Interactions of Freshwater Cyanobacteria with Bacterial Antagonists

Omneya Ahmed Osman,a Sara Beier,b Manfred Grabherr,c Stefan Bertilssona
Department of Ecology and Genetics, Limnology, and Science for Life Laboratory, Uppsala University, Uppsala, Swedena; Leibniz Institute for Baltic Sea Research, Warnemünde, Germanyb; Department of Medical Microbiology and Biochemistry, Bioinformatics Infrastructure for Life Sciences, and Science for Life Laboratory, Uppsala University, Uppsala, Swedenc

ABSTRACT Cyanobacterial and algal mass development, or blooms, have severe effects on freshwater and marine systems around the world. Many of these phototrophs produce a variety of potent toxins, contribute to oxygen depletion, and affect water quality in several ways. Coexisting antagonists, such as cyanolytic bacteria, hold the potential to suppress, or even terminate, such blooms, yet the nature of this interaction is not well studied. We isolated 31 cyanolytic bacteria affiliated with the genera Pseudomonas, Stenotrophomonas, Acinetobacter, and Delftia from three eutrophic freshwater lakes in Sweden and selected four phylogenetically diverse bacterial strains with strong-to-moderate lytic activity. To characterize their functional responses to the presence of cyanobacteria, we performed RNA sequencing (RNA-Seq) experiments on coculture incubations, with an initial predator-prey ratio of 1:1. Genes involved in central cellular pathways, stress-related heat or cold shock proteins, and antitoxin genes were highly expressed in both heterotrophs and cyanobacteria. Heterotrophs in coculture expressed genes involved in cell motility, signal transduction, and putative lytic activity. L,D-Transpeptidase was the only significantly upregulated lytic gene in Stenotrophomonas rhizophila EK20. Heterotrophs also shifted their central metabolism from the tricarboxylic acid cycle to the glyoxylate shunt. Concurrently, cyanobacteria clearly show contrasting antagonistic interactions with the four tested heterotrophic strains, which is also reflected in the physical attachment to their cells. In conclusion, antagonistic interactions with cyanobacteria were initiated within 24 h, and expression profiles suggest varied responses for the different cyanobacteria and studied cyanolytes.

IMPORTANCE Here, we present how gene expression profiles can be used to reveal interactions between bloom-forming freshwater cyanobacteria and antagonistic heterotrophic bacteria. Species-specific responses in both heterotrophs and cyanobacteria were identified. The study contributes to a better understanding of the interspecies cellular interactions underpinning the persistence and collapse of cyanobacterial blooms.

KEYWORDS coculture interaction, cyanobacteria, metatranscriptome

Many nutrient-rich freshwater ecosystems experience episodic mass development of cyanobacteria. These blooms influence water quality through the sheer quantity of organisms and the release of toxic compounds (1–3). Trophic interactions can lead to cyanobacterial toxins being accumulated in aquatic organisms (e.g., mussels, crayfish, and fish), which are subsequently passed on to humans (4). Microcystis is one of the most widespread and problematic toxin-producing freshwater cyanobacteria. Its toxin, microcystin, can cause liver damage in domestic animals and has a number of severe impacts on lake biota (5). Aphanizomenon flos-aquae is another widespread...
cyanobacterium, with several strains having the capacity to produce and release toxins that cause fish mortality (6, 7).

The control of algal blooms has previously been facilitated with mechanical methods, such as filtration, ultrasound, and electrolysis, or the addition of toxic chemicals, such as copper sulfate or sodium hypochlorites (8). However, these methods are expensive and not practical for the elimination of algal blooms at the ecosystem scale. The use of biological control agents, such as viruses and predatory bacteria, has been proposed as an alternative strategy to counteract the harmful effects of algal blooms, but the efficacy of this approach is yet to be demonstrated.

Several phylogenetically diverse heterotrophic bacteria within the Proteobacteria, Bacteroidetes, Firmicutes, and Actinobacteria have displayed antagonistic activities against various cyanobacteria (9–12). These antagonistic bacteria exert lytic activity via mechanisms, including parasitism, antibiosis after host entrapment, or contact lysis (13). For example, close attachment of Streptomyces neyagawaensis to cyanobacterial cells is known to cause efficient lysis after the production of antimicroalgal compounds (10). Other examples include the endoparasitic strain Bdellovibrio bacteriovorus, which was reported to cause lysis of Phormidium luridum by the secretion of extracellular compounds (12, 14) and heterotrophic bacterial production of the antimicroalgal compound β-cyanoalanine (L-CNAla) to control toxin-producing cyanobacteria (15). In the study by Yoshikawa et al. (15), the authors showed that L-CNAla inhibits the growth of some cyanobacterial strains, such as Synechococcus sp. strain CSIRO 94 and Microcystis aeruginosa NIES-298, but not green algae, dinoflagellates, or diatoms.

One powerful approach to map the functional response of microorganisms to environmental cues or interactions is to broadly sequence the combined transcriptome of the full set of interacting organisms. Compared to a metagenomic approach, the major advantage of metatranscriptomic analyses is the possibility to observe actively expressed genes at a certain time point, allowing for the prediction of changes in metabolic pathways and other functional responses (16–18). Recent advances in next-generation RNA sequencing mean that millions of reads can be generated and either be mapped to reference genomes or assembled de novo, making this approach tractable and affordable (19, 20).

We applied a metatranscriptomic approach in the present study, using the input experiment of axenic cultures of Microcystis aeruginosa PCC 7941 and Aphanizomenon flos-aquae PCC 7905, challenged with different confirmed freshwater lake antagonistic or lytic bacteria. The combined mRNA pools, expressed by the heterotrophic antagonistic bacteria cocultured with Microcystis aeruginosa PCC 7941 or Aphanizomenon flos-aquae PCC 7905, were characterized by RNA sequencing. The aim was to uncover the mechanisms underlying the cyanolytic bacterial interactions while at the same time identifying the cyanobacterial response to such pressures.

RESULTS

Isolation of cyanolytic bacteria. Most of the cyanolytic bacteria were from Lake Ekoln and Lake Erken, while only eight strains were from Lake Funbosjön. All of the 31 isolates selected for further characterization propagated well on BG11 medium supplemented with 0.2% Casitone. Twenty of these isolates had high 16S rRNA identity to Pseudomonas species (99 to 100% identity), while 11 isolates showed an equal level of identity to other taxa (Delftia, Stenotrophomonas, Acinetobacter, Marinobacter, and Limnobacter). The 31 isolates displayed high-to-moderate lytic activity with either defined or diffuse lytic zones (see Table S2 in the supplemental material).

Genomic features of experimental cyanolytic bacteria. The 16S rRNA gene analysis of the four cyanolytic bacteria selected for further experiments matched with 100% identity to Stenotrophomonas rhizophila (1,527-bp alignment), Pseudomonas putida (1,144 bp), and Acinetobacter beijerinckii (1,518 bp) and 99% identity to Delftia sp. (1,519 bp). The genome statistics and metabolisms of the four strains are summarized in Tables 1 and 2. Inspection of the genomes revealed that the four cyanolytic bacteria are all heterotrophs, with auxotrophic requirement for specific amino acids. S. rhizophila...
**TABLE 1** Genome statistics of *Acinetobacter beijerinckii* F107, *Delftia* sp. strain F45, *Pseudomonas putida* EK59, and *Stenotrophomonas rhizophila* EK20

| Genome statistics | Acinetobacter beijerinckii F107 | Delftia sp. F45 | Pseudomonas putida EK59 | Stenotrophomonas rhizophila EK20 |
|-------------------|-------------------------------|----------------|-------------------------|-------------------------------|
| No. % of total    | No. % of total                | No. % of total | No. % of total          | No. % of total               |
| Total DNA bases   | 2,304,418 100                  | 5,792,729 100  | 4,767,657 100           | 3,665,436 100                |
| DNA-coding bases  | 2,029,954 88.09                | 5,216,926 90.06 | 4,311,136 90.42        | 3,319,715 90.57             |
| DNA G+C bases     | 885,232 38.41                  | 3,871,767 66.84 | 2,962,115 62.13        | 2,437,651 66.50             |
| DNA scaffolds      | 594 100.00                     | 391 100.00     | 530 100.00              | 384 100                      |
| Total genes       | 2,624 100.00                   | 5,381 100.00   | 4,730 100.00            | 3,583 100                    |
| Protein-coding genes | 2,578 98.25                | 5,304 98.57    | 4,614 97.55             | 3,518 98.19                  |
| With function prediction | 1,881 71.68          | 4,337 81.60    | 3,708 78.39             | 2,762 77.09                  |
| Without function prediction | 697 26.56               | 967 17.79      | 906 19.15               | 756 21.10                    |
| With enzymes      | 545 20.77                     | 1,167 21.69    | 1,079 22.81             | 820 22.89                    |
| Connected to KEGG pathways | 530 20.20 | 1,387 25.78 | 1,246 26.34 | 906 25.29 |
| Connected to KEGG Orthology | 1,033 39.37 | 2,465 45.27 | 2,312 48.88 | 1,638 45.72 |
| Connected to transporter classification | 272 10.37 | 920 17.10 | 691 14.61 | 406 61.76 |
| With COGs         | 1,349 51.41                   | 3,669 42.98    | 3,078 65.07             | 2,213                        |
| With KOGs         | 409 15.59                     | 914 16.99      | 796 16.83               | 618 17.25                    |
| With Pfam         | 1,994 75.99                   | 4,516 83.92    | 3,946 83.42             | 2,903 81.02                  |
| With TIGRfam      | 743 28.32                     | 1,472 27.36    | 1,491 31.52             | 1,076 30.03                  |
| With InterPro     | 1,271 48.44                   | 3,039 56.48    | 2,593 54.82             | 1,883 52.55                  |
| Genes in biosynthetic clusters | 38 1.45 | 165 3.07 | 155 3.28 | 38 1.06 |
| Fused protein-coding genes | 41 1.56 | 118 2.19 | 122 2.58 | 87 2.43 |
| Protein-coding genes coding signal peptides | 211 8.04 | 603 11.21 | 479 10.13 | 530 14.79 |
| Protein-coding genes coding transmembrane proteins | 581 22.14 | 1,222 22.71 | 1,050 22.20 | 888 24.78 |

EK20, *Delftia* sp. strain F45, and *A. beijerinckii* F107 are auxotrophs for 12 to 17 amino acids (such as L-lysine, L-histidine, and L-tyrosine) and prototrophs for 3 to 4 amino acids (such as glycine and glutamate). *Pseudomonas putida* EK59 appeared to be auxotrophic for only five amino acids. In each of the four assemblies, about 98% of the genome consists of protein-coding genes, 70 to 80% of which could be functionally annotated as Clusters of Orthologous Groups (COGs) or protein families (Pfams). Most of the basic cellular and metabolic functions were successfully annotated (Tables 1 and 2). For *Delftia* sp. F45, *P. putida* EK59, and *S. rhizophila* EK20, >100 genes involved in cell motility and extracellular structures were identified, while few genes coding for cell motility and extracellular structures were identified in the *A. beijerinckii* F107 genome (20).

The *Delftia* sp. F45, *P. putida* EK59, and *S. rhizophila* EK20 genomes contain 70 to 76 genes predicted to be involved in defense mechanisms, while 39 genes with such functions were present in the *A. beijerinckii* F107 genome. All four heterotrophic genomes contain peroxiredoxin, and most of them (except for *A. beijerinckii* F107) encode the osmotically inducible protein OsmC.

**Coculture experiment.** Here, cocultures are abbreviated miA, miD, miP, miS, apA, apD, apP, and apS, with the first two letters describing the autotroph genus, followed by the first letter of the heterotroph genus (mi, *Microcystis aeruginosa* PCC 7941; ap, *Aphanizomenon flos-aquae* PCC 7905; P, *Pseudomonas putida* EK59; S, *Stenotrophomonas rhizophila* EK20; D, *Delftia* sp. F45; A, *Acinetobacter beijerinckii* F107). Microscopic observations were first used to describe interactions between the four cyanolytic bacterial strains, and either *M. aeruginosa* PCC 7941 or *A. flos-aquae* PCC 7905. During the interaction with *M. aeruginosa* PCC 7941, all four heterotrophic bacteria exhibited very similar visual patterns: first, the heterotrophs adhered to the cyanobacterial cells at 6 h of incubation. Second, the formation of aggregates surrounding *M. aeruginosa* PCC 7941 was apparent after 24 h of incubation time. Third, many *M. aeruginosa* PCC 7941 cells featured abnormal and irregular shapes, and extracellular protein structures could be visualized with NanoOrange after 96 h (Fig. 1).

During the coculture with the nitrogen-fixing and filamentous cyanobacterium *A. flos-aquae* PCC 7905, the heterotrophs *P. putida* EK59 and *S. rhizophila* EK20 attached...
to the heterocyst of the filaments after 6 h of incubation. The amount of cells adjacent to the filament had increased after 24 h of contact time, and the disruption of filaments was only starting to become visible after 96 h of incubation. In contrast, *Delftia* sp. F45 and *A. beijerinckii* F107 attached mainly to the photosynthetic cells of *A. flos-aquae* PCC 7905 and increased in numbers between 6 and 24 h of contact time. Abnormal filament shapes were observed after 96 h (Fig. 2).

Despite the apparent strong visible effect of the heterotrophic bacteria on cyanobacterial cell morphology, microscopic cell counts revealed that the growth of the two cyanobacterial strains was not significantly affected (Table S3). The count of heterotrophic bacteria increased during the first 24 h of the experimental incubation and remained stable or slightly increased thereafter. However, as the bacterial population forming aggregates and biofilms adjacent to the cyanobacterial cells could not be accurately counted, these numbers may be underestimates. Irregular shapes of cyanobacterial cells were not counted due to the difficulty in estimating their number in aggregates.

### Highly expressed genes in heterotrophs

Three COG functional categories, (i) amino acid transport and metabolism, (ii) translation, ribosomal structure, and biogenesis, and (iii) energy production and conversion, were among the most highly expressed genes in all heterotroph cocultures (Table 3). For representative genes of each COG functional category, see Table S4.

Other functional categories involved in cell growth and survival; carbohydrate, coenzyme, and inorganic ion transport and metabolism; protein turnover; and chaperones were also among the most highly expressed genes in all heterotrophs (Table 3). A considerable number of putative defense mechanisms were expressed in most of the
heterotrophs, particularly in the *S. rhizophila* EK20 cocultures. Most of the heterotrophs featured high expression of the CspA family cold shock protein (3% for *A. beijerinckii* F107 and *S. rhizophila* EK20, 0.8% for *P. putida* EK59), known to protect bacterial cells from damage due to low temperature. *S. rhizophila* featured high expression of 5-formyltetrahydrofolate cyclo-ligase, which is involved in folate metabolism, and the L,D-transpeptidase catalytic domain involved in peptidoglycan cross-linking.

**Highly expressed genes in cyanobacteria.** In both *A. flos-aquae* PCC 7905 and *M. aeruginosa* PCC 7941, genes coding for ribosomal proteins and amino acid synthesis, as well as energy production and conversion required for cell growth, were highly expressed (Table 4). Genes involved in cell wall and membrane/envelope biogenesis, protein turnover, chaperone and carbohydrate transport, and metabolism were also equally highly expressed in both cyanobacteria. Several highly expressed genes (19 and 25, respectively) are involved in defense mechanisms and signal transduction (Table S5). Photosystem q(b) protein and Lhc-like protein Lhl4 (3.7 to 5% and 1 to 3.9%, respectively) were the most abundantly expressed genes in the *M. aeruginosa* PCC 7941 cocultures, while gas vesicle structural protein and allophycocyanin were highly abundant in the *A. flos-aquae* PCC 7905 cocultures (10 to 12% and 2 to 2.5%, respectively).

**Differential expression gene profiles in *P. putida* EK59 and *S. rhizophila* EK20.** Overall clustering of expression profiles of the heterotrophs revealed clear species-specific patterns under all coculture and monoculture conditions. The monoculture control transcriptomes of *P. putida* EK59 and *S. rhizophila* EK20 were distinct but clustered with their respective coculture transcriptomes (Fig. 3). Differential expression
analyses identified 37 COG homologs and an additional 7 genes annotated as being significantly upregulated in at least one of the coculture treatments (Fig. 4a). Thirty-two COG homologs and 7 annotated genes were downregulated in at least one of the coculture treatments compared to monocultures (Fig. 4b). Expression of some COG homologs was specific to either *P. putida* EK59 or *S. rhizophila* EK20, but a few contradictory trends were observed. On the whole, most genes with significant differential expression in one of the cocultures were consistently up- or downregulated across species (Fig. 4a and b).

Among transcripts coding for energy production and conservation, two key enzymes that indicate the activity of the glyoxylate shunt (isocitrate lyase and malate synthetase) were consistently upregulated in cocultures (Fig. 4a). In contrast, a number of other genes coding for enzymes involved in the citrate cycle or reoxidation of reduced NADH (respiration chain) were consistently downregulated in the cocultures, although only statistically significantly in the *S. rhizophila* EK20 cocultures (Fig. 4b). Other genes involved in the respiratory chain, such as cytochromes, were inconsistently regulated (Fig. 4a and b). A simplified metabolic map with the transcriptional response of these pathways is given in Fig. 5.

A catalase and alkyl hydroperoxide oxidase protein, as well as a DNA-break repair enzyme, were consistently upregulated in both heterotroph cocultures (Fig. 4a). For both heterotrophs, some transcripts coding for translation and ribosomal structure were consistently downregulated in coculture (Fig. 4b).

Genes involved in formation of flagella were consistently upregulated in both cocultured heterotrophs. Outer or integral membrane and a surface protein containing fasciclin repeats (COG3188, COG0670, and COG2335, respectively) were significantly upregulated, while the outer membrane protein (COG3047) was downregulated in *P. putida* EK59 cocultures. The 5-formyltetrahydrofolate cyclo-ligase gene involved in folate metabolism was upregulated in most of the cocultured heterotrophs with significant signal in ap5.

Among the genes that were significantly upregulated in cocultures was one enzyme with putative extracellular lytic activity. This may have caused the observed cyanolytic activity with L,D-transpeptidase, which was upregulated in *S. rhizophila* EK20. However, a number of outer membrane candidate genes with potential lytic activities seemed to be
| COG category | Function                                                                 | apP1 | apP2 | miP1 | miP2 | apS1 | apS2 | miS1 | miS2 | apD1 | apD2 | miD1 | miD2 | apA1 | apA2 | miA1 | miA2 |
|--------------|--------------------------------------------------------------------------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|
| E            | Amino acid transport and metabolism                                      | 67   | 67   | 80   | 72   | 81   | 81   | 85   | 72   | 45   | 44   | 42   | 35   | 59   | 58   | 55   | 51   |
| S            | Function unknown                                                         | 62   | 58   | 69   | 56   | 69   | 71   | 76   | 64   | 34   | 23   | 18   | 24   | 18   | 41   | 41   | 42   | 41   |
| J            | Translation, ribosomal structure, and biogenesis                        | 60   | 67   | 75   | 61   | 91   | 93   | 99   | 87   | 62   | 49   | 46   | 46   | 61   | 59   | 65   | 57   |
| C            | Energy production and conversion                                         | 52   | 59   | 57   | 54   | 59   | 57   | 62   | 55   | 42   | 37   | 38   | 34   | 39   | 39   | 43   | 37   |
| R            | General function prediction only                                         | 47   | 45   | 55   | 50   | 59   | 71   | 74   | 58   | 28   | 24   | 27   | 24   | 27   | 42   | 43   | 46   | 42   |
| M            | Cell wall/membrane/envelope biogenesis                                   | 40   | 53   | 57   | 43   | 64   | 67   | 71   | 60   | 39   | 32   | 22   | 24   | 35   | 38   | 33   | 34   |
| G            | Carbohydrate transport and metabolism                                   | 35   | 31   | 34   | 28   | 49   | 50   | 55   | 47   | 19   | 15   | 14   | 11   | 16   | 14   | 16   | 13   | 13   |
| O            | Posttranslational modification, protein turnover, chaperones             | 35   | 36   | 43   | 36   | 52   | 53   | 56   | 47   | 36   | 32   | 27   | 28   | 28   | 26   | 26   | 27   | 28   |
| P            | Inorganic ion transport and metabolism                                   | 34   | 31   | 35   | 27   | 41   | 40   | 50   | 38   | 30   | 22   | 14   | 18   | 14   | 18   | 35   | 33   | 38   | 28   |
| H            | Coenzyme transport and metabolism                                        | 32   | 33   | 39   | 33   | 48   | 50   | 54   | 40   | 21   | 15   | 10   | 10   | 27   | 26   | 25   | 23   | 23   |
| K            | Transcription                                                            | 29   | 29   | 34   | 25   | 36   | 42   | 34   | 26   | 21   | 21   | 21   | 28   | 28   | 28   | 29   | 27   | 27   |
| L            | Signal transduction mechanisms                                           | 26   | 24   | 29   | 24   | 31   | 33   | 33   | 33   | 24   | 17   | 17   | 17   | 17   | 12   | 16   | 15   | 15   |
| I            | Lipid transport and metabolism                                           | 21   | 21   | 31   | 26   | 51   | 54   | 60   | 49   | 19   | 19   | 16   | 13   | 26   | 30   | 24   | 23   | 23   |
| T            | Replication, recombination, and repair                                    | 20   | 24   | 23   | 20   | 36   | 35   | 36   | 33   | 18   | 14   | 9    | 13   | 27   | 23   | 26   | 26   |
| N            | Cell cycle control, cell division, chromosome partitioning               | 16   | 15   | 16   | 11   | 13   | 14   | 13   | 14   | 13   | 7    | 4    | 5    | 14   | 12   | 15   | 11   | 11   |
| V            | Cell motility                                                            | 15   | 15   | 15   | 14   | 18   | 19   | 20   | 12   | 13   | 13   | 6    | 6    | 1    | 0    | 2    | 0    | 0    |
| Q            | Defense mechanisms                                                       | 9    | 9    | 10   | 9    | 17   | 17   | 17   | 15   | 8    | 8    | 4    | 5    | 8    | 11   | 10   | 8    | 8    |
| U            | Secondary metabolite biosynthesis, transport and, catabolism             | 7    | 6    | 12   | 7    | 7    | 9    | 11   | 5    | 10   | 4    | 2    | 7    | 7    | 8    | 6    | 6    | 6    |
| N            | Intracellular trafficking, secretion, and vesicular transport            | 7    | 9    | 10   | 9    | 16   | 12   | 16   | 12   | 8    | 7    | 9    | 6    | 6    | 5    | 5    | 5    | 5    |
| HR           | Coenzyme transport and metabolism                                        | 4    | 4    | 4    | 4    | 4    | 4    | 5    | 3    | 2    | 2    | 2    | 2    | 2    | 2    | 2    | 2    | 2    |
| TK           | Signal transduction mechanisms                                           | 4    | 5    | 4    | 4    | 4    | 4    | 3    | 4    | 6    | 5    | 4    | 6    | 3    | 2    | 2    | 2    |
| NT           | General function prediction only                                         | 4    | 5    | 4    | 4    | 5    | 4    | 5    | 4    | 3    | 2    | 3    | 2    | 2    | 2    | 2    |
| X            | Cell motility                                                            | 4    | 5    | 4    | 4    | 5    | 4    | 5    | 4    | 3    | 2    | 3    | 2    | 2    | 2    | 2    |
| GM           | Signal transduction mechanisms                                           | 4    | 7    | 4    | 6    | 4    | 5    | 8    | 5    | 4    | 2    | 2    | 3    | 1    | 0    | 0    | 0    |
| GM           | Carbohydrate transport and metabolism                                    | 3    | 1    | 3    | 3    | 3    | 4    | 4    | 4    | 1    | 0    | 1    | 1    | 2    | 1    | 0    |
| KT           | Cell wall/membrane/envelope biogenesis                                   | 3    | 3    | 5    | 3    | 2    | 2    | 2    | 2    | 3    | 2    | 1    | 0    | 2    | 1    | 1    |
| EH           | Cell motility                                                            | 2    | 3    | 4    | 4    | 4    | 5    | 5    | 4    | 3    | 3    | 2    | 1    | 3    | 4    | 4    |
| IQ           | Signal transduction mechanisms                                           | 2    | 3    | 4    | 4    | 4    | 5    | 5    | 4    | 3    | 3    | 2    | 1    | 3    | 4    | 4    |
| MN           | Signal transduction mechanisms                                           | 2    | 3    | 4    | 4    | 4    | 5    | 5    | 4    | 3    | 3    | 2    | 1    | 3    | 4    | 4    |
| GM           | Cell wall/membrane/envelope biogenesis                                   | 2    | 2    | 2    | 1    | 1    | 1    | 1    | 1    | 1    | 0    | 0    | 0    | 0    | 1    | 1    |
| KT           | Cell motility                                                            | 2    | 2    | 2    | 1    | 1    | 1    | 1    | 1    | 1    | 0    | 0    | 0    | 0    | 1    | 1    |

mi, Microcystis aeruginosa PCC 7941; ap, Aphanizomenon flos-aquae PCC 7905; P, Pseudomonas putida EK59; S, Stenotrophomonas rhizophila EK20; D, Delftia sp. F45; A, Acinetobacter beijerinckii F107.
| COG category | Function | miA1 | miA2 | miD1 | miD2 | miP1 | miP2 | miS1 | miS2 | apA1 | apA2 | apD1 | apD2 | apP1 | apP2 | apS1 | apS2 |
|-------------|---------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|------|
| J           | Translation, ribosomal structure, and biogenesis | 106 | 87 | 60 | 64 | 84 | 88 | 108 | 83 | 112 | 107 | 102 | 91 | 87 | 85 | 118 | 121 |
| R           | General function prediction only | 91 | 82 | 54 | 59 | 78 | 78 | 107 | 70 | 113 | 105 | 107 | 78 | 80 | 66 | 118 | 116 |
| E           | Amino acid transport and metabolism | 85 | 69 | 51 | 45 | 63 | 64 | 94 | 68 | 90 | 91 | 68 | 63 | 65 | 53 | 97 | 97 |
| H           | Coenzyme transport and metabolism | 64 | 52 | 34 | 34 | 56 | 60 | 75 | 42 | 74 | 71 | 57 | 55 | 46 | 43 | 89 | 87 |
| S           | Function unknown | 60 | 44 | 29 | 30 | 49 | 42 | 69 | 44 | 80 | 67 | 63 | 52 | 50 | 46 | 91 | 91 |
| C           | Energy production and conversion | 58 | 49 | 44 | 42 | 49 | 52 | 60 | 49 | 68 | 64 | 56 | 55 | 49 | 48 | 48 | 67 |
| P           | Inorganic ion transport and metabolism | 58 | 42 | 35 | 38 | 35 | 44 | 57 | 41 | 71 | 57 | 52 | 57 | 40 | 42 | 77 | 69 |
| M           | Cell wall/membrane/envelope biogenesis | 54 | 50 | 34 | 39 | 47 | 45 | 61 | 51 | 44 | 67 | 55 | 52 | 47 | 48 | 48 | 70 |
| O           | Posttranslational modification, protein turnover, chaperones | 50 | 41 | 35 | 38 | 40 | 48 | 55 | 44 | 61 | 61 | 55 | 52 | 48 | 45 | 69 | 66 |
| G           | Carbohydrate transport and metabolism | 51 | 41 | 36 | 38 | 47 | 54 | 42 | 56 | 51 | 53 | 47 | 43 | 40 | 59 | 56 |
| L           | Replication, recombination, and repair | 40 | 31 | 26 | 26 | 33 | 32 | 49 | 27 | 60 | 48 | 42 | 38 | 34 | 33 | 60 | 61 |
| F           | Nucleotide transport and metabolism | 35 | 26 | 23 | 24 | 29 | 30 | 36 | 24 | 36 | 33 | 26 | 21 | 22 | 43 | 40 |
| I           | Lipid transport and metabolism | 24 | 25 | 17 | 17 | 20 | 20 | 26 | 22 | 31 | 26 | 21 | 20 | 22 | 18 | 31 | 31 |
| V           | Defense mechanisms | 22 | 21 | 13 | 14 | 20 | 19 | 25 | 16 | 22 | 23 | 20 | 17 | 16 | 15 | 23 | 26 |
| T           | Signal transduction mechanisms | 20 | 19 | 15 | 18 | 19 | 19 | 26 | 20 | 31 | 32 | 25 | 19 | 23 | 35 | 32 |
| K           | Transcription | 18 | 15 | 9 | 12 | 16 | 15 | 20 | 11 | 26 | 27 | 23 | 19 | 15 | 18 | 33 | 29 |
| U           | Intracellular trafficking, secretion, and vesicular transport | 9 | 10 | 7 | 8 | 10 | 11 | 9 | 15 | 13 | 13 | 12 | 9 | 7 | 15 | 16 |
| D           | Cell cycle control, cell division, chromosome partitioning | 10 | 9 | 6 | 7 | 8 | 9 | 10 | 6 | 14 | 12 | 14 | 11 | 10 | 9 | 15 | 13 |
| Q           | Secondary metabolite biosynthesis, transport, and catabolism | 10 | 8 | 7 | 6 | 9 | 7 | 8 | 8 | 11 | 10 | 11 | 7 | 8 | 6 | 17 | 16 |
| X           | Mobiome: prophages, transposons | 9 | 6 | 8 | 6 | 8 | 7 | 9 | 8 | 17 | 15 | 14 | 10 | 10 | 12 | 15 | 16 |
| IMG/GenBank | Photosynthesis | 32 | 32 | 18 | 18 | 31 | 31 | 26 | 26 | 18 | 18 | 17 | 17 | 18 | 18 | 18 |
| IMG/GenBank | Gas vesicle | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 2 | 1 | 1 | 1 | 1 | 2 | 2 |
| EH          | Amino acid transport and metabolism | 6 | 6 | 3 | 4 | 6 | 6 | 4 | 4 | 4 | 4 | 4 | 4 | 4 | 4 | 4 |
| NW          | Coenzyme transport and metabolism | 6 | 6 | 3 | 4 | 6 | 6 | 4 | 4 | 4 | 4 | 4 | 4 | 4 | 4 | 4 |
| HR          | Cell motility | 4 | 4 | 3 | 4 | 4 | 4 | 3 | 4 | 2 | 5 | 6 | 3 | 4 | 5 | 3 | 7 |
| TK          | Extracellular structures | 4 | 4 | 2 | 4 | 3 | 4 | 4 | 2 | 6 | 4 | 5 | 4 | 5 | 3 | 6 | 6 |
| EF          | General function prediction only | 4 | 3 | 4 | 3 | 4 | 4 | 4 | 4 | 4 | 3 | 3 | 3 | 3 | 4 | 3 |
| N           | Signal transduction mechanisms | 3 | 3 | 1 | 2 | 2 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 |
| NT          | Transcription | 3 | 3 | 1 | 2 | 2 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 |
| PR          | Nucleotide transport and metabolism | 3 | 3 | 1 | 2 | 2 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 |
| HI          | Coenzyme transport and metabolism | 2 | 2 | 1 | 2 | 1 | 1 | 1 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 |
| EQ          | Lipid transport and metabolism | 2 | 2 | 1 | 2 | 1 | 2 | 1 | 3 | 3 | 3 | 3 | 3 | 3 | 3 | 3 |
| IMG/GenBank | Secondary metabolite biosynthesis, transport, and catabolism | 1 | 0 | 1 | 0 | 1 | 0 | 2 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 1 |

Note: mi, Microcystis aeruginosa PCC 7941; ap, Aphanizomenon flos-aquae PCC 7905; P, Pseudomonas putida EK59; S, Stenotrophomonas rhizophila EK20; D, Delftia sp. F45; A, Acinetobacter beijerinckii F107.
upregulated, albeit this effect was not statistically significant. Three outer membrane-coding genes, murein \textit{\text{\text{DD-endopeptidase (COG0739), a phospholipase (COG2829), and a}} penicillin V acylase/amidase (COG3049) (at log2 fold changes in miS/apS of 1.9/0.9, 1.3/1.7, and 1.6/1, respectively), were differentially expressed in \textit{S. rhizophila EK20 cocultures. In miP, the expression of a muramoyl-tetrapeptide carboxypeptidase (COG1619; log2 fold change, 1.1) was increased, while in apP, the expression of an N-acetyl muramoyl--\text{\text{L-alanine amidase (COG0860; log2 fold change, 1.4) was increased.}} Murein \textit{\text{DD-endopeptidase (score, 9.92) is predicted with a high degree of certainty to be localized in the outer membrane of the bacterial cell and muramoyl-tetrapeptide carboxypeptidase in the cytoplasm (score, 9.26), while the other predicted lytic genes have unknown or multiple localizations in the cell.}}

**Differential expression gene profile of cyanobacteria.** Similar to the expression profiles of the heterotrophic organisms, cyanobacteria were also characterized by clear species-specific patterns (Fig. 6). Unlike the heterotrophs, monoculture control treatments did not cluster separately from coculture treatments (Fig. 6). Differential expression analyses revealed 34 genes that were significantly upregulated in at least one of the coculture treatments and 31 genes that were significantly downregulated in at least one of the coculture treatments compared to the monoculture controls (Fig. 7a and b).
In contrast to the set of differentially expressed genes among the heterotrophic organisms, the expression profiles in the cyanobacteria responded less consistently to the experimental manipulation treatments (Fig. 7a and b).

Some expression patterns indicated a contrasting physiological response in *M. aeruginosa* PCC 7941 and *A. flos-aquae* PCC 7905. Several genes involved in photosynthesis were significantly and consistently downregulated in the *M. aeruginosa* cocultures (photosystem proteins, phycobilisome, and allophycocyanin; Fig. 7b), while the light-harvesting protein Lhl4 was consistently significantly upregulated in miA and miD cocultures (Fig. 7a). In *A. flos-aquae*, protein-coding genes involved in photosynthesis (FoF1-type ATP synthase and RuBisCO) were significantly upregulated in apA and apD cocultures, while the same genes were downregulated when these heterotrophs were cocultured with *M. aeruginosa* (Fig. 7a and b). Such contrasting regulation response was also observed for genes in the COG category translation, ribosomal structure, and biogenesis (Fig. 7a and b). The *A. flos-aquae* gas vesicle structural protein (GenBank accession no. WP_015080752.1) was significantly upregulated in all cocultures involving this organism, while the corresponding gene in *M. aeruginosa* was significantly downregulated in miD. The nitrogenase molybdenum-iron protein, involved in nitrogen fixation, was downregulated in all *A. flos-aquae* cocultures (statistically significant in two cocultures) and three *M. aeruginosa* cocultures (Fig. 7b).

**DISCUSSION**

In this study, we isolated, cocultured, and analyzed four bacterial antagonists, *S. rhizophila* EK20, *P. putida* EK59, *Delftia* sp. F45, and *A. beijerinckii* F107, with either *M. aeruginosa* PCC 7941 or *A. flos-aquae* PCC 7905. The type of interaction between...
different heterotrophs and cyanobacteria in cocultures was revealed by several bacterial responses.

**Heterotrophic responses in coculture.** Outer membrane proteins have been described in *Escherichia coli* as multifunctional proteins that maintain the integrity of cell shape, diffusion, and release of molecules, while also contributing to virulence by inhibiting the host defense system (21). A few outer membrane proteins that were significantly differentially expressed in our experiments corroborate the microscopic observation of direct physical interactions between heterotrophs and cyanobacteria (Fig. 1, 2, and 4a). Additionally, the consistent upregulation of genes coding for the formation of flagella in *S. rhizophila* EK20 and *P. putida* EKS9 cocultures suggests cell-to-cell contact antagonism in which heterotrophs appeared to be actively seeking out and attaching to cyanobacterial targets (Fig. 1 and 4a). This is not surprising, as flagellar proteins have previously been linked to virulence, biofilm formation, and adhesion (22). Furthermore, an important role of the fasciclin I domain (FAS1) in cell adhesion has recently been described in *Mycobacterium tuberculosis* and *Rhodobacter sphaeroides* (23). This protein-coding gene was significantly upregulated in *S. rhizophila* EK20 cocultures, suggesting a role in aggregation and biofilm formation on cyanobacterial surfaces (Fig. 1).

Differential expression of several genes that fall into the COG category energy production and conservation (Fig. 4a and b) implies a switch in organic matter processing through the glyoxylate shunt instead of the full Krebs cycle in *S. rhizophila* EK20 and *P. putida* EKS9 (24, 25). Parameters that have been shown to induce upregulation of the glyoxylate shunt are, for example, the scarcity of high-quality organic matter for cell growth (26) or iron deficiency (27–29). For our study, it seems reasonable to assume that the switch from Casitone-containing growth medium for heterotrophic controls to a medium without such Casitone subsidies may have caused the observed
upregulation of the glyoxylate cycle. While the algal exudates are likely to represent high-quality organic substrates in the coculture treatments, the ambient concentration of such substrates in coculture may not be sufficient to satisfy the heterotrophic carbon demand.

Previous work has shown that *Shewanella* sp. strain W3-18-1 (*Shewanella* W3-18-1) and *Synechococcus* sp. strain 7002 (*Synechococcus* 7002) grown in carbon-limited coculture have an efficient strategy to overcome oxidative stress conditions. This involves an exchange of metabolites and increase in the availability of Fe ions by means of extracellular Fe chelators produced by *Shewanella* W3-18-1, which can then be acquired by *Synechococcus* 7002 for metabolic use (30). This might explain the upregulation of the gene coding for 5-formyltetrahydrofolate cyclo-ligase in one of the *P. putida* EK59 cocultures and all *S. rhizophila* EK20 cocultures (Fig. 4a). This ligase would facilitate the conversion of 5-formyltetrahydrofolate to 5,10-methylenetetrahydrofolate, a product essential for the metabolism of one-carbon compounds likely released by the cyanobacteria (31, 32). In addition, all four heterotrophs featured high expression of detoxification genes, such as catalase and alkyl hydroperoxide reductase (Table S4). The products of these genes are used for protection from radiation-induced cell damage and oxidative stress and have been shown to be highly expressed in *Escherichia coli* and *Deinococcus radiodurans* (33). The high expression of peroxiredoxin, known to protect...
bacteria from toxic peroxides, was another indication that the heterotrophs suffer from stress (34).

Essential functional categories for growth maintenance, such as ribosomal proteins, translation, and transcription-processing factors, and energy production were highly expressed in all four heterotrophs (35) (Table 3), even though some of these genes were downregulated compared to their respective controls (Fig. 4b). Cyanobacteria play an important role in providing heterotrophic bacteria with newly fixed nitrogen and reduced carbon compounds. For example, a mesocosm experiment in the Baltic Sea showed that isotopically labeled fixed nitrogen from *Aphanizomenon* was taken up by attached heterotrophic bacteria (36). This implies that the changes in organic matter supply as well as oxygen stress are likely to have a significant impact on the growth of heterotrophs in coculture.

Stenotrophomonas spp. possess a large number of hydrolytic enzymes which act as antibacterial and antifungal compounds by degrading structural and functional proteins of host cells (37–39). Accordingly, the significant upregulation of *L,D*-transpeptidase in *S. rhizophila* EK20 possibly disrupts the cross-linkages within the peptidoglycan layer of the cocultured cyanobacteria. In addition, the upregulation of a few predicted extracellular hydrolytic genes in both *S. rhizophila* EK20 and *P. putida* EK59 indicate that heterotrophs were probably initiating the production of extracellular enzymes. These could be causing the distortion in cyanobacterial cells that was observed microscopically after 96 h of incubation (Fig. 1 and 2) and supports the complete plaque formation seen after 3 to 4 days of incubation (see Materials and Methods).

**Cyanobacterial responses in coculture.** A comparison of the expression patterns in cyanobacterial monoculture controls to those in cocultures identified a small number of genes that were consistently up- or downregulated (Fig. 7a and b). Several photosynthetic genes were downregulated in the *M. aeruginosa* PCC 7941 cocultures (Fig. 7b), suggesting that photosynthesis was inhibited in this organism in response to the activity of the heterotrophs. It was previously reported that microorganisms minimize energy acquisition and metabolism when exposed to environmental changes (40). In addition, it has been shown that *Bacillus mycoides* B16 attached to *M. aeruginosa* in coculture experiments caused a deterioration in light exposure, resulting in the accu-
mulation of glycogen, poly-beta-hydroxybutyrate, and cyanophycin (41). This indicates that either the presence of heterotrophs directly influences photosynthetic activities of the autotroph organisms or that there is an indirect effect by induction of clumps or aggregate formation limiting light exposure (Fig. 1 and S2). The upregulation of the gene for the alternative light-harvesting protein Lhl4 (Fig. 7a) may be a strategy for M. aeruginosa PCC 7941 to maintain basal photosynthetic activity under adverse conditions.

In contrast, A. flos-aquae PCC 7905 cocultures featured an upregulation of photosynthetic genes and genes contributing to gas vesicles which provide regulation of cellular buoyancy. The effect of buoyancy on A. flos-aquae has been studied in the Baltic Sea, where a 2-fold increase in photosynthesis was observed in response to wind-induced mixing events (42). Moreover, the gene coding for the nitrogenase molybdenum-iron protein, the central gene of nitrogen fixation, was consistently downregulated in all A. flos-aquae PCC 7905 cocultures. This implies that the attachment of S. rhizophila EK20 and P. putida EK59 to heterocysts might have an effect on nitrogen fixation-coding genes in the A. flos-aquae PCC 7905 cocultures. However, the same photosynthesis-coding genes were significantly upregulated in A. flos-aquae PCC 7905-Delftia sp. F45 and A. beijerinckii F107 coculture, even if they were not predominantly attached to heterocysts. This indicates that there is a specific response of A. flos-aquae PCC 7905 to different heterotrophic strains, or it may be that Delftia sp. F45 and A. beijerinckii F107 had a weaker interaction with the photosynthetic cells, at least compared to that with S. rhizophila EK20 and P. putida EK59.

In keeping with the four cocultured heterotrophs, the cyanobacteria also featured high expression of growth-related genes, such as ribosomal proteins, amino acid and coenzyme synthesis, and energy production by the Krebs cycle, for maintenance of growth (Table S5). This suggests that cyanobacteria were actively growing without being critically starved for energy or nutrients.

The cyanobacterial model organism Synechocystis sp. strain PCC 6803 harbors 47 pairs of type II toxin-antitoxin systems (TA systems) that exhibit RNase activity and which may have other additional undiscovered functions (43). The production of antitoxin components is usually induced by stress responses linked to reversible growth inhibition or cell death. For example, in order to adapt to environmental stresses via reversible growth arrest, Anabaena sp. strain PCC 7120 expressed genes for the chromosomal type II toxin-antitoxin systems (44). Both of our studied cyanobacterial strains also expressed high levels of several antitoxin components, likely representing defense mechanisms to maintain their growth under stress conditions (Tables 4 and S5).

Conclusions. Our study provides new information on the biology of heterotrophic bacteria capable of lysing cyanobacteria and, in doing so, identifies diverse physiological traits that likely shape these interactions. The overall transcriptional responses of heterotrophs in coculture with cyanobacteria indicate that their interaction could be divided into four main categories: (i) cell-to-cell contact, (ii) nutrient and space competition with cyanobacteria, (iii) entrapment of cyanobacteria with aggregate formation, and (iv) production of extracellular compounds to disrupt or damage cyanobacterial cells. In contrast, heterotroph aggregate formation interferes with M. aeruginosa PCC 7941 photosynthetic processes, and the preferential attachment of the different heterotrophs to either heterocyst or photosynthetic cells of A. flos-aquae PCC 7905 initiates contrasting antagonistic gene expression related to nitrogen fixation and photosynthesis processes. We conclude that the metatranscriptome analysis of samples collected after 24 h of a coculture experiment was only a snapshot of the antagonistic interaction and did not capture the complete mechanism underlying the antagonistic interaction.

MATERIALS AND METHODS
Cyanobacterial cultures. Two cyanobacterial strains, Microcystis aeruginosa PCC 7941 and Aphanizomenon flos-aquae PCC 7905, were obtained from the Pasteur Culture Collection as experimental model
strains for lytic tests. *M. aeruginosa* PCC 7941 is a unicellular, planktonic, and toxin-producing strain capable of buoyancy regulation with gas vesicles. *A. flos-aquae* PCC 7905 is a filamentous and toxin-producing strain with the ability to fix nitrogen. Both strains represent idealized models of bloom-forming freshwater cyanobacteria of environmental concern. Both strains were cultured and maintained in a cyanobacterial BG11 freshwater medium (45) at 20°C under photosynthetically active radiation (PAR) of approximately 5 μE m⁻² s⁻¹ (IL-1400 radiometer with PAR sensor), with a 12-h light/12-h dark cycle.

**Isolation of cyanolytic bacteria.** Water samples from three eutrophic lakes in south-central Sweden, Erken, Eknol, and Funbojsjön, were collected during a summer period of high cyanobacterial biomass to screen for cyanolytic bacteria. Lytic bacterial strains were isolated by first preparing a cyanobacterial lawn according to the method of Whyte et al. (46). A volume of 15 ml of each of the axenic cyanobacterial cultures (PCC 7941 and PCC 7905) at cell densities of approximately 4.8 × 10⁸ cells · ml⁻¹ was mixed with 0.5 ml of unfiltered lake water. The mixed water samples were then gently filtered onto 0.22-μm-pore-size polyether sulfonate membranes (47 mm diameter; Gelman Supor). Filters were rapidly placed on top of BG11 agar plates in individual petri dishes, sealed with Parafilm, and incubated under the same light and temperature conditions described above. Viral and bacterial plaque formation was monitored for 3 to 4 days using a stereo microscope (Olympus SZ61) (Fig. S1). Bacterial colonies that formed lytic zones in the cyanobacterial lawns were further purified by repeated streaking onto BG11 agar supplemented with 0.2% Casitone.

**Lytic test.** One hundred bacterial isolates were recovered from the cyanobacterial lawns. In order to confirm that the colonies maintained their lytic activity, we regrew them in BG11 liquid medium supplemented with 0.2% Casitone at 25°C for 48 h and then inoculated 10 to 20 μl of pure culture suspension in triplicate on a cyanobacterial lawn. The cyanobacterial lawns were photographed after 24 to 72 h of incubation (Canon G9 zoom lens 6 × 15 fixed to a Kaiser R51 camera stand), and both the colony diameter and the lytic halo were quantified using Image J (public domain image processing and analysis in Java). The enzymatic activity was then reported as the colony diameter divided by the halo diameter ratio, expressed as the enzymatic index (EI) (47).

**16S rRNA sequencing.** Thirty-one pure colonies possessing lytic activity were chosen for 16S rRNA gene sequencing. DNA was extracted using the PowerSoil DNA extraction kit (Mo Bio Laboratories, Inc., CA, USA). PCR primers 341F (5’-CTACGGGNGGCWGCAG-3’) and 805R (5’-GACTACHVGGGTATCTAATC-3’) were used for 16S rRNA amplification. PCR was conducted in a 20-μl volume using 1 U of Taq DNA polymerase (New England BioLabs), 0.25 μM primers, 200 μM dNTP mix, 0.4 μg of bovine serum albumin, and 1 μl of DNA template. The thermal program consisted of an initial 95°C denaturation step for 5 min, a cycling program of 95°C for 40 s, 53°C for 40 s, and 72°C for 60 s, and a final extension step for 7 min for 20 cycles. Amplicons were purified with a Qiagen gel-purification kit (Qiagen, Germany) and quantified with a fluorescent stain-based kit (PicoGreen; Invitrogen), BigDye Terminator version 3.1 (Applied Biosystems, Paisley, UK); primer 341F was used for sequencing reactions, and the samples were subsequently analyzed by capillary electrophoresis on an ABI3730XL DNA analyzer (Applied Biosystems).

**Antagonistic interaction between cyanobacteria and heterotrophic bacterial strains.** Four phylogenetically diverse heterotrophic strains with strong and reproducible cyanolytic activity were selected for further experiments: *Stenotrophomonas rhizophila* EK20, *Pseudomonas putida* EK59, *Delftia* sp. F45, and *Acinetobacter beijerinckii* F107. Prior to performing coculture experiments, the viability and purity of heterotrophs were verified by inoculation in LB agar plates and by microscopic observation. Inoculation in Luria broth was also used to verify that the cyanobacterial cultures were axenic. Heterotrophic bacteria were quantified by flow cytometry (CyFlow space; Partec, Münster, Germany) after mixing the sample with a 1.25 μM final concentration of SYTO13 nucleic acid stain (Invitrogen, Eugene, OR, USA). The sample flow rate was 4 μl/s, and the sheath fluid was Milli-Q water. SYTO13-stained cells were excited by blue laser (488 nm) with gain setting of 335 nm for side scatter (SSC), 450 for green fluorescent light (FL), and 240 for forward-scatter light (FSC). Bacterial cells were identified according to their FSC and green fluorescence patterns. The average relative cell size was estimated using the mean FSC value and referred as individual cell size (ICS). Biovolume (BV) of the samples was calculated by multiplying ICS with respective bacterial abundance (BA) (48). Cyanobacterial abundance was assessed by fluorescence microscopy counts. The initial ratio of heterotrophs to cyanobacteria in the cocultures was set to approximately 1:1. Each heterotrophic strain was cultured in darkness using liquid BG11 medium supplemented with 0.2% Casitone at 25°C. Cells were harvested after 48 h of incubation by centrifugation (10,000 rpm, 15 min, 25°C) and washed and suspended in Casitone-free BG11 medium before mixing with *M. aeruginosa* PC 7941 or *A. flos-aquae* PCC 7905. Cocultures were maintained under photosynthetically active radiation (PAR) conditions of approximately 5 μE m⁻² s⁻¹ (IL-1400 radiometer with PAR sensor) with a 12-h light/dark cycle. One milliliter of sample for cell counts was collected from each culture after 6, 24, and 96 h of incubation and fixed with 2% formaldehyde. Additionally, 25 ml of culture was collected by rapid filtration onto 0.2-μm-pore-size Supor membrane filters (47 mm diameter; Pall Corporation) and was immediately frozen at −80°C for later RNA isolation. Control samples of *M. aeruginosa* PCC 7941 and *A. flos-aquae* PCC 7905 were collected separately under the same light conditions mentioned above. Heterotrophic controls for *S. rhizophila* EK20 and *P. putida* EK59 were grown in darkness at 25°C in liquid BG11 medium supplemented with 0.2% Casitone and were collected after 48 h of incubation.

**Epifluorescence microscopy.** Formaldehyde-preserved samples were filtered onto black polycarbonate membrane filters (0.22-μm-pore-size, 25 mm) placed on top of a support filter to ensure even dispersion of cells across the filter area. 4′,6-Diamidino-2-phenylindole was used for DNA staining at a final concentration of 100 μg/ml, whereas NanoOrange (Life Technologies) was used for protein staining.
at 5.0 μg/ml, according to the manufacturer’s instructions. Filters were incubated for 15 min before excess stain was removed by vacuum filtration and further rinsed with deionized water. Image acquisition for each filter section was done using an AxioPlan II epifluorescence microscope with a digital camera (AxioCam) installed (Carl Zeiss, Germany). Cell concentrations of both heterotrophs and cyanobacteria were calculated from the equation \((C \times A_3)/(n \times V \times A_2)\), where \(C\) is the total number of bacteria counted, \(A_3\) is the effective area of the membrane filter, \(A_2\) is the area of the observed grid, \(n\) is the number of grids counted, and \(V\) is the volume of sample filtered. Two sample replicates were counted, with the number of counted grids ranging from 8 to 15 grids per sample. Rough counts of normal heterotrophs and cyanobacterial cell shapes were counted due to the formation of heterotrophic aggregates after 24 h of incubation.

**RNA isolation and sequencing.** Two biological replicates for each combination of the four heterotrophic strains and the two cyanobacteria (\(M. \text{aeruginosa} \ PCC \ 7941\) or \(A. \text{flos-aquae} \ PCC \ 7905\)) were selected for total RNA isolation and metatranscriptome sequencing. Metatranscriptome analyses were carried out for samples retrieved after 24 h of incubation. RNA extraction was performed with the PowerWater RNA isolation kit, as per the manufacturer (Mo Bio Laboratories, USA). Digestion of contaminating DNA was carried out using the Turbo DNA-free kit (Invitrogen, Life Technologies, Europe BV) and the absence of DNA verified by negative PCR amplification of the 16S rRNA gene. RNA integrity was assessed on an Agilent 2100 Bioanalyzer (Agilent Technologies, Inc., USA). First-strand cDNA was synthesized by RevertAid H Minus first-strand cDNA synthesis kit (Thermo Scientific, USA), followed by second-strand cDNA synthesis using a double-stranded DNA synthesis kit (catalog no. E6111; New England BioLabs). The resulting double-stranded DNA was purified by QiAquick PCR extraction kit (Qiagen) and the concentration was measured using the PicoGreen double-stranded DNA assay (Invitrogen). Sequencing was performed on each of two replicate coculture and control samples of \(Stenotrophomonas \ rhizophila \ KE\,20\), \(Pseudomonas \ putida \ EKS\,9\,\), \(M. \text{aeruginosa} \ PCC \ 7941\), and \(A. \text{flos-aquae} \ PCC \ 7905\) (SNP&SEQ Technology Platform at Uppsala University). Libraries were prepared from 20 ng of cDNA using the ThruPLEX FD library preparation kit (Rubicon Genomics) and used for two full runs on the Illumina MiSeq instrument running in paired-end 2 × 300-bp mode with version 3 chemistry.

**Bacterial reference genomes.** The draft genomes of the four selected heterotrophic strains (\(S. \text{rhizophila} \ KE\,20\), \(P. \text{putida} \ EKS\,9\,\), \(Deltia \ sp. \ F\,45\,\), and \(A. \text{beijerinckii} \ F\,107\)) were sequenced by MiSeq using \(2 \times 300\) chemistry, as described above. Reads were first assembled using the SPAdes genome assembler, with contigs \(\geq 2,000\) bp length with high coverage \(\geq 190\times\) submitted to Integrate Microbial Genomes (IMG) for genome annotation using the IMG pipeline (49). The IMG taxon identification (ID) of each strain is as follows: \(Deltia \ sp. \ F\,45\,\), Gp011381; \(A. \text{beijerinckii} \ F\,107\,\), Gp011397; \(S. \text{rhizophila} \ KE\,20\,\), Gp011470; and \(P. \text{putida} \ EKS\,9\,\), Gp011469.

**RNA processing.** RNA reads were assembled using the de novo assembler Trinity (50), according to the assembly protocol described by Hass et al. (51). The de novo transcriptome assembly was performed by the three constituent components: *Inchworm*, which assembles unique sequences to construct transcripts; *Chrysalis*, which makes groups of transcript contigs form components and generate de Bruijn graphs for each component; and *Butterfly*, which compacts and extracts all probable sequences from each graph. Prior to the mapping of raw RNA reads to the assembly, they were quality trimmed using Sickle (https://github.com/najoshi/sickle) (quality cutoff, 20; minimum read length, 75 bp), and noncoding RNA was removed via the SortMerna software (52). The trimmed paired reads were mapped on the assembled Trinity contigs using the Bowtie 2 aligner (53). BLASTP was used to align and annotate Trinity-assembled contigs to the reference genome amino acid sequences of the 4 heterotrophic bacteria (described above) and a group of *Microcystis* genomes (\(M. \text{aeruginosa} \ PCC \ 7041\,\) \(M. \text{aeruginosa} \ DIAN-CHI905\,\) \(M. \text{aeruginosa} \ TAIH\,U98\,\) \(M. \text{aeruginosa} \ PCC \ 9432\,\) \(M. \text{aeruginosa} \ PCC \ 9717\,\) and \(M. \text{aeruginosa} \ PCC \ 9808\)) as well as the *A. \text{flos-aquae} \ NIES-81* genome (BioProject accession no. PRJNA232534). The BLASTP E value cutoff used was \(e^{-0.2}\). While the genome data provided by IMG are annotated to COG homologs, this is not the case for genome data stored in GenBank. Therefore, a BLASTP search of *A. \text{flos-aquae} \ NIES-81* protein data against the COG reference database was performed with the same E value cutoff provided by other COG annotations \(E < 10^{-0.2}\). RNA reads mapping to the respective reference genome were binned and either assigned to a COG homolog or, if this information was not available, to the overall functional annotation provided by IMG/GenBank for each metagenomic open reading frame. An overview containing the number of obtained reads for each sample is available in Table S1. The localization of lytic genes was predicted by the localization prediction tool PSORTb (http://www.psort.org/psortb/).

**Estimation of highly expressed genes.** Highly expressed genes were extracted from the relative abundance counts (derived after normalization of the raw count data to the contig length) of the expressed genes in heterotrophs and cyanobacteria. For each replicate pair, we sorted genes by their respective functional annotation provided by IMG/GenBank for each metagenomic contig to the reference genome amino acid sequences of the 4 heterotrophic bacteria (described above) and a group of *Microcystis* genomes (\(M. \text{aeruginosa} \ PCC \ 7041\,\) \(M. \text{aeruginosa} \ DIAN-CHI905\,\) \(M. \text{aeruginosa} \ TAIH\,U98\,\) \(M. \text{aeruginosa} \ PCC \ 9432\,\) \(M. \text{aeruginosa} \ PCC \ 9717\,\) and \(M. \text{aeruginosa} \ PCC \ 9808\)) as well as the *A. \text{flos-aquae} \ NIES-81* genome (BioProject accession no. PRJNA232534). The BLASTP E value cutoff used was \(e^{-0.2}\). While the genome data provided by IMG are annotated to COG homologs, this is not the case for genome data stored in GenBank. Therefore, a BLASTP search of *A. \text{flos-aquae} \ NIES-81* protein data against the COG reference database was performed with the same E value cutoff provided by other COG annotations \(E < 10^{-0.2}\). RNA reads mapping to the respective reference genome were binned and either assigned to a COG homolog or, if this information was not available, to the overall functional annotation provided by IMG/GenBank for each metagenomic open reading frame. An overview containing the number of obtained reads for each sample is available in Table S1. The localization of lytic genes was predicted by the localization prediction tool PSORTb (http://www.psort.org/psortb/).

**Differential expression analyses.** The transcripts of each strain grown in coculture were compared to their respective control treatment transcripts using a differential expression analysis performed in the R environment with the package DESeq2 (54). For the two heterotrophic bacteria without a heterotrophic-only control, no differential expression analyses were performed. \(P\) values were corrected for multiple testing according to the procedure of Benjamini and Hochberg (55). Transcribed gene orthologs with a corrected \(P\) value (adjusted) of \(< 0.1\) were classed as significantly differentially transcribed genes.
SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/AEM.02634-16.

SUPPLEMENTAL FILE 1, PDF file, 0.8 MB.

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