EVALUATION OF METHANE-OXIDIZING BACTERIA PHYSIOLOGICAL RESPONSES TO CHANGES IN OXYGEN CONCENTRATION, INTERACTIONS WITH NON-METHANE-OXIDIZING HETEROTROPHIC BACTERIA, AND THE TAXONOMIC CLASSIFICATION OF METHYLOCYSTIS SULFLITAE SP. NOV.

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EVALUATION OF METHANE-OXIDIZING BACTERIA PHYSIOLOGICAL RESPONSES TO CHANGES IN OXYGEN CONCENTRATION, INTERACTIONS WITH NON-METHANE-OXIDIZING HETEROTROPHIC BACTERIA, AND THE TAXONOMIC CLASSIFICATION OF METHYLOCYSTIS SUFLITAE SP. NOV.

A DISSERTATION APPROVED FOR THE SCHOOL OF BIOLOGICAL SCIENCES

BY THE COMMITTEE CONSISTING OF

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As a special note regarding the contents of this dissertation, the first chapter is a published multi-author work (Garner CT, Sankaranarayanan K, Abin CA, Garner RM, Cai H, Lawson PA, et al. Methylocystis suflitae sp. nov., a novel type II methanotrophic bacterium isolated from landfill cover soil. International Journal of Systematic and Evolutionary Microbiology. 2024;74(1):006239.). A detailed overview of the author's contributions is found at the end of the chapter.

Thank you to my parents and to my sister for their unwavering encouragement. Their support has been a source of strength and motivation.

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Table of Contents

LIST OF TABLES .................................................................................................................. VII

LIST OF FIGURES ............................................................................................................. VIII

ABSTRACT ........................................................................................................................... IX

CHAPTER 1 ............................................................................................................................. 1

Description of Methylocystis sulfite sp. nov., a novel type II methanotrophic bacterium
isolated from landfill cover soil ............................................................................................. 1
  Introduction ......................................................................................................................... 2
  Isolation and Ecology ......................................................................................................... 3
  Genome Features and Phylogeny ....................................................................................... 4
  Physiology and Chemotaxonomy ...................................................................................... 11
  Description of Methylocystis sulfite ............................................................................. 18
  Supplement to Chapter 1 .................................................................................................. 20
  References ......................................................................................................................... 23

CHAPTER 2 .......................................................................................................................... 26

Effect of Oxygen Concentration on the Methane-oxidizing Bacteria, Methylosinus
trichosporium OB3b and Methylomonas sp. WSC-7 .............................................................. 26
  Abstract .............................................................................................................................. 26
  Introduction ......................................................................................................................... 27
  Methods .............................................................................................................................. 29
  Results ................................................................................................................................. 33
  Discussion .......................................................................................................................... 44
  References ......................................................................................................................... 47

CHAPTER 3 .......................................................................................................................... 50

Effect of Coculturing the Methane-Oxidizing Bacterium Methylosinus trichosporium OB3b
with Flavobacterium, Cupriavidus, or Pseudomonas Bacteria ................................................ 50
  Abstract .............................................................................................................................. 50
  Introduction ......................................................................................................................... 51
  Methods .............................................................................................................................. 55
  Results ................................................................................................................................. 60
  Discussion .......................................................................................................................... 72
  Conclusion .......................................................................................................................... 76
  References ......................................................................................................................... 77
List of Tables

CHAPTER 1......................................................................................................................1
Table 1: Comparative ANI, AAI, and dDDH.....................................................................8
Table 2: Fatty acid profile ...............................................................................................16
Table 3: Strain characteristics .........................................................................................18
Supplemental Table 1: 16S sequence identity..................................................................20
Supplemental Table 2: Number of unique genes .............................................................21
Supplemental Table 3: NLS-7 unique gene annotation categories.................................22
Supplemental Table 4: DDH values ................................................................................22

CHAPTER 3........................................................................................................................50
Table 1: OB3b differentially expressed genes in co-culture ..............................................65
Table 2: Differentially expressed carbon metabolism genes..........................................67
Table 4: Differentially expressed ROS genes ..................................................................69
Table 4: Differentially expressed nitrogen and amino acid genes..................................71
List of Figures

CHAPTER 1 ................................................................................................................................. 1

Figure 1: Methylocystis UBCG Phylogenetic Tree ................................................................. 7
Figure 2: Methylocystis pmoA Phylogenetic Tree ................................................................. 8
Figure 3: Pangene of Methylocystis – Unique Genes ............................................................ 11

CHAPTER 2 .................................................................................................................................. 26

Figure 1: Methane oxidation rates of MOB in Changing Oxygen Conditions.................. 34
Figure 2: ROS gene counts – Methylosinus trichosporium OB3b ........................................ 36
Figure 3: ROS gene counts – Methylomonas sp. WSC-7 ...................................................... 38
Figure 4: Methylomonas sp. WSC-7 Biofilm Formation in Growth Media......................... 39
Figure 5: Gene counts of motility genes in Methylomonas sp. WSC-7 ................................ 40
Figure 6: WSC-7 Methane oxidation gene counts ............................................................... 41
Figure 7: OB3b Methane oxidation gene ........................................................................... 42
Figure 8: WSC-7 SEM Images ............................................................................................. 43

CHAPTER 3 .................................................................................................................................. 50

Figure 1: Visualization of the HPC Method ........................................................................... 58
Figure 2: Growth of NMOHB in Methane-oxidizing co-culture .......................................... 61
Figure 3: MOR of M. trichosporium OB3b in co-culture and monoculture conditions ....... 63
Abstract

This work describes an in-depth investigation into how the methane-oxidizing bacteria (MOB) respond to changes that allow them to effectively function in their ecological role. This includes understanding their response to unfavorable oxygen concentrations, and how they respond in coculture with other bacteria. This work also presents the characterization of a novel species of MOB; *Methylocystis suflitae* sp. nov. MOB play a crucial role in the global methane cycle, serving as significant actors in biogeochemical cycling. However, their physiological response to changing oxygen concentrations remains incompletely understood. One of the studies presented in this dissertation investigates how two MOB species, *Methylosinus trichosporium* OB3b and *Methylomonas* sp. WSC-7, respond to changing oxygen concentrations and the addition of catalase, a hydrogen peroxide scavenger, in growth media. Through transcriptomics analysis, this work showed that under high oxygen conditions, *M. trichosporium* OB3b upregulates genes involved in reactive oxygen species (ROS) defense, including cytochrome c peroxidase and superoxide dismutase, suggesting a need to deal with elevated ROS levels. Conversely, *Methylomonas* sp. WSC-7 exhibits cell clustering behavior, potentially as a defense mechanism against ROS toxicity. Differential expression of flagellar biosynthesis genes and chemotaxis response genes further supports this adaptive response. Moreover, *Methylomonas* sp. WSC-7 shows reduced expression of soluble methane monooxygenase genes under low oxygen conditions, while *M. trichosporium* OB3b exhibits higher expression under catalase-amended conditions, and that rates of methane oxidation for both strains are impacted by the concentration of oxygen and amending the growth media with catalase. Our findings underscore the importance of oxygen concentration in modulating MOB physiology and suggest potential strategies for optimizing their growth conditions. Additionally, despite the MOB’s crucial role in
biogeochemical cycling, there is still a lot to learn about how the MOB interact with other bacteria in their environment. Another aspect of this study investigates the growth conditions and community dynamics of methane-oxidizing co-cultures, focusing on the interactions between *M. trichosporium* OB3b (MOB) and the non-methane-oxidizing heterotrophic bacteria (NMOHB) of the genera *Flavobacterium*, *Cupriavidus*, and *Pseudomonas*. Heterotrophic plate counts were used to determine whether NMOHB could grow in methane-oxidizing co-cultures. There was growth of all three NMOHB, suggesting carbon substrate utilization derived from methane oxidation. Furthermore, co-cultures with *Pseudomonas chlororaphis* HC exhibited enhanced methane oxidation rates compared to monoculture, indicating a stimulating effect of *P. chlororaphis* HC on methane utilization by MOB. Transcriptomic analysis revealed differential gene expression patterns in both MOB and NMOHB during co-culture and points to potential cross-feeding of C4 dicarboxylates and methanol from the MOB to the NMOHB. Further, NMOHB displayed upregulation of genes involved in ROS remediation, suggesting a response to oxidative stress induced by methane oxidation. Additionally, NMOHB exhibited alterations in nitrogen utilization and amino acid metabolism genes, possibly reflecting adaptations to the co-culture environment. Overall, these findings shed light on the metabolic interactions between MOB and NMOHB in methane-oxidizing systems, highlighting the potential for carbon sharing, nitrogen exchange, and ROS defense mechanisms.
Chapter 1

Description of *Methylocystis sulfurae* sp. nov., a novel type II methanotrophic bacterium isolated from landfill cover soil.

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Abbreviations: TEM – Transmission Electron Microscopy; AAI – Average Amino Acid Identity; ANI – Average Nucleotide Identity; dDDH - digital DNA-DNA hybridization

Repositories: This strain is in the type culture collection of the ATCC with the type culture designation TSD-256 and has been deposited in the DSMZ German Collection of Microorganisms and Cell Cultures GmbH under the number DSM 112294.

Genome GenBank Accession Number: GCA_024448135.1

16S rRNA GenBank Accession Number: ON715489

A bacterial strain designated NLS-7ᵀ, was isolated through enrichment of landfill cover soil in methane-oxidizing conditions. Strain NLS-7ᵀ is a Gram-stain negative, non-motile, rod approximately 0.8 µm wide by 1.3 µm long. Phylogenetic analyses based on 16S rRNA gene sequencing places it within the genus *Methylocystis*, with its closest relatives being *M. hirsuta*, *M. silviterrae*, and *M. rosea*, with 99.9%, 99.7%, and 99.6% sequence similarity respectively.
However, ANI and AAI values below the 95% threshold compared to all the close relatives with
dDDH values between 20.9-54.1 demonstrate that strain NLS-7^T represents a novel species.

Genome sequencing generated 4.31 million reads and genome assembly resulted in the
generation of 244 contigs with a total assembly length of 3,820,957 bp (N50: 37,735 bp; L50:
34). Genome completeness is 99.5% with 3.98% contamination. It is capable of growth on
methane and methanol. It grows optimally at 30 °C between pH 6.5-7.0. Strain NLS-7^T is
capable of atmospheric dinitrogen fixation and can use ammonium (as NH₄Cl), L-aspartate, L-
arginine, yeast extract, nitrate, L-leucine, L-proline, L-methionine, L-lysine, and L-alanine as
nitrogen sources. The major fatty acids are C₁₈:₁ ω8c and C₁₈:₁ ω7c. Based upon this
polyphasic taxonomic study, strain NLS-7^T represents a novel species of the genus Methylocystis,
for which the name Methylocystis suflitae sp. nov. is proposed. The type strain is NLS-7^T (ATCC
= TSD-256, DSMZ = DSM 112294). The 16S rRNA gene and genome sequences of strain NLS-
7^T have been deposited in GenBank under accession numbers ON715489 and
GCA_024448135.1, respectively.

Introduction

Methanotrophs are organisms that can oxidize methane as their primary carbon and energy
source (1). Methane is a potent greenhouse gas (2), and these organisms play a significant role in
the global methane cycle (3). Successful isolation and characterization of methanotrophs is
useful in understanding their total effect on the global methane budget. Methanotrophs are highly
abundant in landfill cover soils due to the high levels of methane produced in anoxic zones
beneath the surface (4). Knief et. al (5) speculate that landfill cover soil hosts a diverse group of
uncultivated methanotrophic species and that methanotrophs of the genus Methylocystis show
habitat specificity for landfill cover soil.
Methylocystis is a genus of type II methanotrophs in the class Alphaproteobacteria within the family Methylocystaceae. To date, the genus encompasses eight species with validly published names. In 1993, Bowman et al. presented a revised methanotroph taxonomy to show the Methylocystis genus containing the species M. parva and M. echinoides (6). Since then, six more isolates have been characterized within the genus. These include: M. rosea (7), M. hirsuta (8), M. heyeri (9), M. bryophila (10), M. silviterrae (11), and M. iwaonis (12). The work done to characterize the M. iwaonis species presents the most recent emendation and description of the Methylocystis genus (12).

Isolation and Ecology

Methylocystis strain NLS-7\textsuperscript{T} was isolated from landfill cover soil of the United States Geological Survey Norman Landfill Environmental Research Site in Norman, Oklahoma, USA (35° 10' 12" N 97° 26' 49" W). The soil sample was obtained at a depth of 2-4 cm. Enrichments were made by incubating 5 g of soil in 120 mL serum bottles fitted with butyl rubber stoppers at 26 °C and amended with methane at 1% of the headspace volume. Methane oxidation was monitored by gas chromatography on a Shimadzu GC-14A gas chromatograph (Shimadzu, Japan). Following depletion of methane, enrichment cultures were reamended three times. Following this initial enrichment, a ten-fold dilution series was prepared to further enrich the most abundant methanotrophs present in the soil. The dilution series was performed in triplicate using 10 mL of modified NMS medium in 28 mL serum tubes amended with methane at 0.2% of the tube’s headspace. The modified nitrate mineral salts (NMS) medium contains (per liter of distilled water) 1 g KNO\textsubscript{3}, 0.27 g KH\textsubscript{2}PO\textsubscript{4}, 1 g MgSO\textsubscript{4} \cdot 7H\textsubscript{2}O, 0.14 g CaCl\textsubscript{2} \cdot 2H\textsubscript{2}O, and 0.23 g Na\textsubscript{2}HPO\textsubscript{4}, 0.5% (v/v) acidic trace element solution (AcTES), 0.02% alkaline trace element solution (AlTES), and 1% vitamin solution. AcTES contains (per liter), 2 g FeSO\textsubscript{4} \cdot 7H\textsubscript{2}O, 0.07
g ZnSO₄ · 7H₂O, 0.5 g MnCl₂ · 4H₂O, 0.12 g CoCl₂ · 6H₂O, 0.01 g NiCl₂ · 6H₂O, 0.5 g CuSO₄, 0.01 g H₃BO₃, 0.06 g LaNO₃, 0.06 g CeNO₃, and 100 mM HCl. The ALTES contains (per liter) 0.11 g Na₂SeO₄, 0.05 g Na₂WO₄, 0.29 g Na₂MoO₄, and 0.04 g NaOH. The vitamin solution contains (per liter) 5 mg 4-aminobenzoic acid, 2 mg D-biotin, 2 mg folic Acid, 10 mg pyridoxine-HCl, 5 mg riboflavin, 5 mg thiamine-HCl x 2H₂O, 5 mg nicotinic acid, 5 mg calcium D-pantothenate, 5 mg thiocetic acid (α-Lipoic Acid), and 0.10 mg vitamin B12.

Methane oxidation was again measured via gas chromatography. Dilutions that showed methane oxidation were reamended with methane following depletion and the process was repeated twice. These cultures were diluted and then plated onto solid modified NMS media containing 2% agar. Petri plates were incubated in airtight ammunition boxes (normally used for storing firearm ammunition) fitted with a rubber stopper through a drilled bore hole. Methane was added to the headspace of the ammunition box at 5% of the box’s interior volume. Colonies that formed on the plates were passaged on solid media up to five times until a pure colony of a single bacterial strain was present. The isolated strain was characterized and maintained in liquid modified NMS media in 160 mL serum bottles with methane added at 1-10% of the bottle’s headspace.

**Genome Features and Phylogeny**

Genomic DNA was extracted from a log-phase culture of strain NLS-7⁷ grown in the modified NMS medium at 28°C using the Qiagen DNeasy Blood and Tissue Kit according to the manufacturer’s instructions. Genomic DNA was fragmented using a QSonica 800R3 sonicator and built into Illumina-compatible shotgun libraries using the KAPA HyperPrep Kit following the manufacturer’s protocol. Shotgun libraries were sequenced on an Illumina MiSeq instrument. Sequence data was processed using fastp (v0.23.2) to remove reads containing uncalled bases.
‘N’), trim low-quality bases (q < 30), and merge overlapping reads (13). The resulting analysis-ready reads were used as input for de novo assembly (SKESA v2.5.1) (14). The quality of genomes was assessed by CheckM (v1.1.3) (15).

Reference genome sequences were obtained for representative bacterial species in the family Methylocystaceae (Methylocystis, Methylophila, Chenggangzhangella, and Hansschlegelia) from the NCBI RefSeq and GenBank databases (16,17). Gene prediction was performed using Prodigal (mRNA, v2.6.3), Barrnap (rRNA, v0.9) and tRNAscan-SE (tRNA, v2.0.9) (18). Annotation of protein-coding genes was performed using the KEGG database as reference (BLASTP, KoFamScan) (19–21). Predicted proteins were used to generate pairwise Average Amino Acid Identity scores (AAI, Enveomics toolbox) for genome-based taxonomic delineation (22). Average Nucleotide Identity (ANI) scores were calculated using the fastANI algorithm (23). Finally, sequences for 92 single-copy core bacterial genes were retrieved using the UBCG pipeline, aligned (MUSCLE), filtered to remove invariant sites, and concatenated (24,25). This concatenated sequence set was used to construct a maximum-likelihood amino acid tree (RAxML, 100 bootstrap replicates, GTR substitution model) to characterize the phylogenetic relationship between strain NLS-7T and other representatives of the family Methylocystaceae (26).

A total of 4.31 million reads (1.17 million merged, 0.98 million paired) were generated following quality filtering. Genome assembly resulted in the generation of 244 contigs with a total assembly length of 3,820,957 bp (N50: 37,735 bp; L50: 34). A total of 3,938 protein-coding, 46 tRNA, and three rRNA genes were identified. Genome completion was estimated at 99.5% based on the recovery of 1,095 single-copy genes (total 1,100) found in representative genomes of members of the Methylocystis genus. Genome contamination is 3.98% (15). Of the 3,938
predicted proteins, 2,957 were assigned function, 662 were hypothetical or had unknown function, and 319 had no significant matches.

Comparisons of the 16S rRNA gene sequence for strain NLS-7T with other validly published species of the genus *Methylocystis* were performed by aligning complete 16S rRNA sequences (1,497bp) using MUSCLE (25), followed by estimation of pairwise sequence similarities using NCBI’s blastn tool (20). Strain NLS-7T shares 99.9%, 99.7%, and 99.6% sequence similarity with *M. hirsuta*, *M. silviterrae*, and *M. rosea*, respectively (Supplemental Table 1). A partial 16S rRNA sequence of strain NLS-7T was also obtained by single gene PCR amplification and Sanger sequencing (Oklahoma Medical Research Foundation, Oklahoma City, USA). The primer set used for the single gene amplification was the 27F (5'-AGAGTTTGATCMTGGCTCAG-3') and 1492R (5'-TACGGYTACCTTGTTACGACTT-3') (27). This partial region matches the 16S rRNA gene from the whole genome sequence. Genome-wide estimates of similarity (ANI, AAI and dDDH, Table 1) also show that strain NLS-7T is most closely related to *M. silviterrae* (93.03% ANI, 93.86% AAI, 53.7% dDDH), *M. hirsuta* (92.63% ANI, 93.65% AAI, 54.1% dDDH), and *M. rosea* (91.52% ANI, 92.04% AAI 46.1% dDDH). These ANI and AAI values are below the 95% threshold used to delineate microbial species, and dDDH is less than 70% ranging from 20.9%-54.1% (Table 1). This indicates that strain NLS-7T is a novel species in the *Methylocystis* genus (28–30). The multi-gene UBCG phylogenetic tree (Figure 1) shows that strain NLS-7T forms a robust cluster with *M. silviterrae*, *M. hirsuta*, and *M. rosea* and is distinct from *M. parva*, *M. heyeri*, and *M. bryophila* (Figure 1). To further show the phylogenetic relationship between strain NLS-7T and other relatives of the family *Methylocystaceae*, phylogenetic trees were constructed based on nucleotide and conserved deduced amino acid sequences of genes (*pmoA*) encoding alpha subunit of particulate methane monooxygenase...
(Figure 2). The pmoA gene is commonly used for ecological studies of methanotrophic bacteria (5,31). This was done by extracting the pmoA gene nucleotide sequences from the NCBI reference genomes (16) by annotation with prokka (v 1.14.6) (32) and then the tree was constructed using FastTree (v2.1) (33).
Figure 2: Phylogenetic tree based on pmoA gene nucleotide sequence phylogeny for classified Methylocystis species and Methylosinus trichosporium OB3b. The phylogeny was computed by FastTree 2.1's Heuristic Neighbor-Joining method with a GTR substitution model and 1,000 resamples (32). Branch lengths numbers represent the proportion of resamples that fitted to that branch length.

Table 1: Comparative ANI, AAI, and dDDH values for characterized strains of species in the genus Methylocystis against strain NLS-7<sup>T</sup>. All ANI and AAI values are less than 95% and all dDDH values are less than 70% demonstrating species level delineation of strain NLS-7<sup>T</sup>.

<table>
<thead>
<tr>
<th>Organism Name</th>
<th>GenBank ID</th>
<th>ANI</th>
<th>AAI</th>
<th>dDDH</th>
</tr>
</thead>
<tbody>
<tr>
<td>M. bryophila</td>
<td>GCF_002117405.1</td>
<td>78.41</td>
<td>64.60</td>
<td>20.9</td>
</tr>
<tr>
<td>M. echinoides</td>
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<td>70.89</td>
<td>22.1</td>
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<td>22.0</td>
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<tr>
<td>M. iwaonis</td>
<td>GCF_027925385.1</td>
<td>79.80</td>
<td>71.98</td>
<td>22.0</td>
</tr>
<tr>
<td>M. rosea</td>
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<td>91.53</td>
<td>46.1</td>
</tr>
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<td>M. heyeri</td>
<td>GCF_004802635.2</td>
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<td>66.45</td>
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</tr>
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</tr>
<tr>
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<td>GCF_013350005.1</td>
<td>93.86</td>
<td>93.03</td>
<td>53.7</td>
</tr>
</tbody>
</table>

A search for common chemotaxonomic markers (membrane polar lipids, peptidoglycan diagnostic diamino acids) identified cardiolipin synthase (EC: 2.7.8.41, K08744), CDP-diaeclylglycerol---serine O-phosphatidyltransferase (EC:2.7.8.8, K17103), phosphatidyserine decarboxylase (EC:4.1.1.65, K01613), and the murE ligase (EC:6.3.2.13, K01928), associated with the biosynthesis of diphosphatidylglycerol (DPG, cardiolipin), phosphatidyserine (PS), phosphatidylenthanolamine (PE), and the diagnostic diamino acid meso-diaminopimelic acid (m-
DPM), respectively (34,35). These genes were observed in the genomes of all *Methylocystis* species (including isolate NLS-7T).

The genetic relationship between strain NLS-7T and related species of the genus *Methylocystis* was further studied by determining the pan-genome of *Methylocystis* using the pan-genome workflow of Anvi’o v7 (36). The program “anvi-gen-contigs-database” was first used to profile genomes of the genus *Methylocystis*, during which Prodigal v2.6.3 with default settings identified open reading frames (37). The program 'anvi-run-kegg-kofams' was used to annotate genes with functions by searching them against the latest KEGG database using blastp (20). The program 'anvi-pan-genome' with the flag ‘--use-ncbi-blast’, and parameters ‘--minbit 0.5’, and ‘--mcl-inflation 8’ was used to run the pan-genomic analysis. This program calculates similarities of each amino acid sequence in every genome against every other amino acid sequence in the group of genomes using blastp (20) and uses the MCL algorithm to identify gene clusters (38). The flag "anvi-display-pan" was used to visualize the distribution of gene clusters across genomes. In total, 508 gene clusters were identified that are unique to strain NLS-7T within the genus *Methylocystis*. This work further supplements our genomic analysis described above, demonstrating that with ortholog clustering and unique gene cluster determination, the proportion of genes unique to NLS-7T is similar to the proportion of unique genes found in other species of *Methylocystis*, especially *M. rosea* and *M. silviterrae* (Figure 3, Supplemental Table 2, Supplemental Table 3). *M. silviterrae* has 334 unique gene clusters, and *M. rosea* has 436 unique gene clusters. *M. echinoides* has the most gene clusters unique to its genome with 1629 clusters followed by *M. heyeri* with 1226 and *M. broyophila* with 1149. The species with high numbers of unique gene clusters in their genome also have AAI values that are low (<77%) compared to every other species in the genus *Methylocystis* (Supplemental Table 4). Strains that
don’t have as many unique gene clusters have relatively higher AAI values in pairwise comparison (Supplemental Table 4). A portion of the genes unique to strain NLS-7T were annotated to be in the coenzyme transport and metabolism, cell wall/membrane/envelope biogenesis, carbohydrate transport and metabolism, and amino acid transport and metabolism COG20 annotation categories (39) though most were unknown or unannotated (Supplemental Table 3).
Figure 3: The pangenome of the genus Methylocystis as determined by the Anvi’o workflow [36]. Overlapping black boxes represent portions of the genome that are shared by any two or more species, while segments of the genome that are unique to a certain species do not overlap with another.

Figure 3: Pangenome of Methylocystis – Unique Genes

Physiology and Chemotaxonomy

Cells for microscopy were grown to mid log-phase in the modified NMS medium in air with 10% CH₄ as the carbon and energy source. All the following growth experiments were at 28 °C unless otherwise indicated. Cell size estimates, motility, and morphological descriptions of strain NLS-7ᵀ were determined through light microscopy of wet mounts of cells grown in liquid culture using an Olympus BX41 microscope (Olympus Corporation, Japan) fitted with a Jenoptik Gryphax Subra camera and PROGRES GRYPHAX imaging software v 2.0.0.66 (Jenoptik Group, Germany). Cells for transmission electron microscopy were grown to mid log phase, harvested by centrifugation, and resuspended in 1% molten agarose. After cooling, the agarose was cut into blocks and pre-fixed with 4% (w/v) glutaraldehyde in 0.2 M cacodylate buffer (pH 7.4) for 1 h at room temperature and then overnight at 4 °C. Cells were then rinsed with 3 changes of cold 0.2 M cacodylate buffer for 10 minutes each and fixed in 2% OsO₄ for 2 hours on ice. Blocks were then rinsed in cold buffer and stored overnight at 4 °C. They were dehydrated in an ethanol series followed by a propylene oxide substitution and embedded in a Spurr epoxy resin. Resin blocks were thin sectioned using an Ultramicrotome (Reichert-Jung Ultracut E). Thin sections (50-70 nm) were collected on formvar-coated size 300 mesh copper
grids and stained with 2% uranyl acetate and 2% lead citrate for 10 minutes each. Thin sections were imaged at 80 kV on a JEOL 2010F TEM, equipped with a cold field emission electron source and a Direct Electron DE-12 (4k x 3k) digital imaging system. TEM imaging was performed at the Oklahoma Medical Research Foundation (Oklahoma City, USA).

Cell growth was measured for carbon and nitrogen source utilization tests, NaCl concentration tolerance, and temperature and pH range by analyzing the OD$_{600}$ of cultures with a Thermo Spectronic 20 D+ spectrophotometer (Thermo Scientific, USA) in the above medium.

Temperature range was determined by measuring growth rates at 4, 15, 20, 25, 30, 35, 40, and 45 °C. The pH range was examined from 5.0–9.0 in 0.5 pH increments and NaCl tolerance measured at concentrations of 0–2% (w/v) in 0.25% increments. To determine substrate utilization abilities of the strain, the following carbon sources were individually added to the modified NMS media at 0.1% w/v: methanol, ethanol, formate, glucose, fructose, lactose, sucrose, maltose, xylose, sorbitol, alanine, serine, glycine, acetate, citrate, oxalate, malate, pyruvate, succinate, urea, methylamine, dimethylamine, dimethylcarbonate, formamide, and methane (10% of tube headspace). Formaldehyde was also tested as a growth substrate, but at 0.01% w/v to compensate for potential toxicity issues and growth was measured using direct cell counts with 0.01% acridine orange stained cells and fluorescence microscopy (40). Cells were counted using the Olympus BX41 microscope described above fitted for fluorescence emission at 480 nm (Olympus Corporation, Japan). For all carbon source utilization experiments, growth was measured weekly for 6 weeks. Nitrogen source utilization was tested by adding at 0.1% w/v, the following nitrogen sources to a nitrate free modified NMS medium in tubes containing air and CH$_4$ in a 90:10 ratio in the headspace. Nitrogen sources included ammonium (as NH$_4$Cl), formamide, methylamine, glycine, hydroxylamine, urea, L-serine, L-aspartate, L-arginine, L-
leucine, L-proline, L-cysteine, L-methionine, L-histidine, L-lysine, L-alanine, L-threonine, glutamine, yeast extract, and nitrite. Each of these nitrogen sources were compared to a no added fixed nitrogen source control. Growth was measured every 1-2 weeks for 5-6 weeks. To determine if strain NLS-7T was able to fix atmospheric N₂, it was grown in a nitrogen free medium with N₂, O₂, and CH₄ gases present in the headspace at a 70:20:10 ratio and compared to a control with He substituted for N₂. This control therefore has a headspace gas composition of He, O₂, and CH₄ at the sample 70:20:10 ratio. Growth was measured weekly for 8 weeks.

Strain NLS-7T was capable of growth solely on methane and methanol. It was able to utilize ammonium (as NH₄Cl), L-aspartate, L-arginine, yeast extract, Nitrate, L-leucine, L-proline, L-methionine, L-lysine, and L-alanine as nitrogen sources. Like all other members of the genus Methylocystis, strain NLS-7T can fix atmospheric N₂. Strain NLS-7T can tolerate up to 0.75% (w/v) NaCl. It grew between 10-35 °C with an optimum at 30 °C and grew from pH 5.5-8.5 with an optimal pH of 6.5-7.0. When growing at 30 °C and pH 6.8 the strain has a specific growth rate of approximately 0.06 h⁻¹. Cells are Gram-stain negative rods and are approximately 0.8 ± 0.14 μm wide by 1.3 ± 0.3 μm long (Figure 4a) when observed by light microscopy. When examining thin sections of the cells with transmission electron microscopy, there appears to be accumulation of poly-β-hydroxybutyrate and the cells appear to have a dumbbell shape (Figure 4b and 4c).
Cells for fatty acid analysis were grown in modified NMS medium at 26 °C, harvested in late log phase from batch culture and the cell material was freeze dried. Phospholipid fatty acid analysis with mass spectrometry of strain NLS-7T was carried out by the Identification Service, Leibniz-Institut – Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH, Braunshweig, Germany.

Fatty acid analysis indicates that strain NLS-7T contains the C\(_{18:1}\) \(\omega8c\), C\(_{18:1}\) \(\omega7c\) as the predominant fatty acids. The C\(_{18:1}\) \(\omega8c\) fatty acid is present in all classified members of the *Methylocystis* genus, and the C\(_{18:1}\) \(\omega7c\) fatty acid is present in all species of *Methylocystis* except for *M. iwaonis* (12) as shown in Table 2. For strain NLS-7T, the C\(_{18:1}\) \(\omega8c\) and C\(_{18:1}\) \(\omega7c\) fatty acids are present at the highest abundance at over 97% of the entire fatty acid profile with all
other fatty acids present only at trace amounts. Although not grow all nine of the species were grown in parallel, very similar growth conditions were used to grow NLS-7T cells as were used for the other data. Specifically the data from the characterization of M. iwaonis which grew cells for fatty acid analysis in NMS medium at 30°C (12).

Based on the data obtained from this polyphasic taxonomic study that included phenotypic and phylogenetic information, strain NLS-7T merits recognition as representing a novel species of the genus Methylocystis for which the name Methylocystis suflitae sp. nov. is proposed. The novel species can be delineated from other members of the genus Methylocystis based on its ANI and AAI values, digital DNA-DNA hybridization, and proportion of gene clusters unique to the species. It can also be distinguished from other species of Methylocystis (except M. silviterrae) by its low diversity of fatty acids that comprise >1% of the fatty acid profile (Table 2).

Additionally, it can be distinguished from its nearest neighbors (M. silviterrae, M. hirsuta and M. rosea) by other means. Compared to M. silviterrae, strain NLS-7T can be observed to have a dumbbell shape (using transmission electron microscopy) and is shorter. The shortest M. silviterrae cells are reported to be longer than the average NLS-7T cell. Also, M. silviterrae has been shown to grow at 4 °C while NLS-7T cannot. Strain NLS-7T can be distinguished from M. hirsuta by NLS-7T’s inability to grow on acetate, and its lack of sMMO. Strain NLS-7T can be distinguished from M. rosea by M. rosea’s ability to form pink colored colonies. The strain’s cellular and physiological characteristics are detailed and summarized in Table 3.
Table 2: Fatty acid profile of strain NLS-7T compared to other members of the genus *Methylocystis*. Numbers represent the relative percentage of the acid. The comparative strains are: 1. *M. heyeri*; 2. *M. rosea*; 3. *M. echinoides*; 4. *M. parva*; 5. *M. hirsuta*; 6. *M. bryophila*; 7. *M. silviterrae*; 8. *M. iwaonis*. The data for strain NLS-7T were obtained in this study with the rest of the data obtained in the work describing the characterization of *M. iwaonis* (12). Trace fatty acids comprising <1% of the total profile are not presented and shown as a dash.

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<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
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16
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<tr>
<th>Cell shape</th>
<th>Rods, dumbbells</th>
<th>Small, curved coccoids, short rods</th>
<th>Straight, polymorphic or regularly curved rod, ovoids</th>
<th>Rods, coccobacilli, rods</th>
<th>Rods, coccobacilli, ovoids</th>
<th>Dumbbells</th>
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<td>0.5–0.8</td>
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<td>1.1–2.5</td>
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<td>Color of colonies</td>
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<td>Light cream</td>
<td>White</td>
<td>Pink</td>
<td>White, buff, pale pink</td>
<td>Diffusible brown, pale pink</td>
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<td>≤1</td>
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<td>sMMO</td>
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<td>+</td>
<td>-</td>
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Description of *Methylocystis suflitae*

*Methylocystis suflitae* (suf.li’tae. N.L. gen. n. *suflitae*) was named to honor Joseph M. Suflita, a contemporary American microbiologist, for his pioneering work in understanding biodegradation at the United States Geological Survey Norman Landfill Environmental Research Site. It is a member of the type II *Alphaproteobacterial* methanotrophic genus *Methylocystis*. It is a Gram-negative, non-motile, rod approximately 0.8 µm wide by 1.3 µm long. When growing on solid media in the presence of methane, it is yellow, circular, and convex with an entire margin and has a diameter of approximately 3 mm. It is capable of growth on methane and methanol. It grows optimally at 30 °C but can tolerate temperatures between 10-35 °C. It also grows optimally between pH 6.5-7.0 but can tolerate pH values between 5.5-8.5. Strain NLS-7ª tolerates NaCl concentrations up to 0.75% (w/v). Strain NLS-7ª is capable of atmospheric dinitrogen fixation and can use ammonium (as NH₄Cl), L-aspartate, L-arginine, yeast extract, nitrate, L-leucine, L-proline, L-methionine, L-lysine, and L-alanine as nitrogen sources. The major fatty acids are C18 : 1 ω8c, C18 : 1 ω7c. The type strain is NLS-7ª (= ATCC TSD-256 = DSM 112294). The 16S rRNA gene and genome sequences of strain NLS-7ª have been deposited in GenBank under accession numbers ON715489 and GCA_024448135.1, respectively.

**Funding information**
This work was supported by the National Science Foundation grant No. 1 736 255 under a subcontract from South Dakota School of Mines and Technology through the EPSCoR RII Track-2 FEC program.

Acknowledgments

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Author contributions

Conceptualization of ideas related to overarching research goals and aims was done by C.T.G. and L.R.K. Biochemical and physiological data were curated, maintained, and formally analysed by C.T.G. and genomic and phylogenetic data were curated, maintained and formally analysed by C.T.G. and H.C. Development and design of methodology was performed by C.T.G., L.R.K., C.A.A., K.S., H.C. and P.A.L. Research and investigation was conducted by C.T.G., R.M.G., C.A.A., K.S. and P.A.L. Management and coordination for the research activities was done by L.R.K. Oversight and leadership responsibility for research activity and execution including mentorship external to the core team was done by L.R.K. and C.A.A. Provision of study materials, reagents, materials, laboratory samples, instrumentation, computing resources and other analysis tools were provided by L.R.K., P.A.L., K.S. and the Oklahoma Medical Research
Foundation. Verification of replication and reproducibility of the experiments and results was done by L.R.K., C.A.A., K.S. and P.A.L. Preparation, creation, and presentation of the published work, specifically with visualization and data, was done by C.T.G., L.R.K., K.S., H.C. and P.A.L. Writing and preparation of the original draft was done by C.T.G., L.R.K., K.S., H.C. and R.M.G. Writing, including review, editing and critical commentary and revision was done by C.T.G., L.R.K., K.S. and P.A.L.

Supplement to Chapter 1

Supplemental Table 1: 16S sequence identity based on NCBI's blastn pairwise comparison of genome extracted 16S rRNA genes for each member of the genus *Methylocystis* and strain NLS-7\(^T\). Percentages above 98% are bolded.

<table>
<thead>
<tr>
<th></th>
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</thead>
<tbody>
<tr>
<td><em>M. bryophila</em> S285 [1]</td>
<td>100</td>
<td>97.71</td>
<td>97.51</td>
<td>97.31</td>
<td>97.44</td>
<td>97.78</td>
<td>97.24</td>
<td>97.64</td>
<td>97.37</td>
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<td><em>M. echinoides</em> LMG27198 [2]</td>
<td>97.71</td>
<td>100</td>
<td>97.24</td>
<td>97.91</td>
<td>98.31</td>
<td>99.19</td>
<td>97.57</td>
<td>98.25</td>
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<td><em>M. heyeri</em> H2 [3]</td>
<td>97.51</td>
<td>97.24</td>
<td>100</td>
<td>97.78</td>
<td>96.49</td>
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<td>97.71</td>
<td>97.57</td>
<td>97.84</td>
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<td><em>M. hirsuta</em> CSC1 [4]</td>
<td>97.31</td>
<td>97.91</td>
<td>97.78</td>
<td>100</td>
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<td><strong>98.31</strong></td>
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<td>96.83</td>
<td><strong>100</strong></td>
<td><strong>98.92</strong></td>
<td>96.49</td>
<td>97.17</td>
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<td>99.19</td>
<td>97.17</td>
<td>97.64</td>
<td><strong>98.92</strong></td>
<td><strong>100</strong></td>
<td>97.30</td>
<td>97.98</td>
<td>97.71</td>
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<td>97.24</td>
<td>97.57</td>
<td>97.71</td>
<td><strong>99.53</strong></td>
<td>96.49</td>
<td>97.30</td>
<td><strong>100</strong></td>
<td><strong>99.33</strong></td>
<td><strong>99.60</strong></td>
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<td><em>M. silviterrae</em> FS [8]</td>
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<td><strong>98.25</strong></td>
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<td><strong>99.66</strong></td>
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<td><em>M. sulfitae</em> NLS-7 [9]</td>
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<td><strong>99.60</strong></td>
<td><strong>99.73</strong></td>
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**Supplemental Table 2:** Total number of gene clusters unique to each species of the genus *Methylocystis*. Genomic analysis done on the NCBI reference genome of each species.

<table>
<thead>
<tr>
<th>Species (NCBI Reference Genome)</th>
<th>Number of Unique Clusters</th>
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<tbody>
<tr>
<td><em>Methylocystis broyophila</em></td>
<td>1149</td>
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<tr>
<td><em>Methylocystis echinoides</em></td>
<td>1629</td>
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<td><em>Methylocystis heyeri</em></td>
<td>1226</td>
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<tr>
<td><em>Methylocystis hirsuta</em></td>
<td>657</td>
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<tr>
<td><em>Methylocystis iwaonis</em></td>
<td>978</td>
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<tr>
<td><em>Methylocystis rosea</em></td>
<td>463</td>
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<tr>
<td><em>Methylocystis parva</em></td>
<td>646</td>
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<tr>
<td><em>Methylocystis silviterrae</em></td>
<td>334</td>
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<tr>
<td><em>Methylocystis suflitae NLS-7</em></td>
<td>508</td>
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</tbody>
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**Supplemental Table 3:** COG20 functional annotation of genes clusters among genes that are unique to *Methylocystis suflitae* strain NLS-7 within the genus *Methylocystis*.

<table>
<thead>
<tr>
<th>COG20 Annotation Category</th>
<th>Number of Gene Clusters</th>
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<tr>
<td>Amino acid transport and metabolism</td>
<td>5</td>
</tr>
<tr>
<td>Carbohydrate transport and metabolism</td>
<td>3</td>
</tr>
<tr>
<td>Cell Cycle Control</td>
<td>4</td>
</tr>
<tr>
<td>Cell wall/membrane/envelope biogenesis</td>
<td>14</td>
</tr>
<tr>
<td>Coenzyme transport and metabolism</td>
<td>10</td>
</tr>
<tr>
<td>Defense mechanisms</td>
<td>9</td>
</tr>
<tr>
<td>Energy production and conversion</td>
<td>5</td>
</tr>
<tr>
<td>Inorganic ion transport and metabolism</td>
<td>6</td>
</tr>
<tr>
<td>General function prediction</td>
<td>10</td>
</tr>
<tr>
<td>Lipid transport and metabolism</td>
<td>4</td>
</tr>
<tr>
<td>Mobilome: prophages, transposons</td>
<td>32</td>
</tr>
<tr>
<td>Posttranslational modification, protein turnover, chaperones</td>
<td>4</td>
</tr>
<tr>
<td>Replication, recombination and repair</td>
<td>24</td>
</tr>
</tbody>
</table>
**Supplemental Table 4**: DDH values calculated by Formula 2 of the DSMZ Genome-to-Genome Distance Calculator 3.0 (37,38). This demonstrates that the DDH value of strain NLS-7 is < 70% compared to each unique species of *Methylocystis* (Reference). Each reference genome shown and corresponding data are being compared to strain NLS-7.

<table>
<thead>
<tr>
<th>NCBI Reference genome</th>
<th>DDH</th>
<th>Model C.I.</th>
<th>Prob. DDH is &gt;= 70%</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Methylocystis echinoides</em></td>
<td>22.1</td>
<td>[19.9 - 24.6%]</td>
<td>0</td>
</tr>
<tr>
<td><em>Methylocystis iwaonis</em></td>
<td>22</td>
<td>[19.7 - 24.4%]</td>
<td>0</td>
</tr>
<tr>
<td><em>Methylocystis broyophila</em></td>
<td>20.9</td>
<td>[18.7 - 23.3%]</td>
<td>0</td>
</tr>
<tr>
<td><em>Methylocystis heyeri</em></td>
<td>20.9</td>
<td>[18.7 - 23.4%]</td>
<td>0</td>
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<tr>
<td><em>Methylocystis hirsuta</em></td>
<td>54.1</td>
<td>[51.4 - 56.8%]</td>
<td>32.02</td>
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<tr>
<td><em>Methylocystis parva</em></td>
<td>22</td>
<td>[19.7 - 24.4%]</td>
<td>0</td>
</tr>
<tr>
<td><em>Methylocystis rosea</em></td>
<td>46.1</td>
<td>[43.5 - 48.7%]</td>
<td>10.11</td>
</tr>
<tr>
<td><em>Methylocystis silviterrae</em></td>
<td>53.7</td>
<td>[51 - 56.4%]</td>
<td>30.48</td>
</tr>
</tbody>
</table>
References


Chapter 2

Effect of Oxygen Concentration on the Methane-oxidizing Bacteria, *Methylosinus trichosporium* OB3b and *Methylomonas* sp. WSC-7

Abstract

Methane-oxidizing bacteria (MOB) play a crucial role in the global methane cycle, serving as significant actors in biogeochemical cycling. However, their physiological response to changing oxygen concentrations remains incompletely understood. This study investigates how two MOB species, *Methylosinus trichosporium* OB3b and *Methylomonas* sp. WSC-7, respond to changing oxygen concentrations and the addition of catalase, a hydrogen peroxide scavenger, in growth media. Transcriptomics analysis demonstrates that under high oxygen conditions, *M. trichosporium* OB3b upregulates genes involved in reactive oxygen species (ROS) defense, including cytochrome c peroxidase and superoxide dismutase, suggesting a need to deal with elevated ROS levels. Conversely, *Methylomonas* sp. WSC-7 exhibits cell clustering behavior, potentially as a defense mechanism against ROS toxicity. Differential expression of flagellar biosynthesis genes and chemotaxis response genes further supports this adaptive response. Moreover, *Methylomonas* sp. WSC-7 shows reduced expression of soluble methane monooxygenase genes under low oxygen conditions, while *M. trichosporium* OB3b exhibits higher expression under catalase-amended conditions, and that rates of methane oxidation for both strains are impacted by the concentration of oxygen and amending the growth media with catalase. Our findings underscore the importance of oxygen concentration in modulating MOB physiology and suggest potential strategies for optimizing their growth conditions.
Introduction

Methane-oxidizing bacteria (MOB) are organisms that can oxidize methane as their primary carbon and energy source. These bacteria oxidize methane to CO₂ using a pathway involving conversion of methane to methanol, and further to formaldehyde and formate where the carbon is assimilated by the ribulose monophosphate pathway or the serine pathways (1). Methane is a potent greenhouse gas (2), and these organisms play a significant role in the global methane cycle (3). Studying this group of organisms is critical for understanding their total impact on biogeochemical cycling. Also, isolating and characterizing novel species of methane oxidizers will be useful in biotechnological applications such as bioremediation and generation of value-added products (4). All of these areas of study require detailed understanding of MOB physiology and conditions for optimized growth.

During aerobic metabolism toxic reactive oxygen species (ROS) are generated including superoxide and hydrogen peroxide (5–9). Because endogenous ROS generation in bacteria is a result of cellular respiration, increases in endogenous production in ROS generated during bacterial respiration is positively correlated with increases in cellular metabolic rate as well as elevated O₂ concentrations (5–7,10–12). This endogenous production of ROS can have harmful effects on cells including damaging proteins, nucleic acids, and lipids (10). In addition to the potential generation of toxic ROS by cellular respiration, aerobic MOB oxidize methane using a class of enzymes called methane monooxygenases. These enzymes catalyze the oxidation of methane to methanol in the presence of dioxygen, and this reaction may contribute to ROS generation (13). Some studies have demonstrated that the enzyme activity of methane
monooxygenases may catalyze the production of hydrogen peroxide, and that monooxygenases are inhibited by the presence of hydrogen peroxide leading to decreased enzyme activity (8,14).

Many species of bacteria, including MOB, have evolved systems to counteract the negative effects of ROS. The primary defense systems employed use ROS scavenging enzymes such as catalase, cytochrome c peroxidases, and superoxide dismutase to decrease ROS concentrations (10,15). The general bacterial response to increases in H₂O₂ accumulation includes the expression of a wide set of peroxidase class enzymes that reduce H₂O₂ to H₂O governed by the OxyR regulon (16). H₂O₂ can also be reduced by cytochrome c peroxidase, and it has even been shown in MOB that cytochrome c peroxidase may serve to remediate peroxide buildup from cellular respiration (15). In bacteria, superoxide is scavenged primarily by the enzyme superoxide dismutase that catalyzes the dismutation of superoxide into H₂O₂ and O₂, governed by the SoxRS regulatory system (16).

In addition to using ROS scavenging enzymes to mitigate ROS buildup, some species of bacteria that are known to be sensitive to elevated oxygen levels even employ the use of cell clustering to defend against oxygen stress (17). Some microaerophilic bacteria have chemotaxis sensors that regulate the motility of cells in the presence of unfavorable oxygen concentrations leading to cell clumping which may protect the cells from elevated oxygen by reducing the surface-to-volume ratio of the cells by limiting oxygen diffusion (17-19). However, despite our understanding of ROS defense systems such as these, the precise way that MOB deal with the consequences of ROS, or even how they respond to changes in oxygen concentration is not well understood.

An experiment using transcriptomics analysis was designed to show how two species of MOB, *Methylosinus trichosporium* OB3b and *Methylomonas* sp. WSC-7, respond to changing oxygen concentrations and how the use of growth media amended with catalase as a hydrogen peroxide
scavenger influences the growth and physiology of the two strains. This experiment demonstrated that amending growth media with catalase leads to an increase in methane oxidation rate for both *M. trichosporium* OB3b and *Methylomonas* sp. WSC-7. Also, *M. trichosporium* OB3b responds to increases in oxygen concentration by overexpressing cytochrome c peroxidase and superoxide dismutase encoding genes, but *Methylomonas* sp. WSC-7 responds to increases in oxygen concentration by cell clustering and decreasing expression of flagella biosynthesis genes and aerotaxis response genes.

**Methods**

**Growth Conditions of *M. trichosporium* OB3b and *Methylomonas* sp. WSC-7**

The objective of this study was to determine if changes in headspace oxygen concentration and/or the addition of catalase to the growth media influenced the physiology, response to ROS toxicity, and methane oxidation rate of two strains of methane-oxidizing bacteria, *M. trichosporium* OB3b and *Methylomonas* sp. WSC-7 grown under methane-oxidizing conditions.

*M. trichosporium* OB3b cells were grown statically in modified nitrate mineral salts medium (NMS) similar to the medium developed by Whitenbury et al. (20) for the growth of methane-oxidizing bacteria in the group’s initial characterization studies, but with the addition of a vitamin solution. The modified NMS medium contains (per liter of distilled water) 1 g KNO₃, 0.27 g KH₂PO₄, 1 g MgSO₄ · 7H₂O, 0.14 g CaCl₂ · 2H₂O, and 0.23 g Na₂HPO₄, 0.5% (v/v) acidic trace element solution (AcTES), 0.02% alkaline trace element solution (AlTES), and 1% vitamin solution. AcTES contains (per liter), 2 g FeSO₄ · 7H₂O, 0.07 g ZnSO₄ · 7H₂O, 0.5 g MnCl₂ · 4H₂O, 0.12 g CoCl₂ · 6H₂O, 0.01 g NiCl₂ · 6H₂O, 0.5 g CuSO₄, 0.01 g H₃BO₃, 0.06 g
LaNO₃, 0.06 g CeNO₃, and 100 mM HCl. The AlTES contains (per liter) 0.11 g Na₂SeO₄, 0.05 g Na₂WO₄, 0.29 g Na₂MoO₄, and 0.04 g NaOH. The vitamin solution contains (per liter) 5 mg 4-amino benzoic acid, 2 mg D-biotin, 2 mg folic acid, 10 mg pyridoxine-HCl, 5 mg riboflavin, 5 mg thiamine-HCl x 2H₂O, 5 mg nicotinic acid, 5 mg calcium D-pantothenate, 5 mg thiocetic acid (α-Lipoic Acid), and 0.10 mg vitamin B12.

*Methylomonas* sp. WSC-7 cells were grown statically in HEPES/phosphate buffered Saddle Mountain Creek medium (HPB-SMC Medium). This contains (per liter of distilled water) 2.38 g HEPES (Free Acid), 0.26 g NaCl, 0.14 g Na₂HPO₄, 0.11 g KCl, 0.12 MgSO₄ · 7H₂O, 0.04 g NaNO₃, 0.04 g CaCl₂ · 2H₂O, 0.004 g H₃BO₃, and 0.007 g SrCl₂ · 6H₂O as well as 0.5% (v/v) AcTES, 0.02% alkaline trace element solution AlTES, and 1% vitamin solution.

Three treatments were evaluated in this study. The strains were grown statically in 500 mL of respective media in 1 L stoppered bottles, leaving 500 mL of available gas headspace. One condition (high oxygen) was grown with a headspace gas composition of 69% N₂, 10% CH₄, and 21% O₂. The second condition (low oxygen) was grown with a headspace gas composition of 85% N₂, 10% CH₄, and 5% O₂. The third condition (catalase) was grown at high oxygen, but the medium was amended with 20 µg/mL of catalase (From Bovine Liver, Thermo Scientific). Each condition was incubated at 28 °C without shaking and methane depletion was measured by gas chromatography on a Shimadzu GC-14A gas chromatograph (Shimadzu, Japan). Methane oxidation rates (MOR) were calculated as \( \text{MOR} = \ln(\Delta[\text{CH}_4]) / \ln(\Delta t) \). This is the slope of the natural log of the methane depletion curve where \([\text{CH}_4]\) is the molar concentration of CH₄ and \( t \) is time elapsed in hours.
RNA Extraction, Illumina Sequencing Library Preparation, and Sequencing

Cells for RNA extraction were grown statically in one-liter bottles with 500 ml of medium and harvested in late log phase by centrifugation for 20 minutes at 4 °C and 4000 x g using an Eppendorf 5910 centrifuge with cell pellets stored at -80 °C. The Zymo Research Quick-RNA Fungal/Bacterial Miniprep kit was used for RNA extraction. Additional DNase treatments were carried out using the Invitrogen DNase I Amplification Grade treatment kit. Nucleic acid extracts were quantified using a Qubit Fluorometer 2.0 with RNA high specificity and DNA broad range specificity reagent kits.

Illumina compatible sequencing libraries were prepared for sequencing with the Zymo-Seq RiboFree Total RNA Library Kit (Zymo Research, USA) according to the manufacturer’s specifications. Prior to sequencing, the cDNA library was concentrated using a Thermo Scientific Savant DNA120 SpeedVac Concentrator, and size selection was carried out using the Agilent 4150 TapeStation System.

Sequencing of the final cDNA library was performed at the Oklahoma Medical Research Foundation Center for Clinical Genomics (Oklahoma City, USA) using Illumina NovaSeq 6000 S4 PE150 chemistry.

Reference Genome Sequencing

Genomic DNA of Methylomonas. sp. WSC-7 from a log-phase grown culture grown in the modified NMS medium at 28 °C was extracted using the Qiagen DNeasy Blood and Tissue Kit. DNA was fragmented using a QSonica 800R3 sonicator and built into Illumina-compatible shotgun libraries using the KAPA HyperPrep Kit. Shotgun libraries were sequenced on an Illumina MiSeq instrument. Sequence data was processed using fastp v0.23.2 (21) to remove
reads containing uncalled bases (‘N’), trim low-quality bases (q < 30), and merge overlapping reads. The resulting analysis-ready reads were used as input for de novo assembly with SKESA v2.5.1 (22). The quality of genomes was assessed by CheckM v1.1.3 (23). The genome for *M. trichosporium* OB3b (GCF_000178815.2) was downloaded from the NCBI RefSeq database (24). Gene prediction was performed using Prodigal v2.6.3 (25) and barrnap v0.9 (26).

**Read Mapping and Quantification**

Raw fastq reads were filtered for high quality reads (q < 30) and the adaptors were trimmed using fastp v0.22.0 (21). Quality filtered pair-end reads were mapped to respective reference genomes using Bowtie 2 v2.2.5 (27). The .bam files generated were parsed for DNA contamination issues by removing reads known to directionally be linked to genomic DNA, then removing an equal number of reads from the RNA pool to account for bias. The precomputed alignments generated by Bowtie 2 were used as input for transcript quantification using Salmon 0.14.2 (28).

**Differential Expression Analysis**

The R program DESeq2 was used for differential expression analysis of quantified reads using the likelihood ratio test method (29). Genes that were differentially expressed (adjusted p-value < 0.05, log2FC < -2 or > 2) were annotated using KEGG BlastKOALA (30).

**Scanning Electron Microscopy of Biofilms**

*Methylomonas* sp. WSC-7 cells for scanning electron microscopy were grown statically to late-log phase in HPB-SMC media with 10% CH$_4$ added to the headspace of air. Cells were filtered onto a 0.2 mm membrane (Whatman) by vacuum filtration and the filtered cells are transferred onto a poly-l-lysine coated slide.
The cells were then fixed in HPB-SMC medium with 4% glutaraldehyde for 17 hours at room temperature and then washed for 10 minutes, three times in HPB-SMC medium to remove excess fixative. After fixation, cells were immersed in a 1% OsO₄ solution in modified NMS medium at 4 °C for 1 hour and washed again with deionized water to remove the excess solution. Cells were then dehydrated by immersion for 10 minutes each in 25%, 50%, 70%, 85%, 95%, and three times in 100% ethanol (in deionized water). Critical point drying occurred using a Tousimis Autosamdri-814 and samples were sputter coated with a thin layer (~ 4 nm) of iridium (Ir) using an EMS Quorum Q150 ES Plus sputter coater. Imaging of the biofilms was performed at the University of Oklahoma Samuel Roberts Noble Microscopy Laboratory using a Zeiss Neon 40 EsB field emission scanning electron microscope at 5 kV.

**Results**

**Effect of Catalase and Oxygen Concentration on *M. trichosporium OB3b* and *Methylomonas* sp. WSC-7 Methane Oxidation Rate**

The methane oxidation rate (MOR) was measured for both *M. trichosporium OB3b* and *Methylomonas* sp. WSC-7 across the three treatments. Both *M. trichosporium OB3b* and *Methylomonas* sp. WSC-7 had a significantly higher MOR under the catalase amended conditions compared to both the high oxygen condition and the low oxygen condition. Also, MOR was higher in the high oxygen condition with no catalase compared to the low oxygen condition for both strains. (Fig. 1).
RNA-seq of *M. trichosporium* OB3b and *Methylomonas* sp. WSC-7

To explain these trends in the MORs and the overall physiological response to oxygen concentration and catalase on the two strains, gene expression profiles were analyzed by DESeq2. The likelihood ratio test (LRT) of our differential gene expression analysis demonstrated that 136 genes were significantly differentially expressed (log2 Fold Change > 2.0 or < -2.0 & adjusted p-value < 0.05) by *M. trichosporium* OB3b across all conditions and that 998 genes were differentially expressed by *Methylomonas* sp. WSC-7. The differentially expressed genes were annotated by KEGG’s BlastKOALA. A total of 55 differentially expressed genes from *M. trichosporium* OB3b and 599 genes from *Methylomonas* sp. WSC-7 were annotated.
M. trichosporium OB3b & Methylomonas sp. WSC-7 ROS Gene Expression

There were notable differences in the ROS gene expression profiles across different conditions. M. trichosporium OB3b overexpressed a cytochrome c peroxidase gene when grown at high oxygen. Expression of this gene decreased when grown with catalase and decreased even more when grown at low oxygen (Fig. 2). The same trend can be observed for the katG gene encoding for catalase-peroxidase in OB3b. However, the change in expression is not statistically significant (adjusted p-value = 0.14). Additionally, in OB3b a gene encoding for superoxide dismutase was underexpressed in the catalase condition compared to both the high oxygen and low oxygen conditions. This same gene was expressed the greatest in the low oxygen condition (Fig. 2). The decreased expression of these genes when catalase is added to the growth medium further implies that these genes are involved in ROS remediation and that the addition of catalase may aid in ROS remediation during growth on methane, reducing the need for OB3b to produce ROS defense enzymes.
Methylophans. sp. WSC-7 also expresses a cytochrome c peroxidase that could be reducing H$_2$O$_2$. However, unlike in *M. trichosporium* OB3b, the highest expression level is in the low oxygen condition with less in the catalase condition, and in the high oxygen condition it is expressed at very low levels if at all (Fig. 3). There is also a superoxide dismutase gene being differentially expressed in sp. WSC-7. This gene is expressed at low oxygen, far less at high oxygen condition, and not at all in the catalase condition (Fig. 3). This trend leads us to hypothesize that under elevated oxygen conditions, *Methylophans* sp. WSC-7 uses a method of ROS defense that is distinct from OB3b. Based on observations of cell clustering under growth
conditions of high oxygen and that this cell cluster is lacking when oxygen is lower (Fig. 4), strain WSC-7 may cluster together to reduce the diffusion of oxygen to the cells as a primary means of dealing with increased aeration. In addition to the observed cell clustering, *Methylomonas* sp. WSC-7 expressed several genes associated with flagellar biosynthesis proteins and flagellar component proteins (*flgBCDEFGHIK*) to a much greater extent in the low oxygen conditions than in high oxygen or catalase amended conditions (Fig. 5). Additionally, the *cheBDRV* chemotaxis protein genes are expressed at higher levels in the low oxygen condition as well (Fig. 5). The increase in the expression of flagellar protein synthesis genes and chemotaxis response genes suggests that there is increased motility in the low oxygen condition, perhaps as an aerotaxis response to facilitate access to oxygen when the concentration is lower.
Figure 3: Normalized gene counts of Cytochrome C Peroxidase and Superoxide Dismutase genes in Methylomonas sp. WSC-7 across high oxygen, low oxygen, and catalase containing conditions. Each condition was incubated with a headspace methane concentration of 10%. Data are shown as the mean +/- standard deviation across triplicate samples for the High Oxygen and Low Oxygen conditions and duplicate samples for the Catalase condition.
Figure 4: Representative samples of Methylomonas sp. WSC-7 growing in high oxygen, low oxygen, and catalase conditions depicting that in conditions where the headspace oxygen is 21% the cells form clumps and when the headspace oxygen is at 5% the culture is turbid and there is no visible clumping.
Figure 5: Gene counts of motility genes in Methylomonas sp. WSC-7, M. trichosporium OB3b & Methylomonas sp. WSC-7 Methane Oxidation Pathway Gene Expression

The WSC-7 pmoABC genes coding for the particulate methane monooxygenase are expressed at much greater levels than the mmoXYC genes in all conditions (Fig. 6). For the pmoABC genes there is not a statistically significant change in expression under any condition. However, the expression of each of the mmoXYC genes is significantly lower in the low oxygen condition. In WSC-7, the methanol dehydrogenase genes xoxF and mxaF are expressed in all conditions with xoxF showing increased expression under low oxygen. In OB3b, there are no statistically
significant changes in MMO protein gene expression levels (Fig. 7). Further, in OB3b, the $mxaF$ and $xoxF$ methanol dehydrogenases are expressed at similar levels in each condition (Fig. 7).

Figure 6: Normalized counts of methane and methanol oxidation genes by *Methylomonas* sp. WSC-7 in catalase (blue), high oxygen (orange), and low oxygen (green), conditions. Data are shown as the +/- standard deviation across triplicate samples for the high and low conditions and duplicate samples for the catalase condition.
Figure 7: Normalized counts of methane and methanol oxidation genes by *M. trichosporium* OB3b in catalase (blue), high oxygen (orange), and low oxygen (green) conditions. Data are shown as the +/- standard deviation across triplicate samples for the high and low conditions and duplicate samples for the catalase condition.
Methylomonas sp. WSC-7 Growth Phenotypes in Changing Oxygen Concentrations and Scanning Electron Microscopy of WSC-7 Biofilms

The cells of Methylomonas sp. WSC-7 were observed to form clumps which floated in the high oxygen conditions, but not at low oxygen (Fig. 4). However, biofilms were observed using electron microscopy under all conditions. Although the cells did not appear to form a biofilm in the bottles when growing in the low oxygen conditions, they appear to have the same cell stacking structure as the cells growing in the high oxygen and catalase conditions when observing the scanning electron micrographs (Fig. 8). This suggests the capacity form clumps was present under all conditions, but only visible at high oxygen.

Figure 8: Representative scanning electron micrographs of Methylomonas sp. WSC-7 cells grown in catalase (A) high oxygen (B) and low oxygen (C) conditions. Images were captured at 5000X zoom and electron high tension of 5 kV.
Discussion

Lower Oxygen Concentrations Leads to Lower Expression of Soluble Methane Monooxygenase in *Methylomonas* sp. WSC-7

It has been determined that genes encoding the soluble methane monooxygenase in the type I MOB WSC-7 are more impacted by decreases in oxygen concentration than those of the genes for particulate methane monooxygenase. WSC-7 in low oxygen had decreased expression of *mmoXYC* genes. Interestingly, the same trend was not present in type II MOB OB3b. In OB3b, there was a slightly higher expression of *mmoX* under the catalase condition, but this change was not statistically significant and gene expression in high and low oxygen conditions was similar. Other studies have demonstrated that in wild-type OB3b and OB3b mutants that are unable to express genes for particulate methane monooxygenase, increases in oxygen concentration from 10% headspace O₂ to 80% headspace O₂ decreased sMMO protein levels and enzyme activity (31). This study did not see a change in expression of the sMMO gene *mmoX* for Ob3b, but ranges of O₂ examined in this study were from 5% to 21% which may account for this discrepancy. Perhaps this observation suggests that oxygen levels impact type I and type II MOB sMMO genes differently.

*Methylosinus trichosporium* OB3b May Remediate Negative Impacts on MOR by Expression of Cytochrome c Peroxidase and Superoxide Dismutase

Changing the headspace oxygen concentration and/or amending the growth medium with 20 µg/mL of catalase impacted the MOR and gene expression profile of methane-oxidizing bacteria when they are growing in pure culture. *M. trichosporium* OB3b had genes related to the remediation of radical oxygen toxicity including cytochrome c peroxidase and superoxide
dismutase (SOD) overexpressed in the high oxygen condition compared to when catalase was used to amend the media, or when there was a lower concentration of headspace O₂.

This observation that when O₂ concentration is higher, the expression of ROS defense genes in OB3b increases is plausible. It has been demonstrated that when aerobically growing cells are exposed to elevated oxygen concentrations, there is a proportional increase in the accumulation of ROS such as superoxide and hydrogen peroxide inside cells (10, 32-34). Specifically, the observation that cytochrome c peroxidase is one of the enzymes that is used by OB3b to mitigate hydrogen peroxide buildup is also supported by previous experiments. It has been demonstrated that cytochrome c peroxidase from *Methylococcus capsulatus* Bath is capable of peroxide reduction. Researchers in that study speculate that because of the peroxide formation induced by monooxygenase O₂ activation, peroxide toxicity remediation could be a reasonable role for the protein (15).

In *M. trichosporium* OB3b, the superoxide dismutase encoding gene is also differentially expressed in the same manner, where at a higher headspace oxygen concentration, the gene is expressed more. This could happen for potentially the same reason, as a means of scavenging toxic superoxide radicals which have been shown to be generated by bacterial aerobic metabolism and cause cellular damage (7,9,35).

*Methylomonas* sp. WSC-7 May Utilize Cell Clumping to Defend Against ROS Toxicity

These data suggest that WSC-7 cells are clustering together as a way to defend against ROS toxicity when exposed to unfavorably high concentrations of O₂. At higher O₂ concentrations the *Methylomonas* sp. WSC-7 cells tend to clump together to form flocs, but when growing in the low O₂ condition, the cells disperse throughout the growth medium (Fig. 4). This correlates with
increased expression of flagellar biosynthesis protein genes in the low oxygen condition. Since the flagellar protein genes are downregulated in the high oxygen condition, this could lead to less motility, and reduction in cell dispersal. Moreover, biofilm formation consists of a transition of motility to non-motility by bacterial cells to facilitate cell-to-cell adhesion (36) so it is plausible that when clustered, expression of flagellar proteins is shut down. The WSC-7 cell aggregation could serve as a defense against elevated oxygen exposure because there is a reduction in the surface-to-volume ratio of the cells by limiting oxygen diffusion (19). Further, it was observed that *Methylomonas* sp. WSC-7 was shown to have higher expression of the *cheB*, *cheD*, *cheR*, and *cheV* genes under conditions of low oxygen. It has been shown that *A. brasilense* mutants that have been unable to express *cheB* or *cheR* genes are shown have an impaired aerotaxis response even though the cells remain motile (37). This may suggest that the differentially expressed *che* genes are important to properly navigating to oxygen. Also, *A. brasilense* cells have been shown to cluster together under elevated oxygen concentrations and *A. brasilense* mutants lacking their ability to express chemotaxis genes clump together more frequently (17,18). This makes sense, as *A. brasilense* has been shown to avoid unfavorable oxygen concentrations that are either too high or too low via aerotaxis (38). Strain WSC-7 likely does the same, where under conditions that are oxygen-rich, the cells clump together, but under lower oxygen conditions, the cells are motile and migrate towards oxygen concentrations that optimize oxygen exposure to the cells. However, this may come with having to compensate for ROS toxicity by expressing peroxidase and SOD genes as observed here.

Some key conclusions arise from these findings. The type II methanotroph *M. trichosporium* OB3b transcribes genes needed to mitigate the toxic effects of ROS more when growing at elevated O$_2$ concentrations, but the type I methanotroph *Methylomonas* sp. WSC-7 appears to
mitigate the toxic effects of ROS differently than OB3b by facilitating cell clumping as a potential defense mechanism from ROS toxicity. While this study is limited to just two different strains of MOB, the fact that there such a drastic distinction in their behavior demonstrates that when trying to optimize growth conditions for the isolation, cultivation and biotechnical implication of MOB, the consideration of oxygen concentration is critical and has an impact on cellular metabolism and physiology.

References


Chapter 3

Effect of Coculturing the Methane-Oxidizing Bacterium *Methylosinus trichosporium* OB3b with *Flavobacterium, Cupriavidus, or Pseudomonas* Bacteria

Abstract

Methane-oxidizing bacteria (MOB) are critical members of methane-containing ecosystems and are highly involved in biogeochemical cycling. Despite their crucial role, there is still a lot to learn about how the MOB interact with other bacteria in their environment. This research investigates the growth conditions and community dynamics of methane-oxidizing co-cultures, focusing on the interactions between the MOB *Methylosinus trichosporium* OB3b and the non-methane-oxidizing heterotrophic bacteria (NMOHB) of the genera *Flavobacterium, Cupriavidus,* and *Pseudomonas.* Heterotrophic plate counts were used to determine whether NMOHB could grow in methane-oxidizing co-cultures. All three NMOHB grew in methane-oxidizing conditions suggesting carbon substrate utilization derived from methane oxidation. Furthermore, co-cultures with *Pseudomonas chlororaphis* HC exhibited enhanced methane oxidation rates compared to monoculture, indicating a stimulating effect of *P. chlororaphis* HC on methane utilization by MOB. Transcriptomic analysis revealed differential gene expression patterns in both MOB and NMOHB during co-culture and points to potential cross-feeding of C4 dicarboxylates and methanol from the MOB to the NMOHB. Further, NMOHB displayed upregulation of genes
involved in ROS remediation, suggesting a response to oxidative stress induced by methane oxidation. Additionally, NMOHB exhibited alterations in nitrogen utilization and amino acid metabolism genes, possibly reflecting adaptations to the co-culture environment. Overall, these findings shed light on the metabolic interactions between MOB and NMOHB in methane-oxidizing systems, highlighting the potential for carbon sharing, nitrogen exchange, and ROS defense mechanisms.

Introduction

Methane (CH₄) is a key greenhouse gas. It is the most abundant reduced compound in the Earth’s atmosphere and its impact on the global warming effect is second only to carbon dioxide (CO₂) (1). Between geochemical and biological sources, it is estimated that 500-600 Tg of methane is introduced into the atmosphere every year and about 60% of that total is anthropogenic (2). Some of the sources of CH₄ include arctic tundra and other wetlands, rice paddies, oceans, lakes, ponds, many different soils and sediments, landfills, and deserts. In each of these ecosystems, the major biological sink for CH₄ is oxidation to CO₂ by methane-oxidizing bacteria (MOB) (also called methanotrophs) (3). Because of their distinct role in CH₄ degradation, methanotrophs are critical components of these ecosystems and in the global carbon cycle.

Most aerobic methanotrophic bacteria are considered either type I or II methanotrophs. The most critical difference is that the type I methanotrophs utilize the ribulose monophosphate (RuMP) pathway for carbon assimilation while type II use the serine cycle (4). Prior to carbon assimilation, CH₄ is oxidized by aerobic MOB to methanol (CH₃OH) via an oxygen dependent methane monooxygenase (3). The CH₃OH is further oxidized to formaldehyde via a methanol dehydrogenase where it assimilated by either the RuMP or serine pathways (3). Remaining
formaldehyde is oxidized to formate and then to CO₂ via formaldehyde and formate dehydrogenases (3). Also, some MOB have been shown to use a complete oxidative TCA cycle (5). It has been shown that intermediate metabolites of CH₄ metabolism are released externally from the cell and made available to other organisms, including metabolites such as CH₃OH, acetate, and other organic acids (6,7).

Methanotrophs exist within microbial communities. There are several environments where CH₄ oxidizing bacteria have been described to cohabitate and be highly interactive with other members of the microbial community. Movile Cave in Romania is an underground cave system with CH₄ present in the atmosphere as well as in groundwater (8). In 2004, Hutchens et al. used a stable isotope probing method to track ¹³C labeled CH₄ in the microbial community and discovered evidence of an active methanotrophic community, as well as cross-feeding of CH₄ derived carbon (8). These results suggest that aerobic methanotrophs actively convert CH₄ into complex organic compounds and help to sustain a diverse community of microorganisms. Similarly, in 2007, a ¹³C labeled CH₄ experiment was carried out in rice paddy soil. It was shown that carbon derived from labeled CH₄ accumulated in a community of methanotrophs and non-methane-oxidizing heterotrophic bacteria (NMOHB) as well as amoebae, ciliates, and flagellates. This suggests that the CH₄ derived carbon can be integrated not only into the prokaryotic community but even into microeukaryotes that act as predators on the bacteria (9). CO₂ derived from microbial CH₄ oxidation has also been shown to be incorporated into in algae in peat bogs (10,11). Experiments utilizing stable isotope probing coupled with phylogenetic analyses such as these have been shown to be effective at critically analyzing carbon tracking through microbial communities (12). So, methanotrophs are critical to sustaining a variety of different ecosystems at many levels.
Some studies have been conducted to further characterize the relationships between MOB and NMOHB by analyzing which co-culture community members provide stimulatory or inhibitory growth effects for the CH₄ oxidizers in the consortia. Iguchi et al. in 2011 demonstrated that when *Methylovulum miyakonense* HT12 was grown in co-culture with one of five bacterial strains related to the genera *Rhizobium, Sinorhizobium, Mesorhizobium, Xanthobacter,* and *Flavobacterium,* there was a positive effect on methanotrophic growth and methane oxidation rate. Through further analysis of the stimulatory mechanism exerted by *Rhizobium* sp. Rb122, they identified cobalamin as the growth-stimulating factor. They observed this stimulatory effect for several other Gammaproteobacterial methanotrophs belonging to the genera *Methylobacter* and *Methylomonas,* but the same effect could not be observed in members of the Alphaproteobacterial methanotrophic genera *Methylocystis* and *Methylosinus* (13).

All this work established that the methane-oxidizing bacteria community is crucial to carbon cycling in a variety of environments and, at times, essential for ecosystem function and sustainability. However, there exist many questions regarding the physiological mechanisms that the microbes in CH₄ oxidizing communities use. Some work has been done detailing the specifics of the physiological, biochemical, and genetic mechanisms at work in these microbial interactions. Previous studies on the physiology of the interactions between methanotrophs and NMOHB focused on cross-feeding of CH₃OH derived from the oxidation of CH₄. In 2017, Krause et al. used a combination of transcriptomics and RT-PCR to show that CH₃OH is the dominant carbon and energy source the methanotroph *Methylobacter tundripaludum* provides to support growth of the methanol using NMOHB *Methylotenera mobilis.* They showed that in the presence of *Methylotenera mobilis,* gene expression of the MOB’s dominant methanol
dehydrogenase (MDH) shifts from the lanthanide-dependent MDH (noxF) to the calcium-dependent MDH (mxAF).

Correspondingly, CH\textsubscript{3}OH is released into the medium only when the methanotroph expresses the MxaF-type MDH. These results suggest a cross-feeding mechanism in which the NMOHB partner induces a change in expression of methanotroph MDHs, resulting in release of methanol for its growth. This work was crucial to our understanding of the carbon transfer mechanisms at work during growth in simple MOB-NMOHB co-cultures (7). In 2019, Takeuchi et al. expanded our understanding of how MOB are affected by coculturing with NMOHB. They used RNA-seq and differential gene expression analysis of the transcriptomes of the MOB Methylocaldum marinum S8 and the methanol using NMOHB Methyloceanibacter caenitepidi Gela4 and showed that cross-feeding between the MOB and NMOHB can occur. They also showed that the NMOHB can assimilate a wide range of substrates in addition to methanol, such as acetate, ethanol, formate, and succinate when grown in co-culture with a methanotroph when CH\textsubscript{4} is the only available carbon source. They discovered that the expression levels of nearly half of the genes of the NMOHB were altered during co-culture growth including genes for acetyl-CoA synthesis, the ethylmalonyl-CoA (EMC) pathway, and TCA cycle that were upregulated in co-culture. Since acetyl-CoA is a starting compound of the EMC pathway and TCA cycle, compounds that can be used for the acetyl-CoA synthesis are cross-fed from the MOB to the NMOHB. They speculate that a possible source of acetyl-CoA in co-culture is acetate since expression of acetyl coenzyme A synthetase increased. Also, they showed that the MOB produced acetate (6). These studies show that CH\textsubscript{3}OH is a possible source of carbon cross-fed in methane-oxidizing systems but not the only possible metabolite since acetate can cross-feed into NMOHB as a precursor of the EMC pathway and the TCA cycle.
These prior studies on carbon cross-feeding to NMOHB have been limited to MOB growing with other methylotrophs. This study aims to further our knowledge of carbon transfer physiology in methane-oxidizing systems by investigating how the MOB *Methylosinus trichosporium* OB3b functions in co-culture with three different NMOHB that were isolated from CH₄ oxidizing enrichment cultures, not commonly considered to be methylotrophs and are able to grow on complex growth media. These are *Flavobacterium* sp. HC, a strain of *Pseudomonas chlororaphis* HC, and a strain of *Cupriavidus* sp. HC. Each of these three bacteria are able to grow on carbon derived from CH₄ oxidation in co-culture with *M. trichosporium* OB3b, with *P. chlororaphis* HC providing a stimulatory growth effect in the system. Further transcriptomic analysis of the culture of OB3b and NMOHB indicates that some genes of the NMOHB TCA cycle as well as alcohol dehydrogenases are upregulated and that the most likely candidates for carbon utilized by NMOHB in the system are C₄ dicarboxylates and CH₃OH. Also, NMOHB play a role in nitrogen cycling in the system and upregulate ROS defense genes in response to growing in co-culture with OB3b.

**Methods**

**Growth Condition and Community Composition of Methane-Oxidizing Cocultures**

Starter cultures of the NMOHB were grown statically in 100 mL of modified nitrate mineral salts (NMS) media with 0.1% peptones. The starter culture of *M. trichosporium* OB3b was grown in 100 mL of modified NMS media with 10% CH₄ in air in a 160 mL serum bottle sealed with a butyl rubber stopper.

The modified NMS medium contains (per liter of distilled water) 1 g KNO₃, 0.27 g KH₂PO₄, 1 g MgSO₄ · 7H₂O, 0.14 g CaCl₂ · 2H₂O, and 0.23 g Na₂HPO₄, 0.5% (v/v) acidic trace element
solution (AcTES), and 0.02% alkaline trace element solution (AlTES). AcTES contains (per liter), 2 g FeSO$_4$ · 7H$_2$O, 0.07 g ZnSO$_4$ · 7H$_2$O, 0.5 g MnCl$_2$ · 4H$_2$O, 0.12 g CoCl$_2$ · 6H$_2$O, 0.01 g NiCl$_2$ · 6H$_2$O, 0.5 g CuSO$_4$, 0.01 g H$_3$BO$_3$, 0.06 g La(NO$_3$)$_3$, 0.06 g Ce(NO$_3$)$_3$, and 100 mM HCl. The AlTES contains (per liter) 0.11 g Na$_2$SeO$_4$, 0.05 g Na$_2$WO$_4$, 0.29 g Na$_2$MoO$_4$, and 0.04 g NaOH.

Starter cultures of the co-cultures were established by adding together a stationary phase culture of one of the NMOHB, either the *Flavobacterium* sp. HC, *Pseudomonas chlororaphis* HC, or *Cupriavidus* sp. HC with a stationary phase culture of *M. trichosporium* OB3b cells in a 1:1 (v:v) ratio in modified NMS at a 1% inoculum. Cells were grown together statically in 100 mL of modified NMS media in 160 mL serum bottles with butyl rubber stoppers and a headspace gas mixture of 10% CH$_4$ in air. Prior to inoculation into the co-culture system, the NMOHB cells were washed 2X with modified NMS media by pelleting cells via centrifugation at 10,000 rpm for 10 minutes, discarding old media, and resuspending the cell pellet in fresh modified NMS media. Starter cultures of the co-cultures were monitored for methane depletion by gas chromatography on a Shimadzu GC-14A gas chromatograph (Shimadzu, Japan) and after 70-80% of the CH$_4$ was used, a 5% inoculum was transferred to 500 mL of modified NMS media in 1 L stoppered bottles in triplicate, leaving 500 mL of available gas headspace with the same headspace gas composition as the starter cultures. Newly established experimental co-cultures were incubated at 28 °C without shaking and methane depletion was measured. Methane oxidation rates (MOR) were calculated as MOR=ln(Δ[CH$_4$])/ln(Δt). This is the slope of the natural log of the methane depletion curve where [CH$_4$] is the molar concentration of CH$_4$ and $t$ is time elapsed in hours. Cells in each condition were harvested in late log phase by centrifugation for 20 minutes at 4 °C and 4000 X g using an Eppendorf 5910 centrifuge. The
MOR was measured and compared to *M. trichosporium* OB3b growing in monoculture under the same conditions. The co-cultures of OB3b and NMOHB, the OB3b monoculture, and each of the NMOHB grown in modified NMS media with 0.1% peptones were further subjected to transcriptomic analysis as described below. The NMOHB grown in peptones was used as a reference for how that strain may alter its growth when growing in a CH$_4$ oxidizing system versus with peptones.

**Heterotrophic Plate Counts to Determine Growth of NMOHB in Methane-Oxidizing Cultures**

Heterotrophic plate counts were used to determine if the NMOHB were growing in co-culture where only CH$_4$ was present as a carbon source, Cells were grown on nutrient agar (for *Pseudomonas* and *Cupriavidus*) or TSA (for *Flavobacterium*). Since these media were not incubated in the presence of CH$_4$, only the NMOHB were able to grow. At the point of inoculation and after 10 days of co-culture growth, 1 mL of co-culture was diluted in a tenfold dilution series, plated onto the respective solid media in triplicate and incubated at 28 °C. Colonies appearing on the media were counted after 4 days of growth. This method is visualized in Figure 1. These plate count data were compared to plate count data from a control where the co-culture was established as described above, but no CH$_4$ was added.
RNA Extraction, Illumina Library Preparation, and Sequencing

Cells for RNA extraction were grown and harvested as described above with cell pellets stored at -80 °C. The Zymo Research Quick-RNA Fungal/Bacterial Miniprep kit was used for RNA extraction. Additional DNase treatments were carried out using the Invitrogen DNase I Amplification Grade treatment kit. Nucleic acid extracts were quantified using a Qubit Fluorometer 2.0 with RNA high specificity and DNA broad range specificity reagent kits.

Illumina compatible sequencing libraries were prepared for sequencing with the Zymo-Seq RiboFree Total RNA Library Kit (Zymo Research, USA) according to the manufacturer’s specifications. Prior to sequencing, the cDNA library was concentrated using a Thermo
Scientific Savant DNA120 SpeedVac Concentrator, and size selection was carried out using the Agilent 4150 TapeStation System.

Sequencing of the final cDNA library was performed at the Oklahoma Medical Research Foundation Center for Clinical Genomics (Oklahoma City, USA) using Illumina NovaSeq 6000 S4 PE150 chemistry.

**Reference Genome Sequencing**

Genomic DNA of *P. chlororaphis* HC from a log-phase grown culture grown in liquid TSB at 28 °C was extracted using the Qiagen DNeasy Blood and Tissue Kit. DNA was fragmented using a QSonica 800R3 sonicator and built into Illumina-compatible shotgun libraries using the KAPA HyperPrep Kit. Shotgun libraries were sequenced on an Illumina MiSeq instrument. Sequence data was processed using fastp v0.23.2 (14) to remove reads containing uncalled bases (‘N’), trim low-quality bases (q < 30), and merge overlapping reads. The resulting analysis-ready reads were used as input for de novo assembly with SKESA v2.5.1 (15). The quality of genomes was assessed by CheckM v1.1.3 (16). The genome for *M. trichosporium* OB3b (GCF_000178815.2) was downloaded from the NCBI RefSeq database (17). Gene prediction was performed using Prodigal v2.6.3 (18) and barrnap v0.9 (19).

**Read Mapping and Quantification**

Raw fastq reads were filtered for high-quality reads (q > 30) and the adaptors were trimmed using fastp v0.22.0 (14). Quality-filtered pair-end reads were mapped to respective reference genomes using Bowtie 2 v2.2.5 (20). The .bam files generated were parsed for DNA contamination issues by removing reads known to directionally be linked to genomic DNA, then removing an equal number of reads from the RNA pool to account for bias. The pre-computed
alignments generated by Bowtie 2 were used as input for transcript quantification using Salmon 0.14.2 (21).

**Differential Expression Analysis**

The R program DESeq2 was used for differential expression analysis of quantified reads using the likelihood ratio test method (22). Genes that were differentially expressed (adjusted p-value < 0.05, log2FC < -2 or > 2) were annotated using KEGG BlastKOALA and differentially expressed genes in pathways were reconstructed using the KEGG Mapper Reconstruct tool (23).

**Results**

**NMOHB Grow on Carbon Substrates Derived from Methane Oxidation by MOB**

Heterotrophic plate counts on nutrient agar were used to quantify the growth of *P. chlororaphis* HC, *Cupriavidus* sp. HC, and the *Flavobacterium* sp. HC in methane-oxidizing conditions. Each of the strains were able to grow in co-culture with *M. trichosporium* OB3b on NMS medium with CH₄ as the only carbon source (Fig. 2). The growth of each of the NMOHB in co-culture with *M. trichosporium* OB3b was compared to a control with no added CH₄, and in this control there was no significant increase in growth, with the *Flavobacterium* species even having a slight decrease in cells over time. Each NMOHB strain was also grown in NMS and 10% CH₄ and none of the strains are able to grow on CH₄ in monoculture. This indicates that *P. chlororaphis* HC, *Cupriavidus* sp. HC, and *Flavobacterium* sp. HC are all able to grow on products produced from the oxidation of CH₄ by *M. trichosporium* OB3b.
Figure 2: Mean heterotrophic plate counts (CFU/mL) of *Cupriavidus* sp. HC, *Pseudomonas chlororaphis* HC, and *Flavobacterium* sp. HC in coculture with *Methylosinus trichosporium* OB3b in methane oxidizing conditions and associated controls grown with no added methane. Error bars represent standard deviation of triplicate samples. Statistically significant differences in means (p-val < 0.05) between 0 (blue) and 10 (orange) days of growth represented by *.
The Presence of *P. chlororaphis* in Coculture with *Methylosinus trichosporium* OB3b

Stimulates the Rate of Methane Oxidation

The MOR of co-cultures of either *P. chlororaphis* HC, *Cupriavidus* sp. HC, or the *Flavobacterium* sp. HC with *M. trichosporium* OB3b was measured and compared to *M. trichosporium* OB3b grown in monoculture to determine if the presence of any of the NMOHB strains stimulated the rate of CH$_4$ utilization by the MOB. When *M. trichosporium* OB3b is grown in co-culture with *P. chlororaphis* the MOR is greater than when *M. trichosporium* OB3b is grown in monoculture. When *M. trichosporium* OB3b is grown in co-culture with either *Cupriavidus* sp. HC or the *Flavobacterium* sp. HC, the MOR does not significantly change compared to the monoculture (Fig. 3). This demonstrates that the presence of *P. chlororaphis* HC, but not the other NMOHB, in culture with OB3b is stimulating the MOR of OB3b.
Figure 3: Mean methane oxidation rates of Methylosinus trichosporium OB3b in coculture with Flavobacterium sp. HC (FLV), Pseudomonas chlororaphis HC (PSD), or Cupriavidus sp. HC (CUP) compared to OB3b growing in monoculture. Error bars represent standard deviation of triplicate samples. There is a statistically significant (p-val < 0.05) difference in the mean methane oxidation rate of OB3b in coculture with PSD compared to OB3b in monoculture (*).
Genes Differentially Expressed by *Methylosinus trichosporium* OB3b in Coculture with NMOHB

Across all conditions, there were 25 genes, differentially expressed by *M. trichosporium* OB3b (Table 1). When *M. trichosporium* OB3b was in co-culture with *P. chlororaphis* HC, 20 genes were differentially expressed of which 8 were annotated. These genes included an arsenate reductase gene (*arsC*) and associated transcriptional regulators, hydroxylamine reductase (*hcp*), and the large subunit of a nitrite reductase (*nasB*). When *M. trichosporium* OB3b was in co-culture with *Cupriavidus* sp. HC, 5 genes were differentially expressed of which 4 were annotated. These included an outer membrane immunogenic protein, flavodoxin I, pyrroline-5-carboxylate reductase (*proC*), and L-ornithine Nalpha-acyltransferase (*olsB*). When *M. trichosporium* OB3b was in co-culture with *Flavobacterium* HC there were no differentially expressed genes.
Table 1: Differentially expressed genes of Methylosinus trichosporum OB3b in co-culture with Pseudomonas chlororaphis HC or Cupriavidus sp. HC compared to OB3b in monoculture. A negative log2FC represents genes that are expressed less in the co-culture condition. There were no differentially expressed genes in co-culture with Flavobacterium sp. HC.

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<td>3.09</td>
<td>2.71E-10</td>
<td></td>
</tr>
<tr>
<td>OB3B.00001_2499</td>
<td>-5.18</td>
<td>0.019841577</td>
<td></td>
</tr>
<tr>
<td>OB3B.00001_2609</td>
<td>-2.68</td>
<td>0.001470256</td>
<td></td>
</tr>
<tr>
<td>OB3B.00001_2610</td>
<td>9.15</td>
<td>0.020488634</td>
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</tr>
<tr>
<td>OB3B.00001_2956</td>
<td>2.36</td>
<td>0.04426312</td>
<td>lept; hydroxylamine dehydrogenase [EC:1.7.99.1]</td>
</tr>
<tr>
<td>OB3B.00001_3242</td>
<td>-2.22</td>
<td>0.02684378</td>
<td></td>
</tr>
<tr>
<td>OB3B.00001_3609</td>
<td>-2.00</td>
<td>0.00112231</td>
<td>ade, adh; chromosome partitioning protein</td>
</tr>
<tr>
<td>OB3B.00001_3786</td>
<td>-2.34</td>
<td>0.02191633</td>
<td>msdA, msdb; nitric oxide [NAD(P)H] reductase [EC:1.7.1.4]</td>
</tr>
<tr>
<td>OB3B.00001_3959</td>
<td>-3.79</td>
<td>0.035235756</td>
<td></td>
</tr>
<tr>
<td>OB3B.00001_4109</td>
<td>-2.99</td>
<td>0.000687722</td>
<td></td>
</tr>
<tr>
<td>OB3B.00001_497</td>
<td>2.63</td>
<td>0.030527272</td>
<td></td>
</tr>
</tbody>
</table>

Genes Involved in Carbon Utilization by Cupriavidus sp. HC

In co-culture with M. trichosporum OB3b, Cupriavidus sp. HC overexpresses fdwB, coding for formate dehydrogenase beta subunit as well as genes in the TCA cycle such as succinate dehydrogenase iron-sulfur subunit (frdB), isocitrate lyase (aceA), fumarate hydratase (fumA), and what is putatively annotated as D-malate dehydrogenase (dmlA). Cupriavidus sp. HC also shows increased expression of an alcohol dehydrogenase gene (adh) in co-culture with OB3b that could potentially be responsible for the utilization of methanol (Table 2).

Genes involved in Carbon Utilization by Pseudomonas chlororaphis HC

In co-culture, P. chlororaphis HC overexpresses an alcohol dehydrogenase gene (alhP) and TCA cycle genes encoding for malate dehydrogenase (maeB), acetyl-CoA synthetase, and fumarate hydratase (fumC) when compared to when P. chlororaphis HC growing on peptones.

Additionally, a set of genes coding for an aerobic C4-dicarboxylate transport protein (dctABD),
is also overexpressed in co-culture with OB3b by *P. chlororaphis* HC, implying that the C4 carbon derived from the oxidation of CH₄ by OB3b could be assimilated by *P. chlororaphis* HC (Table 2).

**Genes Involved in Carbon Utilization by *Flavobacterium* sp. HC**

*Flavobacterium* sp. HC downregulates two genes involved in glycolysis/gluconeogenesis including glyceraldehyde 3-phosphate dehydrogenase (*gapA*) and phosphoenolpyruvate carboxykinase (*pckA*) in co-culture with OB3b compared to the monoculture. Under the same condition, it also downregulates one gene involved in the TCA cycle; succinate dehydrogenase iron-sulfur subunit (*sdhB*). Unlike in *P. chlororaphis* HC and *Cupriavidus* sp. HC, genes related to central carbon metabolism are downregulated, and there is no evidence that an alcohol dehydrogenase is being expressed in co-culture with OB3b. In light of this, it is challenging to pinpoint the pathways used for growth in co-culture with OB3b (Table 2).
**Table 2**: Differentially expressed genes related to carbon metabolism for the NMOHB in co-culture with *Methylosinus trichosporium* OB3b. A positive log2FC represents genes that are expressed more in the co-culture condition compared to the NMOHB grown in NMS + peptones medium.

<table>
<thead>
<tr>
<th><em>Pseudomonas chlororaphis</em> HC with <em>Methylosinus trichosporium</em> OB3b</th>
<th>Gene ID</th>
<th>log2FC</th>
<th>Adj P-Value</th>
<th>Annotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCMA.00014_59</td>
<td>7.42736</td>
<td>0.03155606</td>
<td><em>adhP</em>; alcohol dehydrogenase</td>
<td></td>
</tr>
<tr>
<td>PCMA.00019_12</td>
<td>9.00466</td>
<td>0.03153312</td>
<td><em>maeB</em>; malate dehydrogenase</td>
<td></td>
</tr>
<tr>
<td>PCMA.00001_182</td>
<td>2.86899</td>
<td>0.017978906</td>
<td><em>fumC</em>, fumarate hydratase, class II</td>
<td></td>
</tr>
<tr>
<td>PCMA.00001_49</td>
<td>5.83743</td>
<td>0.013491053</td>
<td><em>dctA</em>; aerobic C4-dicarboxylate transport protein</td>
<td></td>
</tr>
<tr>
<td>PCMA.00065_25</td>
<td>7.98401</td>
<td>0.017077523</td>
<td><em>dctB</em>; C4-dicarboxylate transport sensor histidine kinase</td>
<td></td>
</tr>
<tr>
<td>PCMA.00065_24</td>
<td>8.04993</td>
<td>0.008705995</td>
<td><em>dctD</em>; C4-dicarboxylate transport response regulator</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><em>Cupriavidus</em> sp HC with <em>Methylosinus trichosporium</em> OB3b</th>
<th>Gene ID</th>
<th>log2FC</th>
<th>Adj P-Value</th>
<th>Annotation</th>
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</thead>
<tbody>
<tr>
<td>CUP.00008_11</td>
<td>8.08528</td>
<td>1.36E-05</td>
<td><em>fdwB</em>; formate dehydrogenase beta subunit</td>
<td></td>
</tr>
<tr>
<td>CUP.00008_60</td>
<td>6.77362</td>
<td>0.002740338</td>
<td><em>sdhB</em>; succinate dehydrogenase iron-sulfur subunit</td>
<td></td>
</tr>
<tr>
<td>CUP.00154_2</td>
<td>12.1813</td>
<td>0.022377965</td>
<td><em>aceA</em>; isocitrate lyase</td>
<td></td>
</tr>
<tr>
<td>CUP.00282_4</td>
<td>2.77132</td>
<td>0.005704623</td>
<td><em>fumA</em>; fumarate hydratase, class I</td>
<td></td>
</tr>
<tr>
<td>CUP.00383_2</td>
<td>6.44547</td>
<td>0.002530642</td>
<td><em>dmlA</em>; D-malate dehydrogenase</td>
<td></td>
</tr>
<tr>
<td>CUP.00024_14</td>
<td>9.61713</td>
<td>0.000922184</td>
<td><em>adh</em>; alcohol dehydrogenase</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><em>Flavobacterium</em> sp HC with <em>Methylosinus trichosporium</em> OB3b</th>
<th>Gene ID</th>
<th>log2FC</th>
<th>Adj P-Value</th>
<th>Annotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>FB1.00019_44</td>
<td>-4.4595</td>
<td>0.046665232</td>
<td><em>gapA</em>; glyceraldehyde 3-phosphate dehydrogenase</td>
<td></td>
</tr>
<tr>
<td>FB1.00001_337</td>
<td>-2.697</td>
<td>0.035652999</td>
<td><em>pckA</em>; phosphoenolpyruvate carboxykinase</td>
<td></td>
</tr>
<tr>
<td>FB1.00001_206</td>
<td>-2.2889</td>
<td>0.029635791</td>
<td><em>sdhB</em>; succinate dehydrogenase iron-sulfur subunit</td>
<td></td>
</tr>
</tbody>
</table>
*Cupriavidus* sp. HC Overexpresses ROS Remediation Genes in Coculture with

*Methylosinus trichosporium* OB3b

When growing in co-culture with *M. trichosporium* OB3b, *Cupriavidus* sp. HC overexpresses multiple genes that could remEDIATE toxic ROS including SOD2 Fe-Mn family superoxide dismutase, thioredoxin-dependent peroxiredoxin (*prx*), glutathione-dependent peroxiredoxin (*pgdX*), thioredoxin-dependent peroxiredoxin (*tpx*), and genes for glutathione biosynthesis from glutamate via glutamate-cysteine ligase (*gshA*) and glutathione synthase (*gshB*) which are involved in peroxide remediation (24). This implies that in co-culture with OB3b, the *Cupriavidus* sp. HC must deal with exposure to ROS by utilizing ROS remediation proteins in ways that are likely not needed when growing in NMS + Peptones (Table 3).

*Flavobacterium* HC Underexpresses ROS Remediation Genes in Coculture with

*Methylosinus trichosporium* OB3b

When growing in co-culture with *M. trichosporium* OB3b, *Flavobacterium* sp. HC decreases its expression ROS remediation genes including thioredoxin 1(*trxA*), and lipoyl-dependent peroxiredoxin (*osmC*) (Table 3).

*Pseudomonas chlororaphis* Overexpresses ROS Remediation Genes in Coculture with

*Methylosinus trichosporium* OB3b

When *P. chlororaphis* is growing in co-culture with *M. trichosporium* OB3b, it overexpresses genes related to the remediation of ROS species such as hydrogen peroxide and superoxide. Specifically, the strain overexpresses peroxidases including catalase (*katE*) and a thioredoxin-dependent peroxiredoxin (*prx*) (Table 3).
Table 3: Differentially expressed genes related to ROS remediation for the NMOHB in co-culture with *Methylosinus trichosporium* OB3b. A positive log2FC represents genes that are expressed more in the co-culture condition compared to the NMOHB grown in NMS + peptones medium.

<table>
<thead>
<tr>
<th>Pseudomonas chlororaphis HC with Methylosinus trichosporium OB3b</th>
<th>Gene ID</th>
<th>log2FC</th>
<th>Adj P-Value</th>
<th>Annotation</th>
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</thead>
<tbody>
<tr>
<td>PCMA.00077_28</td>
<td>3.68097</td>
<td>0.004639686</td>
<td>katE; catalase</td>
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<tr>
<td>PCMA.00058_20</td>
<td>22.5344</td>
<td>0.046812146</td>
<td>prx; thioredoxin-dependent peroxiredoxin</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cupriavidus sp HC with Methylosinus trichosporium OB3b</th>
<th>Gene ID</th>
<th>log2FC</th>
<th>Adj P-Value</th>
<th>Annotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>CUP.00038_5</td>
<td>7.62229</td>
<td>3.87E-02</td>
<td>prx; thioredoxin-dependent peroxiredoxin</td>
<td></td>
</tr>
<tr>
<td>CUP.00584_2</td>
<td>3.28403</td>
<td>6.07E-07</td>
<td>prx; thioredoxin-dependent peroxiredoxin [EC:1.11.1.24]</td>
<td></td>
</tr>
<tr>
<td>CUP.00243_10</td>
<td>7.85164</td>
<td>0.017935295</td>
<td>gshA; glutamate-cysteine ligase</td>
<td></td>
</tr>
<tr>
<td>CUP.00282_4</td>
<td>3.63785</td>
<td>1.41122E-20</td>
<td>gshB; glutathione synthase [EC:6.3.2.3]</td>
<td></td>
</tr>
<tr>
<td>CUP.00037_2</td>
<td>8.84614</td>
<td>0.045748896</td>
<td>SOD2; superoxide dismutase</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Flavobacterium sp HC with Methylosinus trichosporium OB3b</th>
<th>Gene ID</th>
<th>log2FC</th>
<th>Adj P-Value</th>
<th>Annotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>FB1.00019_44</td>
<td>-9.2792</td>
<td>0.00024807</td>
<td>arcA; thiorexin 1</td>
<td></td>
</tr>
<tr>
<td>FB1.00004_165</td>
<td>-2.3608</td>
<td>1.057E-05</td>
<td>osmC; lipoyl-dependent peroxiredoxin</td>
<td></td>
</tr>
</tbody>
</table>

**Pseudomonas chlororaphis HC Changes Expression of Nitrogen Utilization Genes Under Methane-Oxidizing Conditions**

When *M. trichosporium* OB3b is growing in co-culture with *P. chlororaphis* HC, OB3b decreases its expression of the *nasBE* nitrite reductase large and small subunits. On the other hand, *P. chlororaphis* HC overexpresses the *narK* nitrate/nitrite transporter and the *nirB* nitrite reductase genes when compared to growing in NMS + peptones medium (*Table 4*), suggesting a possible shift in nitrogen metabolism in the co-culture.
*Cupriavidus* sp. HC Changes Expression of Nitrogen Utilization Genes Under Methane-Oxidizing Conditions

In co-culture with OB3b, *Cupriavidus* sp. HC overexpresses the nitrate/nitrite MFS transporter (narK), assimilatory nitrate reductase catalytic subunit (nasA), assimilatory nitrite reductase large subunit (nasB), and the assimilatory nitrite reductases large and small subunits (nirBD). Additionally, in co-culture with OB3b, the strain overexpresses genes related to the biosynthesis or degradation of amino acids. There is an additionally overexpressed gene related to proline degradation to glutamate via proline dehydrogenase (putA). It seems likely that most of these changes are in response to growing in the absence of peptones (*Table 4*).

*Flavobacterium* sp. HC Changes Expression of Nitrogen Utilization Genes Under Methane-Oxidizing Conditions

When *Flavobacterium* sp. HC is growing in co-culture, it overexpresses the nrtA and nrtB nitrate/nitrite transporter and transport system substrate-binding proteins. Additionally, the strain underexpresses ornithine--oxo-acid transaminase gene (rocD) and tryptophan synthase alpha chain gene (trpA), again suggesting a need for additional nitrogen (*Table 4*).
Table 4: Differentially expressed genes related to nitrogen and amino acid utilization for the NMOHB in co-culture with *Methylosinus trichosporium* OB3b. A positive log2FC represents genes that are expressed more in the co-culture condition compared to the NMOHB grown in NMS + peptones medium.

<table>
<thead>
<tr>
<th>Pseudomonas chlororaphis HC with Methylosinus trichosporium OB3b</th>
<th>Gene ID</th>
<th>log2FC</th>
<th>Adj P-Value</th>
<th>Annotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCMA.00014_59</td>
<td>7.42736</td>
<td>0.03155606</td>
<td><em>narK</em>; nitrate/nitrite transporter</td>
<td></td>
</tr>
<tr>
<td>PCMA.00045_11</td>
<td>12.3807</td>
<td>0.025245203</td>
<td><em>nirB</em>; nitrite reductase</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Cupriavidus sp HC with Methylosinus trichosporium OB3b</th>
<th>Gene ID</th>
<th>log2FC</th>
<th>Adj P-Value</th>
<th>Annotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>CUP.00164_7</td>
<td>8.80157</td>
<td>0.001428358</td>
<td><em>narK</em>; nitrate/nitrite transporter</td>
<td></td>
</tr>
<tr>
<td>CUP.00164_10</td>
<td>2.9744</td>
<td>0.039483739</td>
<td><em>nasB</em>; nitrite reductase [NAD(P)H] large subunit</td>
<td></td>
</tr>
<tr>
<td>CUP.00164_11</td>
<td>3.14536</td>
<td>0.000405005</td>
<td><em>nasA</em>; assimilatory nitrate reductase catalytic subunit</td>
<td></td>
</tr>
<tr>
<td>CUP.00164_8</td>
<td>8.86943</td>
<td>0.007747264</td>
<td><em>nirB</em>; nitrite reductase (NADH) large subunit [EC:1.7.1.15]</td>
<td></td>
</tr>
<tr>
<td>CUP.00164_9</td>
<td>6.03439</td>
<td>0.014278022</td>
<td><em>nirD</em>; nitrite reductase (NADH) small subunit [EC:1.7.1.15]</td>
<td></td>
</tr>
<tr>
<td>CUP.00467_1</td>
<td>3.14531</td>
<td>0.048500913</td>
<td><em>putA</em>; proline dehydrogenase</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Flavobacterium sp HC with Methylosinus trichosporium OB3b</th>
<th>Gene ID</th>
<th>log2FC</th>
<th>Adj P-Value</th>
<th>Annotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>FB1.00001_16</td>
<td>2.0401</td>
<td>0.032168366</td>
<td><em>nrtA</em>; nitrate/nitrite transport system substrate-binding protein</td>
<td></td>
</tr>
<tr>
<td>FB1.00001_17</td>
<td>2.10538</td>
<td>0.000780906</td>
<td><em>nrtB</em>; nitrate/nitrite transport system substrate-binding protein</td>
<td></td>
</tr>
<tr>
<td>FB1.00004_84</td>
<td>-2.0899</td>
<td>0.00398162</td>
<td><em>rocD</em>; ornithine--oxo-acid transaminase</td>
<td></td>
</tr>
<tr>
<td>FB1.00005_80</td>
<td>-6.7143</td>
<td>0.017739174</td>
<td><em>trpA</em>; tryptophan synthase alpha chain</td>
<td></td>
</tr>
</tbody>
</table>
Discussion

Cross-feeding of Methane Derived Carbon to NMOHB in Coculture Conditions

This study shows potential ways that carbon sharing occurs between OB3b and *P. chlororaphis* HC by observing the transcription of genes related to metabolism of metabolites that could be derived from CH$_4$ oxidation such as methanol and C4 dicarboxylates. This study indicates that *P. chlororaphis* HC and *Cupriavidus* sp. HC express alcohol dehydrogenases that could potentially be expressed to oxidize methanol derived from the initial step of CH$_4$ oxidation to methanol by methane monooxygenase in OB3b. It has been shown that in strict methylotrophs, methanol cross-feeding can occur and that the non-methane-oxidizing strains can even cause the MOB to shift expression of methanol dehydrogenase from a high affinity methanol dehydrogenase (MDh) to a low affinity MDh, thereby facilitating the release of methanol by the MOB cells for use by the non-methane-oxidizing bacteria (7). OB3b expresses both the high affinity *xoxF* and low affinity *mxaF* type MDhs in co-culture with each of the NMOHB and in monoculture, but there is no indication that OB3b switches expression to the low affinity MDh in co-culture. But because expression of the *mxaF* type is essential in some MOB for release of methanol from the cell in other MOB such as in the system studied by Krause et al. (7), it is reasonable that some methanol would be accessible to the NMOHB.

It is likely the *P. chlororaphis* HC strain may also be taking up TCA cycle intermediates including fumarate, succinate, or malate derived from CH$_4$ oxidation. This carbon could be assimilated by *P. chlororaphis* HC via the *dctABD* C4 dicarboxylate transport system which is expressed by *P. chlororaphis* HC in co-culture with OB3b and would correlate with the observation that fumarate hydratase and malate dehydrogenase are overexpressed by *P.*
*chlororaphis* HC. The expression of the transporter gene *dctA* is critical to the use of C4 carbon by *P. chlororaphis*. Other studies have shown that in *Pseudomonas chlororaphis* strain O6, the expression of *dctA* is directly linked to its utilization of C4 dicarboxylate carbon utilized in the TCA cycle and mutants of strain O6 lacking in their ability to expresses the *dctA* transport gene do not grow on succinate or fumarate and show reduced growth on malate (25–28). This demonstrates the potential for cross-feeding of carbon in methane-oxidizing systems is not necessarily strictly limited to methanol in this system, and perhaps C4 substrates such as fumarate, succinate, or malate are being directly fed into the TCA cycle. There is no indication that transport of C4 substrates is upregulated, however a number of TCA cycle genes in *Cupriavidus* sp. HC were also upregulated indicating a similar type of interaction within the *Cupriavidis* and *Pseudomonas* co-cultures.

**NMOHB Expression of Nitrogen Utilization Genes in Methane-Oxidizing Conditions**

A likely reason that *Cupriavidus* sp. HC may be increasing its expression of the proline dehydrogenase (*putA*) gene involved in the degradation of proline to glutamate in co-culture with OB3b could be that there is simply a shift in the types of amino acids available for metabolism when changing from growth in media with peptones to growth in a CH₄ oxidizing system. It is also observed that OB3b is downregulating *proC*, which catalyzes the biosynthesis of proline, this may also be a response to changing levels of proline available in the co-culture system compared to the monoculture system. While with these data it is hard to pinpoint the specific interactions and amino acid flux across conditions, the data suggest the cross-feeding of amino acids. It has been previously shown that bacterial community structure impacts the available amino acids in a given environment and cross-feeding of amino acids is a prominent interaction of bacterial co-cultures (29,30).
Also of note is that the upregulation of nitrate/nitrite transport system genes occurs in all NMOHB when in co-culture with OB3b. Expression of these genes is essential to the transport of nitrogen anions (31) and may be an indicator of nitrogen anion exchange in the co-culture system; however, the precise mechanism of exchange is unknown. Also of note is that OB3b decreases its expression of nasBE, the assimilatory nitrite reductase large and small subunits when in co-culture with *P. chlororaphis* HC while *P. chlororaphis* HC increases expression of the soluble nitrite reductase nirB in the same condition. Though with these data alone, it is challenging to elucidate the exact role of each organism in the nitrite metabolism, it has been demonstrated that bacteria within microbial biofilms may cooperate in ways such that aerobic methanotrophs and heterotrophic denitrifiers couple oxidation of methane or methane intermediates with denitrification (32). Since the *nas* genes encoding for assimilatory nitrite reduction in OB3b are being downregulated in co-culture, and *nir* (nitrite reductase) is being overexpressed by *P. chlororaphis* HC in the same condition, perhaps *P. chlororaphis* HC is using electrons from the oxidation of CH$_4$ derived metabolites to reduce nitrite in the methane-oxidizing system.

**NMOHB May Need to Compensate for the Buildup of Toxic ROS When Growing with OB3b**

Each of the NMOHB express ROS defense genes in co-culture with OB3b more than they do when growing in monoculture in NMS + peptones medium. It appears that *P. chlororaphis* HC deals with the effects of growing in co-culture with MOB in a methane-oxidizing system by expressing catalase (*katE*), and thioredoxin-dependent peroxiredoxin (*prx*). *Cupriavidus* sp. HC also overexpresses the SOD2 Fe-Mn family superoxide dismutase in co-culture, as well as thioredoxin-dependent peroxiredoxin (*prx*). But it also overexpresses glutathione-dependent
peroxiredoxin (pgdX), and thioredoxin-dependent peroxiredoxin (tpx) and genes for glutathione biosynthesis from glutamate via glutamate-cysteine ligase (gshA) and glutathione synthase (gshB). Unlike the other two strains, Flavobacterium actually decreases its expression of the ROS genes thioredoxin 1 (trxA) and lipoyl-dependent peroxiredoxin (osmC) in co-culture with OB3b. While OB3b possesses and is expressing in monoculture and co-culture conditions several genes related to ROS remediation such as SOD2 superoxide dismutase, catalase peroxidase (katG), these genes are not being differentially expressed across conditions.

During aerobic metabolism toxic reactive oxygen species (ROS) are generated including superoxide and hydrogen peroxide. (33–37). This endogenous production of ROS can have harmful effects on cells including damaging proteins, nucleic acids, and lipids (37). In addition to potential generation of toxic ROS by cellular respiration, aerobic MOB oxidize methane using methane monooxygenases that catalyze the oxidation of methane to methanol in the presence of dioxygen that may contribute to ROS generation (38). Bacteria possess a wide variety of ROS defense mechanisms including the expression of genes that code for proteins that directly act on ROS to reduce toxic effects. Some of the genes responsible for ROS remediation are being expressed by NMOHB in co-culture with OB3b including catalase (37), SOD (39), thioredoxin-dependent peroxiredoxin (40), thioredoxin 1, and lipoyl-dependent peroxiredoxin (40). When NMOHB shift from growing in peptones to growing on carbon derived from methane oxidation, they seem to alter the way that they deal with ROS by overexpressing certain ROS genes. There were some similarities in that the P. chlororaphis HC and Cupriavidus sp. HC both overexpressed prx genes when growing in co-culture. The observation that Cupriavidus sp. HC overexpresses genes for glutathione biosynthesis from glutamate via glutamate-cysteine ligase (gshA) and glutathione synthase (gshB) as a response to stress from peroxide exposure has also
been demonstrated in strains of *Pseudomonas aeruginosa*, where the *gshA* and *gshB* genes are overexpressed in response to different peroxides introduced into the growth system (24). In general, NMOHB appear to upregulate ROS defense genes as a response to growing in methane-oxidizing conditions compared to growing on peptones, however *Flavobacterium* sp. HC downregulates these genes in co-culture pointing to the variation of ROS response across organisms.

**Conclusion**

The NMOHB *Pseudomonas chlororaphis* HC, *Cupriavidus* sp. HC, and *Flavobacterium* sp. HC all grow in methane-oxidizing conditions when in co-culture with *Methylosinus trichosporium* OB3b and that co-cultures of *P. chlororaphis* HC and *M. trichosporium* OB3b present increased MOR compared to OB3b growing in monoculture. Transcriptomics analysis reveals that carbon sharing between OB3b and the NMOHB is most likely occurring as cross-feeding of methanol and/or C4 dicarboxylates. Also, nitrite and amino acid metabolism are altered in both OB3b and NMOHB when growing in co-culture and ROS defense genes by NMOHB are differentially expressed during growth in co-culture with OB3b. These observations provide insight into how NMOHB that are not strict methylotrophs respond to growth in methane-oxidizing systems.
References


Chapter 4
Conclusion

The whole of the work present in the previous chapters demonstrates the breadth and diversity of research regarding our understanding of the group of organisms known as methane-oxidizing bacteria. The research in this field is so diverse because the impact of these organisms is critical to the carbon flux in a very wide variety of ecosystems, and methane metabolism is often the focus of many biotechnological applications including the generation of value-added products, and remediation of pollutants.

This work summarized the knowledge gaps in the fields of ecology and physiology of methane-oxidizing bacteria including how there was a lack of research into how different concentrations of oxygen or the presence of catalase in growth media as an ROS scavenger may affect the methane oxidation rate and alter the expression of ROS defense genes in MOB. This was despite previous knowledge of the chemical potential for MOB to generate ROS during methane metabolism and the critical role that oxygen flux and balance has on the physiology of any organism. This study demonstrates for the first time that methane-oxidizing bacteria alter their expression of ROS defense genes as a physiological response to increases in oxygen concentration and that some MOB such as *Methylomonas* sp. WSC-7 may even utilize cell clumping as a response to unfavorable oxygen conditions. There are several implications to these findings. Foremost, it should be noted that even given that only two MOB were discussed at depth during our investigation into how oxygen impacts MOB physiology, there were major distinctions. Properly understanding and optimizing oxygen concentrations is critical when trying to utilize the unique abilities of methane oxidizers for biotechnological and bioremediation
purposes. Further, as researchers aim to isolate and characterize more methane oxidizers, careful consideration should be made to how ROS generation may hinder our ability to do so effectively. Consideration of oxygen concentrations in the environments where MOB are isolated should be paramount, as unfavorable oxygen concentrations could potentially lead to decreases in methane metabolism, and certain MOB being outcompeted during the enrichment process by MOB that have a wide variety of means allowing them to scavenge ROS and not be adversely affected by ROS toxicity. To further our understanding of how ROS and changes in oxygen concentration impact MOB, it would be beneficial to understand MOB response across a wider range of MOB and try to correlate physiological response to ROS with MOB ecology and evolution. This would allow for more precise means of refining enrichment conditions for the targeted isolation of a wider variety of MOB.

In addition to presenting a refined overview of how MOB respond to changes in oxygen concentration, this work also presented a novel perspective of how MOB behave in co-culture with a variety of NMOHB that were isolated from methane-oxidizing enrichment cultures. This work helped identify some potential means of carbon flux in our established co-culture systems, and also demonstrated that NMOHB express ROS defense genes when growing in methane-oxidizing systems. Until now, most work done on how MOB and NMOHB interact in co-culture had been limited to carbon flux and cross-feeding. This work helped pinpoint potential modes of cross-feeding in our system, and also expanded the understanding of MOB community interactions by demonstrating the potential for NMOHB to deal with the negative consequences of ROS in co-culture, that there are changes in the expression of nitrite and amino acid utilization genes. It has been established that in some cases, the presence of NMOHB in co-culture with MOB can provide growth stimulating effects to the MOB leading to increased growth rates or
MOR rates in these systems. Further, this work shows that when *Methylosinus trichosporium* OB3b is in co-culture with *Pseudomonas chlororaphis* HC there is a growth stimulating effect. However, the cause of his stimulation of the MOR is unknown. Some future directions for this work may include trying to identify how MOB and NMOHB from the same ecosystem interreact to give insight into more ecologically relevant findings. This study utilized NMOHB that were isolated from methane-oxidizing enrichment cultures, but they may not primarily rely on methane-derived carbon in the natural environment, nor is it assumed that they coexist with species of *Methylosinus*.

Finally, this dissertation presents the isolation and characterization of a novel species of MOB, *Methylocystis suflitae*, named for University of Oklahoma emeritus professor Joseph M. Suflita. The characterization of this species highlights a critical aspect of research in the study of MOB, but also in microbiology research as well. Successful discovery and taxonomic classification of new species of microorganisms is an essential component to understanding microbial ecology and cultivating new organisms for the use of microbes to better our world through biotechnological applications. In light of the other work presented in this dissertation, I am hopeful that considerations regarding oxygen concentration, and how microbes are interacting in their broader community may help to refine methods of enrichment and isolation of novel bacteria, especially in targeted approaches following the identification of the presence of specific microorganisms within a particular environment by metagenomic, functional gene, or 16S rRNA surveys.