Stratified Microbial Structure and Activity in Sulfide- and Methane-Producing Anaerobic Sewer Biofilms

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Simultaneous production of sulfide and methane by anaerobic sewer biofilms has recently been observed, suggesting that sulfate-reducing bacteria (SRB) and methanogenic archaea (MA), microorganisms known to compete for the same substrates, can coexist in this environment. This study investigated the community structures and activities of SRB and MA in anaerobic sewer biofilms (average thickness of 800 μm) using a combination of microelectrode measurements, molecular techniques, and mathematical modeling. It was seen that sulfide was mainly produced in the outer layer of the biofilm, between the depths of 0 and 300 μm, which is in good agreement with the distribution of SRB population as revealed by cryosection-fluorescence in situ hybridization (FISH). SRB had a higher relative abundance of 20% on the surface layer, which decreased gradually to below 3% at a depth of 400 μm. In contrast, MA mainly inhabited the inner layer of the biofilm. Their relative abundances increased from 10% to 75% at depths of 200 μm and 700 μm, respectively, from the biofilm surface layer. High-throughput pyrosequencing of 16S rRNA amplicons showed that SRB in the biofilm were mainly affiliated with five genera, Desulfovibrio, Desulfobulbus, Desulfomicrobium, Desulfoverrups, and Desulfatiferula, while about 90% of the MA population belonged to the genus Methanoseta. The spatial organizations of SRB and MA revealed by pyrosequencing were consistent with the FISH results. A biofilm model was constructed to simulate the SRB and MA distributions in the anaerobic sewer biofilm. The good fit between model predictions and the experimental data indicate that the coexistence and spatial structure of SRB and MA in the biofilm resulted from the microbial types and their metabolic transformations and interactions with substrates.

Sewer biofilms comprise complex multispecies microflora with a typical thickness of only about 1 mm (1). Depending on the electron donors and electron acceptors present in the wastewater, different carbon transformation processes can occur in close proximity in the sewer biofilms. Domestic wastewater normally contains a significant concentration (ca. 100 to 1,000 μM) of sulfate but negligible nitrite and nitrate concentrations (2, 3). Therefore, under anaerobic conditions (which normally occur in pressure sewers fully filled with wastewater), sulfate reduction carried out by sulfate-reducing bacteria (SRB) could be an important terminal electron-accepting process in the sewer biofilms. The sulfate reduction activity in anaerobic sewers is important, as the production of sulfide can be transferred to the gas phase of partially filled gravity sewers and cause extensive corrosion of concrete sewer pipes (4, 5). Also, the emission of sulfide from sewers can cause odor problems for the surrounding area and pose health risks to sewer workers (6, 7). Apart from sulfate reduction, methanogenesis caused by the respiration of methanogenic archaea (MA) could also be a key terminal process in anaerobic sewer biofilms (8, 9). Guisasola and colleagues found that methanogenesis accounted for more than 70% of the chemical oxygen demand (COD) loss in laboratory anaerobic sewer biofilm reactors (9). A recent report suggests that methane emissions from sewers contribute significantly to the total greenhouse gas footprint of wastewater systems (10).

Under anaerobic conditions, both sulfate reduction and methanogenesis can potentially occur in the same system while competing for the same electron donors, primarily hydrogen and acetate. In the presence of adequate sulfate concentrations, SRB typically out-compete MA due to kinetic and thermodynamic advantages (11–13). However, the coexistence of SRB and MA has been observed in anaerobic sewer biofilms in the presence of sulfate. Guisasola et al. (9) hypothesized that the coexistence of SRB and MA in sewer biofilms was due to the penetration limitation of sulfate into the biofilms, resulting in a stratified biofilm structure, with SRB being predominant in the outer zone, nearer the wastewater, while MA inhabit the inner zone, nearer the sewer pipe. However, to date this hypothesis has not been verified. A few studies have investigated the vertical distribution of SRB in oxic-anoxic sewer biofilms (biofilms attached to gravity sewer pipe with the presence of oxygen or nitrate in wastewater), but studies on the SRB distribution in the anaerobic sewer biofilms are scarce (14). In addition, the distribution of MA in sewer biofilms and their interaction with SRB have not yet been explored. Similarly, the phylogenetic diversity of SRB and MA in the anaerobic sewer biofilms has rarely been reported. This fundamental information could provide a better understanding of the sulfate reduction and methanogenesis processes in sewer systems, which would be useful for sewer management. Therefore, the aims of this study were to investigate the community structures of both SRB and MA and to determine their spatial arrangement in anaerobic sewer biofilms.

Both experimental investigations and modeling analyses were conducted to achieve the aims of this study. The experiments were carried out in an annular biofilm reactor fed with real domestic wastewater.
wastewater, mimicking anaerobic sewer conditions. First, micro-electrodes were applied to determine the spatial distribution of in situ sulfide production activity within the biofilms. Although it would have been ideal to determine the distribution of methane production activity using the same method, this was difficult to perform due to the lack of suitable microelectrodes (15). Second, the spatial distributions of the SRB and MA in the biofilms and their abundances at different depths were determined by fluorescence in situ hybridization (FISH) after cryosectioning of the biofilm samples. This method has been used frequently to determine the spatial distributions of microbial communities in biofilms or granules. However, little phylogenetic information is revealed due to the limitations of the oligonucleotide probes used in FISH (16). Therefore, 16S rRNA gene amplicon pyrosequencing was applied to further investigate the phylogenetic diversity. In previous studies of sewer biofilms, the phylogenetic analysis was performed on the entire biofilm, and information on the different genera at different biofilm depths has rarely been reported (2, 14). In this study, we determined the phylogenetic diversity in different layers of the sewer biofilms by innovatively using pyrosequencing combined with cryosectioning. To our knowledge, to date, this method has not been applied in any other studies related to biofilms and granules. Finally, a mathematical model focusing on the interaction between SRB and MA in the sewer biofilm was developed to evaluate and interpret the experimental results.

MATERIALS AND METHODS

Reactor configuration, operation, and monitoring. An annular biofilm reactor made of acrylonitrile butadiene styrene (ABS), one of the typical materials used for sewer pipes, was set up to mimic an anaerobic sewer pipe section (Fig. 1). The reactor consisted of an inner cylinder (height, 295 mm; diameter, 130 mm) enclosed in an outer cylinder (height, 345 mm; diameter, 160 mm). The gap between the two cylinders was filled with wastewater (volume, 3 liters). Biofilms were grown on the walls of both cylinders in contact with the wastewater, resulting in a biofilm-area-to-reactor-volume (A/V) ratio of 119 m⁻¹. Mixing was established by the rotation of the inner cylinder driven by a motor at a speed of 200 rpm. The mixing was expected to create a uniform shear stress on the reactor walls to ensure that biofilms grew relatively evenly on the wall. The average shear stress provided by the mixing was 2.11 N/m², which is typical in sewer systems (17). Eight removable ABS slides of 3-mm width and 200-mm length were mounted in recessed slots inside the outer cylinder. The slides were removable via ports on the top of the reactor for biofilm sampling. The reservoir on the top of the reactor was used to ensure that the reactor was full of wastewater during sampling. The reactor was operated in a temperature-controlled room (20 ± 2°C). Domestic sewage, collected on a weekly basis from a local wet well (Brisbane, Queensland), was used as the feed for the reactor. The sewage compositions varied to a certain extent in terms of sulfate, volatile fatty acid (VFA), and COD concentrations. The sewage typically contained sulfate at concentrations of 10 to 25 mg/liter, sulfide at <3 mg/liter, soluble COD (sCOD) at 200 to 300 mg/liter, 50 to 120 mg COD/liter of VFAs, and approximately 50 mg N/liter of ammonium. Negligible amounts of sulfate, thiosulfate (<1 mg S/liter), nitrate, and nitrite (<1 mg N/liter) were present. The sewage was stored in a cold room (4°C) to minimize biological transformation and was heated to 20 ± 1°C prior to being pumped into the reactor (Fig. 1), which was consistent with the ambient temperature.

The sewage was fed to the reactor intermittently by a peristaltic pump (Masterflex 7520-47) to simulate the typical flow patterns of rising main sewers (main pressure sewers). For easier reactor monitoring, each day was divided into three identical 8-h periods. Figure S1 in the supplemental material shows the pumping patterns applied to the reactor for an 8-h period and the hydraulic retention time (HRT) (a measure of the average length of time that a soluble compound remains in a constructed bioreactor) of sewage in the reactor. Every pumping event lasted for 3 min, delivering one reactor volume (3 liters) of wastewater into the reactor. The HRTs of the wastewater ranged between 30 min and 4 h, which are in the range of the HRTs observed in a typical real sewer pipe (9).

Monitoring of the reactor performance was carried out during the 8-h cycle periods every 2 weeks. Sulfide concentrations during the 8-h cycle were continuously monitored using an S::CAN UV-visible light (UV-VIS) spectrophotometer (Messtechnik GmbH, Austria), as previously described by Sutherland-Stacey et al. (18). In addition, samples were taken from the reactor before and after each pumping event and also at 2.5 h, 5 h, and 6.5 h for the analysis of dissolved methane, sulfate, total COD (tCOD), sCOD, and VFAs, using methods described by Jiang et al. (3). Briefly, dissolved methane and VFA concentrations were measured by gas chromatography. Sulfate concentrations were measured using ion chromatography, tCOD and sCOD were determined by means of COD cuvette tests (Merck) (range, 25 to 1,500 mg liter⁻¹). Detailed studies of the biofilm were carried out when the reactor reached pseudo-steady-state conditions after about 10 months operation, as indicated by the relatively constant sulfide and methane profiles.

Microelectrode analyses. Hydrogen sulfide (i.e., molecular H₂S), pH, and dissolved oxygen levels in the biofilm were measured using micro-electrodes (Unisense A/S; Denmark) with tip diameters of 10 μm, 25 μm,
and 100 μm, respectively. The sensors were calibrated according to the manufacturer’s instructions. Hydrogen sulfide and pH profiles were measured to determine the total dissolved sulfide concentration as described by Kuhl et al. (19). Oxygen profiles were measured to confirm anaerobic conditions.

Before the microelectrode analyses, a length of about 5 cm of the biofilm slide was removed from the reactor and mounted in a flow cell (as described by Gutierrez et al. [20]) containing 140 ml of wastewater that had been filtered using a 0.22-μm-pore-size filter and 20 ml of 300 mM phosphate buffer (added to ensure a stable pH of 7.0 to 7.5). Nitrogen gas (99.99% purity) was bubbled through the flow cell to ensure anaerobic conditions and to provide mixing. Microelectrodes were mounted on a micromanipulator and positioned on the surface of the biofilm using a dissecting microscope. The concentration gradients through the biofilm were obtained by moving the microelectrodes in increments of 25 to 100 μm. Steady-state profiles were obtained by incubating the biofilm for 1 h in the medium before measurements were made.

The local sulfide production rates were calculated from the total sulfide profiles based on Fick’s second law of diffusion. The calculation was carried out by a stepwise procedure as described by Gieseker and de Beer (21). Briefly, the production rate at point \( n \) can be calculated using the following equations:

\[
\begin{align*}
    &J_{n-1,n} = \frac{f_{n-1,n} - f_{n,n+1}}{0.5(x_{n-1} + x_n) - 0.5(x_n + x_{n+1})} \\
    &J_{n-1,n} = D_{\text{eff}} \left( C_{n-1} - C_n \right) / \left( x_{n-1} - x_n \right) = D_{\text{eff}} \left( C_n - C_{n+1} \right) / \left( x_n - x_{n+1} \right)
\end{align*}
\]

where \( f_{n-1,n} \) and \( f_{n,n+1} \) represent the flux between point \( n-1 \) and point \( n \) and the flux between point \( n \) and point \( n+1 \) (expressed in mol/m²/s), \( x_{n-1} \), \( x_n \), and \( x_{n+1} \) represent the depths of points \( n-1 \), \( n \), and \( n+1 \) (m), \( C_{n-1} \), \( C_n \), and \( C_{n+1} \) are sulfide concentrations at points \( n-1 \), \( n \), and \( n+1 \) (mol/m³), and \( D_{\text{eff}} \) represents the effective diffusion coefficient of sulfide in the biofilm (m²/s). The value used in this study was 1.39 × 10⁻⁹ m²/s (19), and this was based on the assumption that the diffusion coefficients within the biofilm were equal to the molecular diffusion coefficients.

**Fluorescence in situ hybridization.** Fluorescence in situ hybridization (FISH) was carried out to determine the distributions of SRB and MA in the biofilm. The sequences of all oligonucleotide probes used in this study are summarized in Table S1 in the supplemental material, and further detailed information is in probeBase (22). Due to a drawback of SRB probes, which can detect other bacteria that are not SRB, the SRB levels in the phylum of the Deltaproteobacteria were determined by the overlapping of the fluorescence signals of probes DELTA495a (CY3), DELTA495b (CY3), and DELTA495c (CY3) with those of probes SRB385 (BS), SRB385DB (CY5), and DABAC357 (CY5). Using this approach, most SRB in the phylum of the Deltaproteobacteria were detected while the non-SRB targeted by these probes were discriminated (16). SRB in other phyla were not detected by 16S rDNA pyrosequencing results, so probes for those phyla were not used. For FISH detection of MA, a combination of probes MSMX860 (CY5), MG1200B (CY5), MB1175 (CY5), MC1109 (CY5), and MC504 (CY5) was used to determine the total MA population in the biofilm. This combination of probes covers a wide range of MA in these ecosystems (23). SRB and MA levels were determined using different samples due to the different formamide concentrations required (35% for the SRB detection and 45% for the MA detection). Probes EUB338mix (fluorescin isothiocyanate [FITC]) and ARCA915 (CY3) were used to determine all bacteria and archaea in the biofilm, respectively.

To conduct FISH analysis, the biofilm sampling slides were removed from the reactor and cut into sections with dimensions of approximately 10 mm by 5 mm. The biofilm samples on the small pieces were fixed with freshly prepared 4% paraformaldehyde solution for 2 h at 4°C. The fixed biofilm sample was then embedded in Tissue-Tek OCT compound (Sakura Finetek, Tokyo, Japan) following the procedures described by Batstone et al. (24). The biofilm samples were then allowed to settle on the base of the OCT molds and were frozen at −20°C. The frozen samples were then sectioned using a research cryostat (Leica CM3050 S) with a knife temperature of −20°C, a cabinet temperature of −18°C, and a section thickness of 10 μm. The samples were divided into two groups and cryosectioned in two different directions. One group of samples was sectioned perpendicularly to the substratum to provide sections to visualize the arrangement of SRB and MA distributed through the depth of the biofilm. The samples in the other group were successively sectioned parallel to the substratum from the surface to the bottom of the biofilm. These samples were used to determine the relative abundances of SRB and MA at different depths within the biofilm. The cryosectioned samples were placed on poly-l-lysine-coated microscopy slides (Polysciences Asia Pacific, Inc.) and air dried for 6 to 10 h. The slides were then dehydrated for 3 min each (50%, 80%, and 98%) in an aqueous ethanol solution.

**All in situ hybridizations were performed at 46°C for 2 to 3 h according to a previously described protocol (25) using hybridization buffer.** The buffer contained 0.9 M NaCl, 20 mM Tris hydrochloride (pH 7.2), 0.01% sodium dodecyl sulfate, and formamide concentrations as previously mentioned. Subsequently, a stringent washing step was performed at 48°C for 15 min in 50 ml of washing solution comprising NaCl at a concentration dependent on the formamide concentration and 20 mM Tris hydrochloride at pH 7.2. The slides were examined and recorded using a Zeiss LSM 510 confocal laser scanning microscope (CLSM) (Carl Zeiss, Jena, Germany) and three excitation channels (488 nm, green emission; 545 nm, red emission; 633 nm, blue emission). The biofilm thickness was estimated by measuring the width of the biofilm sections cut perpendicularly to the substratum. FISH images at eight different depths of the biofilm (0 to 10 μm, 100 to 110 μm, 200 to 210 μm, 300 to 310 μm, 400 to 410 μm, 500 to 510 μm, 600 to 610 μm, and 700 to 710 μm) were analyzed using DAIME version 1.3 (26) to determine the biovolume fractions of SRB and MA. About 20 confocal images of the biofilm sections were analyzed for each sample. The quantification data were calculated based on the average results from two separately analyzed samples.

**16S rRNA gene amplicon pyrosequencing.** 16S rRNA gene amplicon pyrosequencing was conducted to investigate the phylogenetic diversity of SRB and MA at different layers in the biofilm. Biofilms on a piece of a slide (10 mm by 5 mm) were quickly removed from the reactor and embedded in OCT compound and then frozen at −20°C in a OCT mold. The frozen samples were then cryosectioned successively from the surface to the bottom of the biofilm with a section thickness of 15 μm, using the cryostat as described above. The sectioned biofilm samples were then placed separately in 1 ml Eppendorf tubes containing 0.5 ml of phosphate-buffered saline (PBS) (containing 137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, and 2 mM KH₂PO₄) for DNA extraction.

Genomic DNA was extracted using a FastDNA SPIN kit for soil according to the manufacturer’s instructions (Q-Bio gene; Australia). The quantity and quality of the extracted DNA were measured using a NanoDrop ND-1000 spectrophotometer (Nano-Drop Technology, Rockland, DE) and agarose gel (0.8% [wt/vol] electrophoresis. Primers 926F (5’-A AAC TYA AA KGA ATTGACGGGCG-3’) and 1392r (5’-ACGGGCGGTGTGAC-3’) (27) containing multiple identifiers and Libl. adaptor sequences (not shown) were used to generate amplicons. The PCR tube (50 μl) contained 5 μl of 10X PCR buffer, 1 μl of 10 mM deoxynucleoside triphosphates (dNTPs), 1 μl (each) of 10 mM primers, 4 μl of 25 mM MgCl₂, 1.5 μl of 10 mg/ml bovine serum albumin (BSA), 0.2 μl of 5.5 μl/μl Fisher Biotech Taq DNA polymerase, and 2.5 μl of 20 μg/ml DNA sample. The PCR was conducted under the following conditions: 95°C for 3 min, followed by 30 cycles of 95°C for 30 s, 55°C for 30 s, and 74°C for 30 s and a final elongation step at 74°C for 10 min. The pyrosequencing of amplicons was carried out according to Roche 454 protocols using a Roche 454 GS FLX sequencer (Roche, Switzerland). The sequence data were analyzed through the use of the ACE Pyrosequencing Pipeline (https://github.com/minillinni/APP) in a local implementation. First, the sequencing reads were split according to the barcode in QIME v1.8.0 (28). Then, demultiplexed sequences were trimmed to 250-bp lengths and denoised using....
**RESULTS**

**Performance of the anaerobic sewer reactor.** The typical sulfide and methane profiles in the sewer biofilm reactor during an 8-h operation cycle are shown in Fig. 2. Sulfide and methane were produced simultaneously in the reactor, and concentrations of sulfide (13.0 to 18.6 mg/liter) and methane (9.3 to 14.9 mg/liter) at the end of each pumping cycle varied according to the HRT. During the 8-h cycle, the tCOD was decreased by 17% and nearly 86% of the sulfide was reduced. Table 1 shows the average daily transformation of COD, VFAs, sulfur species, and methane at the pseudo-steady state, calculated based on the concentration differences at the beginning and end of each pumping cycle. The tCOD was consumed by 688.2 ± 29.2 mg/day, with productions of sulfide and methane at 123.9 ± 11.1 mg/day and 103.4 ± 3.2 mg/day, respectively. Similar daily sulfate consumption and sulfide production indicated that sulfide was the major product of sulfate reduction. The sCOD and propionate were also consumed in the reactor while acetate accumulated. The COD balance was calculated on the basis of the assumption that all of the hydrogen produced due to fermentation was consumed during the experiment. The COD utilization per gram of sulfide and methane formed was assumed to be 2 g COD/g H₂S-S and 4 g COD/g CH₄, respectively (9). Therefore, sulfidogenesis accounted for 36.0% ± 2.4% of the tCOD loss in the wastewater whereas methanogenesis accounted for 60.0% ± 4.3% (Table 1).

**Distribution of sulfide production within the biofilm.** The microscale sulfide, pH, and oxygen levels were measured throughout the depth of the biofilm (Fig. 3). A significant increase of sulfide concentration is seen from the biofilm surface to ca. 250 μm into the biofilm. The pH remained constant throughout the depth of the biofilm, due to the buffering capacity of the system. Negligible levels of oxygen were detected within the biofilm. The in situ sulfide production rates were calculated based on the sulfide profiles according to Fick’s law of diffusion (Fig. 3), which indicated 4.3% (Table 1).

### Table 1 Daily transformation of COD, VFAs, sulfur species, and methane in the sewer biofilm reactor

<table>
<thead>
<tr>
<th>Compound</th>
<th>Unit</th>
<th>Daily transformation^a</th>
<th>% ΔCOD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total COD</td>
<td>mg COD</td>
<td>-688.2 ± 29.2</td>
<td>-100.0 ± 0.0</td>
</tr>
<tr>
<td>Soluble COD</td>
<td>mg COD</td>
<td>-362.5 ± 12.7</td>
<td></td>
</tr>
<tr>
<td>Acetate</td>
<td>mg COD</td>
<td>+49.4 ± 17.2</td>
<td></td>
</tr>
<tr>
<td>Propionate</td>
<td>mg COD</td>
<td>-76.5 ± 3.0</td>
<td></td>
</tr>
<tr>
<td>Sulfate</td>
<td>mg S</td>
<td>-123.5 ± 12.8</td>
<td></td>
</tr>
<tr>
<td>Sulfide</td>
<td>mg S</td>
<td>+123.9 ± 11.1</td>
<td>36.0 ± 2.4</td>
</tr>
<tr>
<td>Dissolved methane</td>
<td>mg</td>
<td>+103.4 ± 3.2</td>
<td>60.0 ± 4.3</td>
</tr>
</tbody>
</table>

^a +, production; −, consumption.
Spatial distributions of SRB and MA populations as determined by FISH. FISH analysis of the biofilm sections cut perpendicularly to the substratum showed the localization of SRB and MA (Fig. 4A and B). SRB (white in Fig. 4A) were mainly situated at the outer layer (depth of 0 to 300 μm) of the biofilm, while MA (purple in Fig. 4B) were mainly located in the inner layer (below 250 μm). Panels C to F in Fig. 4 show typical FISH images of the biofilm sections cut parallel to the substratum at depths of 100 μm and 700 μm. Accordingly, SRB were detected in much higher abundance in the biofilm section at the depth of 100 μm than at the 700-μm-deep section (compare Fig. 4C to Fig. 4D). In contrast, there were hardly any MA at the depth of 100 μm whereas MA were dominant at the depth of 700 μm (compare Fig. 4E to Fig. 4F). The relative abundances of SRB and MA at different depths showed that SRB accounted for about 20% of the total population at the surface and at 100 μm into the biofilm and that the proportion decreased continuously to lower than 3% at the depth of 400 μm (Fig. 5). This distribution of SRB is consistent with the profile of the in situ sulfide production rate (Fig. 3). In contrast to the SRB distribution, the MA were detected at below 3% abundance at the surface and at the depth of 100 μm and increased in abundance to 10% at the depth of 200 μm, 60% at 500 μm, and then 75% at 700 μm (Fig. 5).

Biofilm community structure as determined by 16S rRNA sequence analysis. The 16S rRNA gene sequence analysis was successively applied to five layers of the biofilm, from the surface to the bottom of the biofilm (layer 1 to layer 5). The sequence reads for each layer are shown in Table S6 in the supplemental material. The thickness of each layer was 150 μm. The results revealed that the SRB detected in the biofilm were mainly affiliated with five genera and that their proportions of the total SRB detected were as follows: Desulfobulbus at 33%, Desulfomicrombium at 19%, Desulfovibrio at 24%, Desulfatitihalitalea at 7%, and Desulforuga at 16%. The heat map (Fig. 6A) displaying the distributions of the predominant SRB in the different layers (layer 1 to layer 5) confirmed that SRB were mainly situated in the outer layer. SRB of the genera Desulfobulbus, Desulfomicrombium, and Desulfovibrio were also observed in inner layers 4 and 5, but the SRB in these inner layers accounted for only less than 10% of the total SRB detected in the biofilm. About 90% of the MA population belonged to the genus of Methanoseta, whose members use acetate rather than hydrogen as the substrate. The other 10% of the MA population mainly belonged to five genera: Methanospirillum, Methanomethylovorans, Methanobrevibacter, Methanobacterium, and Methanomethylovorans. The heat map (Fig. 6B) also demonstrates that MA were mainly located in the inner layer of the biofilm. Interestingly, Methanobrevibacter, Methanomethylovorans, and Methanospirillum actually showed higher abundance in the outer layer than in the inner layer. However, these accounted for about 5% of the total MA population detected in the biofilm and thus had only a minor effect on the overall MA distribution in the biofilm.

Mathematical modeling. Mathematical modeling was performed to describe the microbial distribution and the sulfide concentration profiles within the biofilms. The model-predicted relative abundances of SRB and MA fit well with the experimental results determined by FISH (Fig. 7). The SRB abundance was 19% at the surface and decreased gradually to below 5% at the depth of 400 μm. The abundance of MA was lower than 5% at the surface and at 100 μm, increased to 65% at the depth of 500 μm, and then gradually rose to 80% at 700 μm. These results are consistent with the experimental data. The model-predicted sulfide concentration profiles within the biofilms also matched well with the data measured by microelectrode. The good agreement between the model-predicted results and the experimental data indicated that the spatial structure of SRB and MA in the anaerobic sewer biofilms resulted from the microbial types and their metabolic transformations and interactions with substrates.

DISCUSSION

The distribution of SRB and MA in anaerobic sewer biofilms.

This study investigated the distribution of SRB and MA in sewer biofilms through both experimental analysis and simulation analysis. The stratified distribution of SRB and MA in the biofilm was confirmed and verified using two independent molecular techniques, i.e., FISH and pyrosequencing, as well as microelectrode measurements and mathematical modeling. All the results are highly consistent. The results show that SRB were mainly located in the outer layer of the biofilm whereas MA were mainly situated in the inner layer. The distribution of in situ sulfide production activity was consistent with the distribution of the SRB population. The high sulfide concentration in the inner layer of the bio-

**FIG 3** Profiles of measured total dissolved sulfide, oxygen, pH, and calculated sulfide production rate in the biofilm. Negative depths in the profile represent the distance from the biofilm surface into the wastewater.
film was mainly due to the diffusion transport mechanism. While the sulfide production activity in the inner layer of the biofilm was much lower than that in the outer layer, in the absence of a sulfide sink in this layer, any sulfide produced would accumulate to a level higher than that accumulated in the outer layer, providing a concentration gradient for the sulfide produced to be transferred out of the biofilm.

Under anaerobic conditions, SRB and MA are known to compete for the same substrates (primarily acetate and hydrogen) for metabolism. In sulfate-rich environments, SRB can normally out-compete MA; this is commonly attributed to the different affinities of the two populations for substrates. The affinity constant for hydrogen of SRB is considered to be around five times lower than that of MA (35,36). The difference is even greater in the case of acetate (12, 37). However, coexistence of SRB and MA is observed in some systems under sulfate-limiting conditions or even under non-sulfate-limiting conditions, where other factors play a role. These included mass transfer limitations (38) and differences in microbial colonization and adhesion properties (39, 40) or variable sulfide toxicities (41, 42).

In anaerobic sewers, particularly in networks with relatively short HRTs, sulfate is normally not depleted. The stratified distri-
bution of SRB and MA suggests that mass transfer limitation plays an important role for the coexistence of SRB and MA in sewer biofilms. We used model simulation to determine the average concentrations of sulfate and soluble biodegradable COD in the sewer biofilm (Fig. 8). Sulfate could penetrate into the outer layer of the biofilm. Under these conditions, SRB out-competed MA due to their higher affinity to acetate and hydrogen, resulting in a higher abundance of SRB in the outer layer. However, the modeling result showed that the sulfate was almost totally consumed in the outer layer due to the high sulfate reduction activity and thus could not reach the inner layer (Fig. 8). As a result, SRB activity and growth were limited in the deeper layers of the biofilm. On the other hand, soluble biodegradable COD (including propionate, acetate, or hydrogen and the soluble COD which could be fermented to these products) was not totally consumed by SRB in the outer layer of the biofilm and it was able to penetrate into the inner layers, providing substrate for methanogenesis. Consequently, the coexistence and stratification of these populations are largely a result of the mass transfer of substrates into the biofilm.

The domination of MA in cores of anaerobic granules or at the inner layers of anaerobic biofilms has previously been attributed in some studies to better attachment characteristics of MA (40, 43). However, this cannot be a main reason in the case of anaerobic sewer biofilms. During the startup of the sewer reactor, the sulfate-reducing activity increased much faster than the methanogenic activity in the first several weeks (data not shown), indicating that at the beginning, more SRB were attached on the substrate than MA and that these were the pioneering colonizers of the biofilm. Variations of sulfide toxicities with respect to SRB and MA are also considered a reason for the coexistence of SRB and MA in some studies (41, 42). However, in our system, the sulfide concentration was far below toxic threshold levels for either group of microorganisms. It has been reported that sulfide concentrations of above 300 ppm are required to induce 50% inhibition of the growth of most SRB and MA (44).

The spatial arrangement of SRB and MA in sewer biofilms revealed in this study is of practical importance. Chemicals such as nitrate, oxygen, magnesium hydroxide, and sodium hydroxide are often added to sewers to control the emission of hydrogen sulfide in those sewers (6). As MA mainly inhabit the inner layer of the biofilms, they are likely to be protected from being exposed to

![FIG 5](image-url) The SRB (A) and MA (B) proportions of total microorganisms (bacteria and archaea) detected by FISH within the sewer biofilms. *C. Methanomethylophilus,* "Candidatus Methanomethylophilus."

![FIG 6](image-url) Heat map displaying the distribution of the predominant SRB (A) and MA (B) in different biofilm layers from the biofilm surface to the bottom (Layer 1 to Layer 5).
chemicals added for in-sewer sulfide and methane mitigation. Ji-ang et al. (45) found that sewer biofilms were capable of methanogenesis after nitrate dosing for 4 weeks. To explain this, they suggested that nitrate was not able to fully penetrate into the biofilm and that it failed to reach the MA in the deeper layer. This suggestion is supported by results showing a complete suppression of methane production after they increased the nitrate-dosing rate. Similar results were also observed by Ganigué et al. (46), where they found that methane was produced by the sewer biofilms after oxygen treatment and attributed that production to the partial penetration of oxygen. Consequently, given the spatial distribution of MA in sewer biofilms, full penetration of chemicals into biofilms is required to completely control methane production. This should be an important consideration for methane abatement strategies in sewers. Due to the difficulty in obtaining intact biofilm from real sewers, it remains to be verified whether the biofilm developed in our laboratory reactor fully represents that in real sewers, despite the use of realistic wastewater and shear conditions. Therefore, the implications discussed above need to be verified in real sewer systems.

Phylogenic diversities of SRB and MA and their hypothesized functions. This study innovatively used pyrosequencing coupled with cryosectioning to investigate the phylogenetic diversity of SRB and MA in anaerobic sewer biofilms. Pyrosequencing can provide phylogenetic information that is more detailed than that provided by FISH. Together with cryosectioning, the phylogenetic characteristics at different depths in the biofilms were investigated. However, note that, due to the significant quantity of biomass required for pyrosequencing analysis, the biofilm sections needed for this purpose were much thicker than those needed for FISH (150 μm versus 10 μm in this study). Consequently, the spatial resolution of the method was limited to layers of this size. However, this approach was successful and revealed the microbial diversity of both SRB and MA at five depths of the biofilm, allowing us to attempt to reconstruct the possible metabolic transformations in different regions of the sewer biofilm.

SRB detected in this anaerobic sewer biofilm were mainly affiliated with five genera: Desulfobulbus, Desulfovibrio, Desulfomicrobium, Desulfogulana, and Desulfatiferula. The first four of those genera have also been found in aerobic and anoxic wastewater biofilms, with Desulfobulbus, Desulfovibrio, and Desulfomicrobium appearing in higher abundances (2, 14, 47, 48). Also, Desulfobulbus and Desulfovibrio are reported to be numerically important in anaerobic methanogenic-sulfidogenic aggregates (40). Desulfatiferula is a genus that was newly defined by Cravo-Laureau et al. (49), and members are mesophilic, Gram-negative sulfate-reducing bacteria.

SRB can use many different compounds besides acetate and hydrogen as electron acceptors (37, 50, 51). Among the bacteria in the studied sewer biofilm, Desulfobulbus spp., with 98.4% sequence identity to Desulfobulbus propionicus, are well-known pro-
pionate-utilizing SRB (2, 51, 52). As SRB tend to out-compete acetogenic bacteria for propionate utilization due to their stronger affinity for this carbon substrate (36, 53), the high fraction of Desulfobulbus in the SRB population explained the low propionate concentration (<1 mg COD/liter) in the effluent. Desulfovibrio spp. can use hydrogen, formate, lactate, pyruvate, and many other organic compounds to reduce sulfate (54). It has been suggested that Desulfovibrio is an important member of the hydrogen-utilizing bacteria in wastewater biofilms (2, 48). Desulfomicrobium spp. are also able to use various substrates such as hydrogen, acetate, and lactate (55, 56). It is recognized that hydrogen, acetate, and propionate are important electron donors for sulfate reduction in sewer systems (33, 34). However, in this study, we also observed the proliferation of SRB, which normally use on large-molecule organic substrates rather than on hydrogen, acetate, and propionate for growth. Desulfotartrula and Desulfatiferula are known to use long-chain fatty acids and long-chain alkenes to reduce sulfate (49, 57). Also, some Desulfovibrio spp. are known to use amino acids and many other organic compounds as electron donors (54, 58). It is thought that SRB can be out-competed by very fast-growing fermentative (acidogenic) bacteria for the large-molecule organic substrates (50, 59). However, since fermentable COD or sCOD is abundant in sewer systems and they would not be totally used by fermentative bacteria (Fig. 8), the coexistence of SRB using large-molecule organic substrates with fast-growing fermentative bacteria is possible. From an ecological viewpoint, it would be interesting to understand how different SRB, which use different electron donors, compete for sulfate when it is limiting. However, to date, only a few studies have addressed this competition for sulfate (51). The coexistence of different SRB in our biofilm seems to indicate that their affinities to sulfate are similar.

Though SRB mainly inhabited the outer layer of the sewer biofilm, small amounts of Desulfobulbus, Desulfomicrobium, and Desulfovibrio were also observed in the inner layers (Fig. 6). Since sulfate did not penetrate here, the SRB in the inner layers probably grew by fermenting organic matter. Desulfobulbus species can ferment lactate and ethanol (plus carbon dioxide) to acetate and propionate in the absence of sulfate, and many Desulfovibrio and Desulfomicrobium species grow by fermenting pyruvate to form acetate, carbon dioxide, and hydrogen as products (51, 54, 55). In comparison, Desulfotartrula and Desulfatiferula were not detected in the inner layers, as the ability of these SRB to ferment organic matter is limited (49, 57).

Of the MA, about 90% of the population was Methanoseta, which is an obligate acetoclastic methanogen. Therefore, acetate was likely the main substrate for methanogenesis in anaerobic sewer systems. Since acetate could be simultaneously produced and consumed in the sewer system, the accumulation of acetate is probably due to the production rate of acetate being higher than its consumption rate under the tested conditions. Currently, the only genera known to use acetate for methanogenesis are Methanoscina and Methanoseta. However, Methanoscina failed to inhabit in the anaerobic sewer biofilms, which is consistent with the finding that usually only one acetoclastic methanogen dominates such anaerobic environments (60). It is likely that Methanoseta out-competed Methanoscina due to differences in their affinities for acetate. Methanoseta is a superior acetate utilizer in that it can use acetate at concentrations as low as 5 to 20 μM, while Methanoscina requires a minimum concentration of about 1 mM (61). The acetate concentration in the wastewater was about 0.6 mM and would therefore not favor the growth of Methanosaricina.

MA that use other substrates such as hydrogen or methylated compounds accounted for only less than 10% of the total MA population in the sewer biofilm. The hydrogen-utilizing MA mainly belonged to genera Methanobrevibacter, Methanospirillum, and Methanobacterium. Theoretically, the relative contributions of acetate and hydrogen in methanogenesis are close to 2:1, given the fact that the fermentation of hexose yields 4 H2, 2 acetate, and 2 CO2 and that 4 H2 are required to reduce CO2 to methane (62). One possible reason for the low abundance of hydrogentrophic MA in the sewer biofilms was the low hydrogen concentration in the system. This can be explained by the fact that hydrogentrophic MA were out-competed by the hydrogen-utilizing SRB, which have higher affinity and lower threshold values for hydrogen (35, 36). In addition, at 20°C, homoacetogenesis might occur, which could also out-compete methanogenesis for hydrogen (63). It is interesting that, although there were more SRB in the outer layer of the biofilm, the hydrogen-utilizing MA were more abundant in the outer layer than in the inner layers (Fig. 6). Though hydrogen was largely consumed by SRB, still more hydrogen was available in the outer layer, due to H2-producing bacteria having a higher abundance in the outer layer (data not shown). Methanomethylovorans and “Candidatus Methanomethylphilus” are known to use methylated compounds such as methanol, methanethiol, and dimethyl sulfide for methanogenesis (64–66). Their low abundance could be explained by the relatively low concentrations of these substrates in the wastewater (67, 68).

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