lt;sup&gt;T&lt;/sup&gt;</td>
<td>Saline cyanobacterial mat</td>
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<td><em>Methanobium evestigatum</em> Z-7303&lt;sup&gt;T&lt;/sup&gt;</td>
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<td>Sivash Lagoon</td>
<td>31, 35</td>
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<sup>a</sup> OCM, Oregon Collection of Methanogens, Oregon Graduate Institute, Beaverton.
<sup>b</sup> DSM, Deutsche Sammlung von Mikroorganismen, Braunschweig, Germany.
<sup>c</sup> Data from references 31 and 35.
<sup>d</sup> Data from reference 31.

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We collected samples from pond 20 at the solar salterns of the Leslie Salt Company in south San Francisco Bay (from which strain SF-2 was isolated). The water in the lake contained 220 g of dissolved solids per liter, and its pH was 7. Decomposing bodies of brine shrimp (*Artemia salina*) were present on the surface of the sediment; brine shrimp tissue contains high concentrations of trimethylamine oxide, a precursor of tri-methylamine in anoxic environments. Strains FDF-1<sup>T</sup> and FDF-2 were isolated from sediments from two ponds of a salinarian at Figueira da Foz, Portugal. The water of the ponds was slightly pink (presumably because of the presence of phototrophic halophilic microbes), and the water was saturated with NaCl (crystals appearing to be NaCl were present in the sediments). The water contained 300 g of dissolved solids per liter, and the pH was 7. Strain Ret-1 was isolated from sediments from Retba Lake, Senegal. Retba Lake is hydraulically fed by marine groundwater from the nearby Atlantic Ocean. The high evaporation rate in the lake results in year-round saturation with NaCl. At the time of sampling, the lake water contained 450 g of dissolved solids per liter, and the pH was 7.3. The lake water was pink, and gas bubbles collected from the sediments contained methane. Bernard Ollivier collected sediments from the Casamance River in Senegal in June 1986. Strain Cas-1 was isolated from sediments from the Casamance River, a major river draining southern Senegal. During the dry season, freshwater flow is low, and saltwater intrusion extends inland up to 200 km. High rates of evaporation result in a net flow up-river in the lower part of the river, resulting in hypersaline conditions. This site differs from the others in that it has a lower concentration of dissolved solids (190 g/liter), and it is the only site which contained significant quantities of decaying macrophytes.

**Culture methods.** The Hungate roll tube methods were used with serum tubes and bottles closed with butyl rubber stoppers (7). Additions to media and inoculations were made with glass syringes flushed with O₂-free gas. To isolate predominant methanogens from sediments, the samples were serially diluted into enrichment medium, and enrich-
ment cultures were maintained from the highest dilutions which formed methane upon incubation. Organisms were isolated from such enrichment cultures by serial dilution, inoculation of roll tube medium, and incubation until colonies formed. Methanogenic colonies were tentatively identified by observing the roll tubes by epifluorescent microscopy with a Zeiss Universal microscope equipped with a type O2 filter set. Colonies were picked with a modified Pasteur pipet (7) and transferred to sterile medium in an open tube flushed with O2-free gas. The tube was aseptically closed with a stopper, serial dilutions were made in sterile medium, and roll tubes were inoculated from the dilutions. When methanogenic colonies were visible, we repeated the picking, dilution, and roll tube culturing procedures until a single colony type remained (except that subsurface colonies differed in appearance from surface colonies) and then repeated these procedures at least two additional times to ensure purity. The resulting pure cultures were deposited in the Oregon Collection of Methanogens.

For enrichment and isolation, cultures were incubated statically at 37°C. Otherwise, cultures were grown statically at 42°C, except that cultures containing added H2 were incubated on a shaker.

**Culture media.** Enrichment media were prepared by using site water and (per liter) 1 g of NH4Cl, 0.4 g of KH2PO4, 3 g of MgCl2, 6H2O, 0.5 g of CaCl2, 2H2O, 10 ml of a trace mineral solution (6), and 0.2 g of yeast extract. The media were dispensed into individual vessels along with a gas mixture containing N2 and CO2 (4:1), the vessels were closed with stoppers, and the preparations were autoclaved. After the media cooled and 1 h to 18 h before inoculation, we added catabolic substrates from concentrated anoxic stock solutions, as well as 2.5 ml of a sodium sulfide solution (100 g of Na2S · 9H2O per liter) per liter of medium. The final pH was 7.2. For strains FDF-1T and FDF-2 the enrichment medium was modified by diluting the site water 1:1 with deionized water before the medium was prepared.

The media used for isolations in roll tubes were the same as enrichment medium except that they were prepared from a 1:1 dilution of site water and had 20 g of purified agar added per liter of medium. Most characterization studies were done with HS medium (17); the exceptions were the studies for vitamin requirements, which were done with MH medium (3) modified by eliminating Trypticase peptone, yeast extract, and coenzyme M. To determine optimal conditions for growth, we adjusted the pH by varying the CO2 content of the gas (to give pH values between 6.4 and 8.3) or by adding 1 M NaOH to medium with N2 gas or 1 M HCl to medium with CO2 gas to give more extreme pH values.

**Determination of specific growth rates.** Specific growth rates were calculated from exponential increases in accumulated methane levels, taking into account the methane formed by the inoculum culture prior to its transfer (24). Methane was quantified by gas chromatographic analysis of samples collected at the pressure of the vessels, thereby giving values for amount of methane per volume of headspace rather than per volume of gas under standard conditions. Samples were collected in pressure lock syringes and were detected by thermal conductivity (1) or by a sampling loop with flame ionization detection (12). Specific growth rates were calculated by least-squares analysis of the rate of increase of the logarithm of accumulated methane level. Whenever the specific growth rate of a culture was much lower than that of its inoculum, we transferred this culture during late logarithmic phase to medium under the same conditions and measured growth again. Also, the shapes of the growth curves from which specific growth rates were calculated always indicated logarithmic growth.

**Guanine-plus-cytosine content of DNA.** Cells were lysed by adding 0.1 g of sodium dodecyl sulfate per liter of cell pellet, and DNA was extracted with chloroform and isoamyl alcohol (15). Buoyant density was measured in CsCl gradients immobilized after centrifugation by polymerization of acrylamide (25). The guanine-plus-cytosine content was calculated from the buoyant density (27) by comparison with DNAs from Clostridium perfringens (p was assumed to be 1.691 g/cm3; Sigma Chemical Co., St. Louis, Mo.) and “Micrococcus lysodeikiticus” (p was assumed to be 1.731 g/cm3; Sigma).

**Antibiotic susceptibility.** Anoxic, filter-sterilized solutions of antibiotics were added to cultures in mid-logarithmic phase, giving a final concentration of 100 mg/liter. Methane formation was measured for 7 days. The apparent specific growth rates were compared with the growth rates of control cultures grown without antibiotics. Inhibited cultures were always transferred to fresh medium without antibiotics to determine whether any cells remained viable, and in these cases growth always occurred.

**Organic nutrient requirements.** In order to determine the minimum organic nutrients required for growth, we measured the specific growth rates of cultures grown for at least four transfers (2% inoculum by volume) in mineral medium to which 20 mM trimethylamine was added as a catabolic substrate. For strains which grew substantially more slowly in the mineral medium than in the complex medium, we tested the effect of a vitamin mixture (6) or one or more of its constituent vitamins on the specific growth rates of these cultures.

**Electrophoretic analysis of whole-cell proteins.** Whole-cell proteins were extracted from cell pellets lysed by adding 0.5 g of sodium dodecyl sulfate per liter. Approximately 25 μg of protein was added per well of a polyacrylamide gel, and proteins were separated as described previously (11). The gel was stained with Coomassie blue (11). The proteins bands were analyzed by measuring the position of each band and noting its density (light, medium, or dark). The level of similarity between a pair of organisms was calculated by the following algorithm: for each light protein band from the first organism, 1 point was added if the second organism had any band within 0.5 mm of the position of this light protein band; for each medium band from the first organism, 2 points was added for a medium or dark band within 0.5 mm, or 1 point was added for any band within 1.5 mm. The percentage of protein similarity was calculated by dividing the actual points by the total possible points. For each pair of organisms, each organism was compared with the other, and the percentage of similarity for the pair was the average of the reciprocal determinations. From these binary percentages of similarity, a tree (see Fig. 5) was formed in the same way that the DNA reassociation tree was formed.

**DNA reassociation.** DNA was purified from cell pellets by the Marmur method (15) and was labeled with 32P-cytosine by extension of random primers (13). DNA reassociation experiments were done by using the S1 nuclease protocols (32) developed by Johnson (8) and modified by Maestrojuan et al. (13). For each interstrain comparison, each member of the pair was labeled and tested with the cold DNA of the other strain, and the interspecies sequence similarity values
given below are the averages of reciprocal determinations. In the comparison between Methanohalophilus mahii SLP\textsuperscript{T} and Methanohalophilus halophilus Z-7982\textsuperscript{T}, each reciprocal determination was repeated. The tree shown in Fig. 5 was developed by grouping the most closely related organisms first, and subsequently the level of phylogenetic relationship between organisms or a group and another group of organisms was determined as the average of all of the individual similarity values for each pair between the two groups.

**Microscopy.** Epifluorescence was determined microscopically by using a type O2 filter set (Carl Zeiss, Inc., Thornwood, N.Y.); this provided an excitation spectrum with a peak at 365 nm, a cutoff at 395 nm, and a 420-nm long-pass barrier filter.

**RESULTS**

Isolation and characterization of strain SD-1. A sediment suspension from a saltern in San Diego, Calif., was inoculated into enrichment medium supplemented with 20 mM trimethylamine and enrichment medium supplemented with 50 mM sodium acetate (0.1 ml of suspension in 50 ml of culture medium). Cultures in medium containing trimethylamine produced detectable methane within 3 days and completed methane production within 15 days (the quantity of methane formed was near the amount stoichiometrically expected). No significant quantities of methane were formed in cultures in medium containing acetate within 30 days. An enrichment culture was developed from the trimethylamine-containing culture by 5% transfers when methanogenesis was complete; we isolated strain SD-1 from this enrichment culture and deposited it in the Oregon Collection of Methanogens as strain OCM 134.

Surface colonies were 0.5 mm in diameter within 7 days, tannish yellow, round, convex, smooth, and shiny and had entire edges and grainy interiors. Subsurface colonies were similar but had a lenticular shape. The colonies fluoresced brightly when they were examined by epifluorescence microscopy, and the fluorescence did not fade. Cells in the exponential growth phase were refractile, irregular cocci (diameter, 1 μm) which occurred singly, in pairs, and in small regular tetragonal clumps. The cells stained gram negative and fluoresced when they were examined by epifluorescence microscopy, and the fluorescence did not fade. Cells in the exponential growth phase were refractile, irregular cocci (diameter, 1 μm) which occurred singly, in pairs, and in small regular tetragonal clumps. The cells stained gram negative and fluoresced when they were examined by epifluorescence microscopy, and the fluorescence did not fade. Cells in the exponential growth phase were refractile, irregular cocci (diameter, 1 μm) which occurred singly, in pairs, and in small regular tetragonal clumps. The cells stained gram negative and fluoresced when they were examined by epifluorescence microscopy, and the fluorescence did not fade. Cells in the exponential growth phase were refractile, irregular cocci (diameter, 0.8 to 1.2 μm) and slightly refractile and stained gram negative. Cells occurred individually, in pairs, and in irregular clumps. Cells lysed in hypotonic medium and in the presence of 50 mg of sodium dodecyl sulfate per liter.

Strain SD-1 grew in medium containing trimethylamine, dimethylamine, monomethylamine, or methanol as the catabolic substrate. No growth occurred in medium supplemented with 50 mM acetate, 100 kPa of H\textsubscript{2} plus 20 kPa of CO\textsubscript{2}, or 5 mM dimethyl sulfide as the catabolic substrate. When 100 kPa of H\textsubscript{2} plus trimethylamine was added as the catabolic substrate, cultures formed the same quantity of methane and grew at the same rate as controls in medium containing only trimethylamine. With trimethylamine as the catabolic substrate, the cells grew fastest at 42°C (Fig. 1), at pH 7.8 (Fig. 2), and in the presence of 0.9 to 3.5 M Na\textsuperscript{+} (Fig. 3). Cells grew rapidly (specific growth rate, 0.015 h\textsuperscript{-1}) in mineral medium with no organic compound other than trimethylamine added; vitamins did not stimulate growth.

Chloramphenicol inhibited the growth and methanogenesis of cells. Tetracycline, ampicillin, carbenicillin, cycloserine, erythromycin, and penicillin had no effect at a concentration of 100 mg/liter.

**FIG. 1.** Effect of temperature on specific growth rates of halophilic methanogens.

The guanine-plus-cytosine content of the DNA was 41 mol%.

Isolation and characterization of strains FDF-1\textsuperscript{T} and FDF-2. The sediment suspension from the saltern at Figueira da Foz was serially diluted in enrichment medium containing 10 mM trimethylamine and incubated at 37°C. Cultures inoculated with 10 μl or more of inoculum formed stoichiometric quantities of methane within 2 weeks. A culture which had received 10 μl of inoculum was transferred to enrichment medium, and two methanogenic strains (strains FDF-1\textsuperscript{T} and FDF-2) were isolated. The following description applies to both strains. Colonies were 0.5 mm in diameter within 7 days of incubation at 37°C. Surface colonies were circular, tannish yellow, convex, shiny, and opaque and had entire margins; subsurface colonies were lenticular. Cells were irregular cocci (diameter, 0.8 to 1.2 μm) and slightly refractile and stained gram negative. Cells occurred individually, in pairs, and in irregular clumps. Cells lysed in hypotonic medium and in the presence of 50 mg of sodium dodecyl sulfate per liter.

Cells of strain FDF-1\textsuperscript{T} and FDF-2 grew in medium con-
irregular refractile cocci (diameter, 1 μm) which stained gram negative. Cells occurred mainly individually. Cells margins; subsurface colonies were lenticular. Cells were tannish yellow, convex, shiny, and opaque and had entire incubation was at 37°C. Surface colonies were circular, Colonies were 0.5 mm in diameter within 7 days when medium supplemented with 200 g of NaCl per liter supplemented with 150 g of NaCl per liter (2.7 M Na+). inoculum was maintained as an enrichment culture in HS medium supplemented with 20 mM trimethylamine and suspension from Retba Lake was serially diluted in enrich- ment culture was maintained as an enrichment culture in HS agar medium containing 1.5 M NaCl.

Strains FDF-1T and FDF-2 both grew well in mineral medium to which trimethylamine was added as the sole organic substrate (specific growth rates, 0.027 h⁻¹ for FDF-1T and 0.026 h⁻¹ for FDF-2). Growth was almost as rapid in this medium as in MH medium, which contained yeast extract. The growth rate of these cultures in MH mineral medium supplemented with trimethylamine was not strongly stimulated by the presence of 5 mM glycine betaine.

Chloramphenicol inhibited the growth of cells. Tetracycline, ampicillin, carbenicillin, cycloserine, erythromycin, and penicillin had no effect at a concentration of 100 mg/liter. The guanine-plus-cytosine content of the DNA was 43 mol%.

Isolation and characterization of strain Ret-1. The sediment suspension from Retba Lake was serially diluted in enrichment medium supplemented with 20 mM trimethylamine and incubated at 37°C. Cultures inoculated with 0.5 μl or more of inoculum formed stoichiometric quantities of methane within 3 weeks. Other cultures containing methanol as the catabolic substrate and inoculated in the same way also formed methane, but cultures in media containing acetate, H₂, formate, propionate, butyrate, lactate, or cellulose did not form methane. The culture in medium containing trimethylamine as the substrate and inoculated with 0.05 μl of inoculum was maintained as an enrichment culture in HS medium supplemented with 200 g of NaCl per liter (3.5 M Na+), and strain Ret-1 was isolated in HS agar medium supplemented with 150 g of NaCl per liter (2.7 M Na⁺). Colonies were 0.5 mm in diameter within 7 days when incubation was at 37°C. Surface colonies were circular, tannish yellow, convex, shiny, and opaque and had entire margins; subsurface colonies were lenticular. Cells were irregular, refractile coci (diameter, 1 μm) which stained gram negative. Cells occurred mainly individually. Cells lysed in hypotonic medium and in the presence of 100 mg of sodium dodecyl sulfate per liter.

Strain Ret-1 grew in medium containing trimethylamine, dimethylamine, monomethylamine, or methanol as the substrate, but not in medium supplemented with acetate, H₂, or dimethyl sulfide. The presence of H₂ had no effect on methanogenesis from trimethylamine.

Cells grew most rapidly in the presence of 1.4 M Na⁺ (Fig. 3), at 37°C (Fig. 1), and at pH 7.1 (Fig. 2).

Strain Ret-1 required biotin and p-aminobenzoate for good growth. It grew very poorly in MH mineral medium supplemented with trimethylamine (specific growth rate, 0.0057 h⁻¹), but addition of the two vitamins together resulted in good growth (specific growth rate, 0.025 h⁻¹). Growth was still not as rapid as as growth in MH medium (containing yeast extract and peptones; specific growth rate, 0.046 h⁻¹).

Chloramphenicol and tetracycline inhibited the growth of cells. Ampicillin, carbenicillin, cycloserine, erythromycin, and penicillin had no effect at a concentration of 100 mg/liter. The guanine-plus-cytosine content of the DNA was 43 mol%.

Isolation and characterization of strain Cas-1. The sediment suspension from the Casamance River was inoculated into HS medium containing 80 g of NaCl per liter (1.5 M Na⁺) and 20 mM trimethylamine, and the preparation was incubated at 37°C. Sediments inoculated into medium containing 120 g of NaCl per liter and 20 mM trimethylamine did not form methane. Methanogenesis from the enrichment culture containing 80 g of NaCl per liter was complete within 7 days, and an enrichment culture was maintained by transfer in the same medium. Strain Cas-1 was isolated from this enrichment culture in HS agar medium containing 1.5 M NaCl.

Colonies were 1 mm in diameter within 7 days when incubation was at 37°C. Surface colonies were circular, tannish yellow, convex, shiny, and opaque and had entire margins; subsurface colonies were lenticular. Cells were cocci (diameter, 1 μm) which stained gram negative and occurred mainly individually. Cells lysed in hypotonic medium and in the presence of 100 mg of sodium dodecyl sulfate per liter.

Strain Cas-1 grew in medium containing trimethylamine, dimethylamine, monomethylamine, or methanol as the substrate, but not in medium containing acetate, H₂, or dimethyl sulfide. The presence of H₂ had no effect on methanogenesis from trimethylamine.

Cells grew most rapidly in the presence of 0.6 to 1.5 M Na⁺ (Fig. 3), at 42°C (Fig. 1), and at pH 6.4 to 7.8 (Fig. 2).

Chloramphenicol inhibited the growth of cells. Tetracycline, ampicillin, carbenicillin, cycloserine, erythromycin, and penicillin had no effect at a concentration of 100 mg/liter. The guanine-plus-cytosine content of the DNA was 42 mol%.

Isolation and characterization of strain SF-2. The sediment suspension from the saltern in San Francisco Bay was serially diluted and inoculated into enrichment medium supplemented with 20 mM trimethylamine. Within 10 days bottles inoculated with the equivalent of 10 ml of sediment slurry produced significant quantities of methane. This culture was maintained as an enrichment culture for several transfers before a cocccoid methanogen (strain SF-2) was isolated from it.

Surface colonies were 0.5 mm in diameter within 7 days, tannish yellow, round, convex, smooth, and shiny and had entire edges and grainy interiors. Subsurface colonies were...
similar but were lenticular. The colonies fluoresced brightly when they were examined by epifluorescence microscopy, and the fluorescence did not fade. Cells in the exponential growth phase were refractile, irregular cocci (diameter, 1 μm) which occurred singly, in pairs, and in small regular tetragonal clumps. Cells stained gram negative and fluoresced when they were examined by epifluorescence microscopy, but the fluorescence faded after about 30 s. Cultures stored in serum bottles at room temperature retained viability for at least 6 months. Cells were sensitive to lysis by osmotic shock or by the presence of 50 mg of sodium dodecyl sulfate per liter.

Strain SF-2 grew in medium containing trimethylamine, dimethylamine, monomethylamine, or methanol as the catabolic substrate. No growth occurred in medium containing 50 mM acetate, 100 kPa of H2 plus 20 kPa of CO2, or 5 mM dimethyl sulfide as the catabolic substrate. When 100 kPa of H2 plus trimethylamine was added as the catabolic substrate, cultures formed the same quantity of methane and grew at the same rate as controls in medium containing only trimethylamine. Cells in mineral medium containing trimethylamine as the only organic compound grew at a specific growth rate of 0.025 h⁻¹, and growth was not stimulated by vitamins.

Vitamin requirements of other moderately halophilic methanogens. Strains Z-7401, Z-7404, Z-7302, Z-7982T, and SLP² were all able to grow in MH mineral medium containing trimethylamine as the substrate. Strains Z-7401 and Z-7404 grew rapidly in this medium (nearly as rapidly as in MH medium, which contains yeast extract and peptones), but the other strains grew very poorly. Strains Z-7982T and SLP² grew well in MH mineral medium containing trimethylamine when biotin was added. Strains Z-7302 and Ret-1 grew well only when mineral MH medium containing trimethylamine was supplemented with both biotin and p-aminobenzoate.

Growth of Methanohalobium evestigatum on medium containing methanol. Methanohalobium evestigatum Z-7303T was not able to grow on medium containing 20 mM methanol or in medium containing 20 mM methanol plus 40 mM trimethylamine. However, cells grew on medium containing 5 mM methanol.

Analysis of whole-cell proteins. We used electrophoresis in denaturing gels to analyze the whole-cell proteins of a number of halophilic methanogens (Fig. 4). Our results revealed a high degree of diversity among these organisms in their proteins, although we distinguished two groups whose members exhibited similarities with each other. One group comprises strains FDF-1T, FDF-2, Z-7401, and Z-7404, and the other consists of strains Z-7302, Ret-1, and SF-2.

We prepared other protein gels to compare strain Cas-1 with the type strains of Methanolobus and Methanococcoides species (data not shown); because strain Cas-1 is more similar to members of these genera in its optimal salt concentration than it is to Methanohalophilus strains. However, the proteins of strain Cas-1 were more similar to those of the Methanohalophilus strains included in this study than to the proteins of Methanolobus or Methanococcoides strains.

DNA reassociation studies. The results of the DNA reassociation experiments are shown in Fig. 5. The DNA reassociation tree is consistent with the tree derived from a protein analysis (Fig. 5).

**DISCUSSION**

The methanogens investigated in this study, which include most of the known available moderately halophilic methanogens from pH-neutral environments, were isolated from a wide variety of sources (Table 1). Yet all of these methanogens appear to be similar physiologically, having an identical substrate range and exhibiting similar growth responses to different pH, salinity, and temperature conditions. These characteristics suggest that all of these strains, with the possible exceptions of strains Cas-1 and SD-1, should be classified as members of the genus Methanohalophilus. Strains Cas-1 and SD-1 differed from Methanohalophilus strains physiologically in some respects (strain Cas-1 grew more rapidly and was less halophilic, and strain SD-1 was slightly halophilic), and DNA reassociation studies revealed a phylogenetic separation (Fig. 5). We believe that the taxonomic assignment of these two strains should await rRNA sequence analysis.

Figure 5 also shows that Methanobacterium strain GS-16 and Methanohalobium evestigatum Z-7303T were not closely related to the genus Methanohalophilus. Assignment of strain GS-16 to the genus Methanobacterium rather than the genus Methanohalophilus is suggested by the results of a 16S rRNA sequence analysis (26).

**Taxonomy of Methanohalophilus mahii and Methanohalophilus halophilus.** Methanohalophilus mahii and Methanohalophilus halophilus are physiologically very similar (22, 23, 34, 35), and these names could be considered subjective.
were phylogenetically distant from the other strains included. Halophilus strains can be divided into 60% between-group DNA sequence similarity, all of the strains are very similar physiologically and morphologically. Be lost unless it was conserved by an opinion of the Judicial Commission. Therefore, in the interest of taxonomic stability, we continue to consider these taxa two separate species.

**Taxonomy of other moderately halophilic methanogens.** DNA reassociation and whole-cell protein analyses (Fig. 5) showed that strains Cas-1 and SD-1, Methanobacterium sp. strain GS-16, and Methanobacterium evestigatum Z-7303 were phylogenetically distant from the other strains included in this study. We propose that strains FDF-1, FDF-2, Z-7302, Z-7401, Z-7404, Z-7405, SF-1, SF-2, and Ret-1 should be placed in a single species. Even though these strains can be divided into two groups that exhibit less than 60% between-group DNA sequence similarity, all of the strains are very similar physiologically and morphologically.

**Description of Methanohalophilus portucalensis** sp. nov.

*Methanohalophilus portucalensis* (por.tu.cal.en’s is. L. adj. *portucalensis*, from Portugal) cells are irregular, nonmotile, coccoid cells (diameter, 0.6 to 2 μm) and are gram negative. The coccoid cells occur singly, in pairs, or, less commonly, in small aggregates. Strictly anaerobic. Cells are sensitive to lysis by 0.2 g of sodium dodecyl sulfate per liter. The catabolic substrates include methanol and methylamines, but not H₂ plus CO₂, formate, acetate, methane thiol, or dimethyl sulfide. Cells grow in medium containing minerals plus trimethylamine as the sole organic substrate, but some strains require p-aminobenzoate, biotin, or both.

Growth is most rapid in medium containing 0.5 to 2 M NaCl, at pH 6.5 to 7.5, and at temperatures of about 40°C.

The guanine-plus-cytosine content of the DNA is 43 to 44 mol%.

The type strain is strain FDF-1 (= OCM 59), which was isolated from sediments from a salinarium in Figiera da Foz, Portugal.

**ACKNOWLEDGMENTS**

We thank Bernard Ollivier (INRA, ORSTOM, Marseilles, France) for collecting samples and for helpful discussions, Shuinsong Ni (Oregon Graduate Institute) for technical help and discussions, and Thomas O. MacAdoo (Virginia Polytechnic Institute and State University) for checking the etymology of the epithet.

This study was funded in part by the Program in Energy from Biomass, funded by the Gas Research Institute and the University of Florida (grant IFAS-GRI-FIA-MCS 2171), and by the U.S. Geological Survey, Department of the Interior (grants 14-08-0001-G1636 and 14-08-0001-G2098).

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