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Metabolic Network Analysis and Metatranscriptomics Reveal Auxotrophies and Nutrient Sources of the Cosmopolitan Freshwater Microbial Lineage acl

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ABSTRACT An explosion in the number of available genome sequences obtained through metagenomics and single-cell genomics has enabled a new view of the diversity of microbial life, yet we know surprisingly little about how microbes interact with each other or their environment. In fact, the majority of microbial species remain uncultivated, while our perception of their ecological niches is based on reconstruction of their metabolic potential. In this work, we demonstrate how the “seed set framework,” which computes the set of compounds that an organism must acquire from its environment (E. Borenstein, M. Kupiec, M. W. Feldman, and E. Ruppin, Proc Natl Acad Sci USA 105:14482–14487, 2008, https://doi.org/10.1073/pnas.0806162105), enables computational analysis of metabolic reconstructions while providing new insights into a microbe’s metabolic capabilities, such as nutrient use and auxotrophies. We apply this framework to members of the ubiquitous freshwater actinobacterial lineage acl, confirming and extending previous experimental and genomic observations implying that acl bacteria are heterotrophs reliant on peptides and saccharides. We also present the first metatranscriptomic study of the acl lineage, revealing high expression of transport proteins and the light-harvesting protein actinorhodopsin. Putative transport proteins complement predictions of nutrients and essential metabolites while providing additional support of the hypothesis that members of the acl are photoheterotrophs.

IMPORTANCE The metabolic activity of uncultivated microorganisms contributes to numerous ecosystem processes, ranging from nutrient cycling in the environment to influencing human health and disease. Advances in sequencing technology have enabled the assembly of genomes for these microorganisms, but our ability to generate reference genomes far outstrips our ability to analyze them. Common approaches to analyzing microbial metabolism require reconstructing the entirety of an organism’s metabolic pathways or performing targeted searches for genes involved in a specific process. This paper presents a third approach, in which draft metabolic reconstructions are used to identify compounds through which an organism may interact with its environment. These compounds can then guide more-intensive metabolic reconstruction efforts and can also provide new hypotheses about the specific contributions that microbes make to ecosystem-scale metabolic processes.

KEYWORDS freshwater microbial ecology, metabolism, metagenomics, metatranscriptomics, physiology, systems biology

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Natural microbial communities have central roles in the biosphere, ranging from mediating nutrient cycling to influencing human health and disease (1, 2). However, the majority of microbial species remain uncultivated, a state of affairs that poses a significant challenge to our understanding of their physiology and metabolism. Recent advances in sequencing technology and bioinformatics have enabled assembly and analysis of reference genomes for a wide range of hitherto-uncultured community members from diverse environments (3) that can be used to reconstruct an organism’s metabolism.

Common approaches to metabolic reconstruction involve the comprehensive reconstruction of an organism’s metabolic pathways (4) or a targeted search for genes involved in processes of interest (5). These reconstructions can then be analyzed using manual methods or computational approaches such as flux-balance analysis (FBA) (6). However, FBA-based approaches require a comprehensive understanding of an organism’s growth requirements and biomass composition, information which is often unavailable for uncultivated microorganisms. An alternative approach is to compute an organism’s seed set, representing the set of compounds that the organism cannot synthesize on its own and must exogenously acquire from its environment (e.g., its growth requirements) (7). These compounds may represent both auxotrophies, i.e., the essential metabolites for which biosynthetic routes are missing, and nutrients, i.e., the compounds for which degradation routes but not synthesis routes are present in the genome. The seed set framework offers potential advantages over other reconstruction-based approaches, as identification of seed compounds facilitates a focused analysis by identifying those compounds through which an organism interacts with its environment.

In the present report, we present a computational pipeline to predict seed compounds using metabolic network reconstructions generated from KBase (8). We apply this pipeline to a collection of 36 metagenome-assembled genomes (MAGs) and single-cell genomes (SAGs) from the abundant and ubiquitous freshwater actinobacterial lineage acl, which is thought to have a central role in nutrient cycling in diverse freshwater systems (9–18). The seed compounds predicted by our analysis are in agreement with previous experimental and genomic observations (19–27), confirming the ability of our method to predict an organism’s auxotrophies and nutrient sources.

In particular, we found that members of the acl lineage are auxotrophic for essential vitamins and amino acids and may consume as nutrients a wide array of N-containing compounds (including ammonium, branched-chain amino acids, polyamines, and diand oligopeptides) as well as mono-, poly-, and oligosaccharides. To complement these predictions, and to understand which pathways dominate active metabolism of acl in its natural environment, we conducted an in situ metatranscriptomic analysis of gene expression in the acl lineage. This analysis revealed that the members of the acl lineage express a diverse array of transporters for auxotrophies, nutrients, and other compounds that may contribute to their observed dominance and widespread distribution in a variety of aquatic systems.

RESULTS

Phylogenetic affiliation of acl genomes. We identified 17 SAGs and 19 MAGs from members of the acl lineage (see Table S1 in Data Set S1 in the supplemental material) in a larger set of reference genomes derived from our long-term study sites. A phylogenetic tree of these genomes built using a concatenated alignment of single-copy marker genes is shown in Fig. 1. Previous phylogenetic analyses using 16S rRNA gene sequences showed that the acl lineage can be grouped into 3 distinct monophyletic clades (acl-A, acl-B, and acl-C) and 13 so-called “tribes” (28). In this study, the phylogenetic tree also identified three monophyletic branches, enabling MAGs to be classified to the clade and tribe levels based on the taxonomy of SAGs within each branch (as determined by the 16S rRNA gene sequences that had been either PCR amplified or assembled from the single cell). Note that three MAGs formed a mono-
phyletic group separate from clades acl-A and acl-B; we assume that these genomes belong to clade acl-C as no other acl clades have been identified to date.

Estimated completeness of tribe- and clade-level composite genomes. We constructed composite genomes from multiple SAGs and/or MAGs to partially alleviate the limitations presented by incomplete genomes. To do this, we first estimated the completeness of tribe- and clade-level composite genomes using CheckM (29), which uses lineage-specific marker genes organized into collocated sets to obtain a robust estimate of genome completeness. This allowed us to determine the finest level of taxonomic resolution at which we could confidently compute seed compounds, using genome completeness as a proxy for metabolic reaction network completeness (see Fig. S2 in the supplemental material). We deemed genomes to be nearly complete if they contained 95% of the lineage-specific marker genes. With the exception of tribe acl-B1, the tribe-level composite genomes were estimated to be incomplete (Fig. S2A). At the clade level, the genomes of clades acl-A and acl-B are estimated to be nearly complete, while the acl-C composite genome remains incomplete, as it contains only

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**FIG 1** Phylogenetic placement of the genomes used in this study within the acl lineage. The tree was built using RAxML (41) from a concatenated alignment of protein sequences from 37 single-copy marker genes (40). The order Actinomycetales forms the outgroup. Vertical black bars indicate groups of genomes belonging to defined tribes/clades within the acl lineage, as determined using 16S rRNA gene sequences (for SAGs and bin FNEF8-2 bin_7 acl-B only) and a defined taxonomy (28). SAGs are indicated with italic text. Figure S1 shows the position of the acl lineage relative to other orders within the class Actinobacteria.
75% of the 204 marker genes (Fig. S2B). As a result, seed compounds were calculated for composite clade-level genomes, with the understanding that some true seed compounds for the acI-C clade will not be predicted.

**Computation and evaluation of potential seed compounds.** Metabolic network reconstructions for each genome were built using KBase. Composite metabolic network graphs were then constructed for each tribe and clade by merging metabolic network reconstructions of individual genomes. Seed compounds for each clade were then computed from that clade’s composite metabolic network graph using a custom implementation of the seed set framework (Fig. 2). A total of 125 unique seed compounds were identified across the three clades (Table S2 in Data Set S1).

Because KBase is an automated annotation pipeline, the predicted set of seed compounds is likely to contain inaccuracies (e.g., due to missing or incorrect annotations). As a result, we screened the set of predicted seed compounds to identify those that represented biologically plausible auxotrophies and nutrients and manually curated this subset to obtain a final set of auxotrophies and nutrient sources. Of 125 unique compounds, 31 (24%) were retained in the final set of proposed auxotrophies and nutrients. Tables S3 and S4 in Data Set S1 contain this final set of compounds for clades acl-A, acl-B, and acl-C, and Fig. 3 shows the auxotrophies and nutrients that these compounds represent.
Making sense of seed compounds via protein clustering and metatranscriptomic mapping. For seed compounds representing nutrient sources, genes associated with the consumption of these compounds should be expressed. To test this, we collected and sequenced four metatranscriptome samples from Lake Mendota (Dane County, WI, USA). However, because seed compounds were computed from each clade’s composite metabolic network graph, genes associated with the consumption of seed compounds may be present in multiple genomes within the clade. To facilitate the linkage of metatranscriptome measurements to seed compounds, we used OrthoMCL \(^\text{(30)}\) to identify clusters of orthologous groups (COGs) in the set of acI genomes, merged metatranscriptome reads from all four samples, and mapped the reads to COGs within each clade.

Sequencing of cDNA from all four rRNA-depleted metatranscriptome samples yielded approximately 160 million paired-end reads. After merging, filtering, and further \textit{in-silico} rRNA removal, approximately 81 million, or 51\%, of the reads remained (Table S5 in Data Set S1). We then used BBMap \((https://sourceforge.net/projects/bbmap/)\) to map metatranscriptome reads to our reference genome collection. After mapping the metatranscriptomes to our acI genomes, we calculated the average coverage of each genome in our reference collection. Within each clade, the most abundant genome was detected with at least 16-fold coverage (Table S6 in Data Set S1).

Finally, we calculated gene expression for each COG on the basis of the number of reads per kilobase per million (RPKM) \((\text{Fig. 2})\). OrthoMCL identified a total of 5,013 protein clusters across the three clades (Table S7 in Data Set S1) with an average confidence of 84\% in annotation for COGs containing more than one gene. The COGs were unequally distributed across the three clades, with clade acl-A genomes containing 3,175 COGs (63\%), clade acl-B genomes containing 3,459 COGs (69\%), and clade acl-C genomes containing 1,365 COGs (27\%). Of these, 525 COGs were expressed in clade acl-A, 661 in clade acl-B, and 813 in clade acl-C (Table S8 in Data Set S1). Among

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**FIG 3** Seed compounds of members of the acl lineage. (A) Auxotrophies and nutrient sources, not including peptides and glycosides. (B) Peptides and glycosides. These compounds represent those inferred from genome annotations rather than the seed compounds. In panel B, the intensity of the color indicates the log\(_2\) fold change relative to the median (FC Rel. to Med.) of the encoding gene cluster. For compounds acted upon by multiple gene clusters, the percentile of the most highly expressed cluster was chosen.
the expressed genes, the median log₂ RPKM values were 31.1 in clade acI-A, 32.0 in clade acI-B, and 69.4 in clade acI-C. Due to differing RPKM values in each clade, we report gene expression values for each clade relative to the median log₂ RPKM value for that clade.

**Auxotrophies and nutrient sources of the acl lineage.** Seed set analysis yielded seven auxotrophies that could be readily mapped to ecophysiological attributes of the acl lineage (Fig. 3A and Table S3 in Data Set S1). In all three clades, beta-alanine was identified as a seed compound, suggesting an auxotrophy for pantothenic acid (vitamin B5), a precursor to coenzyme A formed from beta-alanine and pantoate (Table S9 in Data Set S1). In bacteria, beta-alanine is typically synthesized via aspartate decarboxylation, and we were unable to identify a candidate gene for this enzyme (aspartate 1-decarboxylase; EC 4.1.1.11) in any acl genome. Pyridoxine 5’-phosphate and 5’-pyridoxamine phosphate (forms of the enzyme cofactor pyridoxal 5’-phosphate [vitamin B6]) were also predicted to be seed compounds, and genes encoding numerous enzymes involved in the biosynthesis of these compounds were not found in the genomes (Table S9 in Data Set S1).

Clades within the acl lineage also exhibited distinct auxotrophies. Clade acl-A was predicted to be auxotrophic for the cofactor tetrahydrofolate (THF [vitamin B9]), and numerous enzymes for its biosynthesis were missing (Table S9 in Data Set S1). This cofactor plays an important role in the metabolism of amino acids and vitamins. In turn, clade acl-B was predicted to be auxotrophic for adenosylcobalamin (vitamin B12), containing only four reactions from its biosynthetic pathway (Table S9 in Data Set S1). Finally, acl-C was predicted to be auxotrophic for the nucleotide UMP (used as a monomer in RNA synthesis) and the amino acids lysine and homoserine. In all cases, multiple enzymes for the biosynthesis of these compounds were not found in the acl-C genomes (Table S9 in Data Set S1).

A number of seed compounds were also predicted to be degraded by members of the acl lineage (Fig. 3B; Table S3 in Data Set S1). Both clade acl-A and clade acl-B were predicted to use d-altronate and trans-4-hydroxy proline as nutrients, and acl-B was additionally predicted to use glycine betaine.

Finally, all three clades were predicted to use dipeptides and the sugar maltose as nutrients. Clades acl-A and acl-C were also predicted to consume the polysaccharides stachyose, mannotriose, and cellobiose. In all cases, these compounds were associated with reactions catalyzed by peptidases or glycoside hydrolases (Tables S10 and S11 in Data Set S1), which may be capable of acting on compounds beyond the predicted seed compounds. Thus, we used these annotations to define nutrient sources, rather than using the predicted seed compounds themselves. Among these nutrient sources were di- and polypeptides, predicted to be released from both cytosolic and membrane-bound aminopeptidases. As discussed below, we identified a number of transport proteins capable of transporting these released residues. In Lake Mendota, clades acl-B and acl-C expressed two aminopeptidases, one of which was expressed at nearly 175% of the median gene expression levels (Table S10 in Data Set S1). Clade acl-A expressed a third aminopeptidase at a lower level (40%, the median gene expression level) (Table S10 in Data Set S1).

All three clades were predicted to encode an alpha-glucosidase, which in Lake Mendota was expressed only in clades acl-B and acl-C, at nearly 60% of the median gene expression level (Table S11 in Data Set S1). All three clades also encode a beta-glucosidase, but it was not expressed in our samples. Furthermore, all three clades encode an alpha-galactosidase and multiple maltodextrin glucosidases (which free maltose from maltotriose), but these were expressed only in clades acl-A and acl-C. The alpha-galactosidase had a log₂ RPKM expression value of 1.5 times the median in clade acl-C, while the maltodextrin glucosidases were expressed at approximately 30% of the median (Table S11 in Data Set S1) in both clade acl-A and clade acl-C.

**Compounds transported by the acl lineage.** Microbes may be capable of transporting compounds that are not strictly required for growth, and comparing such...
compounds to predicted seed compounds can provide additional information about an organism’s ecology. Thus, we used the metabolic network reconstructions for the acI genomes to systematically characterize the transport capabilities of the members of the acI lineage.

All acI clades encode and were found to express a diverse array of transporters (Fig. 4; Data Set S1; Text S1). Consistent with the presence of peptidases, all clades contained numerous genes for the transport of peptides and amino acids, including putative oligopeptide and branched-chain amino acid transporters, as well as putative transporters for the polyamines spermidine and putrescine. All clades also contained a putative transporter for ammonium. The ammonium, branched-chain amino acid, and oligopeptide transporters had expression values above the median, with expression values for the substrate-binding protein (of the ATP-binding cassette [ABC] transporters) ranging from 1.7 to 411 times the median (Table S13 in Data Set S1). In contrast, while all clades expressed some genes from the polyamine transporters, only clade acI-B expressed the binding protein, at a level approximately 27.8 times the median (Table S13 in Data Set S1). Finally, clades acI-A and acI-B also contain a putative transporter for glycine betaine, which was expressed only in clade acI-A, at approximately 9.6 times the median (Table S13 in Data Set S1). However, we cannot rule out the possibilities that the expression of these transporters changes with space and time and that all three clades may express these enzymes under different conditions.

All clades also expressed transporters consistent with the presence of glycoside hydrolases, including transporters annotated as putative maltose, xylose, and ribose ABC-type transporters, which may indicate that acI bacteria are capable of transporting sugars, including both disaccharides (maltose) and monosaccharides (xylose and ribose). Of these, the putative maltose transporter was most highly expressed (but only in clades acI-A and acI-B), with expression values for the substrate-binding protein in a range in excess of 40 times the median (Table S13 in Data Set S1).

Representatives from the acI lineage were also found to encode and express a number of transporters that do not have corresponding seed compounds, including potential nucleobase transporters and purine/pyrimidine transporters (annotated as a uracil and a xanthine/uracil/thiamine/ascorbate family permease, respectively). Both of

FIG 4 Transports that are actively expressed by members of the acI lineage, as inferred from consensus annotations of genes associated with transport reactions present in metabolic network reconstructions. The intensity of the color indicates the log_2 fold change relative to the median value determined for the encoding gene cluster. For multisubunit transporters, the RPKM of the substrate-binding subunit was chosen (see Table S13 in Data Set S1). For some transporters, consensus annotations have been replaced with broad metabolite classes. Such metabolite classes are indicated with superscripts, and the original annotations are as follows: 1, spermidine and putrescine; 2, maltose; 3, xylose; 4, ribose; 5, uracil; 6, cytosine/purine/uracil/thiamine/allantoin; 7, xanthine/uracil/thiamine/ascorbate.
these are expressed in all three clades, with expression values ranging from 4.7 to 46 times the median (Table S13 in Data Set S1). Clades acl-A and acl-B also contained a second potential purine/pyrimidine transporter (annotated as a cytosine/purine/uracil/thiamine/allantoin family permease), which was expressed only in clade acl-B (Table S13 in Data Set S1). These transporters may be responsible for the uptake of the seed compounds UMP (a pyrimidine derivative) and vitamin B1 (also known as thiamine). In addition, clade acl-A contained but did not express a putative transporter for cobalamin (vitamin B12), and both clade acl-A and clade acl-B contained but did not express transporters for thiamine (vitamin B1) and biotin (vitamin B7) (Table S13 in Data Set S1).

Finally, all three clades expressed actinorhodopsin, a light-sensitive protein that is expected to function as a proton efflux pump (31). In all clades, actinorhodopsin was among the top 10 most highly expressed genes (Table S7 in Data Set S1), with expression values in excess of 84 times the median in all three clades (Table S7 in Data Set S1). Given that many of the transport proteins are ABC transporters, we speculate that actinorhodopsin may facilitate maintenance of the proton gradient necessary for ATP synthesis. Coupled with high expression levels of diverse transporters, this result strongly suggests that acl functions as a photoheterotroph. However, it remains to be seen if this behavior is a general feature of acl physiology or if it is restricted to the specific conditions of the lake and our sampling period.

DISCUSSION

This study used high-throughput metabolic network reconstruction and the seed set framework to predict auxotrophies and nutrient sources of uncultivated microorganisms from incomplete genome sequences. The computational approach easily scales to hundreds of metabolic reconstructions and enables a targeted analysis by identifying those compounds through which an organism interacts with its environment. However, predicted seed compounds are sensitive to the metabolic network structure, and analyzing the results requires significant manual curation of the metabolic reconstruction and accurate interpretation of the underlying gene annotations. As a consequence, the seed set framework is not as high throughput as was initially envisioned but is nevertheless suitable for analysis of microorganisms with high-quality metabolic network reconstructions.

Our predictions of the substrate use capabilities of the acl lineage are largely congruent with previous genome-centered studies based on smaller but manually curated genome collections (22, 25, 27), indicating that the use of automatic metabolic network reconstructions yields predictions similar to metabolic reconstruction efforts that are based on more extensively manual methods, while being both more high throughput and more focused on an organism’s substrate utilization capabilities. In particular, this study predicted that the consumption of N-rich compounds is a universal feature of the acl lineage, with all three clades predicted to consume ammonium, branched-chain amino acids, polyamines, and di- and oligopeptides. These findings agree with the results of microautoradiography-fluorescent in situ hybridization (MAR-FISH) studies that confirm the ability of acl bacteria to consume a variety of amino acids (20, 23). Furthermore, the presence of alpha- and beta-glucosidases is consistent with observations that acl bacteria consume glucose (19, 23), even though no obvious glucose transport system was found in the genomes. Because transport proteins are often capable of acting on multiple substrates, one of the putative sugar transporters may be responsible for glucose uptake activity.

However, our approach failed to recapitulate other genomic and experimental observations, including the uptake of N-acetylglucosamine (NAG) (32–34), the deoxynucleoside thymidine (23, 35), and acetate (19) as well as the potential to hydrolyze the cyanobacterial peptide cyanophycin via the enzyme cyanophycinase (22, 25). Inspection of these discrepancies reveals some important limitations of the seed set framework and automatic metabolic reconstructions. First, the seed set framework identifies only the compounds that the metabolic network must obtain from its environment and
fails to identify compounds that the organism can acquire from its environment but can also itself synthesize. Thymidine and acetate fall into this category. Second, automatic metabolic network reconstructions may not fully capture an organism’s metabolic network (e.g., due to missing or incorrect genome annotations). Manual inspection of the product of the previously identified cyanophycinase gene revealed that KBase annotated this putative enzyme as a hypothetical protein. As biochemical characterization of hypothetical proteins and automatic gene and protein annotation are active areas of research, we anticipate that advances in these fields will continue to improve the accuracy of automatic metabolic network reconstructions.

This study also suggested that auxotrophies for some vitamins may be universal features of the acl lineage, as we predicted all clades to be auxotrophic for pantothenic acid and pyridoxal 5'-phosphate (vitamins B5 and B6). We also predict new auxotrophies within the acl lineage, including THF (clade acl-A), adenosylcobalamin (vitamin B12; clade acl-B), and lysine, homoserine, and UMP (clade acl-C). However, with the exception of adenosylcobalamin, we did not identify transporters for any of these compounds. This negative result may reflect our limited knowledge of transport proteins; transporters for these compounds may yet be present in the genomes, or one or more of the predicted transporters may act on these compounds. Furthermore, because the acl-C composite genome was estimated to be around 75% complete, we cannot rule out the possibility that the missing genes might be found in this clade when additional genomes are recovered. Nonetheless, these results provide additional support for the hypothesis that distributed metabolic pathways and metabolic complementarity may be common features of freshwater bacterial communities.

Combined, these results suggest that the members of the acl lineage are photoheterotrophs and that their survival depends on the availability of a diverse array of N-rich compounds, saccharides, and light. The acl lineage does not appear to be metabolically self-sufficient and may participate in the turnover of high-molecular-weight dissolved organic compounds, such as starch, glycogen, and cellulose. Metatranscriptomic analysis showed that transport proteins were among those most highly expressed in the acl genomes, and expression of multiple putative amino acid transporters may facilitate uptake of these labile compounds. We also observed differences in the relative levels of expression of these transporters, which may point to clade-specific differences in affinities for these substrates. Finally, the actinorhodopsin protein was highly expressed and may facilitate synthesis of the ATP needed to drive acl’s many ABC-type transporters.

Finally, the fragmented and incomplete nature of SAGs and MAGs required us to construct composite genomes for individual acl clades by leveraging multiple genomes from closely related populations. Such an approach limits the resolution of predictions, as we cannot make predictions at the level of tribes, smaller populations, or individual cells. Thus, metabolic diversification at these taxonomic levels would be missed. Constructing composite genomes may also overestimate the metabolic capabilities of a clade or group; for example, if a complete pathway is present in a clade but is distributed among different tribes, the clade would be able to carry out the activity of the entire pathway in situ only if all tribes were present in close enough proximity to exchange pathway intermediates. Nonetheless, the seed set approach provides a framework that can be used to generate new hypotheses about the substrates used by members of a defined phylogenetic group, provided that multiple closely related genomes are available. As metagenomic assembly and binning techniques and single-cell sequencing methods improve and complete genomes become available, we anticipate our approach being applied to individual microbial genomes.

**MATERIALS AND METHODS**

A freshwater reference genome collection. This study relied on an extensive collection of freshwater bacterial genomes containing MAGs obtained from two metagenomic time series from two
Wisconsin lakes (27, 39) as well as SAGs from three lakes in the United States (21). Additional information about this genome collection can be found in Text S1.

**Metatranscriptome sampling and sequencing.** This study used four metatranscriptomes obtained as part of a larger study of gene expression in freshwater microbial communities. Additional information about these samples can be found in the Text S1. All protocols and scripts for sample collection, RNA extraction, rRNA depletion, sequencing, and bioinformatic analysis can be found on Github ([https://github.com/McMahonLab/OMD-TOIL](https://github.com/McMahonLab/OMD-TOIL) [DOI: 10.5281/zenodo.839851]). Metadata for the four samples used in this study can be found in Table S6 in Data Set S1 in the supplemental material, and the raw RNA sequences can be found on the Sequence Read Archive (SRA) of the National Center for Biotechnology Information (see below).

**Identification of acI SAGs and actinobacterial MAGs.** The members of the acI were previously phylogenetically divided into 3 clades (acI-A, acI-B, and acI-C) and 13 tribes on the basis of their 16S rRNA gene sequences (28). The acI SAGs were identified within a previously published genome collection (21) and classified to the tribe level using partial 16S rRNA genes and a reference taxonomy for freshwater bacteria, as described in Text S1. Actinobacterial MAGs were identified within two metagenomic time series (27, 39) using taxonomic assignments from a subset of conserved marker genes, as described in Text S1. Phylogenetic analysis of acI SAGs and actinobacterial MAGs was performed using a concatenated alignment of single-copy marker genes obtained via Phylosift (40). Maximum likelihood trees were generated using RAxML (41), the automatic protein model assignment option (PROTGAMMAAUTO), and 100 bootstraps.

**Genome annotation, metabolic network reconstruction, and computation and evaluation of seed compounds.** In the seed set framework, an organism’s metabolism is represented via a metabolic network graph, in which nodes denote compounds and edges denote enzymatically encoded biochemical reactions linking substrates and products (42). Allowable biochemical transformations can be identified by drawing paths along the network, in which a sequence of edges connects a sequence of distinct vertices. In our implementation of the seed set framework, metabolic network graphs were generated as follows:

- Genome annotations were performed and metabolic network reconstructions were built using KBase. Contigs for each genome were uploaded to KBase and annotated using the “Annotate Microbial Contigs” method with default options, which uses components of the RAST toolkit for genome annotation (43, 44).
- Metabolic network reconstructions were obtained using the “Build Metabolic Model” app with default parameters, which relies on the Model SEED framework (45) to build a draft metabolic model. To ensure that the reconstructions contained only reactions with genomic evidence, no gap filling was performed. These reconstructions were then pruned of currency metabolites (compounds used to carry electrons and functional groups) and highly connected compounds and converted to metabolic network graphs (see Fig. S3 and Text S1 in the supplemental material). Many of the individual acI genomes are incomplete. Therefore, to increase the accuracy of seed identification by means of the use of a more complete metabolic network, composite metabolic network graphs were constructed for each tribe and clade (Fig. S4; Text S1).

- Formally, the seed set of the network is defined as the minimal set of compounds that cannot be synthesized from other compounds in the network and whose presence enables the synthesis of all other compounds in the network (7). Seed compounds for each composite metabolic network graph were calculated using a new Python implementation of the seed set framework (7) (Fig. S5 and Text S1). Because seed compounds are computed from a metabolic network, it is important to manually evaluate all predicted seed compounds to identify those that may be biologically meaningful and that do not arise from errors in the metabolic network reconstruction. Compounds involved in fatty acid and phospholipid biosynthesis pathways were removed during curation, as these pathways are often organism specific and unlikely to be properly annotated by automatic metabolic reconstruction pipelines. Seed compounds related to currency metabolites were also removed, as data corresponding to reactions for the synthesis of these compounds may have been removed during network pruning.

- Text S1 contains a series of brief vignettes explaining why selected compounds were discarded based on the aforementioned considerations and provides examples of additional curation efforts applied to biologically plausible compounds. For a plausible auxotrophy, we screened the genomes for the canonical biosynthetic pathway(s) for that compound and retained those compounds for which the biosynthetic pathway was incomplete. For identification of a plausible nutrient source, we screened the genomes for the canonical degradation pathway(s) for that compound and retained those compounds for which the degradation pathway was complete.

- All computational steps were implemented using Python scripts, freely available as part of the reverseEcology Python package developed for this project ([https://pypi.python.org/pypi/reverseEcology/](https://pypi.python.org/pypi/reverseEcology/) [DOI: 10.5281/zenodo.839856]).

**Identification of transported compounds.** For each genome, we identified all transport reactions present in its metabolic network reconstruction. Gene-protein-reaction associations (GPRs) for these reactions were manually curated to remove unannotated proteins, to group genes into operons (if applicable), and to identify missing subunits for multisubunit transporters. These genes were then mapped to their corresponding COGs and grouped accordingly. Finally, the most common annotation for each COG was used to identify likely substrates for each of these groups. Only transporters with >50% confidence in the substrate-binding subunit were retained. Because identification and annotation of transport proteins are active areas of research (36), substrates for each transporter are described as putative and acting on molecular classes (e.g., saccharide, amino acid) instead of on...
specific compounds, in order to better reflect the promiscuity of transport proteins and the ambiguity of their annotation.

**Protein clustering, metatranscriptomic mapping, and clade-level gene expression.** OrthoMCL (30) was used to identify clusters of orthologous groups (COGs) in the set of acI genomes. Both OrthoMCL and BLAST were run using default options (46). Annotations were assigned to protein clusters by choosing the most common annotation among all genes assigned to the respective cluster and a confidence score assigned to each COG (representing the fraction of genes having the most common annotation). Trimmed and merged metatranscriptomic reads from each of the four biological samples were then pooled and mapped to a single reference fasta file containing all acI genomes using BBMap with the ambig/random and minid=0.95 options. The 95% identity cutoff was chosen as this represents a well-established criterion for identifying microbial species using average nucleotide identity (ANI) (47), while combining the ambig option with competitive mapping using pooled acI genomes as the reference ensured that the reads mapped to only a single genome. These results were then used to compute the expression of each COG in each clade.

Next, HTSeq-Count (48) was run using the intersection_strict option to count the total number of reads that map to each gene in our acI genome collection. After mapping, the list of counts was filtered to remove those genes that did not recruit at least 10 reads. Using the COGs identified by OrthoMCL, the genes that correspond to each COG were then identified.

Within each clade, gene expression was computed for each COG on the basis of the number of reads per kilobase per million (RPKM) (49), while also accounting for different gene lengths within a COG and numbers of mapped reads for each genome within a clade. That is, the RPKM value for a single COG represents the sum of RPKM values for each gene within that COG, normalized to the appropriate gene length and total number of mapped reads. RPKM counts were then normalized to the median level of gene expression within that clade. Finally, the expression data (mapping of transcript reads to genes) were visualized to ensure that the RPKM calculations were based on continuous transcription of each gene.

**Accession number(s).** The raw RNA sequences can be found in the Sequence Read Archive (SRA) of the National Center for Biotechnology Information under BioProject accession no. PRJNA362825.

**Data availability.** All genomic and metatranscriptomic sequences are available through IMG and NCBI, respectively. A reproducible version of the manuscript is available at https://github.com/joshamilton/Hamilton_acI_2017 (DOI: 10.5281/zenodo.839858).

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mSystems.00091-17.

- **TEXT S1,** DOCX file, 0.02 MB.
- **FIG S1,** EPS file, 2.6 MB.
- **FIG S2,** EPS file, 1.3 MB.
- **FIG S3,** EPS file, 2.8 MB.
- **FIG S4,** TIF file, 0.3 MB.
- **FIG S5,** EPS file, 1.7 MB.
- **FIG S6,** EPS file, 2.3 MB.
- **DATA SET S1,** XLSX file, 1.3 MB.

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