2020

Investigation of microbial community interactions between Lake Washington methanotrophs using genome-scale metabolic modeling

Mohammad Mazharul Islam
Tony Le

Follow this and additional works at: https://digitalcommons.unl.edu/chemengall
Investigation of microbial community interactions between Lake Washington methanotrophs using genome-scale metabolic modeling

Mohammad Mazharul Islam¹, Tony Le², Shardhat R. Daggumati¹ and Rajib Saha¹

¹ Department of Chemical and Biomolecular Engineering, University of Nebraska—Lincoln, Lincoln, NE, United States of America
² Department of Biochemistry, University of Nebraska—Lincoln, Lincoln, NE, United States of America

ABSTRACT

Background. The role of methane in global warming has become paramount to the environment and the human society, especially in the past few decades. Methane cycling microbial communities play an important role in the global methane cycle, which is why the characterization of these communities is critical to understand and manipulate their behavior. Methanotrophs are a major player in these communities and are able to oxidize methane as their primary carbon source.

Results. Lake Washington is a freshwater lake characterized by a methane-oxygen countergradient that contains a methane cycling microbial community. Methanotrophs are a major part of this community involved in assimilating methane from lake water. Two significant methanotrophic species in this community are Methylobacter and Methylomonas. In this work, these methanotrophs are computationally studied via developing highly curated genome-scale metabolic models. Each model was then integrated to form a community model with a multi-level optimization framework. The competitive and mutualistic metabolic interactions among Methylobacter and Methylomonas were also characterized. The community model was next tested under carbon, oxygen, and nitrogen limited conditions in addition to a nutrient-rich condition to observe the systematic shifts in the internal metabolic pathways and extracellular metabolite exchanges. Each condition showed variations in the methane oxidation pathway, pyruvate metabolism, and the TCA cycle as well as the excretion of formaldehyde and carbon di-oxide in the community. Finally, the community model was simulated under fixed ratios of these two members to reflect the opposing behavior in the two-member synthetic community and in sediment-incubated communities. The community simulations predicted a noticeable switch in intracellular carbon metabolism and formaldehyde transfer between community members in sediment-incubated vs. synthetic condition.

Conclusion. In this work, we attempted to predict the response of a simplified methane cycling microbial community from Lake Washington to varying environments and also provide an insight into the difference of dynamics in sediment-incubated microcosm community and synthetic co-cultures. Overall, this study lays the ground for in silico systems-level studies of freshwater lake ecosystems, which can drive future efforts of
understanding, engineering, and modifying these communities for dealing with global warming issues.

**Subjects** Bioinformatics, Computational Biology, Microbiology, Molecular Biology, Freshwater Biology

**Keywords** *Methylomonas* sp. LW13, Microbial community, Genome-scale metabolic modeling, Methane-cycling community, Lake Washington, *Methylobacter tundripaludum* 21/22

**INTRODUCTION**

The accelerated rise in worldwide average temperature in recent years is posing a serious threat to the environment, terrestrial ecosystems, human health, economy, and the ultimate survival of the planet earth. About 20 percent of global warming is caused by methane and it is expected to be 86 times more potent than carbon di-oxide in warming the earth over the next two decades (Houghton, Jenkins & Ephraums, 1990; IPCC, 2013). The impacts of the rapid increase in atmospheric methane (Nisbet et al., 2019) are compounded as higher temperatures are associated with an increase in methane production from wetlands and lakes (Yvon-Durocher et al., 2014). Aerobic methanotrophs are mostly gram-negative Proteobacteria that are an integral part of the global carbon cycle (Hanson & Hanson, 1996). They exist in diverse environments such as wetlands, lakes, and the tundra and use the enzyme methane monooxygenase (MMO) to oxidize methane as their sole source of carbon (Hanson & Hanson, 1996). Methane-oxidizing Verrucomicrobia found in geothermal acidic environments are also involved in methane oxidation while solely using carbon dioxide in the Calvin cycle (Carere et al., 2019; Mohammadi et al., 2019; Van Teeseling et al., 2014). The anaerobic methanotrophic archaea are also an important sink of methane that couple methane oxidation with sulfate reduction mediated by sulfate-reducing bacteria (Cui et al., 2015). Thus, methanotrophs act as the primary biological sink for methane (Hanson & Hanson, 1996), consuming up to 90 percent of the methane produced in soil/sediments in addition to the atmospheric methane (Whalen & Reeburgh, 1990; Krause et al., 2017). Methanotrophs have also shown the ability to produce various useful products such as single-cell proteins, biodiesels, biopolymers, and osmo-protectants (Strong, Xie & Clarke, 2015).

Lakes act as major sources and sinks for methane and account for 6 to 16 percent of biologically produced methane (IPCC, 2013; Yvon-Durocher et al., 2014). Lake Washington is a freshwater lake characterized by a methane-cycling community where methanotrophs are one of the important functional microbial groups involved in methane oxidation (Yu et al., 2016). It contains a steep vertical counter-gradient of methane and oxygen, and is separated into oxic and anoxic layers where methane production and consumption occur, respectively (Yu et al., 2016). Hence, it can be a model system to better understand methane-cycling communities in lakes and their role in the global methane cycle.

Understanding the metabolic interactions in these communities will aid in developing methods to reduce the amount of methane emitted from lakes. A diverse array of microbes exist in the Lake Washington community, where Proteobacteria comprise 33% of the
community and includes a major subtype of methanotrophs, *Methylococcaceae* at a 10% abundance level (Beck et al., 2013). The genus *Methylobacter* is the most dominant player in *Methylococcaceae* group at 47.7%, while other major players include *Crenothrix, Methylomonas*, and *Methylomicrobium* at 30.0%, 10.8%, and 7.4%, respectively (Beck et al., 2013). Other members include cyanobacteria, bacteriodetes, acidobacteria, and chloroflexi (Beck et al., 2013).

Understanding the physiological dynamics and interactions in natural methane-cycling communities, such as the Lake Washington, are crucial to addressing problems concerning methane’s role in global warming and leveraging methanotrophs’ possible functions in bioremediation and bioproduction. Omics-based techniques and high-throughput sequencing can elucidate important features of the community such as taxonomic information, community composition, and presence of functionally important genes (Temperton & Giovannoni, 2012). However, it is difficult to assign functionality to members of the community and decipher the roles of individual players due to the complexity of the community and the data involved (Zengler, 2009; Zengler & Palsson, 2012). On the other hand, synthetic communities were proven to be efficient models to provide insight into metabolic capabilities and interactions (De Roy et al., 2014). Simple representative community structures can be made to lower complexity, achieve more consistent results, and efficiently elucidate inter-species interactions (De Roy et al., 2014). Certain Lake Washington community members are easier to cultivate in a laboratory setting than other members. For instance, *Methylomonas* and *Methylosinus* species previously shown ease of cultivation in the laboratory (Auman et al., 2000). However, *Methylobacter* species were difficult to isolate and demonstrated poor growth compared to *Methylomonas* and *Methylosinus* (Yu et al., 2016). A synthetic community comprising 50 different Lake Washington microbes belonging to 10 methanotrophic, 36 methylotrophic, and 4 non-methanotrophic heterobacteria showed that *Methylobacter* was outperformed by *Methylomonas* in the community (Yu et al., 2016). Similar results about the dominance of *Methylomonas* in pure cultures and in standard conditions were observed in their later experiments (Yu, Beck & Chistoserdova, 2017). These observations were inconsistent with previous stable isotope probing studies that found *Methylobacter* is the dominant *Methylococcaceae* species among microbes from Lake Washington when grown on methane (Beck et al., 2013; Kalyuzhnaya et al., 2008). These inconsistencies indicate that the complexities of biological systems often make it challenging to understand the functions and interactions within and among organisms in synthetic communities via in vitro and in vivo studies.

In silico evaluation and analysis utilizing mathematical relation-based modeling allow for a high-resolution understanding of the biological processes in a microbial community. The availability of genome-scale metabolic network models combined with biological constraints provide multiple methods to analyze, perform in silico experiment, develop hypotheses, and redesign biological systems at a genome-level (Zomorrodi & Maranas, 2012; Zomorrodi, Islam & Maranas, 2014; Maranas & Zomorrodi, 2016; Islam & Saha, 2018; Alsiyabi, Immethun & Saha, 2019; Islam et al., 2020; Schroeder, Harris & Saha, 2020; Schroeder & Saha, 2019). To develop effective multi-species community models, significant
and comprehensive knowledge of inter-species interactions and experimental data must be utilized. Compartmentalized community metabolic models were used to model simple microbial consortia involved in bioremediation, synthetic auxotrophic co-growth, human gut microbiome, and soil bacterial ecosystems (Stolyar et al., 2007; Bizukoje et al., 2010; Lewis et al., 2010; Zhuang et al., 2011; Shoaie & Nielsen, 2014; Henry et al., 2016). On the other hand, community modeling frameworks incorporating the trade-offs between species- and community-level fitness successfully modeled steady-state and dynamic behavior in naturally occurring and synthetic soil microbial communities, synthetic co-cultures for bioproduction, human gut microbiome, and very recently to understand the microbial interactions in bovine rumen and viral influences (Zomorrodi & Maranas, 2012; Zomorrodi, Islam & Maranas, 2014; Chan, Simons & Maranas, 2017; Islam et al., 2019). Other novel methods were also proposed for modeling of such communities involving elementary mode analysis, evolutionary game theory, nonlinear dynamics, and stochastic processes (Vallino, 2003; Lehmann & Keller, 2006; Shou, Ram & Vilar, 2007; Borenstein & Feldman, 2009; Freilich et al., 2009; Frey, 2010; Nadell, Foster & Xavier, 2010; Magnúsdóttir, Heínken & Kutt, 2017). While there have been multiple studies in recent years involved in the model development of various methanotrophs (Akberdin et al., 2018; Bordel et al., 2019a; Bordel, Rojas & Munoz, 2019b; De la Torre et al., 2015; Lien et al., 2018; Naizabekov & Lee, 2020), an integrated community level analysis of freshwater methan utilizing ecosystems have not been performed yet.

In this work, we developed a simplified community metabolic model with two representative and functionally important strains of Lake Washington, namely, Methylobacter tundripaludum 21/22 (hereafter, Methylobacter) and Methylomonas sp. LW13 (hereafter, Methylomonas) as representative organisms of the methane-oxidizing microbes in the Lake Washington ecosystem. These species were chosen because of their availability in Lake Washington sediments, the ability to mitigate common pollutants, and produce desirable biological products, and the availability of genome-annotation. Draft models of these species were reconstructed followed by careful curation to ensure proper representation of the species. Metabolic pathways and individual reactions that are fundamental to the growth of these organisms such as the ribulose monophosphate pathway, pentose phosphate pathway, methane metabolism, amino acid synthesis and utilization, serine cycle, and Coenzyme B12 biosynthesis were manually scrutinized and then integrated into the models. The metabolites exchanged by the two species models were established by referring to literature and known transporter information (Boden et al., 2011; Caspi et al., 2016; Elbourne et al., 2017; Henry et al., 2010; Kanehisa, 2008; Nguyen et al., 2018; Orata, Kits & Stein, 2018; Svenning et al., 2011; Szklarczyk et al., 2017; UniProt Consortium, 2018; Wartiainen et al., 2006). The curated models of Methylobacter (704 genes, 1,329 metabolites, and 1,404 reactions) and Methylomonas (658 genes, 1,378 metabolites, and 1,391 reactions) were then utilized to develop a community model using a multi-level optimization framework, which was used to estimate biologically feasible metabolite secretion profiles and community compositions (Zomorrodi & Maranas, 2012; Zomorrodi, Islam & Maranas, 2014; Islam et al., 2019). The community was placed under carbon, oxygen, and nitrogen-limiting as well as nutrient-rich environments to study
the changes in intracellular carbon and nitrogen metabolism and metabolite excretion profiles. The community composition of carbon-limited environments predicted a shift in the carbon metabolism of both species. The community also demonstrates conservative metabolism under oxygen and carbon-limited environments and produce less carbon-di-oxide. Under these conditions, the mutualistic behavior involving formaldehyde transfer between *Methylobacter* and *Methylomonas* is rarely observed. Our results also indicate metabolic reprogramming in TCA cycle and pyruvate metabolism, which can help generate new hypotheses for in vivo experiments. We also simulated the observed binary compositions in a sediment-incubated community and a synthetic co-culture to predict the changes in intra- and extracellular metabolic fluxes. Overall, our results enhance the mechanistic understanding of the Lake Washington methane-cycling community, which can drive further engineering efforts for efficient rerouting of carbon and nitrogen as well as mitigation of methane emission from freshwater ecosystems globally.

**MATERIALS & METHODS**

**Metabolic model reconstruction**

The draft genome-scale metabolic models of *Methylobacter* and *Methylomonas* were developed and downloaded using the ModelSEED database (Henry et al., 2010). The models included reactions for glycolysis/gluconeogenesis, citrate cycle, pentose phosphate pathway, steroid biosynthesis, nucleotide metabolism, and various amino acid biosynthesis. Flux Balance Analysis (FBA), a mathematical approach for analyzing the flow of metabolites through a metabolic network, was utilized for model testing and analyzing flux distributions throughout the work (Orth, Thiele & Palsson, 2010). FBA implements the following optimization framework.

\[
\begin{align*}
\text{Maximize} & \quad v_{\text{biomass}} \\
\text{subject to} & \quad \sum_{j \in I} S_{ij} \cdot v_j = 0 \quad \forall i \in I \\
& \quad LB_j \leq v_j \leq UB_j \quad \forall j \in I
\end{align*}
\]

In the framework, \( I \) and \( J \) represent the sets of metabolites and reactions in the metabolic model, respectively. \( S_{ij} \) represents the stoichiometric coefficient of metabolite \( i \) in reaction \( j \). The flux value of each reaction \( j \), \( v_j \), must be within the parameters of the minimum, \( LB_j \), and maximum, \( UB_j \), biologically allowable fluxes. \( v_{\text{biomass}} \) is the flux of the biomass reaction which simulates the yield of cellular growth in the model (Orth, Thiele & Palsson, 2010).

The biomass composition from recently published methanotroph model *Methylomicrobium buryatense* strain 5G(B1) (De la Torre et al., 2015) was adopted in this study with slight modification in the lipid macromolecular stoichiometry to account for different lipid macromolecules in the models (see Data S1). Non-growth associated ATP maintenance
flux was set to 21.6 mmol/gDCW.hr according to calculated values in the closely related organism *Methylomicrobium alcaliphilum* by Akberdin et al. (2018). All of the three modes of electron transfer during Methane oxidation (redox arm, direct coupling and uphill electron transfer) have been included in the models, since they are all possible and there is no definitive conclusion of which organism prefers what mode of methane oxidation.

**Metabolic model curation**

Draft models for both *Methylobacter* and *Methylomonas* underwent an extensive manual curation process, including chemical and charge-balancing, elimination of thermodynamically infeasible cycles, and ensuring network connectivity. Reactions in the draft model reaction set imbalanced in protons were checked with their appropriate protonation states and corrected by adding and deleting proton(s) on either side of the reaction equation. The remaining imbalanced reactions were stoichiometrically inaccurate and required the atoms on both sides of the reaction equation to be balanced. The metabolites consumed and produced by models were established by referring to literature and known transporter information (Boden et al., 2011; Caspi et al., 2016; Elbourne et al., 2017; Henry et al., 2010; Kanehisa, 2008; Nguyen et al., 2018; Orata, Kits & Stein, 2018; Svenning et al., 2011; Szklarczyk et al., 2017; UniProt Consortium, 2018; Wartiainen et al., 2006).

Genome annotations of methanotrophs indicate that formaldehyde assimilation can happen through the oxidative Pentose Phosphate Pathway as well as the formaldehyde/formate dehydrogenases and the tetrahydrofolate-associated pathways (Cai et al., 2016; Khmelenina et al., 2018). However, there have been extensive evidences that the Ribulose Monophosphate (RuMP) cycle is the major route for formaldehyde assimilation (Fu, Li & Lidstrom, 2017; He et al., 2020; Peyraud et al., 2011). Also, multiple studies indicate that formaldehyde assimilation through the Serine cycle is insignificant (De la Torre et al., 2015; Fu, Li & Lidstrom, 2017; He et al., 2020). Based on transcriptomic data and metabolic flux measurements (De la Torre et al., 2015; Fu, Li & Lidstrom, 2017), we restricted the distribution of formaldehyde assimilation between the RuMP cycle and the tetrahydrofolate-associated pathways. In addition, the oxygen stoichiometry in pathways related to direct coupling methane oxidation and the coupling with cytochrome c oxidase have been set (1 mol Oxygen/mol pyrroloquinoline quinone and 0.5 mol Oxygen/mol Cytochrome c) according to literature (De la Torre et al., 2015; Fu, Li & Lidstrom, 2017; Lieven et al., 2018; Sugioka et al., 1988).

While applying mass balance constraints to genome-scale metabolic models can display the net accumulation and consumption of metabolites within each microbial model, it fails to account for the regulation of reaction fluxes. The limitation of this constraint is better elucidated when focusing on reaction cycles that do not consume and produce metabolites. Because of the absence of metabolite consumption and production, the overall thermodynamic driving force of the cycles become zero and the cycle is incapable of supporting any net flux, and thus deemed thermodynamically and biologically infeasible (Schellenberger, Lewis & Palsson, 2011). These thermodynamically infeasible cycles in our models were identified by inhibiting all nutrient uptakes to the cell and utilizing the
optimization framework, Flux Variability Analysis (FVA), which maximizes and minimizes each reaction flux within the model based on the mass balance constraints (Mahadevan & Schilling, 2003). Reactions whose fluxes reached the defined lower and upper bounds were determined to be unbounded reactions, group together based on stoichiometry, and systematically corrected. These cycles were eliminated by removing duplicate reactions, turning off lumped reactions, fixing reaction directionality, or selectively turning reactions on or off based on cofactor specificities found from literature (Ishida et al., 1969; Dekker, Lane & Shapley, 1971; Chen, Lee & Chang, 1991; Schomburg & Stephan, 1995; Achterholt, Priefert & Steinbuchel, 1998; Hadfield et al., 1999; Hutter & Singh, 1999; Kai, Matsumura & Izui, 2003; Feist et al., 2007; Dean, 2012; Flamholz et al., 2012; Lin et al., 2014; Christensen et al., 2017). The FVA optimization algorithm is shown below.

\[
\text{Maximize}/\text{minimize}(v_j) \quad v_j
\]

subject to

\[
\sum_{j \in I} S_{ij} v_j = 0 \quad \forall i \in I
\]

\[
LB_j \leq v_j \leq UB_j \quad \forall j \in I
\]

\[v_{\text{biomass}} = v_{\text{app}} - \text{obj.threshold}\]

Both metabolic models were checked for the ability to produce biomass and metabolites they were known to produce (Svenning et al., 2011; Kalyuzhnaya et al., 2015). The metabolic functionalities of the models were ensured by identifying and manually adding reactions from biochemical databases, such as KEGG (Kanehisa & Goto, 2000) and Uniprot (UniProt Consortium, 2018), to each model. Fully developed models of related organisms such as Methylococcus capsulatus and Methanomonas methanica for both Methylobacter and Methylomonas were utilized in help pinpoint absent enzymatic activity within our models. The addition of these reactions was confirmed using the bioinformatic algorithm, BLAST, which compares the genes of our organisms and related organisms and determines whether they are found to be orthologous. It was then tested whether these reactions would increase the number of thermodynamically infeasible cycles and promote for further curation of the models.

**Community model formation**

Following the curation of each individual microbial model, both were implemented to form a community model using the bi-level multi-objective optimization framework OptCom (Zomorrodi & Maranas, 2012). OptCom simultaneously optimizes each individual community member’s flux balance analysis problem as an inner-level optimization problem and the community model objective as an outer-level optimization problem. The maximization of the community biomass was set as the community objective function. The mathematical framework of the OptCom procedure is the following.
Maximize (or minimize) \( z = \) Community level objective subject to,

\[
\begin{align*}
\text{maximize} & \quad v^k_{\text{biomass}} \\
\text{subject to,} & \quad \\
\sum_{j \in J} x_{ij}^k v_j^k &= 0 \quad \forall i \in I^k (1) \\
\text{LB}_j^k & \leq v_j^k \leq UB_j^k \quad \forall j \in J^k (2) \\
v_{\text{uptake},i}^k &= v_{\text{uptake},i}^k \quad \forall i \in I_{\text{uptake}}^k (4) \\
v_{\text{export},i}^k &= v_{\text{export},i}^k \quad \forall i \in I_{\text{export}}^k (5) \\
\sum_k r_{\text{uptake},i}^k + c_i^c &= \sum_k r_{\text{export},i}^k + u_i^c \quad \forall i \in I_{\text{shared}} (6) \\
r_{\text{uptake},i}^k, r_{\text{export},i}^k, c_i^c, u_i^c & \geq 0 \quad \forall i \in I_{\text{shared}}, \quad \forall k \in K
\end{align*}
\]

The inner-level optimization problem(s) represents the steady-state FBA problem for each community member \( k \) and limits the exchange (uptake and export) fluxes of shared metabolites between the individual species using the outer-level optimization problem parameters \( r_{\text{uptake},i}^k \) and \( r_{\text{export},i}^k \) respectively. The outer-level optimization problem constraint characterizes the mass balance for every shared metabolite in the extracellular environment within the shared metabolite pool. Metabolic interactions, with constraints, were confirmed with prior experimental research involving both community members or one member with a related organism found in the Lake Washington community \((Yu \ et \ al., \ 2016)\).

**Formaldehyde inhibition constraint**

Formaldehyde production can be inhibitory towards the growth of *Methylomonas* and *Methylobacter* \((Bussmann, \ Rahalkar \ & \ Schink, \ 2006)\). *Methylobacter* is able to uptake formaldehyde excreted from *Methylomonas* to alleviate the inhibitory effects on *Methylomonas* growth. A constraint based upon the minimum and maximum inhibitory concentrations of formaldehyde was implemented to simulate formaldehyde’s inhibitory effects on *Methylobacter* \((Hou, \ Laskin \ & \ Patel, \ 1979; \ Costa, \ Dijkema \ & \ Stams, \ 2001)\). While formaldehyde production is not forced in the model simulations, this constraint accounts for the resulting growth inhibition if there is any amount of formaldehyde excreted. The minimum formaldehyde concentration required for growth inhibition was found to be 1 mM \((Hou, \ Laskin \ & \ Patel, \ 1979)\) and the formaldehyde concentration required for total growth inhibition (maximum formaldehyde concentration) was found to be 7 mM \((Costa, \ Dijkema \ & \ Stams, \ 2001)\).

\[
\begin{align*}
v_{\text{biomass}}^k \text{Methylobacter} & \leq v^k_{\text{biomass} \text{maxbiomass}} \times (1 - R \times v^k_{\text{Methylobacteruptake}}) \\
\end{align*}
\]

The parameter \( R \) was derived by finding the rate of change of biomass growth by the change in formaldehyde concentration.
| Models          | Methylobacter | Methylomonas |
|-----------------|---------------|--------------|
| Genes           | 704           | 658          |
| Reactions       | 1,404         | 1,391        |
| Metabolites     | 1,329         | 1,378        |
| Blocked reactions | 660         | 672          |

**Table 1** Model statistics for *Methylobacter* and *Methylomonas*.

**Bridging metabolic network gaps**

Automated draft reconstructions are limited as many reaction networks possess gaps due to missing reactions and blocked reactions. These are defined as reactions that lack production/consumption of its reactants/products. Major metabolic pathways were added based upon the literature of each organism (*Kalyuzhnaya et al., 2015*). Gaps were filled by referencing genetically related organisms to find missing metabolic capabilities. The presence of these possible reactions in the models were validated by cross referencing the relevant amino acid sequences between the reference organism and our models via pBlast. The reactions were then checked for the formation of thermodynamically infeasible cycles before being accepted.

**Computational resources**

The General Algebraic Modeling System (GAMS) version 24.8.5 with IBM CPLEX solver was utilized to run the optimization algorithms. The optimization frameworks were scripted in GAMS and then run on a Linux-based high-performance cluster computing system at the University of Nebraska-Lincoln. The downloaded models from ModelSEED with curations were parsed from System-Biology Markup Language (SBML) level 2 documentation using general-purpose programming language Python to generate the input files necessary for GAMS and subsequent manual curations.

**RESULTS**

**Genome-scale metabolic models of Methylobacter and Methylomonas**

The draft genome-scale models of *Methylobacter* and *Methylomonas* are reconstructed using the ModelSEED database (*Henry et al., 2016*). The manual curation process ensures that there is no chemical and charge imbalance present in either of the models and there is no reaction with unrealistically high fluxes (infeasible reactions) without any nutrient uptake. The manual curation also reconnects a significant number of blocked metabolites to the network in both models (*i.e.*, 107 metabolites for *Methylobacter* and 109 metabolites for *Methylomonas*). This enhancement of network connectivity is performed using available literature pertaining to major metabolic pathways that are known to be present in both the microbes (*Kanehisa & Goto, 2000; Kalyuzhnaya et al., 2015*). The draft models were lacking some reactions in the important metabolic pathways *i.e.*, the methane oxidation, pentose phosphate pathway, nitrogen fixation, cofactor, and amino acid production, which are curated at this stage. The model statistics are shown in Table 1. Data S2 and S3 contain the model files for *Methylobacter* and *Methylomonas*, respectively.
| Condition               | Methane uptake (mmol/gDCW.hr) | Oxygen uptake (mmol/gDCW.hr) | Nitrogen uptake (mmol/gDCW.hr) |
|------------------------|-------------------------------|-----------------------------|-------------------------------|
| A (high C, high O, high N) | 100                           | 100                         | 100                           |
| B (high C, limiting O, high N) | 100                           | 50                          | 100                           |
| C (high C, high O, limiting N) | 100                           | 100                         | 40                            |
| D (limiting C, high O, high N) | 20                            | 100                         | 100                           |

Community dynamics under variable environmental conditions
The individual models are integrated into a community model using available multi-objective computational optimization framework (Zomorrodi & Maranas, 2012). The metabolic interactions between Methylobacter and Methylomonas are described using inter-organisms flow constraints. The community, as a whole, consumes methane, oxygen, and nitrogen, which is then shared between Methylobacter and Methylomonas. In addition, Methylobacter consumes formaldehyde produced by Methylomonas under certain conditions, which alleviates the formaldehyde toxicity on Methylomonas growth (Bussmann, Rahalkar & Schink, 2006). At the same time, both Methylobacter and Methylomonas export carbon di-oxide to the environment. The community is simulated under four conditions in which methane, oxygen, and nitrogen are supplied to the community at various levels. These conditions are denoted as A, B, C, and D in Table 2 and correspond to the panels in Fig. 1. It should be noted that the amount of each nutrient consumed by the members are not necessarily equal to the amount of nutrient supplied as only one of the nutrients acts as a limiting reagent in each limiting condition. The community model is illustrated in Fig. 1.

The community biomass flux is the highest when all the nutrients are highly abundant (Fig. 1A). Methylomonas dominates the community in nutrient-rich condition. It is observed that a limited uptake of oxygen also imposes some restriction on the carbon and nitrogen utilization by the community, which results in reduced biomass fluxes for both Methylomonas and Methylobacter (Fig. 1B). When oxygen uptake is limited, methane utilization by Methylomonas is still higher compared to Methylobacter despite an overall decrease in methane consumption by the community. An overall conservative nature of metabolism is observed in the community, as manifested by the least amount of carbon-di-oxide production and no formaldehyde production. In nitrogen-limited growth (Fig. 1C), Methane uptake by Methylomonas decreases further and a high rate of respiration is observed, with the highest production of carbon-di-oxide. In Methane-limited condition (Fig. 1D), while the metabolism in Methylobacter remains mostly unaffected, Methylomonas growth is severely hindered by the scarcity of Methane. In all the nutrient-limited conditions, Methylobacter is observed to dominate the community. The observations from Optcom simulations are consistent with the shifts in possible flux ranges under different conditions, as estimated by Flux Variability analysis. These observations indicate that the inherent degeneracy of metabolic fluxes in Flux Balance Analysis or Optcom does not affect the conclusions obtained in this work. The detailed results are presented in Data S4.
The total community biomass and community composition under methane and oxygen gradients are simulated to model the methane-oxygen counter gradient that exists in Lake Washington (Yu et al., 2016). In general, the total community biomass is observed to increase with both oxygen and methane uptake, as is expected from the increased abundance of nutrients. The community is completely dominated by Methylobacter under low methane/low oxygen conditions. An increase of methane at low oxygen conditions does not change the dominance of Methylobacter. Furthermore, the increase of oxygen under low methane conditions has minimal impact in changing the community composition, as can be seen from very slowly increasing ratio of Methylomonas to Methylobacter across the entire range of oxygen uptake at low methane uptake condition (Fig. 2). The biggest charge in community composition is observed under high carbon and high oxygen conditions in which the community biomass is composed of 32% Methylobacter and 68% Methylomonas, as compared to 99% Methylobacter and 1% Methylomonas in the low methane/low oxygen condition (Fig. 2). The switch in community composition is consistent with the observations in nutrient rich condition. The complete numerical results are included in Data S4.

In nutrient-rich condition (shown in Fig. 3), Methylobacter is the dominant community member and consumes the major portion of all of the shared the metabolites i.e., methane, oxygen, and nitrogen. Methylomonas consumes the major share of methane (80%) and most of it is accumulated in biomass with a small amount of formaldehyde (0.1 mmol/gDC.hr)
being produced. *Methylomonas* also has a more active serine cycle converting formaldehyde into the metabolites in the central carbon metabolism. While the TCA cycle in both *Methylobacter* and *Methylomonas* is active in nutrient-rich condition, the activity of alpha-ketoglutarate dehydrogenase was very low.

Under the oxygen limiting condition (Fig. 4), the flux through the methane oxidation pathway decreases by 20% in *Methylobacter*. *Methylomonas* consumes the majority (60%) of the total methane uptaken by the community under the oxygen limiting condition (Fig. 1B). *Methylomonas* diverts central carbon compounds to produce pyruvate via the assimilation of carbon di-oxide and acetaldehyde under the oxygen limited condition (Fig. 4B). A small fraction of carbon di-oxide downstream of this reaction is secreted into the environment, which is the lowest among all the conditions. In this condition, *Methylomonas* has a complete TCA cycle activity.

The methane oxidizing pathway and TCA cycle reactions in *Methylobacter* do not show any significant change in nitrogen limited condition (Fig. 5A). *Methylobacter* excretes more carbon di-oxide than nutrient-rich condition while consuming more oxygen than *Methylomonas*. On the other hand, the activity of the methane oxidation pathway of

---

**Figure 2.** The community composition and total biomass under varying methane and oxygen conditions. The size of each pie chart represents the total community biomass.

Full-size DOI: 10.7717/peerj.9464/fig-2
Methylomonas decreases (the lowest in any non-carbon limiting conditions). Furthermore, succinyl-CoA in Methylomonas is produced from alpha-ketoglutarate in the TCA cycle, similar to oxygen limited conditions (Fig. 5B). Methylomonas, similar to Methylobacter, also excretes significantly high amount of carbon-di-oxide in nitrogen-limited condition.

In the carbon limited growth condition, Methylomonas takes up very small amount of methane, while Methylobacter takes up most of the methane supplied to the community (Fig. 6). The central metabolism of Methylobacter is not altered under this carbon limited condition (Fig. 6A). However, in Methylomonas, carbon di-oxide is scavenged by assimilating succinyl-dihydrolipoamide and carbon di-oxide to succinyl-CoA (Fig. 6B). This reaction is inactive in the oxygen limiting condition. Methylomonas also displays minimal activity in its serine cycle under carbon limiting condition.

**Dynamic shifts in metabolism under sediment incubated microcosm and synthetic co-culture composition**

Previous studies have shown inconsistencies between the microcosm incubated from Lake Washington sediments and synthetic community cultured in the lab. In the natural community (microcosm incubated from the lake sediments), Methylobacter has been shown to be dominant under both low methane/high oxygen and high methane/low
oxygen conditions (Beck et al., 2013). However, Methylococcus outcompeted Methylobacter in both low methane/high oxygen and high methane/low oxygen conditions in synthetic co-cultures (Yu et al., 2016). To elucidate the metabolic flux distributions and the extent of inter-species interactions that gives rise to the observed community composition in the two conditions, the experimentally observed species abundance ratio was imposed on the growth rates of Methylobacter (MB) and Methylococcus (ML) as a constraint in the community optimization framework. For the sediment-incubated community, a MB:ML ratio of 9.3:1.0 and for the synthetic community, a MB:ML ratio of 0.05:1.0 were used.

The vast majority of reactions in Methylococcus has lower flux values in the sediment-incubated community compared to the synthetic community (Fig. 7). This occurs because Methylococcus constitutes a smaller portion of the total community biomass in the sediment-incubated community. There are alternate pathways to produce pyruvate which has increased flux (Fig. 7A). These pathways produce pyruvate by assimilating carbon di-oxide and acetaldehyde and by the assimilation of cysteine and mercaptopyruvate. However, Methylococcus produces less pyruvate overall even with the increased flux in these pathways, since other pyruvate producing pathways decrease in flux. Primarily, the flux through L-serine and ammonia assimilation to produce pyruvate is high in the
Figure 5 Central carbon metabolism fluxes in *Methylobacter* (A) and *Methylomonas* (B) under nitrogen limited condition.

synthetic condition but decreases in sediment-incubated condition. Under sediment-incubated conditions, *Methylobacter* uses methane as its primary carbon source and does not consume any formaldehyde produced by *Methylomonas*. On the other hand, *Methylobacter* takes up some of the secreted formaldehyde from *Methylomonas* as a carbon source in addition to methane under the synthetic condition (Fig. 7B).

**DISCUSSION**

Aerobic methane oxidation in freshwater lakes around the world is a key metabolic process that significantly affects the carbon cycle by acting as a major sink of methane. Lake Washington provides an wonderful opportunity to study the methane cycling, with up to 20% of the organic carbon being released as methane through decomposition and consuming up to 10% of the dissolved oxygen in the lake water (Kuivila et al., 1988). With the goal to understand the dynamics of the methane cycling Lake Washington community, we integrated high-quality manually curated and refined genome-scale metabolic models of highly abundant, functionally important, and representative community members using multi-level optimization-based frameworks. While this community have been studied by many researchers in the previous years with in vivo tools like synthetic co-cultures,
metagenomics, and metatranscriptomics (Hernandez et al., 2015; Oshkin et al., 2015; Yu et al., 2016; Krause et al., 2017), an in silico approach like the one used in this work is needed to have a deeper understanding of the underlying mechanisms that govern the inter-species interactions and in turn, the community structure, function and dynamics.
in Lake Washington. As the global carbon transactions are changing and the release of greenhouse gases in the atmosphere is consistently getting worse every year, it is imperative for us to put our best efforts in mitigating the harmful effects. To do that, the use of genome-scale metabolic modeling tools to understand how the microbial communities are involved in these metabolic processes function in natural environments is essential.

To make the community model a good representation of the naturally occurring methane cycling community, we selected two highly abundant and functionally important microbial species. Following the manual curation process of both metabolic models, it was found that *Methylobacter* was more efficient than *Methylomonas* at producing biomass when simulated under the same standard growth environment and biological constraints in the Lake Washington, except at highly nutrient-rich conditions. Since *Methylobacter* and *Methylomonas* are competitors for the sole carbon source, methane, the overall metabolic efficiency is an important factor in the methane utilization ratio of the two species. While there is no direct literature evidence that suggests that one of them is more efficient in utilizing methane for growth compared to the other, it is highly possible that they might be limited by other small molecules that inhibit high methane consumption. For example, the community was tested under carbon, oxygen, and nitrogen limited conditions to observe how its central metabolic pathway and the community composition varied, and we observe the dominance of *Methylobacter* in the community in all of the limited growth conditions (Fig. 4 through 6) but the dominance of *Methylomonas* in the nutrient-rich condition. This is observed even during the oxygen limited growth condition, where *Methylobacter* is unable to consume as much methane as *Methylomonas*. In this condition, *Methylomonas* takes up more methane than *Methylobacter* but *Methylobacter* can still maintain its dominance.

Each of the nutrient limited conditions shows variable differences within the methane oxidation pathway, the serine cycle, and the TCA cycle. While, based on the simulation, we did not observe a noticeable flux through the serine cycle in either *Methylomonas* or *Methylobacter* in all nutrient-limited conditions, it was significantly active in *Methylomonas* in nutrient rich condition. In oxygen limiting condition, the community was also observed to conserve as much resource as possible. For example, while *Methylomonas* excretes some amount of carbon dioxide in all conditions, it routes most of it back to central carbon metabolism through the direction reversal of pyruvate decarboxylase. Although there is currently no experimental studies pointing to this phenomenon, studies in other organisms suggest a high oxygen sensitivity of this enzyme (*Eram & Ma, 2013*). Therefore, a possible explanation of the shifts in pyruvate metabolism is the oxygen sensitivity of this enzyme, which needs to be further studied.

In most of the simulation conditions, *Methylobacter* and *Methylomonas* were observed to assimilate most of the consumed carbon though the Entner-Doudoroff (ED) pathway. This was evident from the high flux of fructose-6-phospahte to 6-phophogluconate and thereafter pyruvate. In contrast, the Embden-Meyerhof-Parnas (EMP) variant of the glycolytic pathway was predicted to be the dominant pathway in some other methanotrophs like *M. buryatense* through model simulations (*De la Torre et al., 2015; He, Fu & Lidstrom, 2019; Kalyuzhnaya et al., 2013*). It should be noted that studies measuring glycolytic fluxes experimentally in *Methylobacter* and *Methylomonas* species are sparse in literature. A
recent $^{13}$C tracer analysis by *Fu, Li & Lidstrom (2017)* suggested that the flux partition between EMP and ED variants is unresolved since they both manifest fully labeled downstream molecules. Another interesting observation from this study is the activity levels of alpha-ketoglutarate dehydrogenase enzyme in all conditions. The presence and expression of alpha-ketoglutarate dehydrogenase in methanotrophs has been a matter of debate for quite some time (*Zhao & Hanson, 1984*; *Theisen & Murrell, 2005*), which is manifested as the inability of methanotrophs to grow on multi-carbon substrates (*Smith, Trotsenko & Murrell, 2010*). In this study we observe very negligible (<2% of the TCA cycle flux) alpha-ketoglutarate dehydrogenase activity in all conditions. At this point it is not straightforward to decipher what exactly might be the regulating factor to this enzyme and warrants further experimentation.

Finally, the community model was simulated with fixed abundance ratios of the two members to reflect the composition in synthetic co-culture and sediment-incubated community. The changes in community composition as oxygen and carbon levels change are more consistent with behaviors in sediment samples than synthetic co-cultures (*Hernandez et al., 2015; Yu et al., 2016*). *Methylobacter* is able to utilize methane to produce biomass at a more efficient level than *Methylomonas*. However, when the synthetic co-culture condition is imposed on the species abundance ratio, *i.e.*, favoring *Methylomonas* biomass, *Methylobacter* takes up formaldehyde produced by *Methylomonas* in addition to methane consumption, which allows it to bypass the oxygen-intensive reaction of oxidizing methane to methanol to some degree. This commensal relationship helps *Methylobacter* to enhance its biomass even when the uptake of the original carbon source (methane) is low while protecting *Methylomonas* from the toxicity and growth inhibitory effects of formaldehyde (*Hou, Laskin & Patel, 1979; Costa, Dijkema & Stams, 2001*). In our optimization formulation, the inhibitory formaldehyde constraint placed on *Methylobacter* makes the consumption of formaldehyde detrimental towards biomass production, but it can simultaneously act as a carbon source and compensates for its inhibitory effects. This is contradictory to what was observed in the co-culture experiments (*Yu et al., 2016*). Similar to formaldehyde, methanol export was also not observed in the simulations although some previous studies have reported ethanol production (*Krause et al., 2017; Zheng et al., 2018*). This lack of excretion of intermediate carbon molecules is often observed in steady-state Flux Balance Analysis models without any enzyme kinetic information, where pathways upstream of biomass consumes most of the intermediates. However, it should be noted that, the synthetic co-culture experiments reported in literature involved other community members, whose metabolic interactions with *Methylobacter* and *Methylomonas* are not well characterized to date. Some studies have indicated co-operative relationships between *Methylobacter* and other microbial species from *Methylophilaceae* family (*Beck et al., 2013*), which might potentially impact that community dynamics in synthetic co-cultures. Moreover, despite the dominance of *Methylomonas* in the synthetic community at a species level (*Soni et al., 1998; Hoefman et al., 2012*), further assessment of community composition at a higher taxonomic level indicated a consistency with naturally observed composition (*Yu et al., 2016*). We hypothesize that this discrepancy is possible, given that there is high functional redundancy present in Lake Washington community, similar to
any naturally occurring microbial ecosystem (Galand et al., 2018; Louca et al., 2018; Islam et al., 2019; Jia & Whalen, 2020).

**CONCLUSION**

In this work, we attempted to enhance our mechanistic understanding of the dynamics in the methane cycling Lake Washington community through genome-scale metabolic modeling of the representative and functionally important community members, *Methylobacter tundripaludum* 21/22 and *Methylomonas* sp LW13. The understanding of this community behavior will be a foundation for future studies that aim at the long-term goal of creating a complex synthetic community capable of carrying out certain desired functions through the consumption of methane, thus mitigating the harmful effects of methane release in the atmosphere. One should be aware of the fact that the in silico results need to be further tested and confirmed through relevant experiments before any engineering strategies can be successfully employed. The community metabolic model, in that regard, should be expanded to include other major players of the Lake Washington community, *i.e.*, members of *Bacteroidetes* and *Proteobacteria* phyla. Including these organisms in our community metabolic model will enable us to explain currently unidentified inter-species metabolite exchanges/interactions that play important role in the cycling of methane as well as other nutrients.

**ADDITIONAL INFORMATION AND DECLARATIONS**

**Funding**

This work was supported by University of Nebraska-Lincoln Faculty Startup Grant (21-1106-4308) to Rajib Saha and University of Nebraska-Lincoln UCARE Program. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Grant Disclosures**

The following grant information was disclosed by the authors:
University of Nebraska-Lincoln Faculty Startup Grant: 21-1106-4308.
University of Nebraska-Lincoln UCARE Program.

**Competing Interests**

The authors declare there are no competing interests.

**Author Contributions**

- Mohammad Mazharul Islam conceived and designed the experiments, performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Tony Le performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
- Shardhat R Daggumati performed the experiments, analyzed the data, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.
Rajib Saha conceived and designed the experiments, prepared figures and/or tables, authored or reviewed drafts of the paper, and approved the final draft.

Data Availability
The following information was supplied regarding data availability:

The simulation results and model files are available in the Supplementary Files.

Supplemental Information
Supplemental information for this article can be found online at http://dx.doi.org/10.7717/peerj.9464#supplemental-information.

REFERENCES

Achterholt S, Priefert H, Steinbuchel A. 1998. Purification and characterization of the coniferyl aldehyde dehydrogenase from pseudomonas sp. strain HR199 and molecular characterization of the gene. Journal of Bacteriology 180:4387–4391 DOI 10.1128/JB.180.17.4387-4391.1998.

Akberdin IR, Thompson M, Hamilton R, Desai N, Alexander D, Henard CA, Guarnieri MT, Kalyuzhnaya MG. 2018. Methane utilization in Methylococcus alcaliphilum 20Z(R): a systems approach. Scientific Reports 8:2512 DOI 10.1038/s41598-018-20574-z.

Alsiyabi A, Immethun CM, Saha R. 2019. Modeling the interplay between photosynthesis, CO2 fixation, and the quinone pool in a purple non-sulfur bacterium. Scientific Reports 9:12638 DOI 10.1038/s41598-019-49079-z.

Auman AJ, Stolyar S, Costello AM, Lidstrom ME. 2000. Molecular characterization of methanotrophic isolates from freshwater lake sediment. Applied and Environmental Microbiology 66(12):5259–5266 DOI 10.1128/AEM.66.12.5259-5266.2000.

Beck DAC, Kalyuzhnaya MG, Malfatti S, Tringe SG, Glavina del Rio T, Ivanova N, Lidstrom ME, Chistoserdova L. 2013. A metagenomic insight into freshwater methane-utilizing communities and evidence for cooperation between the Methylococcaceae and the Methylophilaceae. PeerJ 1:e23 DOI 10.7717/peerj.23.

Bizukojc M, Dietz D, Sun J, Zeng A. 2010. Metabolic modelling of syntrophic-like growth of a 1, 3-propanediol producer, Clostridium butyricum, and a methanogenic archeon, Methanosarcina mazei, under anaerobic conditions. Bioprocess Biosyst Eng 33:507–523 DOI 10.1007/s00449-009-0359-0.

Boden R, Cunliffe M, Scanlan J, Moussard H, Kits KD, Klotz MG, Jetten MS, Vuilleumier S, Han J, Peters L, Mikhailova N, Tshima H, Tapia R, Kyrpides N, Ivanova N, Pagani I, Cheng JF, Goodwin L, Han C, Hauser L, Land ML, Lapidus A, Lucas S, Pitluck S, Woyke T, Stein L, Murrell JC. 2011. Complete genome sequence of the aerobic marine methanotroph Methylocodons methanica MC09. Journal of Bacteriology 193:7001–7002 DOI 10.1128/JB.06267-11.

Bordel S, Rodriguez Y, Hakobyan A, Rodriguez E, Lebrero R, Munoz R. 2019a. Genome scale metabolic modeling reveals the metabolic potential of three Type
II methanotrophs of the genus Methylocystis. Metabolic Engineering 54:191–199 DOI 10.1016/j.ymben.2019.04.001.

Bordel S, Rojas A, Munoz R. 2019b. Reconstruction of a genome scale metabolic model of the polyhydroxybutyrate producing methanotroph methylocystis parvus OBBP. Microb Cell Fact 18:104 DOI 10.1186/s12934-019-1154-5.

Borenstein E, Feldman MW. 2009. Topological signatures of species interactions in metabolic networks. Journal of Computational Biology 16(2):191–200 DOI 10.1089/cmb.2008.06TT.

Bussmann I, Rahalkar M, Schink B. 2006. Cultivation of methanotrophic bacteria in opposing gradients of methane and oxygen. FEMS Microbiology Ecology 56:331–344 DOI 10.1111/j.1574-6941.2006.00076.x.

Cai Y, Zheng Y, Bodelier PL, Conrad R, Jia Z. 2016. Conventional methanotrophs are responsible for atmospheric methane oxidation in paddy soils. Nature Communications 7:11728 DOI 10.1038/ncomms11728.

Carere CR, McDonald B, Peach HA, Greening C, Gapes DJ, Collet C, Stott MB. 2019. Hydrogen oxidation influences glycogen accumulation in a verrucomicrobial methanotroph. Frontiers in Microbiology 10:1873 DOI 10.3389/fmicb.2019.01873.

Caspi R, Billington R, Ferrer I, Foerster H, Fulcher CA, Keseler IM, Kothari A, Krummenacker M, Latendresse M, Mueller LA, Ong Q, Paley S, Subhraveti P, Weaver DS, Karp PD. 2016. The MetaCyc database of metabolic pathways and enzymes and the BioCyc collection of pathway/genome databases. Nucleic Acids Research 44:D471–D480 DOI 10.1093/nar/gkv1164.

Chan SHJ, Simons MN, Maranas CD. 2017. SteadyCom: predicting microbial abundances while ensuring community stability. PLOS Computational Biology 13(5):e1005539 DOI 10.1371/journal.pcbi.1005539.

Chen Z, Lee WR, Chang SH. 1991. Role of aspartic acid 38 in the cofactor specificity of Drosophila alcohol dehydrogenase. European Journal of Biochemistry 202:263–267 DOI 10.1111/j.1432-1327.1991.tb16371.x.

Christensen EM, Patel SM, Korasick DA, Campbell AC, Krause KL, Becker DF, Tanner JJ. 2017. Resolving the cofactor binding site in the proline biosynthetic enzyme human pyrroline-5-carboxylate reductase 1. Journal of Biological Chemistry 292(17):7233–7243 DOI 10.1074/jbc.M117.780288.

Costa CMV, Dijkema C, Stams A. 2001. The effect of oxygen on methanol oxidation by an obligate methanotrophic bacterium studied by in vivo (13)C nuclear magnetic resonance spectroscopy. The Journal of Industrial Microbiology and Biotechnology 26:9–14 DOI 10.1038/sj/jim/7000075.

Cui M, Ma A, Qi H, Zhuang X, Zhuang G. 2015. Anaerobic oxidation of methane: an active microbial process. Microbiologyopen 4:1–11 DOI 10.1002/mbo3.232.

De la Torre A, Metivier A, Chu F, Laurens LM, Beck DA, Pienkos PT, Lidstrom ME, Kalyuzhnaya MG. 2015. Genome-scale metabolic reconstructions and theoretical investigation of methane conversion in Methylomicrobium buryatense strain 5G(B1). Microb Cell Fact 14:188 DOI 10.1186/s12934-015-0377-3.
De Roy K, Marzorati M, Vanden Abbeele P, Van de Wiele T, Boon N. 2014. Synthetic microbial ecosystems: an exciting tool to understand and apply microbial communities. *Environmental Microbiology* 16:1427–1481 DOI 10.1111/1462-2920.12343.

Dean L. 2012. Methylene tetrahydrofolate reductase deficiency. In: Pratt VM, McLeod HL, Rubinstein WS, Scott SA, Dean LC, Kattman BL, Malheiro AJ, eds. *Medical genetics summaries* [Internet]. Bethesda, MD: National Center for Biotechnology Information (US), 419–422.

Dekker EE, Lane RS, Shapley A. 1971. 2-Keto-4-hydroxybutyrate aldolase. Identification as 2-keto-4-hydroxyglutarate aldolase, catalytic properties, and role in the mammalian metabolism of L-homoserine. *Biochemistry* 10:1353–1364 DOI 10.1021/bi00784a013.

Elbourne LD, Tetu SG, Hassan KA, Paulsen IT. 2017. TransportDB 2.0: a database for exploring membrane transporters in sequenced genomes from all domains of life. *Nucleic Acids Research* 45:D320–D324 DOI 10.1093/nar/gkw1068.

Eram MS, Ma K. 2013. Decarboxylation of pyruvate to acetaldehyde for ethanol production by hyperthermophiles. *Biomolecules* 3:578–596 DOI 10.3390/biom3030578.

Feist AM, Christopher SH, Reed JL, Krummenacker M, Joyce AR, Karp PD, Broadbelt LJ, Hatzimanikatis V, Palsson BO. 2007. A genome-scale metabolic reconstruction for *Escherichia coli* K-12 MG1655 that accounts for 1260 ORFs and thermodynamic information. *Molecular Systems Biology* 3:121 DOI 10.1038/msb4100155.

Flamholz A, Noor E, Bar-Even A, Milo R. 2012. eQuilibrator—the biochemical thermodynamics calculator. *Nucleic Acids Research* 40:D770–D775 DOI 10.1093/nar/gkr874.

Freilich S, Kreimer A, Elhanan B, Yosef N, Sharan R, Gophna U, Ruppin E. 2009. Metabolic-network-driven analysis of bacterial ecological strategies. *Genome Biology* 10(6):R61 DOI 10.1186/gb-2009-10-6-r61.

Frey E. 2010. Evolutionary game theory: theoretical concepts and applications to microbial communities. *Physica A: Statistical Mechanics and its Applications* 389:4265–4298 DOI 10.1016/j.physa.2010.02.047.

Fu Y, Li Y, Lidstrom M. 2017. The oxidative TCA cycle operates during methanotrophic growth of the Type I methanotroph Methylomonas bursmarense 5GB1. *Metabolic Engineering* 42:43–51 DOI 10.1016/j.menb.2017.05.003.

Galand PE, Pereira O, Hochart C, Auguet JC, Debroas D. 2018. A strong link between marine microbial community composition and function idea of functional redundancy. *The ISME Journal* 12:2470–2478 DOI 10.1038/s41396-018-0158-1.

Hadfield A, Kryger G, Ouyang J, Petsko GA, Ringe D, Viola R. 1999. Structure of aspartate–β-semialdehyde dehydrogenase from *Escherichia coli*, a Key Enzyme in the Aspartate Family of Amino Acid Biosynthesis. *Journal of Molecular Biology* 289:991–1002 DOI 10.1006/jmbi.1999.2828.

Hanson RS, Hanson TE. 1996. Methanotrophic bacteria. *Microbiological Reviews* 60(2):439–471 DOI 10.1002/0471263397.env316.
He H, Noor E, Ramos-Parra PA, Garcia-Valencia LE, Patterson JA, Diaz de la Garza RI, Hanson AD, Bar-Even A. 2020. in vivo rate of formaldehyde condensation with tetrahydrofolate. Metabolites 10(2):65 DOI 10.3390/metabo10020065.

He L, Fu Y, Lidstrom ME. 2019. Quantifying methane and methanol metabolism of methylotuvimicrobium buryatense 5gb1c under substrate limitation. mSystems 4(6):e00748-19 DOI 10.1128/mSystems.00748-19.

Henry CS, Bernstein HC, Weisenhorn P, Taylor RC, Lee J-Y, Zucker J, Song H-S. 2016. Microbial community metabolic modeling: a community data-driven network reconstruction. Journal of Cellular Physiology 231:2339–2345 DOI 10.1002/jcp.25428.

Henry CS, DeJongh M, Best AA, Frybarger PM, Linsay B, Stevens RL. 2010. High-throughput generation, optimization and analysis of genome-scale metabolic models. Nature Biotechnology 28:977–982 DOI 10.1038/nbt.1672.

Hernandez ME, Beck DAC, Lidstrom ME, Chistoserdova L. 2015. Oxygen availability is a major factor in determining the composition of microbial communities involved in methane oxidation. PeerJ 3:e801 DOI 10.7717/peerj.801.

Hoefman S, Van der Ha D, De Vos P, Boon N, Heylen K. 2012. Miniaturized extinction culturing is the preferred strategy for rapid isolation of fast-growing methane-oxidizing bacteria. Microbial Biotechnology 5:368–378 DOI 10.1111/j.1751-7915.2011.00314.x.

Hou CT, Laskin AI, Patel RN. 1979. Growth and polysaccharide production by methylocystis parvus OBBP on methanol. Applied and Environmental Microbiology 37:800–804 DOI 10.1128/AEM.37.5.800-804.1979.

Houghton JT, Jenkins GJ, Ephraums JJ. 1990. Climate change: the IPCC scientific assessment. Report prepared for Intergovernmental Panel on Climate Change by Working Group I. New York, USA.

Hutter B, Singh M. 1999. Properties of the 40 kDa antigen of Mycobacterium tuberculosis, a functional L-alanine dehydrogenase. Biochemical Journal 343:669–672 DOI 10.1042/bj3430669.

Ishida Y, Shirafuji H, Kida M, Yoneda M. 1969. Studies on the guanosine degrading system in bacterial cell. Agricultural and Biological Chemistry 33:384–390 DOI 10.1080/00021369.1969.10859324.

IPCC. 2013. Summary for policymakers. In: Stocker TF, Qin D, Plattner G-K, Tignor M, Allen SK, Boschung J, Nauels A, Xia Y, Bex V, Midgley PM, eds. Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change. Cambridge, United Kingdom and New York, NY, USA: Cambridge University Press.

Islam MM, Fernando SC, Saha R, Saha R. 2019. Metabolic modeling elucidates the transactions in the rumen microbiome and the shifts upon virome interactions. Frontiers in Microbiology 10:2412 DOI 10.3389/fmicb.2019.02412.

Islam MM, Saha R. 2018. Computational approaches on stoichiometric and kinetic modeling for efficient strain design. New York: Humana Press DOI 10.1007/978-1-4939-7295-1_5.
Islam MM, Thomas VC, Van Beek M, Ahn JS, Alqarzaee AA, Zhou C, Fey PD, Bayles KW, Saha R. 2020. An integrated computational and experimental study to investigate Staphylococcus aureus metabolism. *NPJ Systems Biology and Applications* 6:3 DOI 10.1038/s41540-019-0122-3.

Jia Y, Whalen JK. 2020. A new perspective on functional redundancy and phylogenetic niche conservatism in soil microbial communities. *Pedosphere* 30:18–24 DOI 10.1016/S1002-0160(19)60826-X.

Kai Y, Matsumura H, Izui K. 2003. Phosphoenolpyruvate carboxylase: three-dimensional structure and molecular mechanisms. *Archives of Biochemistry and Biophysics* 414:170–179 DOI 10.1016/s0003-9861(03)00170-x.

Kalyuzhnaya MG, Lamb AE, McTaggart TL, Oshkin IY, Shapiro N, Woyke T, Chistoserdova L. 2015. Draft genome sequences of gammaproteobacterial methanotrophs isolated from lake Washington sediment. *Genome Announcements* 3:e00103-15 DOI 10.1128/genomeA.00103-15.

Kalyuzhnaya MG, Lapidus A, Ivanova N, Copeland AC, McHardy AC, Szeto E, Salamov A, Grigoriev IV, Suciu D, Levine SR, Markowitz VM, Rigoutsos I, Tringe SG, Bruce DC, Richardson PM, Lidstrom ME, Chistoserdova L. 2008. High-resolution metagenomics targets specific functional types in complex microbial communities. *Nature Biotechnology* 26:1029–1034 DOI 10.1038/nbt.1488.

Kalyuzhnaya MG, Yang S, Rozova ON, Smalley NE, Clubb J, Lamb A, Gowda GA, Ratery D, Fu Y, Bringel F, Vuilleumier S, Beck DA, Trotsenko YA, Khmelenina VN, Lidstrom ME. 2013. Highly efficient methane biocatalysis revealed in a methanotrophic bacterium. *Nature Communications* 4:2785 DOI 10.1038/ncomms3785.

Kanchisa M. 2008. KEGG for linking genomes to life and the environment. *Nucleic Acids Research* 36:D480–D484.

Kanchisa M, Goto S. 2000. KEGG: kyoto encyclopedia of genes and genomes. *Nucleic Acids Research* 28:27–30 DOI 10.1093/nar/28.1.27.

Khmelenina VN, Colin Murrell J, Smith TJ, Trotsenko YA. 2018. Physiology and biochemistry of the aerobic methanotrophs. In: Rojo F, ed. *Aerobic utilization of hydrocarbons, oils and lipids*. Cham: Springer International Publishing, 1–25.

Krause SMB, Johnson T, Karunaratne YS, Fu Y, Beck DAC, Chistoserdova L, Lidstrom ME. 2017. Lanthanide-dependent cross-feeding of methane-derived carbon is linked by microbial community interactions. *Proceedings of the National Academy of Sciences of the United States of America* 114(2):358–363 DOI 10.1073/pnas.1619871114.

Kuivila KM, Murray JW, Devol AH, Lidstrom ME, Reimers CE. 1988. Methane cycling in the sediments of Lake Washington. *Limnology and Oceanography* 33:571–581 DOI 10.4319/lo.1988.33.4.0571.

Lehmann L, Keller L. 2006. The evolution of cooperation and altruism—a general framework and a classification of models. *Journal of Evolutionary Biology* 19:1365–1376 DOI 10.1111/j.1420-9101.2006.01119.x.

Lewis N, Schramm G, Bordbar A, Schellenberger J, Andersen M, Cheng J, Patel N, Yee A, Lewis R, Eils R, Konig R, Palsson BO. 2010. Large-scale in silico modeling of
metabolic interactions between cell types in the human brain. *Nature Biotechnology* 28:1279–1285 DOI 10.1038/nbt.1711.

Lieven C, Petersen LAH, Jorgensen SB, Gernaey KV, Herrgard MJ, Sonnenschein N. 2018. A genome-scale metabolic model for methyllococcus capsulatus (bath) suggests reduced efficiency electron transfer to the particulate methane monooxygenase. *Frontiers in Microbiology* 9:2947 DOI 10.3389/fmicb.2018.02947.

Lin X, Tfaily MM, Green SJ, Steinweg JM, Chanton P, Invittaya A, Chanton JP, Cooper W, Schadt C, Kostka JE. 2014. Microbial metabolic potential for carbon degradation and nutrient (nitrogen and phosphorus) acquisition in an ombrotrophic peatland. *Applied and Environmental Microbiology* 80:3531–3540 DOI 10.1128/AEM.00206-14.

Louca S, Polz MF, Mazel F, Albright MBN, Huber JA, O’Connor MI, Ackermann M, Hahn AS, Srivastava DS, Crowe SA, Doebeli M, Parfrey LW. 2018. Function and functional redundancy in microbial systems. *Nature Ecology & Evolution* 2:936–943 DOI 10.1038/s41559-018-0519-1.

Magnúsdóttir S, Heinken A, Kutt L. 2017. Generation of genome-scale metabolic reconstructions for 773 members of the human gut microbiota. *Nature Biotechnology* 35:81–89 DOI 10.1038/nbt.3703.

Mahadevan R, Schilling C. 2003. Models., The effects of alternate optimal solutions in constraint-based genome-scale metabolic. *Metabolic Engineering* 5:264–276 DOI 10.1016/j.ymben.2003.09.002.

Maranas CD, Zomorrodi AR. 2016. *Optimization Methods in Metabolic Networks*. New Jersey, USA: Wiley.

Mohammadi SS, Schmitz RA, Pol A, Berben T, Jetten MSM, Opden Camp HJM. 2019. The acidophilic methanotroph methylacidimicrobium tartarophylax 4ac grows as autotroph on H2 under microoxic conditions. *Frontiers in Microbiology* 10:2352 DOI 10.3389/fmicb.2019.02352.

Nadell CD, Foster KR, Xavier JB. 2010. Emergence of spatial structure in cell groups and the evolution of cooperation. *PLOS Computational Biology* 6:e1000716 DOI 10.1371/journal.pcbi.1000716.

Naizabekov S, Lee YE. 2020. Genome-scale metabolic model reconstruction and in silico investigations of methane metabolism in methylosinus trichosporium OB3b. *Microorganisms* 8(3):437 DOI 10.3390/microorganisms8030437.

Nguyen NL, Yu WJ, Gwak JH, Kim SJ, Park SJ, Herbold CW, Kim JG, Jung MY, Rhee SK. 2018. Genomic insights into the acid adaptation of novel methanotrophs enriched from acidic forest soils. *Frontiers in Microbiology* 9:1982 DOI 10.3389/fmicb.2018.01982.

Nisbet EG, Manning MR, Dlugokencky EJ, Fisher RE, Lowry D, Michel SE, Myhre CL, Platt SM, Allen G, Bousquet P, Brownlow R, Cain M, France JL, Hermansen O, Hossaini R, Jones AE, Levin I, Manning AC, Myhre G, Pyle JA, Vaughn BH, Warwick NJ, White JWC. 2019. Very strong atmospheric methane growth in the 4 years 2014–2017: implications for the paris agreement. *Global Biogeochemical Cycles* 33(3):318–342 DOI 10.1029/2018GB006009.
Orata FD, Kits KD, Stein LY. 2018. Complete genome sequence of methylomonas denitrificans strain FJG1, an obligate aerobic methanotroph that can couple methane oxidation with denitrification. Genome Announcements 6(17):e00276-18 DOI 10.1128/genomeA.00276-18.

Orth JD, Thiele I, Palsson BO. 2010. What is flux balance analysis? Nature Biotechnology 28:245–248 DOI 10.1038/nbt.1614.

Oshkin IY, Beck DA, Lamb AE, Tchesnokova V, Benuska G, McTaggart TL, Kalyuzhnaya MG, Dedysn SN, Lidstrom ME, Chistoserdova L. 2015. Methane-fed microbial microcosms show differential community dynamics and pinpoint taxa involved in communal response. ISME Journal 9:1119–1129 DOI 10.1038/ismej.2014.203.

Peyraud R, Schneider K, Kiefer P, Massou S, Vorholt JA, Portais JC. 2011. Genom-scale reconstruction and system level investigation of the metabolic network of Methylobacterium extorquens AM1. BMC Systems Biology 5:189 DOI 10.1186/1752-0509-5-189.

Schellenberger J, Lewis N, Palsson BO. 2011. Elimination of thermodynamically infeasible loops in steady-state metabolic models. Biophysical Journal 100:544–553 DOI 10.1016/j.bpj.2010.12.3707.

Schomburg D, Stephan D. 1995. 2-Hydroxy-3-oxopropionate reductase. Enzyme Handbook. In: Enzyme Handbook 9. Berlin, Heidelberg: Springer DOI 10.1007/978-3-642-85200-8_60.

Schroeder WL, Harris SD, Saha R. 2020. Computation-driven analysis of model polyextremotolerant fungus Exophiala dermatitidis: defensive pigment metabolic costs and human applications. iScience 24:100980 DOI 10.1016/j.isci.2020.100980.

Schroeder WL, Saha R. 2019. A computational framework to study the primary lifecycle metabolism of Arabidopsis thaliana. bioRxiv 12:44 DOI 10.1101/761189.

Shoaie S, Nielsen J. 2014. Elucidating the interactions between the human gut microbiota and its host through metabolic modeling. Frontiers in Genetics 5:86 DOI 10.3389/fgene.2014.00086.

Shou W, Ram S, Vilar JMG. 2007. Synthetic cooperation in engineered yeast populations. Proceedings of the National Academy of Sciences of the United States of America 104:1877–1882 DOI 10.1073/pnas.0610575104.

Smith TJ, Trotsenko YA, Murrell JC. 2010. In: Timmis KN, ed. Physiology and biochemistry of the aerobic methane oxidizing bacteria BT—Handbook of hydrocarbon and lipid microbiology. Berlin, Heidelberg: Springer Berlin Heidelberg, 765–779 DOI 10.1007/978-3-540-77587-4_58.

Soni BK, Conrad J, Kelley RL, Srivastava VJ. 1998. Effect of temperature and pressure on growth and methane utilization by several methanotrophic cultures. Applied Biochemistry and Biotechnology 70:729–738 DOI 10.1007/BF02920184.

Stolyar S, Van Dien S, Hillesland K, Pinel N, Lie T, Leigh J, Stahl D. 2007. Metabolic modeling of a mutualistic microbial community. Molecular Systems Biology 3:92 DOI 10.1038/msb4100131.

Strong PJ, Xie S, Clarke WP. 2015. Methane as a resource: can the methanotrophs add value? Environmental Science and Technology 49:4001–4018 DOI 10.1021/es504242n.
Sugioka K, Nakano M, Naito I, Tero-Kubota S, Ikegami Y. 1988. Properties of a coenzyme, pyrroloquinoline quinone: generation of an active oxygen species during a reduction-oxidation cycle in the presence of NAD(P)H and O2. *Biochimica et Biophysica Acta/General Subjects* **964**:175–182 DOI 10.1016/0304-4165(88)90164-x.

Svenning MM, Hestnes AG, Wartiainen I, Stein LY, Klotz MG, Kalyuzhnaya MG, Spang A, Bringel F, Vuilleumier S, Lajus A, Médigue C, Bruce DC, Cheng J-F, Goodwin I, Ivanova N, Han J, Han CS, Hauser L, Held B, Land ML, Lapidus A, Lucas S, Nolan M, Pitluck S, Woyke T. 2011. Genome sequence of the Arctic methanotroph Methylobacter tundripaludum SV96. *Journal of Bacteriology* **193**:6418–6419 DOI 10.1128/JB.05380-11.

Szklarczyk D, Morris JH, Cook H, Kuhn M, Wyder S, Simonovic M, Santos A, Doncheva NT, Roth A, Bork P, Jensen LJ, Mering Cvon. 2017. The STRING database in 2017: quality-controlled protein-protein association networks, made broadly accessible. *Nucleic Acids Research* **45**:D362–D368 DOI 10.1093/nar/gkw937.

Temperton B, Giovannoni S. 2012. Metagenomics: microbial diversity through a scratched lens. *Current Opinion in Microbiology* **15**:605–612 DOI 10.1016/j.mib.2012.07.001.

Theisen AR, Murrell JC. 2005. Facultative methanotrophs revisited. *Journal of Bacteriology* **187**:4303–4305 DOI 10.1128/JB.187.13.4303-4305.2005.

UniProt Consortium. 2018. UniProt: a worldwide hub of protein knowledge. *Nucleic Acids Research* **47**:D506–D515 DOI 10.1093/nar/gky1049.

Vallino JJ. 2003. Modeling microbial consortia as distributed metabolic networks. *The Biological Bulletin* **204**:174–179 DOI 10.2307/1543554.

Van Teeseling MC, Pol A, Harhangi HR, Van der Zwart S, Jetten MS, Opden Camp HJ, Van Niftrik L. 2014. Expanding the verrucomicrobial methanotrophic world: description of three novel species of Methylacidimicrobium gen. nov. *Applied and Environmental Microbiology* **80**:6782–6791 DOI 10.1128/AEM.01838-14.

Wartiainen I, Hestnes AG, McDonald IR, Svenning MM. 2006. Methylobacter tundripaludum sp. nov. a methane-oxidizing bacterium from Arctic wetland soil on the Svalbard islands, Norway (78 degrees N). *International Journal of Systematic and Evolutionary Microbiology* **56**:109–113 DOI 10.1099/ijs.0.63728-0.

Whalen SC, Reeburgh WS. 1990. Consumption of atmospheric methane by tundra soils. *Nature* **346**:160–162 DOI 10.1038/346160a0.

Yu Z, Beck DAC, Chistoserdova L. 2017. Natural selection in synthetic communities highlights the roles of methylococcaceae and methylphilaceae and suggests differential roles for alternative methanol dehydrogenases in methane consumption. *Frontiers in Microbiology* **8**:2392 DOI 10.3389/fmicb.2017.02392.

Yu Z, Krause SMB, Beck DAC, Chistoserdova L. 2016. A synthetic ecology perspective: how well does behavior of model organisms in the laboratory predict microbial activities in natural habitats? *Frontiers in Microbiology* **7**:946 DOI 10.3389/fmicb.2016.00946.
Yvon-Durocher G, Allen AP, Bastviken D, Conrad R, Gudasz C, St-Pierre A, Thanh-Duc N, Del Giorgio PA. 2014. Methane fluxes show consistent temperature dependence across microbial to ecosystem scales. *Nature* **507**:488–491 DOI 10.1038/nature13164.

Zengler K. 2009. Central role of the cell in microbial ecology. *Microbiology and Molecular Biology Reviews* **73**:712–729 DOI 10.1128/MMBR.00027-09.

Zengler K, Palsson BO. 2012. A road map for the development of community systems (CoSy) biology. *Nature Reviews Microbiology* **10**:366–372 DOI 10.1038/nrmicro2763.

Zhao S-J, Hanson RS. 1984. Isolate 761M: a new type i methanotroph that possesses a complete tricarboxylic acid cycle. *Applied and Environmental Microbiology* **48**:1237–1242 DOI 10.1128/AEM.48.6.1237-1242.1984.

Zheng Y, Huang J, Zhao F, Chistoserdova L. 2018. Physiological Effect of XoxG(4) on Lanthanide-Dependent Methanotrophy. *Mbio* **9**(2):e02430-17 DOI 10.1128/mBio.02430-17.

Zhuang K, Izallalen M, Mouser P, Richter H, Risso C, Mahadevan R, Lovley DR. 2011. Genome-scale dynamic modeling of the competition between Rhodoferax and Geobacter in anoxic subsurface environments. *ISME Journal* **5**:305–316 DOI 10.1038/ismej.2010.117.

Zomorrodi AR, Islam MM, Maranas CD. 2014. D-OptCom: dynamic multi-level and multi-objective metabolic modeling of microbial communities. *ACS Synthetic Biology* **3**(4):247–257 DOI 10.1021/sb4001307.

Zomorrodi AR, Maranas CD. 2012. OptCom: a multi-level optimization framework for the metabolic modeling and analysis of microbial communities. *PLOS Computational Biology* **8**(2):e1002363 DOI 10.1371/journal.pcbi.1002363.