Microbial Community Metabolic Modeling: A Community Data-Driven Network Reconstruction

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Metabolic network modeling of microbial communities provides an in-depth understanding of community-wide metabolic and regulatory processes. Compared to single organism analyses, community metabolic network modeling is more complex because it needs to account for interspecies interactions. To date, most approaches focus on reconstruction of high-quality individual networks so that, when combined, they can predict community behaviors as a result of interspecies interactions. However, this conventional method becomes ineffective for communities whose members are not well characterized and cannot be experimentally interrogated in isolation. Here, we tested a new approach that uses community-level data as a critical input for the network reconstruction process. This method focuses on directly predicting interspecies metabolic interactions in a community, when axenic information is insufficient. We validated our method through the case study of a bacterial photoautotroph–heterotroph consortium that was used to provide data needed for a community-level metabolic network reconstruction. Resulting simulations provided experimentally validated predictions of how a photoautotrophic cyanobacterium supports the growth of an obligate heterotrophic species by providing organic carbon and nitrogen sources.

The role of microbial communities has been increasingly recognized in a multitude of scientific disciplines, including: soil ecology, environmental engineering, agriculture, food safety, and human health. Researchers have been keen to reveal the organizational principles in microbial communities and to predict their response to environmental cues, which requires a mechanistic understanding of interspecies interactions. Such understandings are also essential for designing and engineering microbial ecosystems for controllable outputs (Bernstein and Carlson, 2012; Lindemann et al., 2016). Advancements in metatranscriptomics and -omics analyses have significantly expanded the volume of biological data at hand, as well as our collective biological knowledge. However, experimental determination of diverse forms of microbial interactions in a community still remains a challenge. As a complementary tool, metabolic network analysis provides comprehensive predictive capabilities that can serve as ab initio hypotheses on cross-species metabolite exchanges (Song et al., 2014; Biggs et al., 2015; Cardona et al., 2016).

Reconstruction of reliable microbial community networks is a challenging task. Even in the single-species case, genome-scale metabolic network reconstruction is an iterative process that takes a substantial period of time (Thiele and Palsson, 2010). Community modeling is a more challenging process due to the increased complexities involving interacting, non-independent species. A conventional practice to build community metabolic networks focuses on the reconstruction of high-quality individual networks so that their combination provides quantitative predictions of metabolic interactions and community behaviors (Shoaie et al., 2015). This approach becomes ineffective; however, if sufficient data required for curating individual networks are not available. This is often the case with environmental communities whose member species are not axenically cultivable or do not grow on defined media,
for which we may lack high-quality complete genome sequences.

To address this issue, we proposed an approach that directly
refines metabolic networks at a community level. Community
metabolic network reconstruction is primarily driven by a
community-level data, which is readily collectible even when
member species are not isolable or individually cultivable. For
testing the proposed approach, we considered a binary
consortium composed of a Thermosynechococcus elongatus BP-I
and Microthrix ruber Strain A (hereafter T. elongatus and
M. ruber, respectively) as a model photoautotroph–heterotroph
consortium. We evaluated the qualities of resulting networks
and the transported substrates de
these enzymes comprises the intracellular metabolic network,
and extension of such protocols to the building of models of
microbial communities, using our case study for illustration.

Protocols of network reconstruction from single

Current sequencing methods are unable to read a whole
genome at a time, so all sequencing protocols first shear DNA
into smaller fragments that the sequencer can read. To
reassemble these fragments, many copies of the genome must be
fragmented. That way, the shreds from one copy might overlap
with the shreds from another copy so that the original DNA
sequence can be reconstructed (Lander and Waterman, 1988).
These contiguous stretches of overlapping fragments are called
contigs. If the sequencing is deep enough, then these contigs can
be assembled into one or more scaffolds that cover the full
genome. Next, we need to know where genes are located and
what their functions are. The most important genes to identify
for a metabolic reconstruction are those that function as
enzymes and transporters. The set of all reactions catalyzed by
these enzymes comprises the intracellular metabolic network,
and the transported substrates define the interface with the
extracellular environment. The final step—formulation of a
biomass synthesis equation—creates a testable metabolic
model. If the metabolic network can produce all the essential
compounds necessary for biomass from a set of extracellular
metabolites, this is a prediction that the organism can grow in
that nutrient condition. If we remove a reaction or transporter
from the network, and it is no longer capable of producing
biomass, then this is a prediction that the gene encoding that
function is essential. By proceeding in this manner, metabolic
network predictions of biomass production can be validated
against growth or no-growth phenotype observations.

Procedures for network reconstruction of single genomes were
summarized in Table S1.

Strategies for community network construction

Community networks can be constructed from individual
species’ genomes in many different forms. At the simplest level,
we can take a mixed-bag (or gene-soup) approach by treating a
microbial community as a single supra-organism (Abubucker
et al., 2012; Faria et al., 2016). The metabolic pathways and
transmembrane transport reactions, resulting from all
members, are combined by ignoring species boundaries. The mixed-bag network has one cytosolic compartment and one
extracellular compartment, analogous to a single species
prokaryotic network. The primary usage of the mixed-bag
approach is to analyze environment-community interactions.
Construction of a mixed-bag network requires the full genome
sequences of all member species or a deep metagenome sequence. The growth conditions of the microbial community
are also useful but species level resolution is not required.

Prediction of cross-species metabolic interactions requires
species-resolved network modeling. This can be achieved by
treating species networks as compartments of a community,
the structure of which is similar to those of eukaryotic
metabolic networks where inter- and intra-compartmental
activities are assumed to be in a quasi-steady state. To predict
non-steady-state microbial interactions, we require multi-
species dynamic models that account for kinetics of nutrient
uptake and metabolite production for individual species. By
overlaying species-resolved kinetic expressions we can
interrogate the dynamic interplay between members in a
community. Multi-species metabolic modeling (for a
compartmentalized or kinetically connected model) starts with
the construction of metabolic models for the individual species
comprising the community. The methods of obtaining species-
resolved networks depend on the nature of the microbial
community involved. For a defined community composed of
isolated and cultivable microbes, we can use a single-organism
pipeline like the ModelSEED (Henry et al., 2010), RAVEN
(Agren et al., 2013), or COBRA (Thiele and Palsson, 2010)
to produce a separate model of each species. In contrast, for an
undefined community derived from natural environment, we
need a process to build species-level metabolic models from
the metagenomics data by separating out the assembled contigs
into species-level bins and subsequently performing gene
calling, annotation, and model building on each bin. In Figure 1,
we illustrated the three alternative approaches for building
microbial community metabolic models described in this
section.

A binary consortium to model

photoautotroph–heterotroph communities

With a focus on steady state analyses (i.e., methods shown in
Fig. 1B and C), we will explore the advantages and
disadvantages of the various strategies for constructing
microbial community models through the use of a model
microbial consortium, comprised of the photoautotrophic
cyanobacterium T. elongatus and an obligate aerobic
heterotroph M. ruber. Cyanobacteria are essential
contributors to primary productivity in natural ecosystems and
are frequently associated with dependent populations of
heterotrophic bacteria. These phototrophic-heterotrophic
communities are of keen interest to biologists seeking to
uncover mechanisms of interaction that drive global
biogeochemical cycles (e.g., involving C, N, and O2). They are
also attractive bioprocess hosts because of their inherent
robustness, diverse metabolisms, and fast growth rates. We
selected these specific taxa to demonstrate community
modeling because of the obligate dependency of M. ruber on
T. elongatus, when cultured under autotrophic conditions.
Depending on the environment, M. ruber relies upon
cyanobacterial derived organic-carbon, fixed nitrogen,
vitamins, and O2.

T. elongatus BP-I is a thermophilic, unicellular cyanobacterium
that has been studied extensively (Onai et al., 2004; Zhang et al.,
2005; Araki and Kino, 2008; Abed et al., 2009; Eberly and Ely,
2012). Hence, it is a model cyanobacterium and has a completely
sequenced and annotated genome (Nakamura et al., 2002). T.
elongatus was isolated from a cyanobacterial mat environment
near Beppu, Japan (Yamaoka et al., 1978). M. ruber Strain A is an
aerobic, heterotrophic, thermophile that was isolated from an
enrichment culture originally sampled from the cyanobacterial
mat inhabiting Octopus Spring in Yellowstone National Park
(WY, USA) (Thiel et al., 2015). It shares 98.6% nucleotide
identity to the 16S rRNA gene sequence of M. ruber DSM 1279
(Logino et al., 1984) and displays strong functional relatedness
with regard to central carbon and energy metabolism genes
(Thiel et al., 2015). Similar to M. ruber DSM 1279, Strain A lacks
transportable metabolites, and gap conditions has been common in both organisms (orange). Solid and open circles indicate extracellular and intracellular metabolites; solid line boxes represent compartments outside of which the quasi-steady state assumption may not hold, and dashed line boxes represent compartments outside of which the steady-state assumption continues to hold. In A, \( r_1 \) s and \( r_2 \) s denote exchange rates that are kinetically modeled. The nitrate assimilatory pathway. Hence, it depends upon reduced N-sources produced by \( T. \) elongatus when growing in an environment that is depleted of fixed/reduced nitrogen.

Reconstruction of single species metabolic models using the KBase platform

Before we can construct a community model of \( T. \) elongatus and \( M. \) ruber, we first need to construct individual metabolic models for each of these species. The KBase platform offers two alternative pipelines for the generation and refinement of metabolic models from sequence data as described in the following. One pipeline enables the direct reconstruction of a new draft model from a genome sequence, consisting of four steps: (i) import of the genome into KBase; (ii) structural and functional annotation of the genome using the RAST annotation algorithm (Aziz et al., 2008); (iii) reconstruction of a draft metabolic model using the ModelSEED algorithm (Henry et al., 2010); and (iv) gapfilling the model on a specified growth condition using optimization-based approaches (Dreyfuss et al., 2013; Latendresse, 2014). The final gapfilling step can be performed iteratively on multiple media conditions if a capacity for growth under multiple conditions has been confirmed experimentally.

We applied this pipeline to build a model of the heterotrophic member from our example community, \( M. \) ruber (https://narrative.kbase.us/narrative/ws.13807.obj.1). We created three versions of our \( M. \) ruber model: (i) an ungapfilled version (\( dMr729 \)); (ii) a version gapfilled on a rich LB media in which \( M. \) ruber is known to grow (\( lbMr729 \)); and (iii) a version gapfilled on glucose minimal media (\( mmMr729 \)), on which it is known not to grow.

Table I shows the number of reactions, genes, transporters, transportable metabolites, and gapfilled reactions contained in each model. Flux balance analysis (FBA) of the model gapfilled on LB media predicted that this species lacks a capacity to synthesize several vitamins/cofactors (niacin, pantothenate, riboflavin, and heme) and amino acids (proline, valine, isoleucine, lysine, and histidine). These results are largely (albeit not totally) consistent with experimental data on the nutrients required for culturing \( M. \) ruber Strain A. While this result is encouraging, it is also well-established that draft models constructed directly from genome sequence data typically require substantial curation before they can be truly predictive (Thiele and Palsson, 2010).

Fortunately, it is not always necessary to start from scratch when building a model from a genome sequence. KBase offers an alternative pipeline in cases where a curated model already exists for a close relative of the genome of interest. This pipeline consists of five steps: (i) import the genome; (ii) annotate the genome; (iii) import a curated model and genome of a closely related species; (iv) perform bi-directional all-versus-all BLAST comparison of the genes in the primary genome and the well-curated relative; (v) optionally gapfill the model on a specified growth conditions using optimization-based approaches. We applied this second pipeline in KBase to build a model of \( T. \) elongatus based on the previously published \( jN678 \) model of \( S. \) elongatus (Nogales et al., 2012) (https://narrative.kbase.us/narrative/ws.13806.obj.1). As it is known that \( T. \) elongatus is autotrophic, we only generated two versions of this model (Table I): (i) an ungapfilled model (\( dTe583 \)) and (ii) a model gapfilled on autotrophic media (\( aTe583 \)).

Reconstruction of consortium metabolic models using the KBase platform

Now that we have single species models constructed for both of the members of our example consortium, we can combine these models into either a mixed-bag community model (Fig. 1C) or a compartmentalized community model (Fig. 1B). As before, we applied the KBase platform to build both model types. When constructing a mixed-bag model, the reactions from each single-species model are merged together into a

Table I. Statistics on all individual and community metabolic models constructed

| Model       | Reactions | Genes | Transporters | Transported metabolites | Gapfilled reactions |
|-------------|-----------|-------|--------------|-------------------------|---------------------|
| \( dMr729 \) | 1163      | 729   | 68           | 61                      | 0                   |
| \( lbMr729 \) | 1253      | 729   | 76           | 71                      | 95                  |
| \( mmMr729 \) | 1257      | 729   | 71           | 65                      | 98                  |
| \( dTe583 \) | 889       | 583   | 88           | 86                      | 0                   |
| \( aTe583 \) | 917       | 583   | 89           | 87                      | 28                  |
| \( mbMrTe1312 \) | 1707      | 1312  | 124          | 93                      | 74                  |
| \( preMrTe1312 \) | 2174      | 1312  | 160          | 91                      | 126                 |
| \( mixMrTe1312 \) | 2175      | 1312  | 168          | 100                     | 128                 |
| \( postMrTe1312 \) | 2168      | 1312  | 164          | 98                      | 123                 |
single intracellular compartment. Gene associations are preserved from the original models, and they are merged in the cases where the same reaction appears in multiple models. The biomass objective functions from all individual models are combined together into a single consortium biomass objective function by summing the stoichiometric coefficients from each individual biomass reaction, and rescaling the coefficients by the number of models being combined. Like individual models in KBase, mixed-bag models can be gapfilled in a specified growth condition. We applied this capability in KBase to combine our ungapfilled M. ruber and T. elongatus models into a mixed-bag model (https://narrative.kbase.us/narrative/ws.13838.obj.1), which we subsequently gapfilled in autotrophic media. We call this model mbMrTe1312 (Table I). Here we see that the mixed-bag model has fewer total reactions than the sum of the two constituent models. Additionally, the mixed-bag model required more gapfilled reactions than either single model, but fewer than both models combined.

Next we applied tools in KBase to construct a compartmentalized model of our model consortium. When constructing a compartmentalized community model in KBase, the two individual models are still combined into a single model, but in this case, the reactions from each individual model are integrated into distinct cellular compartments in the new consortium model. These two compartments share a single common extracellular compartment that permits the exchange of metabolites between the species models. As with all models in KBase, compartmentalized community models can be gapfilled. In this case, gapfilling is distinctive in that the entire database of candidate reactions is added to each separate species compartment in the consortium model, meaning the number of candidate reactions considered by the gapfilling scales with the number of species models combined in the consortium model, which in turn results in a greater complexity to the gapfilling formulation and a longer run-time in finding an optimal solution. However, this formulation offers a decisive advantage over gapfilling of individual models, as the algorithm now considers the possibility that metabolic functions may be delegated among the species comprising the consortium, and as such, will propose solutions that involve interactions between species in the cases where such interactions result in the greatest parsimony. This is particularly valuable for gapfilling models of species that cannot be grown in isolation, because the only phenotypic data available for these species involves multiple organisms operating in concert. We now explore the impact of applying a range of gapfilling strategies when constructing a compartmentalized model of a microbial consortium.

Alternative gapfilling strategies in reconstruction of consortium metabolic models

Because it is possible to gapfill models on a variety of growth conditions, and because it is possible to gapfill models both before and after they have been merged together into a compartmentalized consortium model, we are presented with a wide range of potential strategies for constructing a consortium model. We summarize the many potential options with three fundamental alternative approaches: (i) pre-gapfilling individual models on minimal media prior to merging into a community model (Fig. 2A); (ii) merge ungapfilled individual models into a community model, then gapfill the community model (Fig. 2B); or (iii) pre-gapfill individual models on rich media, merge into a community model, then post gapfill the community model (Fig. 2C).

We applied all three alternative strategies in KBase (https://narrative.kbase.us/narrative/ws.13838.obj.1), resulting in the development of three alternative compartmentalized models of our M. ruber and T. elongatus consortium: (i) pre-gapfilled preMrTe1312; (ii) post-gapfilled postMrTe1312; and (iii) gapfilled mixMrTe1312 (Table II). We then simulated the growth of these three consortium models in autotrophic media using FBA and identified the interspecies interactions predicted by each model (Fig. 3). As expected, the preMrTe1312 model predicts the fewest interactions with only 9 metabolites exchanged, while postMrTe1312 predicts the most interactions, with 15 metabolites exchanged. The mixMrTe1312 model splits the difference, predicting 13 metabolites exchanged. Thus, we see that typically more interactions are predicted when more of the gapfilling is conducted after the individual models have been merged into a community model. This trend occurs because gapfilling conducted on merged community models will consider solutions that involve trophic interactions between species, and where such trophic interactions may actually be occurring, these solutions will typically be lower cost than the non-interactive alternative of adding all biosynthetic pathways to constituent species. We also highlight that the pre-gapfilling approach may be ineffective in a typical situation where some of the members are cultivable only on rich media, because choosing to gapfill their networks on rich media alone may be insufficient to identify microbial interactions that enable community growth.

Of our three compartmentalized community models, the interactions predicted by the postMrTe1312 model also best match our experimental understanding of the dependencies between M. ruber and T. elongatus growing on autotrophic media, although it is worth noting that the pantothenate

![Fig. 2. Alternative gapfilling strategies when constructing compartmentalized consortium metabolic models: (A) individual gapfilling (igf), (B) community-level gapfilling (c.gfl), (C) combination of A and B. N_1 and N_2 denote draft networks of species 1 and 2, respectively.](https://narrative.kbase.us/narrative/ws.13838.obj.1)
TABLE II. Consistency between reaction flux and gene expression in various model versions

| +Flux, +Exp | -Flux, -Exp | +Flux, -Exp | -Flux, +Exp | Accuracy for active genes (%) | Accuracy for inactive genes (%) | Overall accuracy (%) |
|------------|-------------|-------------|-------------|-------------------------------|-------------------------------|---------------------|
| mMr729     | 16.1        | 37.0        | 17.5        | 29.5                          | 35.3                         | 67.9                | 53.1                |
| bMr729     | 25.5        | 15.9        | 5.6         | 53.0                          | 32.5                         | 74.0                | 41.4                |
| aTe583     | 54.5        | 14.9        | 9.4         | 21.2                          | 72.0                         | 61.2                | 69.4                |
| nMrTe1312  | 20.5        | 35.2        | 20.4        | 23.9                          | 46.2                         | 63.3                | 55.7                |
| preMrTe1312| 17.5        | 37.1        | 25.0        | 20.4                          | 46.2                         | 59.7                | 54.6                |
| mixMrTe1312| 17.4        | 37.0        | 25.2        | 20.5                          | 45.9                         | 59.5                | 54.4                |
| postMrTe1312| 17.6       | 37.1        | 25.0        | 20.4                          | 46.3                         | 59.7                | 54.6                |

exchange predicted by the mixMrTe1312 is more physiologically reasonable than the exchange of CoA predicted by postMrTe1312.

Evaluation of community networks in comparison with gene expression profiles

In addition to evaluating our models based on the number and type of gapfilled reactions and the predicted trophic interactions, we can also evaluate the capacity of our models to accurately predict which pathways will be active in our organisms while they are growing together in a consortium. To accomplish this, we gathered RNA-seq data for our M. ruber and T. elongatus consortium growing in autotrophic conditions. We mapped the reads to our individual M. ruber and T. elongatus genes and loaded the expression profile into KBase. We also called each gene “active” or “inactive” based on its expression level (see the next section). Finally, we ran transcriptional FBA (Seaver et al., 2015) to fit the flux predictions made by our models to gene activity computed from our RNA-seq data. This approach attempts to force reactions associated with inactive genes off, while forcing reactions associated with active genes on, without preventing the model from producing biomass. We thus compared our models’ flux predictions with active/inactive gene activity based on gene expression data for enzymes catalyzing reactions in our models.

We ran this analysis on our two gapfilled M. ruber models: mMr729 and bMr729 (Table II). The mMr729 model performed better at predicting active reactions, correctly predicting 35% of reactions associated with actively expressed genes as active (vs. 33% predicted by the model in LB media). However, the bMr729 model performed much better at predicting inactive reactions, correctly predicting 74% of reactions associated with poorly expressed genes as inactive (vs. 68% predicted by the model in glucose minimal media). These results show that minimal media activates too much of the metabolic network, while LB media activates too little. However, these results also show that both models suffer from inaccuracies in their capacity to produce flux profiles that are consistent with observed patterns in gene expression. Much of this inaccuracy is likely due to comparing the flux of a single species model with the gene expression data generated from a community in which it is metabolically dependent on another member. However, some inaccuracy will typically be the case from draft models constructed directly from genome sequence, which was unavoidable in the case of M. ruber, as no published curated model currently exists for this species or any close relatives of this species.

We conducted the same analysis on our T. elongatus aTe583. For this model, we saw significantly improved accuracy compared with the M. ruber model (Table II). Our T. elongatus model correctly predicted 72% of reactions associated with highly expressed genes as active; and it correctly predict 61% of reactions associated with poorly expressed genes as inactive. This result clearly shows the value of starting analysis with a curated model whenever one is available. In this case, the gene expression data are also more directly comparable to the case of a single genome model as T. elongatus is not dependent upon M. ruber and can grow in isolation on the media condition used to generate the expression profile.

Finally, we ran the analysis with our gapfilled mixed-bag model, as well as all three compartmentalized models (Table II). The mixed-bag model performed the best of our community models, but this model type provides no information on potential interactions among the species in our consortium. Among our compartmentalized models, the postMrTe1312 performed the best, although all three compartmentalized models produced very similar levels of performance. We can also see a pathway-by-pathway breakdown of the agreement between model flux and expression data (Fig. 4). This analysis shows that the riboflavin and quinone pathways showed the poorest agreement, indicating potential flaws in the biomass equation of the M. ruber model.

Fig. 3. Interactions between M. ruber and T. elongatus predicted by consortium models.
percentile as the threshold for calling genes “on” or “off.” Genes with expression values higher than this threshold are called “on,” and genes with expression values lower than this threshold are called “off.”

Summary and future directions

Here, we provided a new approach that uses community-level data for microbial community network reconstruction, and we demonstrated tools that implement this approach in a user-friendly manner in the DOE Systems Biology Knowledgebase. Development of this approach was motivated by: (i) community data provides key information on microbial interactions, which are not necessarily obtainable from axenic cultures and (ii) the conventional approaches that require high-quality individual networks are ineffective when member species in a community cannot be sufficiently characterized in isolation. In the case study of a binary consortium, metabolic networks reconstructed using this method led to predictions of phototroph–heterotroph interactions that are consistent with experimental data. Manual curation is a necessary next step for more accurate predictions. Further developments using KBase are in progress to extend the proposed method to more complex communities beyond simple binary consortia.

Acknowledgments

This work was supported by the Genomic Science Program (GSP), Office of Biological and Environmental Research (OBER) and US Department of Energy (DOE) under contract DE-AC02-06CH11357 as a part of the DOE Knowledgebase project (CSH, PW) and as a contribution of the Pacific Northwest National Laboratory (PNNL) Foundational Scientific Focus Area (HCB, RCT, JYL, JZ, HSS). Author HCB is also grateful for support from the Linus Pauling Distinguished Postdoctoral Fellowship, a laboratory directed research and development program at PNNL.

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