Dynamics of Different Bacterial Communities Are Capable of Generating Sustainable Electricity from Microbial Fuel Cells with Organic Waste

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(Received October 1, 2013—Accepted February 18, 2014—Published online April 30, 2014)

The relationship between the bacterial communities in anolyte and anode biofilms and the electrochemical properties of microbial fuel cells (MFCs) was investigated when a complex organic waste-decomposing solution was continuously supplied to MFCs as an electron donor. The current density increased gradually and was maintained at approximately 100 to 150 mA m⁻². Polarization curve analyses revealed that the maximum power density was 7.4 W m⁻³ with an internal resistance of 110 Ω. Bacterial community structures in the organic waste-decomposing solution and MFCs differed from each other. Clonal analyses targeting 16S rRNA genes indicated that bacterial communities in the biofilms on MFCs developed to specific communities dominated by novel Geobacter. Multidimensional scaling analyses based on DGGE profiles revealed that bacterial communities in the organic waste-decomposing solution fluctuated and had no dynamic equilibrium. Bacterial communities on the anolyte in MFCs had a dynamic equilibrium with fluctuations, while those of the biofilm converged to the Geobacter-dominated structure. These bacterial community dynamics of MFCs differed from those of control-MFCs under open circuit conditions. These results suggested that bacterial communities in the anolyte and biofilm have a gentle symbiotic system through electron flow, which resulted in the advance of current density from complex organic waste.

Key words: population dynamics, community structure, microbial fuel cell, Geobacter, selective enrichment

Technologies that convert organic waste to energy are needed for the construction of a sustainable society. Renewable organic waste accounted for approximately 60% (3.2×10⁴ tons year⁻¹) of the total amount of waste produced in Japan in 2008 (40). Although chemical and biological approaches to sustainable energy production, such as methane, ethanol, and hydrogen, have been developed, many of these approaches have encountered technical and economical hurdles (12, 14). Microbial fuel cells (MFCs) represent an alternative strategy capable of directly converting organic waste to electricity (34, 35).

MFCs are devices that exploit microorganisms as “biocatalysts” to generate electric power from organic matter. MFC systems have been researched as a method of recovering energy from organic waste in the form of electrical power (17, 34, 35, 43) and generating power from aquatic sediments on the bottom of the ocean (37, 53), rice paddy field soil (25, 27), wheat straw biomass (56) or cellulose (45). However, the current densities produced have been too low for practical use.

Many studies have been conducting to improve the potential of MFCs, including the development of a harness, electrode and proton exchange membrane (7, 21, 24, 32, 33, 46) or the analysis of exoelectrogens in pure cultures (15, 57), especially Geobacter and Shewanella strains (2, 4, 44). Two kinds of extracellular electron transfer (EET) mechanisms have been identified from a bacterial cell to an anode; Geobacter spp. use indirect EET through mediator compounds (38). Although Geobacter strains are known as high electricity-producing bacteria, the electricity-producing activities of complex or mixed cultures were shown to be higher than those of pure Geobacter cultures (25). These findings indicated that it may be possible to produce higher current density by controlling the complex microbial ecosystem in the anode chamber. To practically use MFCs, complex organic matter (i.e., food garbage and wastewater) has to be supplied as electron donors, while complex microbial communities have to simultaneously develop in the anode chamber. However, strategies to control microbial ecosystems capable of efficiently producing electricity have yet to be developed. The aim of the present study was to characterize the relationship between bacterial community structures and their electricity-producing properties, and also to evaluate the performance of MFCs continuously supplied with real organic waste.

Materials and Methods

MFC configuration and operation

A mediator-less and air-cathode MFC was used to examine power generation with organic waste, in which paddy field soil was inoculated into the MFC as the initial inoculum. Fig. 1 shows the experimental system used in this study. A carbon paper electropolated with platinum (0.5 mg cm⁻²) on one side was used as the cathode electrode (Chemix, Sagamihara, Japan), providing a total projected cathode surface area (on one side) of 16 cm². A proton exchange membrane (Nafion 117, DuPont, Delaware) was placed between the anode and cathode. A total of 135 pieces of cubic (125 mm³) graphite felt (Sohgoh-C, Yokohama, Japan) were packed into the anode chamber (36 mL in capacity) and the total projection surface area of the anode was 0.02025 m². Twenty of these pieces were directly connected to the platinum wires (0.3 mm; AlfaAesar). A
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The total of 0.4 g of paddy field soil was inoculated into the MFC with 20 mM lactate as the initial carbon and energy sources, and the electrodes were then connected with an external resistor (10 Ω) 5 days after the inoculation. As a control, an MFC was run under an open circuit condition (control MFC) that was also constructed with the same materials. Organic waste was collected from the canteen and 20 g or 40 g of this was placed directly in a bottle (organic waste). Organic waste was collected from the anode chamber in MFCs. Twenty pieces of cubic graphite felt were directly connected with the platinum wire. Panel (B); A total of 115 pieces of cubic graphite felt were set in the anode chamber. Panel (C); This shows the MFCs used in this study.

**Bacterial community analyses**

The analytic culture (1.0 mL) was directly sampled from the anode compartment of MFCs and cells were collected by centrifugation for 5 min at 4°C and 20,000×g. Pieces of the anode were collected and maintained at −20°C until DNA extraction. DNA was digested with *Escherichia coli* genomic DNA (Promega, Madison, WI, USA) according to the manufacturer’s recommendations. Clones were produced into competent DH5α cells using the Target CloneTM-Plus-kit (FMC Bioproducts) recommended by the manufacturer. Cloned genes were sequenced with an ABI PRISM BigDye Terminator version 3.1 cycle sequencing kit and analyzed with an ABI PRISM 3100-Avant genetic analyzer (Applied Biosystems). Sequence data were analyzed for chimeras with the CHIMERA_CHECK program deposited in databases using the BLAST homology search system (10). Sequencing data were compared with those deposited in databases using the BLAST homology search system. The multiple alignments of sequences and calculation of the nucleotide substitution rate by Kimura’s two-parameter model (31) were performed using the CLUSTAL W program (54). Distance matrix analysis, based on the Bray-Curtis index, was used to analyze the dynamics of the bacterial community structure because this index has been recognized as one of the most useful methods for evaluating differences among populations (8, 13). The following equation was used to calculate the Bray-Curtis index.

\[
\delta_{AB} = \left( \frac{\sum (n_A + n_B)}{\sum (n_A - n_B)} \right) 0 \leq \delta_{AB} \leq 1
\]

where \( \delta_{AB} \) means the dissimilarity index between communities A and B, \( n_A \) and \( n_B \) mean the intensities of DGGE bands in clusters of A and B, respectively (3, 18, 39). MDS analysis and cluster analysis were conducted using the R software program v2.12.1 (The R Project for Statistical Computing; http://www.r-project.org/; University of Tsukuba, Japan: http://cran.md.tsukuba.ac.jp/) (51). Commands used in the R software program v2.12.1 were shown in supplemental datum 1. The 3D graph was constructed using RINEARN Graph 3D v5.2.0.

**Clonal analyses**

Bacterial community structures were examined using 16S rRNA clone library analysis. The paddy field soil used as the inoculum was analyzed as the sample at day 0. Cells were collected from the anode liquid and anode biofilm of MFCs on day 34 and day 168. The DNA fragments of 16S rRNA genes were amplified using the primers 5'-AGAGTTTGATCCTGGCTCAG-3' (corresponding to *Escherichia coli* 16S rRNA gene positions 8 to 27) [5] and 5'-AAGGAGGTTGATCCAGCC-3' (corresponding to *Escherichia coli* 16S rRNA gene positions 1525 to 1542). Amplification was performed with the thermal cycler PC320 (ASTEC, Osaka, Japan) by using 50 μL of the mixture containing 0.5 U of KOD FX DNA polymerase (Toyobo, Osaka, Japan), buffer solution attached with the PCR kit, each deoxynucleoside triphosphate at a concentration of 400 μM, 15 pmol of each primer, and 50 ng of template DNA. The PCR conditions were 2 min to activate the polymerase at 94°C, then 25 cycles of 1 min at 94°C, 1 min at 53°C, and 1 min at 72°C, and finally 10 min of extension at 72°C. The PCR products were assessed by electrophoresis on a 1.5% (w/v) agarose gel in TAE buffer (48) and stained with GelRedTM (Wako, Osaka, Japan). Cloned genes were sequenced with an ABI PRISM BigDye Terminator kit and analyzed with an ABI PRISM genetic analyzer. Cloned genes were sequenced with an ABI PRISM BigDye Terminator kit and analyzed with an ABI PRISM genetic analyzer.

**Sequencing and phylogenetic analyses**

Cloned genes were sequenced with an ABI PRISM BigDye Terminator version 3.1 cycle sequencing kit and analyzed with an ABI PRISM 3100-Avant genetic analyzer (Applied Biosystems). Sequence data were compiled with the GENETYX-MAC program (GENETYX, Tokyo, Japan). 16S rRNA gene sequence data were analyzed for chimeras with the CHIMERA_CHECK program version 2.7, and compared with those retrieved from Ribosomal Database Project II (10). Sequencing data were compared with those deposited in databases using the BLAST homology search system. The multiple alignments of sequences and calculation of the nucleotide substitution rate by Kimura’s two-parameter model (31) were performed using the CLUSTAL W program (54). Distance matrix trees were constructed by the neighbor-joining method (47), and the topology of the trees was evaluated by bootstrapping with 1,000 resamplings (16).

**MDS analyses**

Bacterial community structures were also analyzed by denaturing gradient gel electrophoresis (DGGE) analysis that targeted 16S rRNA genes. The variable region, V3 of bacterial 16S rRNA genes (corresponding to positions 341 to 534 in the *Escherichia coli* sequence) was amplified using the primers P2 and P3 (containing a 40 bp GC clamp [41]) and thermal cycler PC320, as described previously (20). A Dcode DGGE system (Bio-Rad Laboratories, Hercules, CA, USA) was used for electrophoresis as recommended by the manufacturer. A total of 10 μL of the PCR-amplified mixture was subjected to electrophoresis in a 10% (w/v) polyacrylamide gel at 200 V for 3.5 h at 60°C. Gel gradients used for separation, which were applied in parallel to the electrophoresis direction, were 35% to 55%. After electrophoresis, the gel was stained with SYBR Green I (FMC Bioproducts) for 30 min as recommended by the manufacturer.

The intensities of bands in the DGGE gel were measured by Gel Doc XR+ (Bio-Rad Laboratories). Multidimensional scaling (MDS) analysis was performed with these band intensities. Since DGGE analysis does not necessarily completely reproduce the same results, all the intensities and locations of the DGGE bands used in the MDS analysis were compensated for by comparing the intensities and locations of common samples in different DGGE gels. MDS analysis, based on the Bray-Curtis index, was used to analyze the dynamics of the bacterial community structure because this index has been recognized as one of the most useful methods for evaluating differences among populations (8, 13). The following equation was used to calculate the Bray-Curtis index.

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\]

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Real-time PCR to monitor Geobacter strains

Real-time PCR assays were performed on genomic DNA to measure the 16S rRNA gene copy number of Geobacteraceae. Template DNAs were prepared as described above. The Geobacter spp.-specific primers used for real-time PCR were Geobacteraceae-494f (5’-AGGAGAGGACCGGCTAATCC-3’) (22) and Geo285r (5’-TACCCGCTACACTATG-3’) (1). Standard DNA fragments were produced using cloned DNA affiliated with the genus of Geobacter. The PCR profile consisted of preheating at 95°C for 10 min, followed by 40 cycles of denaturation at 95°C for 10 s, annealing at 65°C for 5 s, and extension at 72°C for 15 s. The fluorescence signal was detected at 72°C in each cycle, and a melting curve was obtained by heating the product to 95°C and cooling to 40°C. The novel Geobacter clade-specific primers used for real-time PCR were designed by aligning the Geobacter 16S rRNA sequences obtained from this experiment and GenBank; Novel Geobacter-f (5’-GAGGCCTCTGATATGCTTCTGTA-3’) and Novel Geobacter-r (5’-AGCATAACGGGTATTAACCCG-3’) (Fig. S4). Standard DNA fragments were produced using a cloned DNA affiliated with the novel Geobacter clade. Real-time PCR was conducted under the same conditions used for Geobacteraceae. The reaction was performed using a LightCycler FastStart DNA Master SYBR GREEN I kit (Roche Molecular Biochemicals, Indianapolis, IN, USA) and LightCycler system (Roche Diagnostics, Mannheim, Germany) according to the manufacturer’s instructions. The copy number of amplicons was calculated using LightCycler software version 3.52.

Electrochemical analyses

Voltage across the external resistor (10 Ω) was automatically monitored every 5 min using a data logger (GL200A, Graphitec, Tokyo, Japan) connected to a personal computer. To evaluate the performance of MFCs, a polarization curve was measured using a potentiostat (HAV-110, HOKUTO DENKO, Tokyo, Japan) at 2 mV min⁻¹ of a slope range at an approximate interval. Cell-performance indices (open-circuit voltage [Voc], short-circuit current density per volume of the anode harness [Isc], maximum power density per volume of the anode harness [P max], and internal resistance [R int]) were calculated from the slopes of the polarization curves. In some tests, an Ag/AgCl reference electrode (0.199 V versus standard hydrogen electrode [SHE], HX-R6, Hokuto Denko) was placed into the anode compartments to determine individual electrode potentials. Coulombic efficiency was obtained by calculating the ratio of total recovered coulombs by integrating the current over time to the theoretical amount of coulombs that could be produced from organic waste (see Chemical analysis). Detailed information can be found in a previous study (32).

Chemical analyses

Liquid samples including small particles were collected from the effluent solution of the tank to measure the redox potential, pH, and COD cr by a colorimetric standard method (5220D. Closed Reflux, Colorimetric Method). The redox potential and pH were measured by the electrode (TPX-999Si, Teko Chemical Lab., Tokyo, Japan). COD cr has primarily been used in MFCs (50) because the amount of COD cr removed by anode microbial metabolism can be used to calculate the number of electrons released from organics (1 g of COD cr was shown to be equivalent to 125 mmol of electrons [42]). These liquid samples were also filtered with a syringe filter (Millipore LG [pore size; 0.2 μm, diameter; 13 mm], EMD Millipore, Billerica, MA, USA) and organic acids were then analyzed by HPLC equipped with a Shodex HRPacked KC-811 column (300×8.0 mm) (Showa Denko, Kanagawa, Japan) and UV detector. The column oven was set at 50°C. Samples were eluted with 0.1% H3PO4 solution at a flow rate of 1.0 mL min⁻¹ and elutes were monitored at 210 nm. Formate, pyruvate, lactate, butyrate, and acetate were identified according to the retention time, and the concentration was determined by comparing the peak area with that of its respective standard sample.

Accession numbers

The nucleotide sequences reported here have been deposited in the DDBJ under accession numbers AB716976 to AB717136, AB720077, AB721108 to AB721264, AB721266 to AB721283, AB724116 to AB724215, AB724352 to AB724383, AB731904 to AB731955, and AB844727 to AB844796.

Results

Decomposition of organic waste and performance of MFCs

Organic waste was directly added to the tank [Fig. 1 (b)] at every 20 to 40 d interval (Fig. S1) and physicochemical parameters were monitored (Fig. 2, Fig. S2 and S3). Current generation was observed at approximately day 20, and

Fig. 2. (A) Electricity generation from MFCs with 10 Ω of external resistance. (B) Monitoring of organic acid concentration in the effluent from the organic waste-decomposing tank. Arrows mean the time of addition of organic waste into the organic waste-decomposing tank. Red Diamond: lactate, green square: acetate, black circular: butyrate, purple triangle: propionate, blue bar: formate. Black bars mean the time when organic acid concentrations were not analyzed. (C) Enumeration of Geobacteraceae population density by real-time PCR with specific primers. Black square and solid line: the MFC (closed circuit), open circular and broken line: the control-MFC (open circuit).
Table 1. Electricity-producing properties of MFCs used in this study

<table>
<thead>
<tr>
<th>Incubation time (d)</th>
<th>Voc (mV)</th>
<th>Maximum current density (mA m(^2) anode)</th>
<th>Maximum power density (W m(^{-2}))</th>
<th>Internal resistance (Ω)</th>
</tr>
</thead>
<tbody>
<tr>
<td>27</td>
<td>160</td>
<td>4</td>
<td>0.12</td>
<td>2300</td>
</tr>
<tr>
<td>34</td>
<td>500</td>
<td>44</td>
<td>4.9</td>
<td>370</td>
</tr>
<tr>
<td>69</td>
<td>300</td>
<td>140</td>
<td>5.3</td>
<td>160</td>
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<tr>
<td>105</td>
<td>475</td>
<td>63</td>
<td>6.7</td>
<td>200</td>
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<tr>
<td>167</td>
<td>470</td>
<td>200</td>
<td>6.9</td>
<td>140</td>
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<tr>
<td>182</td>
<td>500</td>
<td>220</td>
<td>7.4</td>
<td>110</td>
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</tbody>
</table>

current density then increased to approximately 80 to 120 mA m\(^{-2}\) between day 55 and day 85 (Fig. 2A). After the generation of current declined slightly between day 90 and day 125, current-generating activity developed and reached approximately 180 mA m\(^{-2}\) between day 165 and day 185 (Fig. 2A). Electrochemical analyses revealed that the maximum current and power densities also increased from 4 mA m\(^{-2}\) and 0.12 W m\(^{-2}\) to 220 mA m\(^{-2}\) and 7.4 W m\(^{-2}\), respectively, while Voc was stable at 470 to 500 mV (Table 1). Internal resistance significantly decreased from 2300 Ω to 110 Ω.

Lactate, acetate, propionate, formate, and butyrate were detected and their concentrations varied from several mM to 30 mM (between day 45 and day 200). In the initial stage (between day 0 and day 45), lactate was detected after the addition of organic waste, and the maximum concentration was approximately 30 mM. On the other hand, acetate, propionate, and butyrate were detected after the initial stage, and their concentrations varied from approximately 5 mM to 30 mM (between day 45 and day 200) (Fig. 2B). In the middle stage (between day 95 and day 150), the proportion of acetate detected in organic acids was higher than that in the other stages. The redox potential and pH ranged from −200 mV to −450 mV and from 5.6 to 7.8, respectively (Fig. S2). COD\(_a\) and coulombic efficiency ranged from 375 mg L\(^{-1}\) to 11,000 mg L\(^{-1}\) and from 0.3% to 46%, respectively (Fig. S3).

Bacterial community structure

Clonal analyses targeting the 16S rRNA gene were performed to investigate the bacterial community structure in paddy filed soil, the tank, control-MFCs, and MFCs (Fig. 3). The results of sequence analyses were summarized in Supplemental Table S1. Sequence analysis revealed that *Firmicutes*, α- and β-proteobacteria (denoted the Faβ-group) dominated over 90% of all communities in the tank and anolytes of both the control-MFCs and MFCs on day 34 (Fig. 3. Lanes T34, C34A, M34A). The community structure of soil was more diverse than those of the tank, the control-MFCs and the MFCs in which the proportion of the Faβ-group to total soil communities was 76% (Fig. 3. Lane S). Although the Faβ-group also dominated 92% of all biofilm communities in control-MFCs on day 34 (Fig. 3. C34B), it only dominated 55% of those in MFCs (Fig. 3. M34B).

Forty-eight out of the 100 clones analyzed were affiliated with *Acetobacterium* sp. HAAP-1 (98% identity) in *Firmicutes*, which was most frequently detected in paddy field soil. The clones related to *Firmicutes bacterium BV9-3a* (94% identity), belonging to *Firmicutes*, and those related to *Holophaga foetida* (89% identity), belonging to *Betaproteobacteria*, were the second and third most frequently detected, respectively (Fig. 3. Lane S). Forty of the 46 clones belonging to β-proteobacteria were *Microvirgula aerodenitrificans* (over 99.8% identity), which were most frequently detected in the tank on day 34 (Fig. 3. Lane T34).

These results indicated that the anolytic bacterial community structures in control-MFCs and MFCs were similar to those in the tank on day 34, (Fig. 3. Lane T34, C34A, and M34A). However, their biofilm community structures were different (Fig. 3. Lane T34, C34B, and M34B). On the other hand, differences were observed in bacterial community structures among the tank and anolytes in control-MFCs and MFCs on day 168. For example, the percentage of α-proteobacteria in the anolytic bacterial community in MFCs (33%) was significantly higher than those in the tank.
Dynamics of a microbial ecosystem in MFCs

and control-MFCs (7%) (Fig. 3. Lanes T168, C168A, and M168A). Furthermore, the biofilm bacterial community structure in MFCs was significantly different from that in control-MFCs (Fig. 3. Lanes C168B and M168B). \( \delta \)-proteobacteria in particular dominated 81% of the biofilm bacterial communities in MFCs on day 168 (Lanes M168B), and only consisted of Geobacter spp. (Table S1). On the other hand, \( \delta \)-proteobacteria dominated half of the biofilm bacterial communities in MFCs on day 34 (Fig. 3. Lane M34B). These results indicated that bacterial communities adapted to the respiration process by using anode electrodes as electron acceptors.

Phylogenetic analysis and enumeration of Geobacteraceae populations

Since Geobacter species are known to be high electricity-producing bacteria, improvements in MFCs are important for analyzing and monitoring the communities in MFCs. Clones closely related to Geobacteraceae were detected in paddy field soil (2 clones), the biofilm in control-MFCs on day 34 (2 clones), the anolyte and biofilm in control-MFCs on day 168 (2 clones and 2 clones, respectively), and the biofilm in MFCs on day 34 and day 168 (17 clones and 46 clones, respectively).

Phylogenetic analysis revealed that Geobacter communities were grouped into four clusters consisting of the Geobacter metallireducens clade, G. subsurface clades I, II, and a novel clade (Fig. 4). Most Geobacter spp. from the biofilm in MFCs on day 34 belonged to the G. metallireducens clade (10 clones out of 16 shared 97% identity with Geobacter hydrogenophilus), while those on day 168 belonged to the novel clade (42 clones out of 46...
shared 94% identity with *Geobacter lovleyi* SZ). This result indicates that the *Geobacter* community was markedly changed and converged to a specific genotype. Clonal analysis targeting the 16S rRNA gene showed that clones closely related to *Geobacter* spp. dominated the biofilm population, approximately 50% and 80% of all analyzed clones on day 34 and day 168, respectively (Fig. 3).

Real-time PCR analyses revealed that the population density of *Geobacteraceae* in the anode biofilm in MFCs increased from approximately $6.3\pm0.25\times10^6$ copies cm$^{-2}$ on day 34 (current density was 28 mA m$^{-2}$) to $2.4\pm0.12\times10^7$ copies cm$^{-2}$ on day 74 (74 mA m$^{-2}$). Population density decreased to $2.8\pm0.10\times10^6$ copies cm$^{-2}$ on day 168 (164 mA m$^{-2}$), but increased again to $6.4\pm0.22\times10^7$ copies cm$^{-2}$ on day 198 (7.5 mA m$^{-2}$) (Fig. 2). The population density of the novel *Geobacter* clade was approximately $5.3\pm0.86\times10^6$ copies cm$^{-2}$ on day 74 and $1.4\pm0.05\times10^7$ copies cm$^{-2}$ on day 168. On the other hand, population density in the anode biofilm in control-MFCs increased from approximately $3.0\pm0.12\times10^5$ copies cm$^{-2}$ on day 34 to $1.0\pm0.08\times10^6$ copies cm$^{-2}$ on day 74, but decreased to $5.3\pm0.21\times10^5$ copies cm$^{-2}$ on day 198 [Fig. 2 (C)].

**Bacterial community dynamics**

MDS analyses based on DGGE profiles were performed to understand the dynamics of the bacterial community in the MFC system. All stress values were less than 0.20, which indicated that these data were valuable statistically. The bacterial community structure in the tank always changed with no dynamic equilibrium (Fig. 5A). On the other hand, the community structures of analytic bacteria in both control-MFCs and MFCs had dynamic equilibrium with fluctuations, and also differed from each other (Fig. 5D). The biofilm-community structures in both MFCs did not have dynamic equilibrium and developed different structures from those of the anolytic community structures (Fig. 5B–D). The dissimilarity index value of the tank changed within the scope from 0.22 to 0.96 and the average value was 0.71±0.18. The dissimilarity index values of the anolyte bacterial community in control-MFCs varied from 0.35 to 0.96 and the average was 0.72±0.14, while these values in MFCs were from 0.32 to 0.84 and the average was 0.62±0.14. These results indicated that the anolyte-bacterial community structure in MFCs fluctuated less than that in control-MFCs. The dissimilarity index values of the biofilm bacterial community in control-MFCs varied from 0.81 to 1.00 and the average was 0.90±0.10, while these values in MFCs were from 0.71 to 1.00 and the average was 0.83±0.15. These results indicated that selective pressure on the surface of the anode was stronger in MFCs than in control-MFCs.

**Discussion**

Many studies have been conducted on MFCs, and bacterial metabolism was shown to be of significant importance due its effect on the efficiency of electricity production (52). The metabolic processes of microorganisms under anaerobic conditions connect with each other, and this is known as

![Fig. 5. Multidimensional scaling (MDS) analyses based on DGGE profiles. (a); the organic waste-decomposing tank, (b); biofilm (orange) and anolyte (red) in MFCs, (c); biofilm (purple) and anolyte (green) in control-MFCs, (d); All plots were shown under the same scales. The number beside the plots means the sampling date and “B” beside the number means the biofilm sample.](image-url)
interspecies hydrogen transfer. Bacterial community structures are known to be affected by substrates supplied to the anode of MFCs (6, 28, 49). A clear understanding of the anaerobically complex microbial ecosystem in the anode of MFCs is important when attempting to improve MFCs for practical use. The results obtained in the present study revealed the relationship between anolyte and anode biofilm bacterial communities and the electrochemical properties of the MFCs used, and also provided an insight into why the performance of MFCs was increased and maintained during this experiment [Fig. 2 (A) and Table 1] regardless of the different components of organic waste supplied as electron donors (Fig. S1).

The MFCs used in this study had a power density of approximately 1.1 W m\(^{-3}\) (or 2.0 W m\(^{-2}\)) with electricity production of 100 mA m\(^{-2}\). Although these MFCs were the middle to high performance type (28, 49, 56), the development of MFCs, in which performance is increased 1000-fold, is needed for their practical application. pH was almost stable at approximately 6.5 irrespective of diverse and complex organic waste (Fig. S1 and S2), which contributed to the sustainable production of electricity. On the other hand, no obvious positive correlation was observed between current density and redox potential [Fig. 2(A) and Fig. S2]. There appeared to be a negative correlation between COD and coulombic efficiency (Fig. S3). However, since the current density was almost stable during this experiment, it was suggested that the change of coulombic efficiency was pretense.

Although bacterial communities in the tank had been always fed into control-MFCs and MFCs, MDS analyses demonstrated that the dynamics of bacterial community structures differed between the tank, control-MFCs, and MFCs (Fig. 3 and Fig. 5D). The anolytic bacterial community structure in MFCs fluctuated less than that in control-MFCs, and the dynamic equilibrium of anolytic bacterial community structures in both MFCs differed from each other (Fig. 5D). Furthermore, the difference observed in the biofilm bacterial community structure between control-MFCs and MFCs was larger than that in anolytic bacterial community structures (Fig. 5D). These results demonstrated that the anode electrode as an electron acceptor was directly involved in the dynamics of biofilm bacterial communities on the anode and also indirectly influenced the dynamics of anolytic bacterial communities. Our results suggested that the relationship between the anolytic and biofilm bacterial communities has a gentle symbiotic system through the electron flow. Furthermore, the anaerobic microbial ecosystem may also be controlled by using the anode electrode as an (terminal) electron acceptor. A few studies have also suggested the presence of electric symbiosis between fermenters in an anolyte and exoelectrogens on an anode surface (26, 30, 36).

By combining the results of clonal and MDS analyses, we showed that the dynamics of the biofilm bacterial community in MFCs converged on the selective enrichment of Geobacter (Fig. 3 and Fig. 5D). Geobacteraceae has been classified into three clades: the Geobacter metallireducens clade, and subsurface clades 1 and 2 (23). The key factors for the selective enrichment of Geobacter are diverse. Kato et al. reported that diverse Geobacter spp. were enriched in the soil layer of a batch culture supplied with (semi)conductive iron-oxide species as the terminal electron acceptor and sodium acetate as the electron donor (29). Other studies demonstrated that the addition of acetate stimulated the growth of Geobacter in field and pure culture (Geobacter sulfurreducens PCA) experiments, resulting in high electroactivity (23, 52). However, a positive correlation was not necessarily observed between the population density of Geobacter and acetate concentration in the tank in the present study (Fig. 2). A previous study demonstrated that complex organic acids alleviated the metabolic constraints derived from assimilation or dissimilation (52). Furthermore, lactate, butyrate, and propionate produced in the tank were typically difficult to use as carbon and energy sources, but were identified as electron donors for Geobacter (9). Therefore, these complex organic acids may have stimulated the growth and electroactivity of Geobacter in MFCs. These processes enabled the enrichment of Geobacter on the anode surface, resulting in an increase in the production of electricity. Phylogenetic analysis showed that the unique phylotype of Geobacter, which shared 94% identity with Geobacter lovleyi SZ, was enriched in the biofilm in MFCs on day 168 (Fig. 4). Since real-time PCR results demonstrated that the population of the novel Geobacter clade was 5-fold higher than that of Geobacteraceae detected with previous primer sets (Geobacteraceae-494f/Geo825r) on day 169, the specific primer set for the novel Geobacter clade designed here was useful for understanding the dynamics of Geobacteraceae. Commault et al. reported that closely related strains of Geobacter have different electron transfer capabilities (11), which suggested that the specific Geobacter species adapted to complex organic acids and was enriched on the anode in the present study. By combining the results of electrochemical properties (Table 1), we propose that the enrichment of the novel Geobacter clade on the anode may have contributed to the establishment of efficient electron transfer from cells to the anode, resulting in the production of a high current density.

Conclusion

In conclusion, although it has been suggested that the smooth flow of electrons from the electron donor to the electrode through microorganisms is important for efficient electricity production, information regarding the dynamics of the microbial ecosystem in the anode of MFCs supplied continuously with complex substrates remains sparse. This study characterized electron flow in the anode chamber standing on the sight of microbial ecosystem and the results presented herein demonstrated that anolytic and biofilm bacterial communities developed interdependency. Geobacter was enriched selectively and naturally on the surface of the anode. It is inherently important to establish how to control the microbial ecosystem. These subjects are currently under investigation in our laboratory.

Acknowledgements

We thank Prof. Yuichi Saeki for MDS analyses. This study was carried out as a part of grant KAKENHI(B) 22310045, Japan. It was also supported partially by ALCA project, Japan Science and Technology Agency.
References


