Meganema perideroedes gen. nov., sp. nov., a filamentous alphaproteobacterium from activated sludge

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An industrial wastewater treatment plant at Grindsted, Denmark, has suffered from bulking problems for several years caused by filamentous bacteria. Five strains were isolated from the sludge by micromanipulation. Phylogenetic analysis of the 16S rRNA gene sequences showed that the strains formed a monophyletic cluster in the Alphaproteobacteria, and they were phenotypically different from their closest relatives and from all hitherto known filamentous bacteria described (closest relative Brevundimonas vesicularis ATCC 11426T, 89·8 % sequence similarity). In pure culture, the cells (1·5–2·0 μm) in filaments are Gram-negative and contain polyphosphate and polyhydroxyalkanoates. The optimum temperature for growth is 30°C and the strains grow in 2 % NaCl and are oxidase- and catalase-positive. Ubiquinone 10 is the major quinone. The major fatty acid (C18:1ω7c) and smaller amounts of unsaturated fatty acids, 3-hydroxy fatty acids with a chain length of 16 and 18 carbon atoms and small amounts of 10-methyl-branched fatty acids with 18 carbon atoms (C18:0 10-methyl) affiliated the strains with the Methylobacterium/Xanthobacter group in the Alphaproteobacteria. The G+C content of the DNA is 42·9 mol% (for strain Gr1T). The two most dissimilar isolates by 16S rRNA gene comparison (Gr1T and Gr10; 97·7 % identical) showed 71·5 % DNA–DNA relatedness. Oligonucleotide probes specific for the pure cultures were designed for fluorescence in situ hybridization and demonstrated that two filamentous morphotypes were present in the Grindsted wastewater treatment plant. It is proposed that the isolates represent a new genus and species, Meganema perideroedes gen. nov., sp. nov. The type strain of Meganema perideroedes is strain Gr1T (= DSM 15528T = ATCC BAA-740T).

Filamentous bacteria affiliated to the Alphaproteobacteria were isolated from activated sludge biomass from a wastewater treatment plant (WWTP) in Grindsted, Denmark. The treatment plant has carbon removal, the mean cell residence time (sludge age) is 8–10 days, the annual mean temperature is 15–25°C and the salinity is relatively high, with a conductivity between 15 and 23 mS cm−1 and a sodium concentration around 5–7 g l−1. The plant has been affected by bulking problems for several years, mainly caused by Thiothrix spp. (Nielsen et al., 2000) and other bacteria similar to the isolates reported in this study. Five strains (Gr1T, Gr2, Gr5, Gr10 and Gr28) were isolated by micromanipulation using a Skerman micromanipulator as described by Blackall (1991) and grown on R2A agar medium (Reasoner & Geldreich, 1985). Sliding motility, Gram- and Neisser-staining (for Gram-reaction and polyphosphate inclusions, respectively; Eikelboom & van Buijsen, 1983) and poly-β-hydroxybutyrate (PHB) inclusions as visualized by Nile blue staining (Ostle & Holt, 1982) were investigated. Only Gr1T and Gr10 were characterized phenotypically, but they were not able to grow on minimal media (Williams & Unz, 1985), so a full characterization of the growth requirements was not possible. Biolog analysis was done according to the manufacturer’s description (http://www.biolog.com/main.html). However, reproducible results could not be obtained, so the results are therefore

Abbreviations: FISH, fluorescence in situ hybridization; PHB, poly-β-hydroxybutyrate; WWTP, wastewater treatment plant.

The GenBank/EMBL/DDBJ accession numbers for the 16S rRNA gene sequences of strains Gr1T, Gr2, Gr5, Gr10 and Gr28 are AF180468 and AY170117–AY170120.

Interstrain 16S rRNA gene sequence similarities, confocal laser scanning micrographs of FISH of activated sludge from Grindsted WWTP and a phase-contrast micrograph of filaments of strain Gr1T are available as supplementary material in IJSEM Online.
not described here. The temperature range and salt tolerance for growth were also studied. The results obtained are given in the species description below.

Phylogenetic analysis was performed on the 16S rRNA gene as described by Blackall (1994). Nearly complete 16S rRNA gene sequences (Gr1T, 1383 bp; Gr2, 1284 bp; Gr5, 1208 bp; Gr10, 1234 bp; Gr28, 1214 bp) were compiled and aligned using the automatic nucleic acid aligner in the ARB software package (http://www.arb-home.de) and alignments were refined manually. A total of 1208 unambiguously aligned nucleotides from position 70 to 1277 were used for calculation of trees by distance matrix, parsimony and maximum-likelihood approaches using default settings in the ARB software. The methods used generated congruent tree topologies and a maximum-likelihood tree is shown in Fig. 1. The five isolates always formed a monophyletic cluster within the Alphaproteobacteria, and the 16S rRNA gene sequence similarity in this cluster ranged between 97-7 and 99-6 % (for individual values see Supplementary Table S1 in IJSEM Online). Strains Gr1T and Gr10 and Gr1T and Gr2 were the most dissimilar strains by 16S rRNA gene comparison, showing 97-7 % identity. The closest relative was Brevundimonas vesicularis LMG 2350T, which showed 89-8 % identity with strain Gr1T. Levantesi et al. (2004) reported on other filamentous alphaproteobacteria. They are so far described only by 16S rRNA gene sequences from uncultured bacteria or from isolates that have not yet been described. These are all closely related to the novel strains.

The detailed results of chemotaxonomic analysis are given in the species description. The following analytical procedures were performed. Respiratory quinones and cellular fatty acids were analysed at the DSMZ according to Tindall (1990a, b) and Sasser (1990). Ubiquinone 10 (Q-10) was the major quinone component in strain Gr1T and Gr10, supporting the affiliation of Gr1T to the Alphaproteobacteria, where the majority of the species have Q-10 as the major quinone (Lechner et al., 1995; Yokota et al., 1992). Both strains synthesized mainly straight-chain saturated and unsaturated fatty acids and 3-hydroxy fatty acids with a chain length of 16 and 18 carbon atoms. Small amounts of 10-methyl-branched fatty acids with 18 carbon atoms (C19:0 10-methyl) were also found. According to the fatty acid patterns, the two strains were assigned to the Methylbacterium/Xanthobacterium group within the Alphaproteobacteria.

The G+C content was determined at the DSMZ according to a standard protocol (Mesbah et al., 1989; Tamaoka & Komagata, 1984). The G+C content of DNA of strain Gr1T was 42.9 mol%. DNA–DNA hybridization was carried out at the DSMZ as described by De Ley et al. (1970), with the modifications described by Huß et al. (1983) and Escara & Hutton (1980). Strain Gr1T and Gr10 had high DNA–DNA relatedness (71-5 %), supporting the conclusion that the two strains can be regarded as members of one species.

Oligonucleotide probes specific for the isolated filaments were designed using the probe design/match tools of the ARB software package (http://www.arb-home.de; Hugenholtz et al., 2002). Fluorescence in situ hybridization (FISH) was carried out as described by Amann (1995) and visualized with a Zeiss LSM 510 (Carl Zeiss) confocal laser scanning microscope. Oligonucleotide probes were 5′-labelled with 5(6)-carboxyfluorescein-N-hydroxy-succinimide ester (FLUOS) or with sulfoindocyanine dyes (Cy3 and Cy5) (Thermo Hybaid). Probe-binding profiles of the designed probes (Meg983 and Meg1028) for FISH were determined using image analysis and with hybridization buffers containing 0 to 60 % formamide at 5–10 % increments (Amann, 1995), initially employing the isolates (perfect probe matches) as probe target organisms and Streptococcus salivarius ATCC 7073T (one central mismatch with probe Meg983) and Escherichia coli ATCC 25922 (more than seven mismatches with Meg983 and Meg1028) as non-target organisms. Two probes were designed for the strains isolated in this study: S-Ss-Meg-983-a-A-19 (Meg983), 5′- CCGGATGTCAAAAGGTGG-3′, and S-Ss-Meg-1028-a-A-19 (Meg1028), 5′-CTGTACCGAGTCCCTTGC-3′. Both probes targeted the novel strains perfectly, as well as the strains described by Levantesi et al. (2004). Probe binding profiles of hybridized pure cultures (Gr1T, Gr2 and Gr10) and Grindsted sludge samples showed probe Meg983 hybridized well up to 35 % formamide, while the intensity of probe Meg1028 decreased above 50 % formamide. Only one clone, clone oc48 (GenBank accession no. AY491589), was found using ARB and RDP (http://rdp.cme.msu.edu/html) with a weighed mismatch below 1.5 to probe Meg1028, indicating that the probe is very specific. Probe Meg983 was less specific so, for in situ identification, it is recommended to use the two probes in combination with two different fluorochromes at 35 % formamide or to use

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**Fig. 1.** Maximum-likelihood tree based on 16S rRNA gene sequences of strains Gr1T and Gr10 and related organisms calculated using default settings in the ARB software. Bar, 10 % estimated sequence divergence.
probe Meg1028 only (at 45 % formamide). *E. coli* ATCC 25922 was used as a non-target organism for both probes, and positive hybridization signals were never observed. Micrographs from FISH of activated sludge from Grindsted WWTP are available as Supplementary Fig. S1 in IJSEM Online. The application of the specific 16S rRNA-targeted oligonucleotide probes allowed the monitoring of sludge samples from Grindsted WWTP for the novel organism (*Meganema perideroedes*). In all samples from the past 3 years, *Meganema perideroedes* filaments predominated in the sludge. *Thiothrix* and other filamentous bacteria were present in moderate numbers. Morphological observations, staining reactions and FISH of the Grindsted sludge sample showed the presence of two variations of *Meganema perideroedes*. Filaments of type A were about 1-5 μm wide, some filaments were Gram-positive, others were Gram-negative and filaments containing both Gram-positive and Gram-negative cells were also found. Some of the filaments were Neisser-negative (no purple granules inside red cells), while others contained Neisser-positive granules (purple inclusions) and thus contained polyphosphate. The cells contained large PHB granules according to Nile blue staining. Type B filaments were slightly wider than those of type A and they showed Gram-variable staining similar to type A. PHB granules were smaller in type B filaments than in type A filaments. The isolated strains were very similar to type A filaments from Grindsted. Thus, the isolated strains look like the Gram-variable filamentous bacteria in activated sludge with the ‘*Nostocoida limicola*’ type II morphology described by Eikelboom & van Buijsen (1983). This morphotype was also observed in previous studies (Snaird et al., 2002; Schade et al., 2002; Blackall et al., 2000; Levantesi et al., 2004). Our results confirm the observation that different filaments from the ‘*N. limicola*’ type II morphotype are phylogenetically heterogeneous, emphasizing that accurate identification in situ is impossible without the application of gene probes. Nevertheless, the specific gene probes designed for the *Meganema perideroedes* sequences were not able to differentiate between the type A and type B morphotypes observed in sludge from Grindsted WWTP.

**Description of Meganema gen. nov.**

*Meganema* (Me.ga.ne’ma. Gr. nom. neut. adj. mega big; Gr. nom. neut. n. *nema* thread; N.L. neut. n. *Meganema* large thread-like micro-organism).

Filamentous bacteria. Cells stain Gram-negative and contain some Neisser-positive granules and PHB granules. The major quinone is Q-10. Fatty acid profiles are dominated by C18:1ω7c; characteristic hydroxy acids are C14:0 3-OH and C18:0 3-OH. *Meganema* belongs to the *Alphaproteobacteria* and the type species is *Meganema perideroedes*.

**Description of Meganema perideroedes sp. nov.**

*Meganema perideroedes* (pe.ri.de.ro’ed.es. Gr. nom. neut. adj. *perideroedes* necklace-like).

The species shares the characteristics listed in the genus description. The DNA G+C content of the type strain is 42.9 mol%. Habitat: mixed liquors of activated sludge WWTP. Although all isolates grow on R2A medium, some (Gr1T, Gr2 and Gr10) grow well, taking only a few days to produce macroscopically visible colonies, which are dense and white. Storage of strains Gr5 and Gr28 was unsuccessful and the strains were lost after sequencing of their 16S rRNA genes. Strains Gr1T, Gr2 and Gr10 demonstrate a very similar morphology in that the cells in all filaments are irregularly disc-shaped and are approximately 1.5–2 μm in diameter (see Supplementary Fig. S2); they show no gliding motility. Gram- and Neisser-staining shows that they are Gram-negative and contain Neisser-positive granules. Only strains Gr1T and Gr10 have been characterized phenotypically; they are not able to grow on minimal media, so a full characterization of the growth requirements has not been possible. They are not able to grow anaerobically with nitrate as an electron acceptor. Starch and tributyrin are not hydrolysed. Oxidase and catalase activities are present. The temperature range for growth is 15–35 °C and the optimum temperature for growth is 25–30 °C. Both strains are able to grow in the presence of NaCl concentrations up to 2 %, but do not grow in the presence of 3 % NaCl. The fatty acid profile of the type strain Gr1T is composed of C18:1ω7c (86–4 %), C18:0 (3–8 %), summed feature 2 (C14:0 3-OH) (2–4 %), C16:0 (2–9 %), C18:0 3-OH (2–3 %) and C19:0 10-methyl (1–1 %). The profile of strain Gr10 is almost identical.

The type strain is strain Gr1T (=DSM 15528T = ATCC BAA-740T), isolated by micromanipulation from sludge of an industrial WWTP at Grindsted, Denmark. There are currently three extant strains in the species.

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**References**


