Schineria larvae gen. nov., sp. nov., isolated from the 1st and 2nd larval stages of Wohlfahrtia magnifica (Diptera: Sarcophagidae)

Erika Tóth, Gábor Kovács, Peter Schumann, Attila L. Kovács, Ulrike Steiner, András Halbritter† and Károly Márialigeti†

Four bacterial strains were isolated from the fly larvae of an obligate parasitic fly, Wohlfahrtia magnifica (Diptera: Sarcophagidae). These isolates were characterized by a polyphasic approach and represent a new lineage of \( \gamma \)-Proteobacteria as their closest relative is Xylella fastidiosa (87.1% 16S rDNA similarity). The four strains are identical at the 16S rDNA level, the level of similarity between them, based on DNA–DNA hybridization, is high (97–102.5%) and they are similar in their physiological and biochemical characteristics, although they differ in their utilization of different sole carbon sources. All produce chitinase. They are obligately aerobic; no growth is detected under anaerobic conditions, even in the presence of \( \text{NO}_3^- \) as terminal electron acceptor. Their predominant respiratory quinone is Q-8. The G+C content of their DNA is 42 mol%. Their cell membrane contains phosphatidylglycerol, diphosphatidylglycerol, phosphatidylethanolamine, phosphatidylserine and two unknown polar lipids. Their main fatty acids are C₁₈:1ω9c, C₁₆:0, C₁₄:0 and C₁₆:1ω6c. To accommodate these bacteria, a new genus, Schineria gen. nov., with the type species Schineria larvae sp. nov., is proposed.

Keywords: Schineria larvae, myiasis, fly, \( \gamma \)-Proteobacteria, taxonomy

INTRODUCTION

It is generally accepted that one of the basic resources for bacterial species diversity is the world of insects. They are speculated to harbour thousands of novel bacterial symbionts (parasites, mutualistic partners or protoco-operants). This supposition seems to be corroborated by numerous publications (Cruden & Markovetz, 1984; Ishikawa, 1989; Groombridge, 1992; Tanada & Kaya, 1993; Cazemier et al., 1997) and recent investigations on traumatic myiasis support this opinion (Khoga, 1994; Tóth et al., 1998). Myiasis is a disease of vertebrate animals caused by different fly larvae. The obligate parasitic fly (developing on/in the tissues of living animals, like sheep, goats, camels, geese, turkeys, etc.) Wohlfahrtia magnifica (Diptera: Sarcophagidae) (Schiner, 1862) is responsible for serious losses in animal husbandry in Eurasia (Martinez et al., 1987, 1991; Hall & Wall, 1995; Farkas et al., 1997; Hall, 1997; Valentin et al., 1997). There is scant information about the aetiology of the disease. During studies in the field, detailed bacteriological investigations have been carried out on the different developmental stages of the fly (1st, 2nd, 3rd stage larvae, pupae and imago). Among the bacterial strains isolated from the fly larvae, some were found to strongly utilize chitin and represent a separate phylotypic line of descent. The isolation, characterization and taxonomic delineation of strains L1/68T, L1/57, L1/58 and L2/11 are described in this paper. We propose a new genus, Schineria gen. nov. and the species Schineria larvae sp. nov. to accommodate these bacteria.
METHODS

Isolation and cultivation. Strains L1/68\textsuperscript{r}, L1/57 and L1/58 were isolated from whole larval magerate of 1st stage larvae and L2/11 was isolated from 2nd stage larvae of *Wohlfahrtia magnifica* (Schiner, 1862) as follows. Larvae were collected from the vulval wound of Romney breed sheep at Mezőfalva State Farm (18° 40' E, 46° 50' N), Hungary, in 1997. They were identified both to species level and for developmental stage by stereomicroscopic investigation. They were washed three times in sterile 0.025 M sodium phosphate buffer solution (pH 6.8) and then homogenized in 0.025 M sodium phosphate buffer under aseptic conditions. All bacterial strains were isolated, grown and later subcultured and maintained on King B agar medium (King, 1954).

Morphology. Colony morphology was tested on King B agar medium by direct and stereomicroscopic observations of single colonies. Cell morphology and motility were studied by phase-contrast microscopy of native preparations. Gram staining was performed according to Claus (1992), spore staining according to Smibert & Krieg (1994) and capsules were visualized by negative staining according to Duguid (1951). Bacteria were prepared for ultrastructural study by transmission electron microscopy (TEM) as follows. Cells were grown on King B agar plates for 24 h at 28 °C, then fixed in 1% glutaraldehyde solution buffered with sodium cacodylate (0.1 M, pH 7.2) for 2 h at room temperature. The fixed samples were subsequently embedded in 2% agar and washed three times in sodium cacodylate. Post-fixation was carried out in cacodylate buffered 0.5% OsO\textsubscript{4} solution for 1 h. After staining with uranyl acetate (1% in distilled water, 30 min) the samples were dehydrated in a graded ethanol series (50, 70, 90, 96, 100%) followed by propylene oxide and embedded in Durcupan (Fluka). After polymerization the resin-embedded samples were cut with a Reichert ultramicrotome, the sections stained with lead-citrate and examined in a JEM100CX II microscope (JEOL).

Physiological and biochemical characterization. Growth in the presence of oxygen was tested using slant agar cultures on King B medium incubated in an anaerobic chamber (Forma Scientific). Oxidase activity was checked by the method of Tarrand & Gröschel (1982), and catalase production and Voges–Proskauer reaction were demonstrated by the standard methods of Cowan & Steel (1974). Growth on synthetic medium was tested by using the medium of Droop (1969) and on semi-synthetic medium using PCa agar containing 2 g peptone, 0.2 g MgSO\textsubscript{4}, 7H\textsubscript{2}O, 0.15 g CaCl\textsubscript{2}. Acid production from different carbohydrates was studied by the OF test according to Hugh & Leifson (1953) and using API CH50 test strips (CHE inoculation fluid; bioMérieux). Growth at different temperatures (4, 28, 37 and 45 °C), NaCl tolerance (2-5 %, w/v) and pH tolerance (2, 4, 6, 8, 9, 10 and 11) were determined using King B agar or broth. Urease activity, reduction of NO\textsubscript{2} to NO\textsubscript{3}, aesculin and starch hydrolysis, indole production from tryptophan, casease, gelatinase, phosphatase activity and H\textsubscript{2}S production from cysteine were studied according to Smibert & Krieg (1994). Haemolysin production was tested on ox-blood agar plates (Elek & Levy, 1954). Utilization of citrate was checked according to Simmons (1926) and hydrolysis of chitin was studied by the modified method of The-Holding & Collée (1971) (instead of a nutrient basal medium, King B agar was used). Utilization of 95 different carbon sources as sole carbon source was tested on BIOLOG GN-plates. To evaluate the results System Release 3.5 was used, following the manufacturer’s instructions. Mannose and arabinose utilization as sole carbon source was tested according to Gordon & Mihm (1957). Acetylene reduction activity was studied with cappled cultures of bacteria grown on semisolid RBA medium (DSMZ-Medium 441) without yeast extract. One-tenth volume of acetylene was added to the atmosphere 36 hours after inoculation and the amount of ethylene was measured immediately after the addition of acetylene, then 3, 6, 12 and 24 h later using a Hewlett Packard 5890A gas chromatograph equipped with a flame ionization detector and a packed metal column [6 ft x 1/8 in (1.83 m x 3.2 mm)] stainless steel, 80/100 HayeSepN. The flow rate of the carrier gas (nitrogen) was 20 ml min\textsuperscript{-1}. Injectors, column and detector temperatures were 100, 100 and 165 °C, respectively (Hardy et al., 1973).

Chemotaxonomic characterization. Cells for chemotaxonomic analysis were grown in liquid rich medium (Yamada & Komagata, 1972) to the middle of their exponential phase on a rotary shaker at 28 °C. Isoprenoid quinones were extracted as described by Collins et al. (1977) and the profile was analysed by HPLC (Groth et al., 1997). Cellular fatty acids were extracted according to Stead et al. (1992) and analysed by GC (Groth et al., 1996). Polar lipids were determined according to the method described by Minnikin et al. (1979).

Determination of the G+C content of DNA. DNA was isolated and its G+C content was determined by HPLC analysis of derived deoxyribonucleosides as described by Groth et al. (1996).

DNA–DNA hybridization. DNA–DNA hybridizations between each of the four strains were performed as described by Maszenan et al. (1999). 16S rDNA sequence determination and phylogenetic analysis. Genomic DNA extraction and PCR-mediated amplification of the 16S rDNA of the selected bacterial strains were carried out by the method of Rainey et al. (1996). The amplified PCR products were purified and concentrated by using a Prep-A-Gene kit (Bio-Rad). Cycle sequencing was performed in a Gene-Amp 2400 PCR machine (Perkin Elmer) with the Big Dye Terminator Cycle Sequencing kit, according to the protocol of the manufacturer. Base sequences were determined in an Applied Biosystems model 310 Genetic Analyser (Perkin Elmer). The 16S rDNA sequences of L1/68\textsuperscript{r}, L1/57, L1/58 and L2/11 were aligned against the ARB-formatted RDP Release 7.1 (Maidak et al., 1997) using the ARB programme package (Strunk & Ludwig, 1995). To check for more recent updates an additional BLAST search (Altschul et al., 1997) was also performed. The methods of Jukes & Cantor (1969) and the Kimura 2-parameter method were used to calculate evolutionary distances. Phylogenetic analysis was carried out applying various treeing algorithms (neighbour-joining, maximum-lihood, parsimony and DeSote tree fit) integrated into the ARB program. The consensus phylogenetic dendrogram was reconstructed from Jukes–Cantor evolutionary distances based on the neighbour-joining method using Bacterillus subtilis as root.

Nucleotide accession numbers. The strain designations and accession numbers of nucleotide sequences used in the phylogenetic analyses are as follows: *Bacterillus subtilis* D26185; *Cardiobacterium hominis* ATCC 15826\textsuperscript{r}, M53014; *Comamonas testosteroni* ATCC 11996\textsuperscript{r}, M11224; *Desulfovibrio desulfuricans* ATCC 27774\textsuperscript{r}, M34113; *Escherichia coli* K-12, U00096; *Frateria aurantia* IFO 3245\textsuperscript{r}, A010481; *Lysobacter antibioticus* DSM 2077\textsuperscript{r}, AB019582; *Proteus vulgaris* IFAM 1731, X07652; *Rhodanobacter lindanilasticus* LMG 18385\textsuperscript{r}, AF039167; *Rhodospirillum rubrum*, X87278; *Sphingomonas paucimobilis* ATCC 29837\textsuperscript{r}.
RESULTS

Culture characteristics and colony morphology

Growth of the bacterial strains isolated from the larvae of *Wohlfahrtia magnifica* was optimal on King B agar medium under aerobic conditions, while growth was weak on the semi-synthetic PCa medium and no growth appeared on synthetic medium (Droop, 1969). Isolates did not grow on media containing mannose or arabinose as sole carbon source (Gordon & Mihm, 1957), although the utilization of the aforementioned carbon sources on the BIOLOG microplate was

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**Table 1** Differences in the main biochemical and physiological characteristics of *Wohlfahrtia* strains

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>L1/68&lt;sup&gt;T&lt;/sup&gt;</th>
<th>L1/57</th>
<th>L1/58</th>
<th>L2/11</th>
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<tbody>
<tr>
<td>Oxidase</td>
<td>+</td>
<td>+</td>
<td>−</td>
<td>+</td>
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<tr>
<td>Catalase</td>
<td>±</td>
<td>+</td>
<td>±</td>
<td>+</td>
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<td>Growth at:</td>
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<tr>
<td>4 °C</td>
<td>±</td>
<td>+</td>
<td>±</td>
<td>+</td>
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<tr>
<td>45 °C</td>
<td>±</td>
<td>+</td>
<td>±</td>
<td>+</td>
</tr>
<tr>
<td>Phosphatase</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>H&lt;sub&gt;2&lt;/sub&gt;S from cysteine</td>
<td>±</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Tween 80 hydrolysis</td>
<td>−</td>
<td>±</td>
<td>−</td>
<td>−</td>
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<tr>
<td>Chitinase activity</td>
<td>+ + +</td>
<td>+</td>
<td>+</td>
<td>+ + +</td>
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</tbody>
</table>

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*Fig. 1.* Longitudinal section of bacteria from strain L1/68<sup>T</sup> as depicted by TEM. Cells are regularly rod-shaped with typical Gram-negative cell wall structure. Note invaginations of the inner membrane and the ring-like structure in the longer cell. Bar, 0.5 µm.

*Fig. 2.* Transmission electron micrograph of a dividing strain L1/68<sup>T</sup> cell. Note the deep invagination in the upper and ring-like structure in the lower daughter cell. Bar, 0.5 µm. The insert clearly shows that the inner membrane is continuous with the membrane of the invagination. Bar, 0.1 µm.
variable among the examined strains. All strains formed bright, easily spreadable, entire, convex, translucent, small colonies (1–3 mm diameter) on the surface of King B agar medium. No distinctive colony colour or diffusible pigments were observed.

Cell morphology, staining properties and ultrastructure

On King B medium 24-h-old cultures of strains L1/68T, L1/57, L1/58 and L2/11 were composed of regular rods (2–3 µm x 0.8–0.9 µm) occurring singly. In ageing cultures (48 h–1 week) no characteristic shape shifts were detected. Cells stained Gram-negative and were non-motile. Endospores were not formed. Capsules were detected. Electron micrographs of thin sections of strain L1 showed the typical Gram-negative cell wall structure. Some of the invaginations in appropriate sections may appear as finger-like intrusions. Fig. 2 and the insert clearly show that the membrane of a deep invagination is continuous with the cytoplasmic membrane. The frequently occurring ring-like structures (Figs 1 and 2) are most likely to be cross sections of the finger-like intrusions.

Physiological and biochemical characteristics

Bacterial strains L1/68T, L1/57, L1/58 and L2/11 grew under strictly aerobic conditions. No growth was observed for all strains except strain L1/57, where a weakly positive reaction could be detected. This latter trait was also proven by the results of Tween 80 utilization as sole carbon source in the BIOLOG GN test. Differences in the basic biochemical characteristics of the strains are summarized in Table 1. The results of utilization of different sole carbon sources showed that there were 22 different carbon sources which always gave negative reactions in all strains (α-cyclodextrin, glycerogen, d-arabitol, cellubiose, i-erythritol, gentiobiose, α-D-lactose, α-D-lactose-lactulose, xylitol, d-galacturonic acid lactone, d-glucosaminic acid, α-ketovaleric acid, propionic acid, quinic acid, D-saccharic acid, sebacic acid, L-threonine, Dl-carnitine, phenylethylamine, putrescine, 2-amino-ethanol and 2,3-butanediol). In the case of 14 carbon sources (monomethylsuccinate, cis-aconitic acid, citric acid, γ-hydroxybutyric acid, α-ketogluutaric acid, succinic acid, L-asparagine, L-glutamic acid, L-threonine, Dl-carnitine, phenylethylamine, putrescine, 2-amino-ethanol and 2,3-butanediol). In the case of 14 carbon sources (monomethylsuccinate, cis-aconitic acid, citric acid, γ-hydroxybutyric acid, α-ketogluutaric acid, succinic acid, L-asparagine, L-glutamic acid, L-threonine, Dl-carnitine, phenylethylamine, putrescine, 2-amino-ethanol and 2,3-butanediol). In the case of 14 carbon sources (monomethylsuccinate, cis-aconitic acid, citric acid, γ-hydroxybutyric acid, α-ketogluutaric acid, succinic acid, L-asparagine, L-glutamic acid, L-threonine, Dl-carnitine, phenylethylamine, putrescine, 2-amino-ethanol and 2,3-butanediol). In the case of 14 carbon sources (monomethylsuccinate, cis-aconitic acid, citric acid, γ-hydroxybutyric acid, α-ketogluutaric acid, succinic acid, L-asparagine, L-glutamic acid, L-threonine, Dl-carnitine, phenylethylamine, putrescine, 2-amino-ethanol and 2,3-butanediol). In the case of 14 carbon sources (monomethylsuccinate, cis-aconitic acid, citric acid, γ-hydroxybutyric acid, α-ketogluutaric acid, succinic acid, L-asparagine, L-glutamic acid, L-threonine, Dl-carnitine, phenylethylamine, putrescine, 2-amino-ethanol and 2,3-butanediol). In the case of 14 carbon sources (monomethylsuccinate, cis-aconitic acid, citric acid, γ-hydroxybutyric acid, α-ketogluutaric acid, succinic acid, L-asparagine, L-glutamic acid, L-threonine, Dl-carnitine, phenylethylamine, putrescine, 2-amino-ethanol and 2,3-butanediol). In the case of 14 carbon sources (monomethylsuccinate, cis-aconitic acid, citric acid, γ-hydroxybutyric acid, α-ketogluutaric acid, succinic acid, L-asparagine, L-glutamic acid, L-threonine, Dl-carnitine, phenylethylamine, putrescine, 2-amino-ethanol and 2,3-butanediol).
methylpyruvate, acetic acid, formic acid, D-galacturonic acid lactone, D-gluconic acid, D-glucuronic acid, α-hydroxybutyric acid, β-hydroxybutyric acid, p-hydroxyphenylacetic acid, itaconic acid, α-ketobutyric acid, DL-lactic acid, malonic acid, bromosuccinic acid, succinic acid, glucuronamide, alaninamide, D-alanine, L-alanine, L-alanyl-glycine, L-aspartic acid, hydroxy-L-proline, L-ornithine, L-proline, D-serine, L-serine, urocanic acid, inosine, uridine, thymidine, glycerol, DL-glycerol phosphate, glucose 1-phosphate, glucose 6-phosphate). Some strains lost the ability to utilize some carbon sources after 2 years of maintenance in the laboratory. Using the API CH50 system acid production was negative in all cases. No nitrogenase activity could be detected, although the strains grew slightly on RBA medium.

Phylogenetic analysis
The results of the phylogenetic analysis are shown in Fig. 3. (Jukes–Cantor phylogenetic distance calculations, neighbour-joining treeing). Almost complete 16S rDNA sequences of strains L1/68T, L1/57, L1/58 and L2/11, composed of 1514, 1514, 1523 and 1518 bp, respectively (>98 % of the complete E. coli sequence; Brosius et al., 1978), were used for analysis. All isolates were completely identical to each other. The highest similarity values were found with members of the Xanthomonas–Rhodanobacter assemblages within the γ-subclass of the Proteobacteria. Similarity ranks for the nearest neighbours of the Xanthomonas–Rhodanobacter line were not significantly different (Xylella fastidiosa ATCC 35871, 86·57%; Lysobacter antibioticus, 86·7%; Frateria aurantia, 86·3%; Rhodanobacter lindaniclasticus, 85·8%; Cardiobacterium hominis, 85·7%; Xanthomonas campestris, 86·5%; Stenotrophomonas maltophilia, 86·9%), although Xylella fastidiosa ATCC 35879 proved to be the closest relative (87·1%). Similarities to representatives of other subclasses of Proteobacteria were also determined: Rhodospirillum rubrum (80·7%), Comamonas testosterone (82·9%), Desulfovibrio desulfuricans (79·3%) and Wolinella succinogenes (79·1%). Similar tree topology was achieved using Jukes–Cantor distance calculations with neighbour-joining and maximum-likelihood treeing and Kimura 2-parameter/neighbour-joining tree generation; however, the bootstrap value on the position of Schineria larvae was only 50%. The position of Schineria larvae within the Rhodanobacter–Xanthomonas assemblage of the γ-Proteobacteria was practically the same using parsimony and De Soete tree fit algorithms.

Chemotaxonomic characterization
Since all strains proved to have identical 16S rDNAs only the fatty acid profiles were examined in all strains, and strain L1/68T was studied in detail chemotaxonically. The G+C content of DNA of this strain was 42 mol%. The predominant respiratory quinone was Q-8, while Q-7 and Q-5 could be detected as minor components. It contained phosphatidylglycerol (PG) and diphosphatidylglycerol (DPG; minor amounts), phosphatidylethanolamine (PE), phosphatidylserine (PS) and two unknown polar lipids. The fatty acid profiles of the four strains are reported in Table 2.

DNA–DNA hybridization
At the DNA level the four examined strains are quite similar as their similarity values demonstrate: L1/57 to L1/58, 98·9%; L1/57 to L2/11, 102·5%; L1/57 to L1/68T, 98·6%; L1/58 to L1/68T, 99·4%; L2/11 to L1/68T, 97·8%.

DISCUSSION
Although all developmental stages of Wohlfahrtia magnifica and bacterial communities of fly-infested wounds were tested for their bacterial partners, bacteria belonging to this lineage of γ-Proteobacteria were detected only in the 1st and 2nd larval stage samples (Tóth et al., 1998). It is interesting to note, however, that the strains examined originating from different specimens of different larval stages of Wohlfahrtia magnifica are totally identical with regard to their 16S rDNA and their DNA–DNA hybridization values are similarly high. At the same time extremely high variations could be detected in the utilization of carbon sources: among 95 examined, only 36 (39%) were uniform at different times of testing. The ability to utilize other carbon sources disappeared or became positive during the 2 years of investigation. It can be speculated that some of these traits were encoded on

Table 2 Fatty acid composition of the bacterial strains

<table>
<thead>
<tr>
<th>Strain</th>
<th>Fatty acid composition (%)</th>
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<tbody>
<tr>
<td></td>
<td>C14:0</td>
<td>C16:0</td>
</tr>
<tr>
<td>L1/68T</td>
<td>20·1</td>
<td>23·1</td>
</tr>
<tr>
<td>L1/58</td>
<td>18·3</td>
<td>25·9</td>
</tr>
<tr>
<td>L1/57</td>
<td>17·7</td>
<td>22·9</td>
</tr>
<tr>
<td>L2/11</td>
<td>22·5</td>
<td>21·2</td>
</tr>
</tbody>
</table>
plasmids and subsequently lost due to subculturing under non-selective pressure.

The temperature range for growth of the strains is not surprising as the larvae grow in the live tissue of warm-blooded animals, where the temperature is usually between 35 and 38 °C. The observation that these bacteria preferred a slightly alkaline pH may result from the feeding habit of the fly larvae on tissues of sheep where the pH is alkaline, due to ammonification performed by a diverse population of wound bacteria (Guerrini et al., 1988; Khoga, 1994). Based on the strong chitinase activity of the strains we can speculate that these bacteria may play a significant role in the metamorphosis of the fly as reported for some Pseudomonas species which have an important function in the moulting of the screwworm Cochliomyia hominivorax by their chitinase enzyme production (Gassner et al., 1983). The slow growth of strains on diazotrophic medium (RBA) indicates that a low nitrogen supply is adequate for reproduction of these bacteria (though the fly larvae live in a nitrogen-rich environment).

Based upon the distinct phylogenetic position a new genus is proposed for the Wohlfahrtia magna species. Low similarity values to related species and the phylogenetic dendrogram shown in Fig. 3 indicate that the isolates represent a new genus as a deeply rooting lineage next to the Rhodanobacter–Lysobacter–Xylella assemblage within the γ-Proteobacteria. The base G+ C content of the 16S rDNA of the isolates is 42 mol% which is not close to the values determined for Xylella (53 mol%), Rhodanobacter (63 mol%) and Lysobacter (66 mol%), but our strains are located far from their closest known relatives. The affiliation of the isolates to the γ-Proteobacteria is supported by the presence of three signature nucleotides in the 16S rDNA (976, 1233 and 1234 – G, G and C respectively; Woese, 1987) distinguishing the members of the γ- and β-subclasses. Furthermore, the presence of ubiquinone Q-8 confirms the membership of the γ-subclass (Collins & Jones, 1981).

All strains are identical in their 16S rDNA sequence and highly similar in biochemical and physiological properties. Their fatty acid composition is quite similar and their hybridization values to each other are also high. As they cannot be differentiated unambiguously by the carbon source utilization tests, we consider these strains as a single species.

**Description of Schineria gen. nov.**

Schineria (sh’i-ner’i-a. N.L. fem. gen. n. Schineria pertaining to Schiner who first described the fly Wohlfahrtia magna in 1862).

Cells are Gram-negative regular rods. Non-motile, endospores are not formed. Capsules may be produced. Aerobic. Catalase reaction is positive (occasionally weak), oxidase reaction is variable. Main fatty acids are C_18:1ω9c, C_{16:1}ω7c, and C_{14:0}:3ω2. Predominant polar lipids are PG, DPG, PE, and PE. Predominant respiratory quinone is Q-8. G+C content of DNA is 42 mol%.

Phylogenetically a member of the γ-subclass of the Proteobacteria. Unequivocal affiliation by primary structure of 16S rDNA. The type species is *Schineria larvae*.

**Description of Schineria larvae sp. nov.**

Schineria larvae (lar’væ. L. gen. fem. larvae pertaining to the origin of the type strain isolated from Wohlfahrtia magna magniffs maggots).

Cells are straight, short, single rods (2–3 μm × 0.8–0.9 μm), non-motile and encapsulated, occurring singly at all ages. On agar media colonies are non-pigmented, small (1–3 mm in diameter), circular with entire margin and smooth surface. Strictly aerobic, NO_3 is not reduced. Optimum growth occurs between pH 6 and 10. Temperature range is wide (4–45 °C), optimum growth at 28–37 °C. Catalase-positive, oxidase-variable. Urea and chitin are hydrolysed. Indole, casein and gelatin are not metabolized. H2S is produced from cysteine. Simmons’s citrate, Voges–Proskauer and aesculin digestion are negative. Not haemolytic. Uses monomethylsuccinate, cis-aconitic acid, citric acid, γ-hydroxybutyric acid, α-ketoglutaric acid, succinic acid, l-asparagine, l-glutamic acid, l-glutamic acid, l-histidine, l-leucine, l-phenylalanine, l-proline, l-proline, and γ-aminobutyric acid as sole carbon source. No utilization of α-cyclodextrin, glycogen, d-arabitol, cellobiose, i-erythritol, gentiobiose, β-lactose–lactulose, xylitol, d-galacturonic acid lactone, d-glucosaminic acid, α-ketoveralac acid, propionic acid, quinic acid, d-saccharic acid, sebacic acid, l-threonine, dL-carnitine, phenylethylamine, putrescine, 2-aminoethanol and 2,3-butanediol. No acid production from carbohydrates in API CH50. The major isoprenoid quinone is Q-8. The major fatty acids are C_18:1ω9c, C_{16:1}ω7c and C_{14:0}:3ω2. Cyclo-C_{18:0} also occurs. Predominant polar lipids are PG, PE and PS. G+C content of DNA is 42 mol%. Isolated from the L1 and L2 stage larvae of an obligate parasitic fly, Wohlfahrtia magna. Type strain is L1/68T (= DSM 13226T = NCAIM B01938T).

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