**Teredinibacter turnerae gen. nov., sp. nov., a dinitrogen-fixing, cellulolytic, endosymbiotic γ-proteobacterium isolated from the gills of wood-boring molluscs (Bivalvia: Teredinidae)**

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**Keywords:** Teredinidae, bivalia, shipworms, symbiont, 16S rRNA phylogeny

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**INTRODUCTION**

Endosymbiotic bacteria are found within the gills of several hundred bivalve species. These species are confined to a relatively small but extremely diverse selection of bivalve taxa, including all examined species within the families Teredinidae, Solemyidae, Vesicomyidae and Lucinidae, most Thyasiridae, and all examined members of the subfamilies Xylophaginiae (family Pholadidae) and Bathymodioliinae (family Mytilidae) (Fisher, 1990).

Much has been inferred about the physiology and phylogeny of bivalve gill endosymbions based on molecular and biochemical investigations. For example, symbions of Solemyidae, Vesicomyidae, Lucinidae and Thyasiridae have been determined to be sulfur-oxidizing chemolithoautotrophs, while those of Bathymodioliinae are either chemolithoautotrophs or methanotrophs (Fisher, 1990). Carbon, fixed by these symbions, provides an important contribution to the host’s diet. Symbions of Teredinidae and Xylophaginiae, on the other hand, play a different role in host nutrition. These symbions are thought to provide cellulolytic enzymes that facilitate the digestion of...
wood by their hosts (Distel & Roberts, 1997; Greene, 1994; Waterbury et al., 1983).

The shipworm isolates reported by Waterbury et al. (1983) and formally described here, however, are the only gill-born bivalve endosymbionts yet cultivated, characterized, or named. The shipworm isolate T7902 has been confirmed by molecular methods (fluorescent in situ hybridization) to be a symbiotic inhabitant of gill tissues of the shipworm *Lyrodus pedicellatus* (Distel et al., 1991) and the morphology and ultrastructure of this symbiosis has been described. Growth characteristics (Greene & Freer, 1986), cellulolytic and proteolytic activities (Greene et al., 1988; Imam et al., 1993), and cellulose binding properties (Imam et al., 1990) have also been described for strain T8201, isolated from *Psiloteredo healdi*. Here we report and summarize the properties, growth characteristics and phylogenetic affiliations of a cellulolytic, dinitrogen-fixing isolate from the shipworm *Lyrodus pedicellatus* and 58 similar isolates from the gills of nine genera and 24 species of teredinid bivalves. A new genus and species, *Teredinibacter turnerae* gen. nov., sp. nov., is proposed with strain T7902T (= ATCC 39867T = DSM 15152T) as the type strain.

**METHODS**

**Culture conditions.** Cells were cultivated in a basal medium (shipworm basal medium, SBM) (Waterbury et al., 1983), containing the following: filtered seawater (750 ml l−1), distilled water (250 mg l−1), KH₂PO₄ (15.3 mg l−1), Na₂CO₃ (10 mg l−1), Na₂MoO₄, 2H₂O (2.5 mg l−1) disodium EDTA (0.5 mg l−1), ferric ammonium citrate (brown crystals) (3 mg l−1), HEPES buffer (pH 8.0) (5.2 g l−1) and A5 trace metals (Rippka et al., 1979) (1.0 ml l−1). Appropriate carbon sources were added with or without (5 mM) ammonium chloride as a source of combined nitrogen. Cells were incubated at temperatures ranging from 15 to 40 °C. To obtain individual colonies, cells were plated on 1% agar, SBM plates supplemented with powdered cellulose (0.2% w/v) (Sigmacell 100) with or without combined nitrogen. Axenic stock cultures were maintained in SBM agar (0.2%) and Sigmacell 100 (0.2%) without combined nitrogen at room temperature. Maintenance cultures were transferred to fresh medium at 30-day intervals. Stock cultures grown in SBM were amended with 5% DMSO and cryopreserved in liquid nitrogen. Liquid cultures, used for nucleic acid extractions, contained SBM supplemented with Sigmacell 100 (0.2%) and ammonium chloride (5 mM) and were incubated with gentle shaking at 35 °C.

**Isolation.** Isolates (Table 1) were obtained from gill tissue of freshly collected bivalve specimens by serial dilution as previously described (Waterbury et al., 1983). Gills were removed, washed and homogenized in sterile seawater. Homogenates were used to inoculate 10 ml serial dilutions (10⁻¹–10⁰) in SBM supplemented with 0.2% agar and powdered cellulose (0.2% w/v, Sigmacell 100) without addition of combined nitrogen. After incubation for 5–10 days at room temperature, cell growth appeared as a lens about 1 cm below the agar surface. Cells were removed from the greatest dilutions in which growth was observed (10⁻¹–10⁰) and streaked on 1% agar SBM plates supplemented

<table>
<thead>
<tr>
<th>Bacterial strain no.*</th>
<th>Host species †</th>
<th>MCZ voucher no. ‡</th>
<th>Host source</th>
</tr>
</thead>
<tbody>
<tr>
<td>T7902T&lt;br&gt;(Quatrafragos)</td>
<td><em>Lyrodus pedicellatus</em></td>
<td>297526</td>
<td>Laboratory Culture, WHOI-ESL. Original specimens from Long Beach, CA, USA</td>
</tr>
<tr>
<td>T7901&lt;br&gt;(Bartsch)</td>
<td><em>Bankia gouldi</em></td>
<td>297551</td>
<td>Laboratory Culture, WHOI-ESL. Original specimens from Duke University Marine Lab, Beaufort, NC, USA</td>
</tr>
<tr>
<td>T8402&lt;br&gt;(Turton)</td>
<td><em>Teredora malleohus</em></td>
<td>297577</td>
<td>Floating wood 190 mi S.E. of Woods Hole, MA (38°18′4″ N, 69°35′6″ W). From the West Indies via the Gulf Stream</td>
</tr>
<tr>
<td>T8602&lt;br&gt;(Wright)</td>
<td><em>Dicyathifer mannii</em></td>
<td>297578</td>
<td>Ferguson Point, near Australian Institute of Marine Science Lab, S. of Townsville, Qld, Australia</td>
</tr>
</tbody>
</table>

* Nomenclature – example of strain no. T7902: T, designates an isolate from the bivalve family Teredinidae; 79, the year of isolation (e.g. 1979); 02, the second isolate of that year.
† Host specimens were collected, identified and dissected by Dr Ruth Turner and C. B. Calloway, Museum of Comparative Zoology, Harvard University, Cambridge, MA, USA.
‡ Voucher specimens are held in the Mollusc Department of the Museum of Comparative Zoology, Harvard University, Cambridge, MA, USA. They consist of the shells, pallets and remaining soft tissue of the shipworms from which shipworm strains were isolated.

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**Table 1. Strain histories of *T. turnerae* isolates**

A complete list of shipworm isolates can be found as supplementary data in IJSEM Online (http://ijsem.sgmjournals.org).
with Sigmacell 100 (0.2%). Individual colonies could be observed on the agar surface after 5 days incubation at room temperature. Colonies were excised with a sterile loop and were used to inoculate fresh liquid medium supplemented with Sigmacell 100 (0.2%). After 5 days at room temperature, these liquid cultures were used to inoculate new serial dilutions. Rounds of serial dilution, plating and inoculation of liquid medium were repeated until axenic isolates were obtained. Host taxa were identified and dissected by Dr Ruth D. Turner or Dr C. B. Calloway and voucher specimens (shells and pallets) are on deposit at the Harvard Museum of Comparative Zoology, Malacology Department, Harvard University, Cambridge, MA, USA (see Table 1). Isolates were named as follows: the letter T indicates isolates from bivalves of the family Teredinidae, the following two digits indicate the year of isolation and the remaining digits chronologically differentiate strains isolated in a given year.

**Phenotypic characterization.** Fourteen isolates from stock cultures maintained as described above were streaked on 1% agar SBM plates supplemented with Sigmacell 100. A single colony was picked from each isolate and restreaked on the same medium, then single colonies from this plate were used to inoculate 25 ml SBM liquid medium supplemented with NH₄Cl (0.025%) with a strip of Whatman no. 1 filter paper (1 x 2 cm) added as a carbon source. Cells grown to mid-exponential phase were then used to inoculate media in glass culture tubes (15 x 100 mm). Each tube contained SBM (10 ml) supplemented with 0.2% agar, NH₄Cl (0.025%), 20 mM HEPES (pH 8.0) buffer (except in the tubes containing cellobiose in which 50 mM HEPES was used), 1.0% phenol red as a pH indicator, and a selected carbon source. Two additional consecutive transfers were performed in the same medium to confirm growth. Cells were observed using a Zeiss compound microscope (1000 x magnification). Each isolate was also inoculated into the same medium lacking an added carbon source. After the third transfer on the indicated carbon source each isolate was regrown on 1% agar SBM plates supplemented with Sigmacell 100 to confirm its identity.

Salt requirements and temperature and pH range and optima for six isolates (T7901, T7902, T7903, T8201, T8202 and T8203) were determined as in (Greene & Freer, 1986) and (Waterbury et al., 1983).

Catalase activity was assayed with 0.3% hydrogen peroxide using cells grown in SBM medium in 0.2% agar stab cultures supplemented with Sigmacell 100 and inoculated from a single colony. Oxidase activity was tested using the Dry Slide Oxidase test kit (Difco). Growth under anaerobic conditions was tested on SBM cellulose in a GasPak system (BBL). Dinitrogenase activity was determined using the acetylene reduction assay on cultures grown in tubes and then capped with rubber stoppers during the assay (Hardy & Holsten, 1977).

**Morphological examination.** The type of flagellation was determined by electron microscopy using cells negatively stained with 0.5% aqueous uranyl acetate. A Philips 300 transmission electron microscope was used to examine samples. Motility, characteristics of cell division, Gram type and staining characteristics were observed by light microscopy using phase-contrast illumination with a Zeiss Standard microscope. Cells for microscopy were grown in SBM supplemented with Sigmacell 100 (0.2%) and ammonium chloride (5 mM) and were incubated with gentle shaking at 35 °C. Cells were harvested during exponential-phase growth. Staining to determine Gram type and presence or absence of PHB (0.3% Sudan black B) or polyphosphate (0.3% methylene blue) was performed as described by Gerhardt (1981).

**DNA isolation.** Residual cellulose powder was removed from 20 ml of exponential-phase liquid culture by centrifugation (1000 g, 2 min). Cells were then pelleted from the supernatant by centrifugation (16000 g, 5 min). Cells were washed by resuspension in sterile seawater (100 µl), pelleted again as above, and resuspended in a final volume of 300 µl STET (0.1 M NaCl; 10 mM Tris/HCl, pH 8.0; 1 mM EDTA, pH 8.0; 5% Triton X-100) containing lysozyme (Sigma) at 10 mg ml⁻¹. Cells were incubated at 37 °C for 1 h, then Proteinase K (Sigma) was added to a final concentration of 50 µg/ml and incubation was continued overnight at 37 °C. After incubation, 200 µl guanidium (iso)thiocyanate (GSCN) lysis medium [50 M guanidinium (iso)thiocyanate, 50 mM Tris/HCl, pH 7.5; 50 mM sodium acetate, pH 7.0; 25 mM EDTA, pH 8.0] was added to the Proteinase K lysate and gently mixed in a Dounce homogenizer. To the homogenate 60 µl sodium lauryl sarcosinate (10% w/v) was added and cellular debris was removed by centrifugation (2 x 5 min at 16000 g). Nucleic acids were precipitated by addition of one volume of cold absolute ethanol to the supernatant followed by centrifugation (2 min at 16000 g). The resulting pellet was dissolved by trituration in 100 µl 6 M guanidine · HCl and undissovled material was removed by centrifugation (2 x 5 min at 16000 g). Nucleic acids were precipitated by centrifugation (2 min at 16000 g) after addition of 2.5 volumes of cold ethanol and 1/10 volume of 3 M sodium acetate. The pellet was washed with 70% ethanol (100 µl), lightly dried under vacuum and dissolved in 50 µl TE buffer (1 mM EDTA; 10 mM Tris/HCl, pH 7.4). Residual protein was removed by organic extraction (Sambrook et al., 1989) with phenol:chloroform:isoamyl alcohol (25:24:1) and chloroform:isoamyl alcohol (24:1) followed by ethanol precipitation and dissolution in TE buffer as above.

**DNA base composition.** DNA base composition was determined by thermal denaturation as in (Herdmann et al., 1979).

**PCR amplification.** Bacterial 16S rDNAs were amplified by PCR (Saiki et al., 1987) from genomic nucleic acid extracted from shipworm isolates using bacterial-domain-specific primers 27f (AGAGTTTGATCATGGCTCAG) and 1492r (GGTTACCTTGTTACGACTT) (Lane, 1991). Reactions were carried out in 50 µl volumes containing 44 µl AmpliTaq buffer (Perkin Elmer) 10 × PCR buffer, 1.8 mM MgCl₂, 25 mM each dATP, dCTP, dGTP and dTTP, 100 nM each primer, 1.25 U AmpliTaq DNA polymerase (Perkin Elmer) and 0.2 ng template nucleic acid µl⁻¹. PCR reactions used the following parameters: presoak: 94 °C, 1 min; 35 cycles of 1 min each at 94, 60 and 72 °C; post-soak: 72 °C, 7 min. Amplification products (~1.5 kb) were visualized by gel electrophoresis (0.8% agarose, 0.5 × TBE, (Sambrook et al., 1989). Reaction products were purified and concentrated by ultrafiltration (Microcon YM100 spin microconcentrator; Amicon) prior to sequencing.

**16S rRNA gene sequencing.** Sequencing reactions were performed using the ABI Prism Dye Terminator Cycle Sequencing Ready Reaction kit (Applied Biosystems) using AmpliTaq FS DNA Polymerase, on an ABI 373A DNA sequencer. Nearly complete double-stranded sequence of the 16S rRNA gene (1453 bp) was determined from PCR products using primers: 27f, 1492r, 357f (CTCCTACGGAGA-GTACCTTGTTACGACTT) (Lane, 1991). Reactions were carried out in 40 µl volumes containing 37.5 µl of PCR product, 0.5 µl AmpliTaq polymerase (Perkin Elmer), 10 mM Tris/HCl, pH 8.0, 50 mM KCl, 20 mM MgCl₂, 25 mM each dATP, dCTP, dGTP and dTTP, 100 nM each primer, 0.25 U AmpliTaq buffer, 1.8 M MgCl₂, 25 mM each dATP, dCTP, dGTP and dTTP, 100 nM each primer, 0.25 U AmpliTaq buffer, 1.8 M MgCl₂, 25 mM each dATP, dCTP, dGTP and dTTP, 100 nM each primer, 25 U AmpliTaq DNA polymerase (Perkin Elmer) and 0.2 ng template nucleic acid µl⁻¹. PCR reactions used the following parameters: presoak: 94 °C, 1 min; 35 cycles of 1 min each at 94, 55 and 72 °C; post-soak: 72 °C, 7 min. Amplification products (~1.5 kb) were visualized by gel electrophoresis (0.8% agarose, 0.5 × TBE, (Sambrook et al., 1989). Reaction products were purified and concentrated by ultrafiltration (Microcon YM100 spin microconcentrator; Amicon) prior to sequencing.
GGCACAGC), 704f (GGTGAATAGGCTAGA), 907r (CCGTCAATTGTTTTRAGTTT), 110r (AGGGTGGC-GCTCGTT) and 690r (TCTACGCATTTCACC) (Lane, 1991).

Phylogenetic analyses. Alignments were designed to maximize agreement with the proposed secondary structure for the Escherichia coli (Gutell, 1994). Agreement with this model was verified in new sequences by the occurrence of compensatory substitutions that preserve secondary structure in conserved internal helices. Gaps were inserted to compensate for length variation in identified loops and helices or to accommodate missing data. No additional gaps were added to increase primary sequence similarity among taxa. Sites within regions of uncertain alignment were identified and eliminated from further analyses. Alignments were edited using SeqLab (GCG version 10.1). The sequence of the T7902T 16S rDNA is on deposit in the GenBank sequence database (accession no. AY028398).

Fifty reference taxa were selected to represent the diversity within the gamma subclass of proteobacteria with emphasis on lineages with highest 16S rDNA sequence identity to the shipworm isolates, as identified through BLAST searches of GenBank and searches of the Ribosomal Database Project (RDP) database using Sequence Match version 2.7 (Maidak et al., 2001). Additional sequences of subjective interest (selected bivalve gill endosymbionts and taxa with known cellulolytic capabilities) were also considered.

Phylogeny reconstruction. Phylogenetic analyses were performed using algorithms contained in PAUP* (version 4.0b8) (Swofford, 1997). Maximum-parsimony, maximum-likelihood and evolutionary distance methods were used to infer tree topology. Maximum-likelihood trees were inferred using the HKY85 substitution model (Hasegawa et al., 1985) with ti:tv = 2, empirical base frequencies and equal rates for all sites. Maximum-parsimony analyses were performed with character states optimized by accelerated transformation (ACCTRAN). Evolutionary distance analyses used Kimura two-parameter distances the minimum evolution as the objective criterion and equal rates for all sites. In all analyses branch swapping was by tree-bisection-reconnection (TBR) with characters weighted equally (weight = 1) and gaps were treated as missing data. Bootstrap analyses were performed using the full heuristic search option with 1000 replicates for maximum-parsimony and evolutionary distance analyses. Bootstrap analyses under maximum-likelihood were limited to 100 replicates on a reduced taxa set (33 taxa) to accommodate the greater computational demands of this technique. Uncorrected pair-wise distance estimates for comparative purposes were estimated using Sequence Match version 2.7 and sequences selected from the Ribosomal Database Project (Maidak et al., 2001).

RESULTS
Phenotypic characteristics of shipworm isolates

Fifty-eight bacterial isolates similar to each other in morphology and growth characteristics were obtained from gill tissues dissected from 24 species of teredinid bivalves collected in various locations [see Table 1, plus additional data in supplementary data in IJSEM Online (http://ijsem.sgmjournals.org)]. All isolates were highly motile, slightly curved, rod-shaped cells of uniform dimensions (0.4–0.6 x 3–6 μm) bearing a single polar flagellum and granular inclusions near each cell apex when observed during exponential-phase growth in SBM medium with cellulose as a sole carbon source (Fig. 1). The granular inclusions are phase- and electron-dense and are consistent with polyphosphate in staining characteristics (staining with methylene blue but not Sudan black) and in appearance in transmission electron micrographs. All fix dinitrogen and display characteristics of microaerophilic growth when inoculated in soft agar stab cultures. All isolates display a distinctive colony morphology when grown on SBM cellulose with growth occurring as a diffuse inverted dome beneath the surface of the agar. Young colonies are translucent and difficult to discern while older colonies may produce a yellow-brown pigment and are surrounded by a broad halo of clearing due to cellulose hydrolysis. Shallow depressions may form on the surface of older colonies. These are most likely due to the removal of solid cellulose from the medium. The depressions do not deepen after clearing of the cellulose and there is no evidence that agar is hydrolysed. No growth is observed on agar in the absence of a suitable carbon source. Cell morphology varied for individual isolates on different carbon sources, ranging from motile rods to non-motile rods, helices, spheres, or pleomorphic forms. DNA G+C content for tested isolates ranged was 49–51 mol%.

Growth of all isolates is supported by cellulose (Whatman no. 1 filter paper or SigmaCell 100 cellulose powder) with or without an added source of combined nitrogen. Fourteen isolates tested (T7901–T7903, T8201–T8203, T8304, T8402, T8505, T8506, T8508, T8509, T8601 and T8602) also showed growth on carboxymethylcellulose (0.5%), cellulose (0.5%), fructose (0.5%), sucrose (0.5%), mannitol (0.5%), and xylan (0.5%). Most tested isolates also exhibit growth on sucrose (0.5%; except T8304), xylose (0.5%; except T8509), yeast extract (0.1%; except T7901), acetate (0.1%; except T7901, T8509 and T8602) and glucose (0.5%; except T8202, T8505, T8506, T8509 and T8601). A yellow pigment was produced by some isolates during growth on SigmaCell 100, Whatman no. 1 filter paper, cellulose and salicin, and acid production was observed by all isolates with growth on sucrose, fructose, xylose, glucose, pectin, salicin, cellulose, SigmaCell 100 and Whatman no. 1 filter paper. No growth was observed on dextrin, trehalose, melezitose, or in controls to which no carbon source was added.

The pH, temperature and salinity optima for growth have been determined for six isolates (T7901, T7902T, T7903, T8201, T8202 and T8203) and are approximately 8·5, 30–35 °C and 0·3 M NaCl respectively. No growth occurs above 39 °C. Specific growth rates measured at 35 °C in the presence and absence of combined nitrogen were 0·12 and 0·05 per hour, respectively. The isolates are marine, i.e. in addition to NaCl they require elevated concentrations of Ca2+ and Mg2+ that reflect the chemistry of seawater (Greene &
Freer, 1986; Waterbury et al., 1983). Strain T7902<sup>T</sup> exhibits catalase and oxidase activity.

**Phylogenetic analyses**

Nearly complete sequence (1453 bp) was determined for the 16S rRNA gene of strain T7902<sup>T</sup>. This sequence was indistinguishable from partial sequences (891 positions examined) determined previously for isolates T7902<sup>T</sup>, T8402, T8602, T7901 (GenBank accession nos M64338–40) by reverse transcriptase sequencing of purified rRNAs (Distel et al., 1991). The T7902<sup>T</sup> sequence was aligned (accounting for rRNA structure) with 50 published bacterial 16S sequences (taxon names, isolate and sequence accession numbers are listed in Fig. 2, and on-line supplementary material). A mask was employed to eliminate sites containing missing or ambiguous data and sites within regions of uncertain alignment, leaving a final character set of 1158 nucleotide positions, 376 of which are parsimony informative. Phylogenetic analyses were conducted using maximum-likelihood, maximum-parsimony and evolutionary distance methods.

A heuristic search by maximum-parsimony retained a single best tree of 2628 steps [consistency index (CI) = 0.3006, homoplasy index (HI) = 0.6994, retention index (RI) = 0.5782, and rescaled consistency index (RC) = 0.1738] (Fig. 2). Evolutionary distance analysis retained a single best tree with minimum evolution score of 210605. Maximum-likelihood analysis retained a single best tree with —ln likelihood = 15893.05033. Best trees inferred by all methods are topologically identical with respect to nodes supported by significant bootstrap proportions.

Trees inferred by all methods include shipworm strain T7902<sup>T</sup> within a clade containing an uncultivated symbiont from the shipworm Bankia setacea (Gen-
Bank no. AF102866) (Sipe et al., 2000), an unordered marine heterotrophic isolate from the Southern California coast (SCB11, GenBank no. Z31658) (Rehnstam et al., 1993), and an uncultivated bacterium from deep-sea sediments in Sugura Bay, Japan (1521 m; 34° 55’, 138° 39’) (BD2-12, GenBank no. AB015541) (Li et al., 1999). This clade (hereafter referred to as the Teredinibacter clade) receives significant bootstrap support (87–98%) by all inference methods. Three additional taxa are placed at the base of this clade in maximum-likelihood and maximum-parsimony analyses. These taxa include strain 2-40 (GenBank no. AF055269) (Gonzalez & Weiner, 2000) isolated from salt marsh cord grass Spartina alterniflora in the Chesapeake Bay, VA; an uncultivated bacterium (NKB4, GenBank no. AB013256) from deep-sea sediments near a cold-seep bivalve community in the Nankai Trough, Japan (3843 m; 33° 39’, 137° 56’) (Li et al., 1999) and Microbulbifer hydrolyticus str. IRE-31 (ATCC 700072, GenBank no. U58338) (Gonzalez et al., 1997) a marine heterotroph isolated from pulp mill effluents inoculated with water from a sea-grass bed (Sapelo Island, GA). The latter same a sister group to the group that contains the Teredinibacter clade+2-40 in evolutionary distance analyses. The groups that include the Teredinibacter clade+2-40 and the Teredinibacter clade+2-40+ NKB4+IRE-31 receive no significant bootstrap support in maximum-parsimony and evolutionary distance analyses and only weak support in the maximum-likelihood analysis. Uncorrected pair-wise identity values estimated for 16S rDNAs of strain T7902T and its closest described relatives, IRE-31 and 2-40, are approximately 89 and 91 % respectively.

Additional analyses using less complete sequences available in GenBank (data not shown) suggest that four additional sequences may be associated with the Teredinibacter clade. This group does not receive significant bootstrap support, however, resolution may be limited due in part to the small number of nucleotide positions determined for these taxa. These sequences include uncultivated symbionts (‘Candidatus Endobugula sertula’ BnSP (GenBank no. AF06606), BnPV (AF06607) and BnTP (AF06608)) from larvae of a bryozoan Bugula neritina, and an uncultivated bacterium (HstpL85.st, GenBank no. AF159682) detected on the surface of sea grass Halophila stipulacea.

**DISCUSSION**

All 58 strains isolated from gill tissue of shipworms were very similar to each other with respect to morphology, physiological characteristics and substrate utilization. Growth rates, temperature ranges and optima for growth, pH and salt requirements and G + C content were similar for all strains tested (T7901, T7902T, T7903, T8201, T8202 and T8203) (Greene, 1989; Waterbury et al., 1983) and 16S rRNA sequences were identical over 891 positions examined for four strains tested (T7901, T7902T, T8402 and T8602) (Distel et al., 1991). All strains were isolated from shipworm gill tissue at dilutions of at least 10⁻⁷–10⁻⁸ and all grow on cellulose as a sole carbon source and fix dinitrogen under microaerobic conditions.

The shipworm isolates are unique in several respects. First, of the many intracellular endosymbionts of marine invertebrates that have been identified by microscopic and molecular methods in the past quarter century, the shipworm isolates remain the only such bacteria to be grown in pure culture. The endosymbiotic status of the shipworm isolates has been confirmed by culture-based (Waterbury et al., 1983) and molecular methods (Distel et al., 1991).

Second, the shipworm isolates are phylogenetically unique. In the 20 years since their cultivation was first reported, no bacterium, cultivated or uncultivated, has been shown to be closely related to the shipworm isolates by molecular phylogenetic methods. While there are currently no accepted criteria for delimiting prokaryotic taxa based on comparative analyses of 16S rRNA sequences, comparisons of pair-wise sequence identities of ribosomal rDNAs are frequently reported and may be informative in cases where sequence differences are extreme. For example, it has been shown that bacteria that share less than 97.5% sequence identity typically do not show greater than 60–70% DNA similarity by a variety of reassociation methods and so are unlikely to be related at the species level (Stackebrandt & Goebel, 1994). Searches of GenBank and RDP databases using BLAST and Sequence Match version 2.7, followed by alignment and estimation of pair-wise identity using Similarity Matrix 1.1 (RDP) demonstrate that the closest cultivated and characterized relative of strain T7902T shares only about 91% identity in its 16S rDNA sequence. For comparison, all 29 species included in the genus Pseudomonas sensu stricto (Anzai et al., 1997, 2000) share greater than 93.3% identity and species of Escherichia, Erwinia, Enterobacter, Citrobacter, Klebsiella, Proteus, Salmonella, Serratia, Yersinia, Xenorhabdus and many other genera within the family Enterobacteriaceae share greater than 91% identity by the same estimation method. Thus, T7902T differs from its nearest described relatives by values substantially exceeding those that separate many well-defined genera.

Third, the shipworm isolates are physiologically unique. Their ability to grow with cellulose as a sole carbon source and to fix nitrogen is a rare combination. Similar capabilities have been reported in several strictly anaerobic Gram-positive bacteria, e.g. Clostridium hungatei (Monserrate et al., 2001). Functional endoglucanase genes have also been identified in dinitrogen-fixing, rhizosphere-associated bacteria from the z-Proteobacteria, e.g. Azospirillum irakense (Faure et al., 2001) and Azorhizobium caulindos (Geelen et al., 1995), and beta proteobacteria, e.g. Azorarcus sp. (Reinhold-Hurek et al., 1993). However, these bacteria do not grow with cellulose as a sole carbon source and probably use their cellulolytic
Teredinibacter turnerae; endosymbiont of shipworms

capabilities as a means to penetrate plant tissues. To our knowledge, the shipworm isolates are the only Gram-negative bacteria described to date that fix dinitrogen and grow with cellulose as a sole carbon source, and the only bacterium that combines these capabilities to grow under (micro)aerobic conditions.

Fourth, the shipworm isolates are unusual in terms of their distinctive habitat. This bacterium has only been cultivated from gill tissues of wood-boring marine bivalves of the family Teredinidae (shipworms) where it exists as an endosymbiont. Attempts to cultivate this bacterium from other tissues of shipworms, from wood surfaces and from seawater surrounding or within shipworm burrows have been unsuccessful (Waterbury et al., 1983). This contrasts with cultivable symbionts of other marine invertebrates, e.g. the Vibrio fischeri and V. logei strains that exist extracellularly in the lumen of the light organ crypts of squid but that also can be cultivated easily from water in which the hosts have been maintained (Nishiguchi, 2000). The ability of T7902T to exist as an endosymbiont in a specific animal group is in itself a highly unusual phenotype.

Of the taxa identified here as potential affiliates of the shipworm isolate T7902T, only two (marine bacterium strain 2-40; ATCC 43961T and Microbulbifer hydrolyticus str. IRE-31 ATCC 700072T) have been cultivated and characterized physiologically. Strain 2-40 is a strictly aerobic marine heterotroph capable of hydrolysing a broad range of complex polysaccharides, including agar, carboxymethylcellulose (but not cellulose), chitin, laminarin and xylans (Gonzalez & Weiner, 2000). The isolation of cellulose-hydrolysing bacteria from gill tissues of wood-boring marine bivalves of the family Teredinidae, we propose a new generic and species, Teredinibacter turnerae gen. nov., sp. nov., with T7902T as the type strain. Three additional strains (Table 1), which are indistinguishable from strain T7902T with respect to 16S rRNA sequence, are also included within the proposed species. Data available to date suggest that the remaining strains, described herein and in supplementary data in IJSEM Online (http://ijsem.sgmjournals.org), will fall within the new species upon more complete characterization.

Description of Teredinibacter gen. nov.

Teredinibacter [Ter.e din.i bac'ter. N.L. fem. pl. n. Teredinidae a family of wood-boring bivalve molluscs (shipworms); N.L. masc. n. bacter, equivalent of bacterium, a staff or rod; N.L. masc. n. Teredinibacter a rod isolated from members of the family Teredinidae].

Cells are rigid, Gram-negative rods of 0.4–0.6 µm in width and 3–6 µm in length during exponential-phase growth on SBM medium with cellulose as a carbon source. Division is by binary fission in a single plane. Motile by means of single polar flagellum. In stationary-phase cultures, cells often become pleomorphic, appearing spiraled or as very long rods. Occur in nature as long rods associated with the gill system of...
Table 2. Characteristics of *Teredinibacter turnerae* and related bacteria

<table>
<thead>
<tr>
<th>Trait</th>
<th><em>Teredinibacter turnerae</em></th>
<th>Strain 2-40</th>
<th><em>Microbulbifer hydrolyticus</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell shape</td>
<td>Rods</td>
<td>Pleomorphic rods</td>
<td>Rods</td>
</tr>
<tr>
<td>Cell length</td>
<td>3–6 µm</td>
<td>1–20 µm</td>
<td>1–1–1·7 µm</td>
</tr>
<tr>
<td>Cell arrangement</td>
<td>Single</td>
<td>Single, chains</td>
<td>Single, chains</td>
</tr>
<tr>
<td>Flagellum</td>
<td>Single, polar</td>
<td>Single, polar</td>
<td>None</td>
</tr>
<tr>
<td>Surface nodules</td>
<td>–</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Salt requirement for growth</td>
<td>Marine</td>
<td>Marine</td>
<td>Marine</td>
</tr>
<tr>
<td>Vitamin requirement</td>
<td>–</td>
<td>B cofactors</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td></td>
<td>stimulatory</td>
<td></td>
</tr>
<tr>
<td>Growth on monosaccharides</td>
<td>+</td>
<td>+</td>
<td>Limited</td>
</tr>
<tr>
<td>Hydrolysis of:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cellulose</td>
<td>+</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>Agar</td>
<td>–</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>Nitrogen fixation</td>
<td>+</td>
<td>?</td>
<td>–</td>
</tr>
<tr>
<td>Habitat</td>
<td>Endosymbionts of teredinid bivalves</td>
<td>Free-living, salt marsh</td>
<td>Free-living, salt marsh</td>
</tr>
<tr>
<td>Mol% G+C</td>
<td>49–51</td>
<td>46·7</td>
<td>57·6</td>
</tr>
</tbody>
</table>

Teredinid molluscs. Aerobic chemoheterotrophs that grow in a simple mineral medium containing seawater and a suitable carbon and energy source (e.g. cellulose, simple sugars and organic acids). Require a source of combined nitrogen if vigorously aerated but will fix dinitrogen when grown in the absence of combined nitrogen under microaerobic conditions. Analysis of their 16S rDNA indicates that the genus belongs to a unique clade of the γ-Proteobacteria. Type species is *Teredinibacter turnerae*.

**Description of *Teredinibacter turnerae* sp. nov.**


In addition to the characteristics of the genus, the species characterization is based on the properties of the type strain (T7902) and three additional strains (T7901, T8402 and T8602<sup>®</sup>) isolated from phylogenetically divergent species and genera of teredinid molluscs collected in distantly separated locations in the Atlantic and Pacific Oceans. All isolates grew on cellulose with or without an added source of combined nitrogen and also demonstrated growth on carboxymethylcellulose, cellobiose, fructose, salicin, glutamate, succinate, pyruvate, pectin, xylan, sucrose, xylose, yeast extract and glucose. All but T8604 also grew on acetate. Acid was produced by all isolates when grown on sucrose, fructose, xylose, glucose, pectin, salicin, cellobiose and cellulose. The temperature range for growth is approximately 20–30 °C, with optimal growth occurring at 30–35 °C. No growth is observed at 15 or 40 °C. The optimum pH for growth is 8·5 and the pH range for growth is 6·0–10·5. The NaCl range for growth is 0·1–0·6 M, and optimal growth occurs at 0·3 M. Growth also requires elevated concentrations of Ca<sup>2+</sup> and Mg<sup>2+</sup> that reflect the chemistry of seawater. Growth factors are not required. T7902<sup>®</sup> exhibits oxidase and catalase activity. No growth was observed under anaerobic conditions on SBM agar with cellulose as the carbon source. Colonies are translucent with all growth occurring beneath the agar surface. As colonies age a yellow-brown pigment is observed and clearing occurs around the colonies due to cellulose hydrolysis. The G+C content of the DNA is 49–51 mol%. The type strain T7902<sup>®</sup> (= ATCC 39867<sup>T</sup> = DSM 15152<sup>T</sup>) was isolated from *Lyrodus pedicellatus* originally collected from Long Beach, CA, USA, but kept in culture at Woods Hole Oceanographic Institution for many years.

**ACKNOWLEDGEMENTS**

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**REFERENCES**


