Draft Genome Sequences of Gammaproteobacterial Methanotrophs Isolated from Marine Ecosystems


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This is contribution 12 from OMeGA.

The genome sequences of Methylobacter marinus A45, Methylobacter sp. strain BBA5.1, and Methylomarinum vadi IT-4 were obtained. These aerobic methanotrophs are typical members of coastal and hydrothermal vent marine ecosystems.

Microbial methane oxidation is one of the key drivers of oxygen consumption in marine sediments and the overlying water column (1). Methanotrophic bacteria are the primary producers of many cold and hot seep ecosystems (2, 3). Here, we report three genome sequences of gammaproteobacterial methanotrophs isolated from three marine ecosystems. Methylobacter marinus A45 (a methanol-adapted strain, formerly Methylomonas methanica A4, ACM 4717) was isolated from sewage outfall sediment near Los Angeles, CA (4). Methylobacter sp. strain BBA5.1 was isolated from the surface layer of estuary sediment collected at low tide near Newport, Bay Estuary (CA) (5). Methylomarinum vadi IT-4 (= JCM 13665T = DSM 18976T) was isolated from a

TABLE 1 General genome statistics and accession numbers

<table>
<thead>
<tr>
<th>Species</th>
<th>Sequencing platform(s)</th>
<th>Genome assembly and annotation</th>
<th>Genome coverage (X)</th>
<th>Genome size (Mb)</th>
<th>No. of scaffolds (no. of contigs)</th>
<th>Core (accessory) metabolic pathways</th>
<th>NCBI accession no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>M. marinus A45</td>
<td>Illumina</td>
<td>Velvet 1.1.05, AllPaths, Phrap</td>
<td>4.24, Prodigal 2.5</td>
<td>1,237</td>
<td>4.99</td>
<td>9 (49)</td>
<td>pMMO, pXmo, Mxa, XoxF1, XoxF2, H4F, H4MPT, FDH, RuMP, EMP, EDD, dPPP, PPP, pSC, TCA</td>
</tr>
<tr>
<td>Methylobacter sp. BBA5.1</td>
<td>Illumina, PacBio RS</td>
<td>AllPaths, Prodigal 2.5</td>
<td>290</td>
<td>5.07</td>
<td>87 (91)</td>
<td>pMMO, pXmo, Mxa, XoxF1, XoxF2, H4F, H4MPT, FDH, RuMP, EMP, EDD, dPPP, PPP, pSC, TCA</td>
<td>JQKS000000000</td>
</tr>
<tr>
<td>M. vadi IT-4</td>
<td>Illumina</td>
<td>Prodigal 2.5</td>
<td>272</td>
<td>4.33</td>
<td>1 (1)</td>
<td>pMMO, Mxa, XoxF, H4F, H4MPT, FDH, RuMP, EMP, EDD, dPPP, PPP, pSC, TCA</td>
<td>JPON000000000</td>
</tr>
</tbody>
</table>

*dPPP, dissimilatory pentose-phosphate pathway; EDD, Entner-Doudoroff pathway; EMP, Embden-Meyerhof-Parnas pathway; FDH, formate dehydrogenases; H4F, folate-linked Ct transfer; H4MPT, methanopterin-linked Ct transfer; Mxa, PQP-linked methanol dehydrogenases; pMOMO, membrane-bound methanol monooxygenase; pSC, partial serine cycle; pXmo, methane/ammonia monooxygenase related proteins of unknown function; PPP, pentose-phosphate pathway; RuMP, assimilatory ribulose monophosphate pathway; Xox, PQP-linked methanol and formaldehyde dehydrogenases (i.e., no evidence for the glyoxylate regeneration pathway was found); TCA, tricarboxylic acid cycle.

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microbial mat of a shallow submarine hydrothermal system near Taketomi Island, Okinawa, Japan (6).

DNA samples from the three strains were prepared using the standard phenol-chloroform method (7). DNA sequence data were obtained at the Joint Genome Institute using a combination of PacBio (8) and Illumina (9) technologies, and draft genome sequences were assembled. The computational tools used for genome sequencing and assembly are listed in Table 1.

All three sequenced marine methanotrophs are obligate methane and methanol utilizers. All three genomes harbor genes typical for type I methanotrophs, including genes encoding particulate methane monoxygenase (pmoCAB), the POR-dependent methanol dehydrogenases (mxaFI and multiple copies of xoxF), genes for tetrahydromethanopterin (H₄MPT) and tetrahydrofolate (H₄F)-dependent C₅-transfer pathways, genes of the ribulose monophosphate pathway, including its phosphoketolase variant (10), and genes encoding a complete tricarboxylic acid (TCA) cycle and a partial serine cycle (10) (Table 1). The genes encoding a complete tricarboxylic acid (TCA) cycle and a partial serine cycle (10) (Table 1).

REFERENCES


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