Diversity of Cultivable Methane-Oxidizing Bacteria in Microsites of a Rice Paddy Field: Investigation by Cultivation Method and Fluorescence in situ Hybridization (FISH)

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The diversity of cultivable methane-oxidizing bacteria (MOB) in the rice paddy field ecosystem was investigated by combined culture-dependent and fluorescence in situ hybridization (FISH) techniques. Seven microsomes of a Japanese rice paddy field were the focus of the study: floodwater, surface soil, bulk soil, rhizosphere soil, root, basal stem of rice plant, and rice stumps of previous harvest. Based on pmoA gene analysis and transmission electron microscopy (TEM), four type I, and nine type II MOB isolates were obtained from the highest dilution series of enrichment cultures. The type I MOB isolates included novel species in the genus Methydomonas from floodwater and this is the first type I MOB strain isolated from floodwater of a rice paddy field. In the type I MOB, two isolates from stumps were closely related to Methydomonas spp., one isolate obtained from rhizosphere soil was most related to Methyloccoccus Methylocaldum-Methylogaeae clade. Almost all the type II MOB isolates were related to Methylocystis methanotrophs. FISH confirmed the presence of both types I and II MOB in all the microsites and in the related enrichment cultures. The study reported, for the first time, the diversity of cultivable methanotrophs including a novel species of type I MOB in rice paddy field compartments. Refining growth media and culture conditions, in combination with molecular approaches, will allow us to broaden our knowledge on the MOB community in the rice paddy field ecosystem and consequently implement strategies for mitigating CH4 emission from this ecosystem.

Key words: Cultivable bacteria, diversity, methane-oxidizing bacteria, FISH, rice paddy field microsite.

Methane (CH4) is an important greenhouse gas ranking second to carbon dioxide, and wetland rice fields are one of the major sources of methane emission, accounting to 5–19% of the global CH4 budget (15). Methane emission from a rice field is the net effect of its production (methanogenesis) and its oxidation (methanotrophy). About 60–90% of the produced CH4 is oxidized in situ before it escapes to the atmosphere (56). Microbial CH4 oxidation driven by methane-oxidizing bacteria (MOB) is the only biological suppression of methane emission from rice fields, and consequently MOB are considered to be important regulators of methane effluxes from this ecosystem (39). MOB are a unique group of bacteria that oxidize CH4 with molecular O2 and use it as a carbon and energy source (9, 26). It is generally recognized that CH4 oxidation occurs at anoxic-anoxic interfaces in rice paddy fields, at the soil-water interface, and in the rhizosphere and rhizoplane of rice plants with available O2 and CH4 (18, 21). Consequently, intermittent water management (flooding/drying) and the resulting significant biogeochemical processes generated (34) may affect CH4 biological oxidation and probably the composition of MOB in the microsites of the rice paddy field ecosystem.

Methanotrophs are mostly classified into the Gammaproteobacteria (type I MOB) and the Alphaproteobacteria (type II MOB) based on their intracytoplasmic membrane structure, carbon-assimilation pathway, phospholipid fatty acid profile, and phylogenetic placement. Both types I and II MOB have been found in rice field bulk soil, rice rhizosphere, soil-water-interface, and on rice roots using phospholipid fatty acid analysis (38, 50), PLFA-stable isotope probing (50), 16S rDNA and pmoA sequencing (18, 20, 21, 32, 35, 40, 50). In contrast with these results revealed through molecular approaches and despite the heterogeneity of microsites in the rice paddy field ecosystem with regard to O2, CH4, and nutrient availability, only type II MOB have been isolated from rice paddy soil and rice roots using cultivation methods (16, 24, 52, 53). However more recently, using a combined molecular and cultivation technique, a mesophilic type I MOB was isolated at the soil-water-interface from a rice paddy field in Uruguay (21, 23), and is the first and sole type I strain isolated from rice paddy fields. Therefore, cultivable MOB diversity associated with microsites in rice paddy field ecosystem remains to be elucidated. In particular, no type I MOB have been isolated from the rice rhizosphere, rice roots, bulk soil or floodwater, although their presence was revealed using molecular techniques as mentioned above. It is commonly accepted that only a small fraction of microbes is cultivable and that molecular approaches always cover a broader spectrum of microbial diversity than cultivation methods (3, 22, 47), although in some cases this seems untrue (17, 36, 45). Therefore, in line with Hengstmann et al. (28),
Oremland et al. (45) and Donachie et al. (17), implementing comprehensive strategies, which include combined improvement of culturing and molecular techniques, may help to gain more insight into cultivable MOB diversity in the rice paddy field ecosystem.

In order to address this problem, we used cultivation techniques and fluorescence in situ hybridization (FISH) in a combined approach to characterize the cultivable MOB inhabiting floodwater, surface soil, rice stumps from the previous harvest, bulk soil, rhizosphere soil, root, and rice stem from a Japanese rice paddy field.

**Materials and Methods**

**Field site**

The studied site was a rice-wheat double cropping paddy field in Aichi-ken Anjo Research and Extension Center, central Japan (latitude 34°48'N, longitude 137°30'E). Principal soil characteristics as described by Watanabe et al. (37) and Jia et al. (32) were as follows: total C, 12.8 g kg⁻¹; total N, 1.1 g kg⁻¹; pH (H₂O), 6.3; amorphous Fe content, 3.76 g kg⁻¹. The soil was classified as Oxyaquic Dystrudept with a clay content of 230 g kg⁻¹. The environment for this study: floodwater (Fw), surface soil (SS), bulk soil (S), rhizosphere soil (RS), total root (homogenate) (Rt), basal stem (Ste), and stump (Stu; only in 2003) of the previous harvest. Floodwater, rhizosphere soil (RS), total root (homogenate) (Rt), basal stem (Ste), and stump (Stu; only in 2003) of the previous harvest. Floodwater, surface soil (10 g) were ground with a sterile mortar and pestle to give a homogenous mixture. The resulting samples were used for cultivation of cells or after enrichment. We focused on seven rice harvest stage (October 10, 2008) from plots E2 (in 2003) and B4 (in 2008), respectively. The samples in 2003 were used for isolation of approx. 20% (v/v) CH₄ in the dark for 3–4 weeks. Enrichment from floodwater was performed similarly with 5 mL sample added to 5 mL autoclaved NMS medium.

**Sampling scheme and rice paddy field microsites**

Samples were taken at the tilling stage (August 4 and 11, 2003), at the flowering stage (September 4, 2003), and at the maturity and harvest stage (October 10, 2008) from plots E2 (in 2003) and B4 (in 2008), respectively. The samples in 2003 were used for isolation of MOB and the samples in 2008 for FISH observation of MOB by extraction of cells or after enrichment. We focused on seven rice paddy field compartments (microsites; abbreviations in parentheses) for this study: floodwater (Fw), surface soil (SS), bulk soil (S), rhizosphere soil (RS), total root (homogenate) (Rt), basal stem (Ste), and stump (Stu; only in 2003) of the previous harvest. Floodwater, surface soil, and bulk soil (soil) between rice plants footpaths, and stumps were collected at the field. All samples were collected from three points randomly distributed in the plot and then mixed. Floodwater (5–10 cm depth) was collected with a 100 mL plastic beaker and poured into sterile plastic bottles. Surface soil (0–0.5 cm) and bulk soil (2–10 cm) were collected with a small trowel and sterile plastic bags. Samples from the other microsites of the rice field were obtained after taking the total rice plant with soil (three cores of approximately 20 cm diameter by 18 cm depth randomly distributed in the plot) to the laboratory. From each core, a rice plant was dug out by hand and shaken to remove loose soil on the roots. The fine layer of soil firmly attached to the roots was released into 100 mL nitrate mineral salt (NMS) medium. The resulting mixed soil solution obtained from the three cores (approximately 1 g [10 mL]⁻¹ was referred to as rhizosphere soil in this study. After removing the rhizosphere soil, the roots were rinsed again repeatedly with sterile distilled water until the water ran clear, before removing the remaining water by pressing between paper. The roots were then cut into small pieces and referred to as roots in this study. Basal stems (0–2 cm) of rice plants and stumps were rinsed with sterile distilled water to remove the remaining soil before they were pressed between paper and cut into small pieces as for roots. Bulk soil and surface soil (10 g) were ground with a sterile mortar and pestle to give a homogenous mixture. The resulting samples were used for enrichment and isolation of MOB.

**Preparation of enrichment cultures and MOB isolation**

All enrichment cultures were prepared using NMS medium (59). Trace elements were added before autoclaving according to Roslev and King (49) to give the following final concentrations: zinc chloride, 2.0 μM; cupric chloride, 2.0 μM; sodium bromide, 1.0 μM; sodium molybdate, 0.5 μM; manganese chloride, 2.0 μM; potassium iodide, 1.0 μM; boric acid, 2.0 μM; cobalt chloride, 1.0 μM; nickel chloride, 1.0 μM; and iron sulphate, 0.1 μM. The pH of the medium was then adjusted to 6.8 by addition of 1 M NaOH. The medium was then autoclaved at 121°C for 20 min.

MOB were isolated from the 2003 samples without enrichment (A), and from enrichment cultures prepared according to Whittenbury et al. (59) (B) and Bowman et al. (10) (C). Some factors (CH₄ pressure, shaking culture) were reported to affect the composition of MOB populations (4, 25, 27, 31, 60); therefore, we performed enrichments under 5 and 25% (v/v) CH₄ phase (D), and under 20% (v/v) CH₄ phase with shaking (E). For enrichment B (59), samples (0.5 g) except floodwater were added to 10 mL sterile NMS liquid medium in 60 mL flasks before the flasks were sealed with butyl rubber stoppers and an aluminium seal. Methane (6 mL) was then passed through a 0.20 μm pore filter with a gas-tight syringe and injected into each flask, giving about 10% (v/v) CH₄ in the head gas-phase. The flasks were incubated statically at 30°C in the dark for 3–4 weeks. Enrichment from floodwater was performed similarly with 5 mL sample added to 5 mL autoclaved NMS medium.

Enrichment C according to Bowman et al. (10) was prepared as follow: Surface soil, soil, root, stem, and stump materials (1 g) were added to 15 mL sterile NMS liquid medium in 34 mL test tubes (18 mm i.d.×180 mm) with glass beads and mixed for 3–5 min. Rhizosphere soil solution with the same concentration as the material was treated similarly. The supernatants of the solutions and floodwater (250 mL) were filtered through 0.20 μm pore filters. The filters were placed into 60 mL flasks containing 10 mL sterile NMS liquid medium. The flasks were then incubated statically at 30°C and 37°C under about 20% (v/v) CH₄ for 3–4 weeks. At the end of the incubation period, pellets formed at the surface and solution were used separately for serial dilutions and isolation. In enrichments under 5 and 25% (v/v) CH₄ (D), floodwater and rhizosphere soil (60 mL), and 60 mL of 5% NMS solutions (w/v) from surface soil, soil, root, stem, and stump samples were placed in sterile 120 mL flasks. The flasks were incubated at approximately 5% and 25% (v/v) CH₄ statically at 30°C for 3–4 weeks. For the enrichment E series, samples (200 or 320 mL) used for enrichment D were placed in 500 and 700 mL flasks. The flasks were incubated under about 20% CH₄ at 30°C with shaking (150 rpm) for 3–4 weeks.

In 2008, enrichments for FISH observation were performed according to Bowman et al. (10) (C), Wise et al. (60) and Auman et al. (5) with respect to type I MOB preferential cultivation. Surface soil, soil, rhizosphere soil, root, and stem (1 g) were mixed in 9 mL NMS medium with glass beads by vortexing vigorously for 3–5 min. Floodwater (250 mL) was centrifuged at 13,000 rpm for 10 min at 4°C before the pellet was resuspended in 9 mL NMS with glass beads by vortexing as above. The obtained suspensions were serially 10-fold diluted (10⁻¹ to 10⁻⁶) in 18 mL NMS medium supplemented or not with 20 μM copper in 121 mL flasks and the headspace was replaced by air containing 10% CH₄, 2% CO₂, (v/v) or 45% CH₄, 5% CO₂ (v/v). The flasks were then incubated at 30°C in the dark under moderate shaking (150 rpm) for 3–4 weeks. Additional enrichment series under 20% (v/v) CH₄ were performed in parallel according to method E described above.

Growth turbidity and CH₄ uptake in the flasks were periodically checked with a gas chromatograph (GC-9; Shimadzu, Kyoto, Japan), using a Porapak N column connected to a flame ionization detector (FID). The oven temperature was 60°C and He served as carrier gas.

**MOB isolation in pure culture**

For isolating MOB in 2003, the enrichment cultures were diluted by serial 10-fold dilution in 9 mL NMS medium in 34 mL test tubes up to 10⁻⁴ (2 dilution series). From the C enrichment process (10), pellets formed on the surface of filters and the culture solution were treated separately. The 10⁻¹ to 10⁻⁶ series were then spread
onto NMS agar (7 mL) slopes in 34 mL test tubes. The tops of the tubes were closed with butyl rubber stoppers and CH₄ (6 mL) was injected into each tube, giving about 18% (v/v) CH₄ in the head gas-phase. Control tubes without CH₄ injection were prepared in duplicate to detect colonies of non-methane-oxidizing contaminants. The tubes were then incubated at 30°C in the dark and observed at 3-day or 1-week intervals over 3–4 weeks. Single colonies formed from positive cultures were then transferred repeatedly onto fresh NMS agar slopes and incubated again for 1–3 weeks. Then, cultures were examined by phase-contrast microscopy using wet mounts with an Olympus BH-80. Subsequent transfers of a single colony and a 2–3-week incubation period at 30°C were determined with a 373S DNA Automated Sequencer or genetic analyzers (PRISM 310 Genetic Analyzer, PRISM Genetic Analyzer 3100 and ABI 3130 Genetic Analyzer; Applied Biosystems) with the protocol of whole cell hybridization was adapted from Amann et al. (2), Bourne et al. (8) and Eller et al. (19). Hybridizations were performed in 8-well Teflon-coated slides. Slides were preclenched by soaking for 1 h in 99.5% ethanol and rinsing in distilled water before they were washed in 1% (v/v) HCl and 70% (v/v) ethanol, and then air-dried. Fixed cell suspension (1–1.5 μL) was added to each well. The slides were dried at 46°C for 10 min in an oven (HB-80; TAITEC, Koshigaya, Japan), subsequently dehydrated by immersing in 50, 80, and 99.5% (v/v) aqueous ethanol for 3 min each, and then air-dried, before wells were covered by 8 μL hybridization buffer (Tris 2.4 g L⁻¹, SDS 2.0 g L⁻¹, EDTA 2.0 g L⁻¹, NaCl 0.9 M, 20% (v/v) formamide, pH 7.4). To each well, 1 μL probe solution (50 ng μL⁻¹) was added and hybridization was carried out for 1 h 30 min (cultures) or 2 h (cells extracted from microsite samples) at 46°C in a water-saturated atmosphere chamber (50 mL Falcon tube containing a piece of a 150 mm 5C-filter paper soaked in 4 mL hybridization buffer) in a hybridization incubator (HB-80; TAITEC). Unbound nucleotides were removed by rinsing the slides with 20 mL washing buffer (Tris 2.4 g L⁻¹, SDS 2.0 g L⁻¹, EDTA 2.0 g L⁻¹, pH 7.4, 225 mM NaCl containing 20% (v/v) formamide concentration in hybridization buffer) and prewarmed to 48°C in a water bath. Subsequently, the slides were washed with 8 μL washing buffer per well for 20 min at 48°C in a water-saturated atmosphere chamber (50 mL Falcon tube containing a piece of a 150 mm 5C-filter paper soaked in 4 mL washing buffer) in the hybridization incubator. They were then rinsed again with 20 μL washing buffer prewarmed to 48°C, air-dried, and then DNA-stained with 10 μL of 50 μL μL⁻¹ DAPI (4',6-diamino-2-phenylindole) solution for 10–15 min at room temperature in the dark. The remaining DAPI solution was removed by

**Phylogenetic analysis of 16S rDNA and pmoA sequences**

Cells of isolates were suspended in 600 μL TESS buffer (25 mM Tris-HCl, 5 mM EDTA 2Na; 50 mM NaCl; 25% [w/v] sucrose) with lysozyme (5 mg mL⁻¹), placed on ice for 30 min, and 30 μL of 10% (w/v) SDS and 20 μL Proteinase K (10 mg mL⁻¹) were added. The preparations were incubated at 50°C for 2 hours and then DNA was extracted with phenol-chloroform-isoamylalcohol and chloroform-isoamylalcohol reagents and by isopropanol and ethanol precipitation. Gene fragments of 16S rRNA and pmoA were amplified by PCR using the following primers: 27f/1492r (58) and A189f /A682r (30) or mb661r (14), respectively. The sequences were determined with a 373S DNA Automated Sequencer or genetic analyzers (PRISM 310 Genetic Analyzer, PRISM Genetic Analyzer 3100 and ABI 3130 Genetic Analyzer; Applied Biosystems) with the global alignment algorithm using the EzTaxon server (http://www.eztaxon.org/; [13]). Phylogenetic trees were constructed using the neighbor-joining method with the ClustalW program based on the web site of the DNA Data Bank of Japan (DDBJ; http://www.ddbj.nig.ac.jp/Welcome-ej.html) by 1,000 replication bootstrap analysis and nj plot software (46).
After rinsing the slides with distilled water.

After air drying, slides were mounted in immersion oil (05; Olympus). Epifluorescence microscopy was performed with an Olympus BX-FLA microscope (Olympus) equipped with a 100-1.25 Oil Ph3 immersion lens and fitted with a 50-W high pressure bulb and an image recorder (ICY-SHOT DXC-S500; Sony, Japan). The following Olympus light filter sets were used: NV (U-MNV) for phase contrast, WIG (U-MWIG) for Cy3 and NUA (U-MNUA) for DAPI.

Detection of MOB inhabiting rice paddy field microsites by FISH

Bacterial cells were extracted from the rice paddy field microsites using a method adapted from Eller et al. (19). Washed roots, surface soil, bulk soil, rhizosphere soil, basal stem (10–11 g), and floodwater (250 mL) were shaken with glass beads (diameter 1 mm) and sterile distilled water (30 mL in 50 mL Falcon tubes) on a horizontal shaker (TS-4N; TAITEC) for 30 min at 200 rpm. Decantation was then performed on ice for 15 min before transferring the supernatant into 15 mL sterile Falcon tubes. The extraction process was repeated and then the resulting mixed supernatants for each sample were centrifuged at 500–1,000 rpm for 2 min at 4°C to precipitate large particles. The resulting supernatants were centrifuged at 12,000–15,000 rpm for 10 min at 4°C and then the pellet (cells) was resuspended in 100 μL (or more according to pellet amount) PBS at pH 7.0 before adding 300 μL of 4% paraformaldehyde (w/v in PBS). After gentle mixing, cells were then left to fix for 2–16 h at room temperature. Whole cell hybridization and microscopy were then performed as described above for pure culture. FISH was performed in parallel on the same microsite samples collected on October 10, 2008 and on the issued enrichment cultures with appreciable CH₄ uptake and turbidity.

Accession numbers

The GenBank/EMBL/DDBJ accession numbers for the 16S rRNA gene and pmoA sequences of MOB strains in the present study are AB669143–AB669172.

Results

Characteristics of MOB isolates

From the enrichment cultures, 13 MOB isolates were obtained from floodwater (two), surface soil (two), bulk soil (two), rhizosphere soil (three), stem (one), and stump (three) (Table 1). The name of isolates was designated with the source, isolate number, enrichment method, and pigment, e.g. Fw12E-Y. All the MOB isolates were motile and stained Gram negative. Exospore or cyst formation was not observed for any MOB isolates. Cell morphology under phase-contrast microscopy varied from rod, short rod, short rod-curve, rod-curve to long rod and long rod-curve among the isolates (Fig. 1A–F). Although all colonies were opaque on agar, pigmentation varied (white, red, yellow to orange, yellow, purple to pink, or pink to purple). All isolates except SS10D-Y-Pr were able to grow with 0.1% (v/v) methanol. Except Stu1B-Pr, Stu5B-P-Pr, and Stu20C-Re, all isolates were able to grow at 37°C on NMS medium; however, none of the MOB isolates could grow at 45°C.

Table 1. Properties of methane-oxidizing bacterial isolates from rice paddy field microsites

| Isolate  | Cell morphology | Motility | Exospore formation | Gram stain | Colony aspect | Growth on 0.1% methanol | Growth at 37°C | Growth at 45°C |
|----------|----------------|----------|--------------------|------------|---------------|------------------------|----------------|----------------|
| Fw12E-Y  | SR             | +        | –                  | –          | Y-Or, O       | +                      | +              | –              |
| Fw1B-WF  | LRC            | +        | –                  | –          | W, O, F       | +                      | +              | –              |
| SS10D-Y-Pr | SR           | +        | –                  | –          | Y, O, G       | –                      | +              | –              |
| SS37A-Re | LR             | +        | –                  | –          | Re, O         | +                      | +              | +              |
| S5B-W    | SR             | +        | –                  | –          | W, O          | +                      | +              | –              |
| S18C-Re  | R              | +        | –                  | –          | Re, O         | +                      | +              | –              |
| RS5A-Re  | R              | +        | –                  | –          | Re, O         | +                      | +              | –              |
| RS6A-Re  | R              | +        | –                  | –          | Re, O         | +                      | +              | –              |
| RS11D-Pr | RC             | +        | –                  | –          | Pr-P, O       | +                      | +              | –              |
| Ste3C-Re | R              | +        | –                  | –          | Re, O         | +                      | +              | –              |
| Stu1B-Pr | LR             | +        | –                  | –          | P-Pr, O, G    | +                      | –              | –              |
| Stu5B-P-Pr | R            | +        | –                  | –          | P-Pr, O, G    | +                      | –              | –              |
| Stu20C-Re | R              | +        | –                  | –          | Re, O         | +                      | –              | –              |

1 Name of isolates was designated with the source, isolate number, enrichment method, and pigment: Fw, floodwater; SS, surface soil; S, bulk soil; RS, rhizosphere soil; Ste, stem; Stu, stump. A–E, enrichment methods. Y, yellow; W, white; Y-Pr, yellow to purple; Re, red; Pr, purple; P-Pr, pink to purple; F, filamentous. Isolates were cultivated at 30°C.

2 R, rod; SR, short rod; RC, rod curved; SRC, short rod curved; LR, long rod; LRC, long rod curved.

3 W, white; Y, yellow; Y-Or, yellow to orange; Re, red; P-Pr, pink to purple; Pr-P, purple to pink. O, opaque; F, filamentous; G, gelatinous.
Table 2. Closest species of methane-oxidizing bacterial isolates from rice paddy field microsites based on 16S rRNA gene sequence

| MOB isolate   | Alignment (Similarity %) | Closest species (Strain number in culture collection) | Taxonomic description | Accession number |
|---------------|---------------------------|-----------------------------------------------------|-----------------------|-----------------|
| Fw12E-Y       | 1271/1339 (94.9)           | Methylomonas scandinavica R5 (VKM B-2140)            | Gammaproteobacteria   | AJ151369        |
| Fw1B-W        | 1385/1402 (98.8)           | Methylocystis parvus OBBP (NCIMB 11129)              | Alphaproteobacteria   | Y18945          |
| SS10D-Y-Pr    | 1365/1390 (98.2)           | Methylocystis parvus OBBP (NCIMB 11129)              | Alphaproteobacteria   | Y18945          |
| SS37A-Re      | 1355/1405 (96.4)           | Methylocystis parvus OBBP (NCIMB 11129)              | Alphaproteobacteria   | Y18945          |
| SS5B-W        | 1365/1395 (97.8)           | Methylocystis parvus OBBP (NCIMB 11129)              | Alphaproteobacteria   | Y18945          |
| S18C-Re       | 1383/1404 (98.5)           | Methylocystis parvus OBBP (NCIMB 11129)              | Alphaproteobacteria   | Y18945          |
| RS5A-Re       | 1353/1394 (97.1)           | Methylocystis parvus OBBP (NCIMB 11129)              | Alphaproteobacteria   | Y18945          |
| RS6A-Re       | 1389/1406 (98.8)           | Methylocystis parvus OBBP (NCIMB 11129)              | Alphaproteobacteria   | Y18945          |
| RS11D-Pr      | 1351/1451 (93.1)           | Methylocaldum szegediense OR2 (NCIMB 11912)         | Gammaproteobacteria   | U89300          |
| Ste3C-Re      | 1382/1405 (98.4)           | Methylocystis parvus OBBP (NCIMB 11129)              | Alphaproteobacteria   | Y18945          |
| Stu1B-Pr      | 1428/1452 (98.3)           | Methylomonas methanica S1 (NCIMB 11130)              | Gammaproteobacteria   | AF304196        |
| Stu5B-P-Pr    | 1428/1452 (98.3)           | Methylomonas methanica S1 (NCIMB 11130)              | Gammaproteobacteria   | AF304196        |
| Stu20C-Pr     | 1389/1408 (98.7)           | Methylomonas parvus OBBP (NCIMB 11129)              | Alphaproteobacteria   | Y18945          |

1 NCIMB, The National Collections of Industrial and Marine Bacteria; VKM, All-Russian Collection of Microorganisms.

Phylogenetic characteristics of the MOB isolates

Sequence analysis of PCR amplification of nearly the full sequence of 16S rRNA genes (1,300–1,400 bp) of MOB isolates was carried out (Table 2). Isolates Stu20C-Re, Ste3C-Re, RS5A-Re, SS37A-Re, SS10D-Y-Pr, and Fw1B-W were most closely related to Methylocystis sp. EB-1 with 97.4–99.7% similarity, while RS6A-Re and S18C-Re, and SS5B-W showed 99.7–99.9% and 98.6% similarity to Methylocystis sp. KSPIII (100%), respectively. The closest species of these isolates was Methylocystis parvus, which showed 96.4–98.8% similarity to type strain RS6A-Re and S18C-Re, and SS5B-W showed 99.7–99.9% and 98.6% similarity to type strains R5 and S1T, which showed 96.4–98.8% similarity to type strain Methylocystis parvus, indicating that the isolates belonged to the genus Methylocystis (type II MOB). Four isolates, Fw12E-Y, RS11D-Pr, Stu1B-Pr, and Stu5B-P-Pr, were closely related to uncultured bacterium clone Er-MS-95 (96.1%), uncultured bacterium clone Er-MS-95 (96.1%), uncultured bacterium clone Er-MS-95 (96.1%), uncultured bacterium clone Er-MS-95 (96.1%), and uncultured bacterium clone Er-MS-95 (96.1%), respectively. The closest species of these isolates was Methylocaldum szegediense (type I MOB), and five within Methylocystis sp. EB-1 with 97.4–99.7% similarity to Methylocaldum szegediense (type I MOB) in OR2, respectively, indicating that the isolates belonged to the genus Methylocaldum (type II MOB). Isolate RS11D-Pr was most closely related to Methylocaldum szegediense OR2 (93.1%) (type I MOB) in OR2, respectively, indicating that the isolates belonged to the genus Methylocaldum (type II MOB).

Fig. 2. Phylogenetic analysis of the deduced amino acid sequences encoded by pmoA genes showing the relationships between the isolates/cultures and other methane-oxidizing bacteria. Bar = 0.1 substitutions per nucleotide sequence position. Bootstrap values more than 50% are shown as closed circles at the branch point. "Methylacidiphilum kamchatkense" Kam1 and "Methylacidiphilum lumidifusorium" SolV were used as an outgroup. Accession numbers of reference sequences are shown in parentheses. Deduced amino acid sequences encoded by pmoA gene analysis derived from the same paddy field are included. PS, paddy soil (32); RS, rice straw (33); MCR, microcrustaceans in floodwater (43); FW, floodwater (Shibagaki-Shimizu et al., unpublished results).
clade, while RS11D-Pr was distantly related to Methylocaldum-Methylococcus-Methylogaea cluster. Type II MOB isolates/cultures except for S1A-W were found closely related to Methylocystis (Methylocystis parvus and Methylocystis echinoides) clade, whereas S1A-W belonged to a distant cluster including Methylosinus sporium. Deduced amino acid sequences encoded by the pmoA gene derived from the same paddy field, i.e. paddy soil (32), rice straw (33), microcrustaceans in floodwater (43), and floodwater (Shibagaki-Shimizu et al., unpublished results) were included in the tree in Fig. 2. Only RS11D-Pr showed relationships to pmoA sequences of the clones.

Morphological observation of MOB isolates by TEM and FISH

Transmission electron micrograph of ultrathin sections of bacterial cells from the MOB isolates was performed. Isolates Fw12E-Y, Stu1B-Pr, and Stu5B-P-Pr showed a typical internal cytoplasmic membrane (ICM) structure of type I MOB, while the ICM structures of isolates S18C-Re and RS6A-Re were of type II MOB (Fig. 3). The highly purified MOB cultures, Fw5B-W, S1A-W (data not shown), and Rt4B-Y-O contained type I and type II MOB cells with the respective ICM structures (Fig. 3). These results showed good agreement with the phylogenetic placements of the MOB isolates, as revealed by 16S rRNA/pmoA gene sequence analysis.

FISH also confirmed the phylogenetic placements of the isolates. Isolate Fw12E-Y, RS11D-Pr, and Stu1B-Pr hybridized with probes My84 and My705, while not hybridizing with probe Mx450. Reversely, the type II MOB isolates, SS37A-Re and RS6A-Re, hybridized well with probe Mx450, and not with the probes for type I MOB. Figure 4 shows the fluorescence micrographs of Stu1B-Pr and RS6A-Re.

Identification of the MOB isolates

We tried to identify the isolates, focusing mainly on type I-related MOB isolates, in further work and in course of the studies we found very few non-MOB cells in some of the cultures, probably due to accidental contamination during preservation or sub-culturing of the isolates. Finally, strain Fw12E-Y was successfully purified and described as Methylomonas koyamae sp. nov. (44). Strain Fw12E-Y represents the second type I MOB isolated from rice paddy field and the first type I MOB strain obtained from a floodwater microsite in a rice paddy field ecosystem to our knowledge.

Cultivable MOB detected by FISH in MOB enrichment cultures

Type I and type II MOB were present in the enrichment cultures obtained from all the rice paddy field microsite samples (floodwater, surface soil, bulk soil, rhizosphere soil, root homogenate, and stem of rice plant) collected in 2008. The floodwater enrichment series appeared to contain abundant populations of type I MOB, although certain dilution series seemed to favor type II MOB (Fig. 5).

Detection of MOB groups in rice paddy field microsites by FISH

Type I and type II MOB were found in all the rice paddy field microsite samples (floodwater, surface soil, bulk soil, rhizosphere soil, root homogenate, and stem of rice plant) collected in 2008, irrespective of the pitfalls of FISH with soil-related samples (1). Type I MOB were found to be abundant in the floodwater, while type II MOB appeared to be more abundant in root and stem microsites of the paddy field in the sampling period (Fig. 6). The ratios of cells of type I MOB to those of type II MOB were about 22, 0.36, and 0.25 for floodwater, root homogenate, and stem of rice plant, respectively. Although accurate counting of bacterial populations revealed by FISH was difficult, especially from soil-related microsite samples (surface soil, bulk soil, rhizosphere soil), due to the autofluorescence of soil particles as reported in several studies (1, 6, 7, 48, 55), the results confirmed the relative abundance of both type I and type II MOB in all the microsites studied, as also indicated by Eller et al. (19).
Discussion

Although the presence of both types I and II MOB in rice paddy field compartments has been reported by many studies (18, 20, 21, 29, 31–33, 35, 38, 40, 50), only two type I strains (Methylogae oryzae [21, 23] and Methylomonas koyamae [strain Fw12E-Y; 44]) have been isolated from this environment so far. What could be the limitations of a culture-dependent technique in isolating type I MOB from rice paddy fields? Some studies underlined that methane pressure or shaking during culture incubation in liquid media could favor the growth of type I or type II MOB in the isolation process in addition to varying the culture media (4, 12, 25, 27, 31, 60). Five kinds of enrichment procedures under various culture conditions, such as the concentration of CH₄, temperature, and shaking or static were used in the present study. While type II MOB were isolated by enrichments A–D, type I MOB (Fw12E-Y, RS11D-Pr, Stu1B-Pr, Stu5B-P-Pr and R+4B-Y-O) were isolated/cultivated by enrichments B, D and E, and in addition we obtained an enrichment culture (SS19A-Pr) containing type I MOB based on the partial 16S rRNA gene sequence from surface soil by enrichment A (data not shown). These findings indicate that culture conditions in the isolation process may have slight influences on the growth preference of type I or type II MOB in the enrichment culture. One reason could also be the difficulty of separating type I MOB cells from non-MOB contaminants, as reported by Ferrando and Tarlera (21) and Geymonat et al. (23), and repeated transfer of colonies on agar slants or plates to liquid media and subsequent sub-culturing may lead to the loss of type I MOB cells. Type I MOB have been found to be preferentially grazed by protozoa in rice paddy fields (41, 42), which may also support the difficulty in isolating type I from this ecosystem. As underlined by Leadbetter (36), this enormous gap between the diversity of cultivated methanotrophs and the diversity of MOB in the rice paddy field ecosystem may partially be linked to our insufficient knowledge or imagination of the chemistry of their extracellular milieu. Therefore, refining and improving the isolation media and conditions may yield more type I MOB and...
consequently lead to deep insight into the true diversity of cultivable MOB in the rice paddy field ecosystem. To obtain the best picture of the methanotroph community present in the rice paddy field ecosystem, it is suitable to combine both molecular and cultivation techniques with regard to the limitation of both methods (17).

In this study, we combined the FISH method and the cultivation technique to characterize the cultivable methanotroph communities present in seven microsites of the paddy field ecosystem, and we compared our data with those obtained from the paddy field using pmoA gene analysis. Floodwater and stump compartments are very poorly studied niches in the rice paddy field ecosystem. The isolates/cultures consisted of 12 type II and five type I MOB. The five type I MOB were issued from floodwater (Fw12E-Y), rhizosphere soil (RS11D-Pr), root (Rt4B-Y-O), and stump (Stu1B-Pr, Stu5B-P-Pr), respectively, and strain Fw12E-Y was found to represent a new species in the genus Methylomonas (44), the first type I MOB isolated from floodwater in a rice paddy field. Type II MOB were isolated from almost all microsites except the root and were affiliated with pmoA sequences of the Methylocystis cluster or Methylomis swirum cluster; the findings were in accordance with previous studies using a cultivation technique (16, 24, 52, 53). Identification of these isolates/cultures is necessary in future work. The isolation of type I MOB in pure culture proved very difficult. As also reported by Ferrando and Tarlara (21), despite repeated streaking on different solid media, phase-contrast microscopic examination of cells from well-isolated colonies revealed the presence of contaminants with typical morphological features of Hyphomicrobium-like cells. In agreement with these authors, repeated transfer of colonies on highly purified agar slants to liquid medium, and subsequent subculturing may allow the isolation of type I methanotrophs from the highest dilution series.

The phylogenetic placement of MOB isolates was in agreement with studies conducted on floodwater, bulk soil, rice straw, and microcrustaceans in the rice paddy field based on pmoA analysis (32, 33, 43, Shibagaki-Shimizu et al. unpublished results). Almost all the type II isolates were associated with Methylocystis parvus or Methylocystis echinoides (Fig. 2); however, S1A-W was related to Methylomis swirum. RS11D-Pr (type I-like MOB isolate) was distantly related to clones from floodwater (FW), rice straw (RS), and microcrustaceans (MCR), together with a novel species Methanogaea oryzae from the soil-water interface in a flooded rice field in Uruguay (21, 23) (Fig. 2). We also obtained an enrichment culture (Fw10D-Pr) showing typical morphological features the same as RS11D-Pr (Fig. 1) from floodwater. Isolates/cultures of Fw12E-Y, Stu1B-Pr, Stu5B-P-Pr, and Rt4B-Y-O were associated with the Methylomonas cluster and did not show any close relationships to pmoA sequences of the clones. That might be related to the different paddy field conditions during both studies; however, the relative abundance found for Methylomonas-related MOB isolates in stomp and floodwater microsites seemed in agreement with the results of Ferrando and Tarlara (21). Using pmoA-base clones libraries analysis, they found for one group (group 3), 44.2% and 4.5% of clone species related to Methylomonas species at the soil-water-interface (SWI) and rhizosphere soil, respectively. The SWI microsite was related to a similar environment to the floodwater and stump microsites in our study from which we obtained the Methylomonas cultures.

Using the FISH technique, type I and type II MOB were found in all the microsites of the paddy field as well in the enrichment cultures. The presence of type I methanotrophs in these cultures may anticipate more type I MOB in pure cultures from the rice paddy field ecosystem. The relative abundance of type I and type II MOB found in different microsites of the rice field overall resembled the results in a rice microcosm system obtained by Eller and Frenzel (18) with the same group-specific probes; they reported that type II MOB predominated over type I MOB in bulk soil, rhizosphere and rhizoplane, but type I MOB also occurred in the compartments, especially at a higher relative proportion in the rhizoplane. In contrast, the proportion of type II MOB seemed to be higher in root homogenate in the present study. One principal reason may be the difficulty in counting cells, with another reason being the differences in paddy fields conditions.

For this study we focused on seven compartments of the rice paddy field ecosystem. Using combined cultivation and FISH techniques, the presence of both type I and type II methanotrophs was revealed. We successfully isolated strain Fw12E-Y from floodwater and obtained four type I MOB cultures from the stump, root and rhizosphere soil, respectively. Although their presence in these microsites has been shown in rice paddy fields using molecular approaches, the isolation of type I MOB has not been reported to our knowledge from these microsites of a rice paddy field. As Leadbetter (36) and Donachie et al. (17) underlined, in agreement with Hengstmann et al. (28) and Oremland et al. (45), by implementing comprehensive strategies that include culture-dependent techniques and molecular approaches, we can identify the full extent of microbial diversity in a given environment. From the present study, we elucidated the extent of the diversity of cultivable methanotrophs in a rice paddy field ecosystem including type I MOB, for which only one isolation has been recently reported so far (23).

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