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## **Actinorhodopsin genes discovered diverse in freshwater habitats and among cultivated Actinobacteria**

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
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## ORIGINAL ARTICLE

# Actinorhodopsin genes discovered in diverse freshwater habitats and among cultivated freshwater *Actinobacteria*

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Microbial rhodopsins are membrane proteins that utilize a retinal chromophore to harvest sunlight for energetic and photosensory functions. Recently, a group of novel rhodopsin sequences named ‘actinorhodopsins’ (ActRs) was hypothesized to exist among uncultured planktonic *Actinobacteria*. ActRs were discovered by mining metagenomic data obtained during the Venter Institute’s Global Ocean Sampling expedition, from a hypersaline lagoon, two estuaries and a freshwater lake. On the basis of these findings, and many studies that show *Actinobacteria* are common inhabitants of lakes, we predicted that ActR genes would likely be present in other freshwater habitats and among the genomes of cultivated *Actinobacteria*. Using degenerate polymerase chain reaction primers, we discovered an ActR gene present in an actinobacterial isolate of the family Microbacteriaceae. Isolate MWH-Uga1 was cultivated prior to this study from a freshwater pond in Uganda and belongs to a group of *Actinobacteria* previously identified in freshwater ecosystems. ActR genes were also discovered present in numerous mixed cultures containing freshwater *Actinobacteria* and among environmental DNA samples obtained from three freshwater sources; a small woodland pond and the Laurentian Great Lakes Superior and Erie. An analysis of small subunit ribosomal RNA genes from metagenomic DNA samples harboring ActR genes suggests that organisms belonging to the *acl* lineage, an uncultured group of *Actinobacteria* commonly present in fresh waters, may utilize rhodopsins. The co-occurrence of an *acl* organism with a specific ActR variant in a mixed culture supports our hypothesis.

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**Subject Category:** microbial ecology and functional diversity of natural habitats

**Keywords:** rhodopsins; actinorhodopsin; freshwater; *Actinobacteria*; Superior; Erie

## Introduction

Microbial rhodopsins are membrane proteins that use a retinal chromophore to harvest solar energy for ion pumping and photoreception. In 2000, Beja *et al.* discovered rhodopsin-encoding genes in the surface waters of the earth’s oceans and designated this novel group of proteins ‘proteorhodopsin’. Their designation was based on the discovery of metagenomic fragments that linked the first discovered of these proteorhodopsin genes to a small subunit ribosomal RNA (SSU rRNA) gene defining the

uncultivated marine SAR86 group of the  $\gamma$ -*Proteobacteria* (Beja *et al.*, 2000). Since then, proteorhodopsin genes have been found in high abundance in the photic zone of the ocean, where they are believed to confer proteorhodopsin-based phototrophy and light-sensing abilities on diverse groups of marine microorganisms (Rusch *et al.*, 2007; Campbell *et al.*, 2008; Fuhrman *et al.*, 2008).

Recently, proteorhodopsin-related genes were discovered through metagenomic analysis of several non-marine aquatic environments sampled during the Venter Institute’s Global Ocean Sampling (GOS) expedition (Rusch *et al.*, 2007). Rusch *et al.* described the presence of these genes among microbes inhabiting the freshwater Lake Gatun (part of the Panama Canal), the estuaries Delaware Bay and Chesapeake Bay, and the hypersaline Punta Cormorant lagoon in Ecuador. In a separate analysis of rhodopsin

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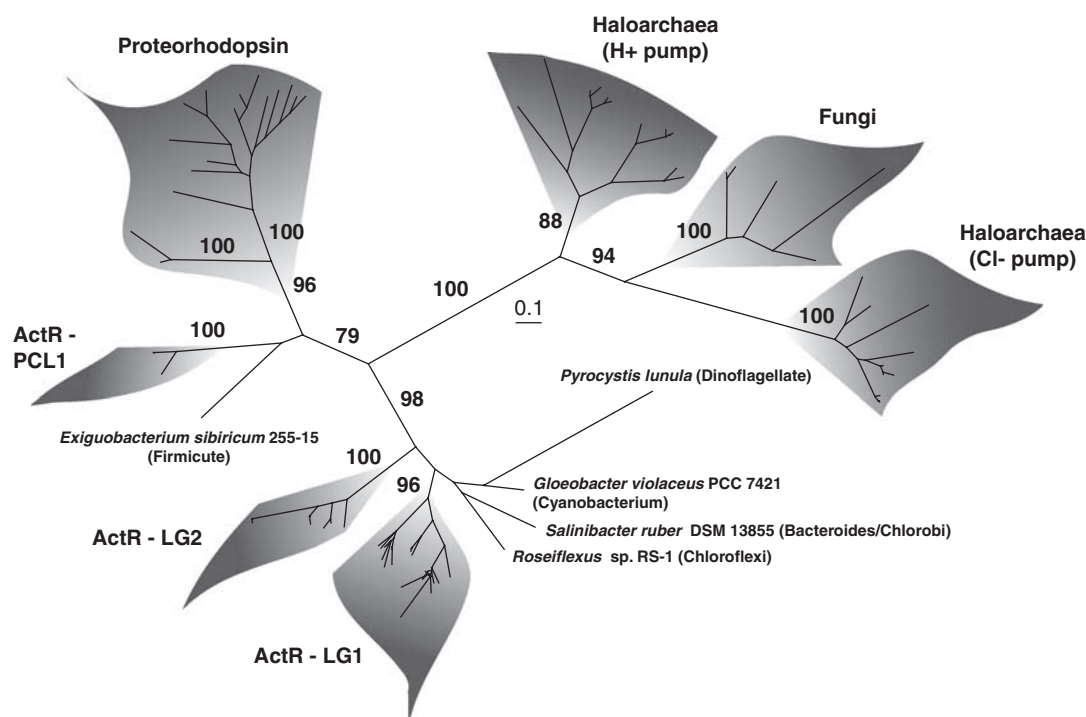
sequences from these non-marine environments in Sharma *et al.* (2008), we described three novel groups of sequences (LG1, LG2 and PCL1, Figure 1) comprising the majority of rhodopsin diversity. Our earlier analyses also consisted of a bioinformatic assessment of GOS-derived (Rusch *et al.*, 2007) metagenomic assemblies containing rhodopsin genes of these three novel clades and found they were almost exclusively linked to open reading frames most closely related to homologues encoded in sequenced actinobacterial genomes. In Sharma *et al.* (2008) we designated these novel clades of rhodopsins collectively as 'actinorhodopsins (ActRs)' based on (1) the aforementioned pattern of gene linkage on GOS metagenomic assemblies, (2) the observation by Rusch *et al.* (2007) that actinobacterial SSU rRNA sequences were abundant in non-marine samples and (3) the knowledge that *Actinobacteria* are common inhabitants of freshwater ecosystems (Warnecke *et al.*, 2004). However in our previous study, we were unable to identify the specific actinobacterial carriers of ActRs from GOS samples. On the basis of our findings in Sharma *et al.* (2008), we predicted that ActR gene sequences would likely be found in other freshwater environments and in the genomes of cultivated *Actinobacteria*. To test these predictions, we designed degenerate primers and used the polymerase chain reaction (PCR) to amplify ActR genes from freshwater metagenomic DNA samples

and *Actinobacteria* cultivated from various freshwater sources.

## Methods

### Sampling and DNA preparation

Point Pleasant Park pond in Halifax, NS was sampled in late summer (early September) 2006. Pond water was collected in a sterile 1 l glass bottle and prefiltered through Whatman paper. Biomass was concentrated onto a 0.2 µm filter using a disposable Nalgene bottle top filter apparatus and the filter was frozen at -20 °C overnight, then the following day DNA was isolated by bead beating a slice (~1/8) of the 0.2 µm filter using the FastDNA SPIN for Soil kit (MP Biomedicals, LLC., Solon, OH, USA). Lake Superior biomass was obtained during a research cruise on the R/V *Blue Heron* in June 2005 during isothermal mixing of the water column. Water was collected from a depth of 25 m at station Castle Danger 1 (CD1; 47.30°N, 91.67°W; see Ivankova *et al.* (2007) for sampling details). Biomass was collected from the central basin of Lake Erie (Environment Canada Station 84, 41° 56' 46" N, 81° 38' 46" W) during a July 2005 cruise aboard the CCGS *Limnos* when the water column was thermally stratified (for sampling details see Wilhelm *et al.*, 2006). DNA for rhodopsin PCR was integrated from



**Figure 1** Best maximum likelihood tree of 81 selected microbial rhodopsin proteins (177 positions). Only selected bootstrap values are displayed. Major clades of rhodopsin sequences are outlined. In Sharma *et al.* (2008), we described three clades of actinorhodopsins (ActR) identified from the Global Ocean Sampling (GOS) expedition's metagenomic samples of non-marine sites (Rusch *et al.*, 2007). In addition to ActR sequences from our earlier study, this tree also contains five full-length LG1 ActR sequences obtained here, from actinobacterial cultures MWH-Uga1, MWH-Dar1, MWH-Dar4, MWH-Ta8, MWH-EgelM2-3. Other sequences were obtained from GenBank searches and the accession numbers for sequences in each group are given in Supplementary file 1.

different sample depths in the Lake Erie water column at 5 m (24 °C), 13.5 m (16.5 °C) and 16.5 m (9 °C). Filters containing Great Lakes biomass were rinsed with 3 ml of TE buffer to remove cells and the resulting suspension was spun down at maximum speed in a table top microcentrifuge at 4 °C. Following centrifugation, the pellet was treated using the bacterial genomic DNA isolation using CTAB protocol of Feil WS and Feil H available on the Joint Genome Institute's recommended protocols page (<http://my.jgi.doe.gov/general/>).

### Rhodopsin PCR

Degenerate PCR primers (Table 1) were designed to amplify partial gene sequences from ActR sequences clades (LG1, LG2 and PCL1) uncovered from our previous study (Sharma *et al.*, 2008). Rhodopsins were amplified from metagenomic DNA samples using either the combined set of three primers targeting ActR clades LG1 and LG2 (Table 1) or a set of three degenerate primers previously designed by Sabehi *et al.* (2005) to target proteorhodopsin. PCR reactions were performed in 20 µl volumes using Eppendorf MasterMix (2.5 ×) containing 1 µl or ~20 ng of DNA, 8 µl of master mix, 1 µl of a 10 µM working solution of each of the three primers (3 µl total) and 8 µl dH<sub>2</sub>O. The amplification steps were: initial denaturation at 95 °C for 4 min, then 40 cycles of 94 °C for 1 min, annealing temperature (46.6 °C for actR primers or 44 °C for pR primers) for 1 min, 68 °C for 30 s, then a final elongation step of 68 °C for 2 min. PCR products of ~330 bp were gel-purified (MiniElute; Qiagen, Valencia, CA, USA), and cloned by using the Invitrogen TOPO-TA kit. Clones were screened by colony PCR using primers flanking the insert in the TOPO vector and amplicons were cleaned using the High Pure PCR Product Purification kit (Roche, Indianapolis, IN, USA) and sequenced using the GenomeLab Dye Terminator Cycle Sequencing with Quick Start kit on a CEQ 8000 genetic analysis capillary sequencer both from Beckman Coulter (Fullerton, CA, USA). Sequences were edited using Sequencher 4.8 from Gene Codes Corporation (Ann Arbor, MI, USA).

**ActR gene amplification from cultivated Actinobacteria**  
All investigated cultures (Supplementary Table S1) were obtained using the filtration-acclimatization method and maintained as outlined in Hahn *et al.* (2004). Genomic DNA was extracted from the cultures using the FastDNA kit and the FastPrep instrument (MP Biomedicals, LLC.) as recommended by the manufacturer. Partial ActR gene sequences were obtained as indicated above and multiple clones were sequenced for each ActR gene fragment amplified and cloned from a single culture. Only culture MWH-EgelM2-3 was discovered to have three ActR sequence variants and the seven remaining cultures were found to harbor only a single actR allele each, which was subsequently verified by direct sequencing of actR PCR products from each culture. A long walk PCR technique designed by Katz *et al.* (2000) was used to obtain full-length ActR gene sequences.

### SSU rRNA PCR

With the exception of the acI strain, SSU rRNA sequences of all other actinobacterial strains were directly sequenced with universal primers (Hahn *et al.*, 2003) or by using the more specific LEAD1 primer (Hahn and Pöckl, 2005). The SSU rRNA gene of the acI actinobacterium from MWH-EgelM2-3 was initially identified with an acI-specific SSU rRNA PCR screen developed by Newton *et al.* (2007) (using AcI.856R 5'-TCGCASAAACCGTGGGAAG-3') and by Warnecke *et al.* (2005). PCR reactions were carried out as above and the amplification cycles were: initial denaturation at 94 °C for 2 min, followed by 30 cycles of 94 °C for 40 s, annealing at 55.5 °C for 40 s, elongation at 68 °C for 30 s and then a single cycle of 68 °C for 2 min. Owing to the low abundance of the acI organism (<0.1% of total bacterial cells) in the culture as revealed by fluorescence *in situ* hybridization (FISH; data not shown), the full-length SSU rRNA gene could not be obtained with universal bacterial primers. The AcI.856.R primer was paired with the Bact.SSU.8F primer (annealing temperature 56.5 °C) in one reaction and the complement of the AcI.856.R primer,

**Table 1** Primers designed to target Actinorhodopsin sequence clades

| ActR clade        | Recovered by PCR | Forward primer(s)  | Reverse primer(s)  | Additional sequence diversity targeted               |
|-------------------|------------------|--|--|--|
| LG1               | Yes              | YRYVDW (5'-TAYMGNTAYGTNGAYTGG-3')<br>RYIDWL <sup>a</sup><br>(5'-MGNTAYATHGAYTGGYT-3')                          | WGVYPI (5'-ATNGGRTANACNCCCCA-3')   | <i>Gloeobacter violaceus</i><br>PCC7421              |
| LG2               | No               | YRYADW (5'-TAYMGNTAYGCNGAYTGG-3')  | WGVYPI (5'-AT NGG RTA NAC NCC CCA-3')  | <i>Roseiflexus</i> sp. RS-1                          |
| PCL1 <sup>b</sup> | No               | RYIDWL <sup>a</sup><br>(5'-MGNTAYATHGAYTGGYT-3')<br>(L/Y/F/Y) RYVDW <sup>a</sup><br>(5'-WWNMGNTAYGTNGAYTGG-3') | NKVGFG (5'-CC RAA NCC NAC YTT RTT-3')<br>GWMIYP (5'-GG RTA DAT CAT CCA NCC-3') | <i>Exiguobacterium sibiricum</i> 255-15 <sup>c</sup> |

<sup>a</sup>Primer sequences that were the same as those from Sabehi *et al.* (2005).

<sup>b</sup>PCL1 primers did not amplify rhodopsin genes from any of the DNA samples obtained in this study.

<sup>c</sup>The rhodopsin sequence of *E. sibiricum* would only be targeted from the top set of PCL1 forward and reverse primers.

AcI.856.F (5'-CTTCCACGGTTTSTGCCA-3') was paired with Bact.SSU.1492R (annealing temperature 49.5 °C) in another reaction to generate two sequence fragments. Concatenation of the two parts into a single longer sequence and analysis by BLAST recover a sequence identical to a previously deposited acI SSU rRNA sequence (EU117876) from Mirror Lake, Wisconsin (Newton *et al.*, 2007). The sequence of the full-length acI SSU rRNA gene was verified by amplification of the entire sequence with match-specific PCR primers and direct sequencing.

#### *Rhodopsin phylogenetic methods*

The tree in Figure 1 was constructed using only full-length sequences. ActR sequences obtained in this study, as well additional ones from the NCBI database representing haloarchaea and fungi, were added to alignments generated in our previous study of ActRs as outlined in Sharma *et al.* (2008), using the program Hmmer (Eddy, 1998). Partial ActR sequences amplified from freshwater metagenomic DNA samples and from cultures are represented in a separate tree in Figure 3. Full-length and partial LG1 sequences identified in Sharma *et al.* (2008) that completely overlapped with those amplified here were also aligned with Hmmer. The site-specific clustering analysis of partial and full-length GOS rhodopsin sequences mapped onto the phylogeny in Figure 3 was performed in Sharma *et al.* (2008). All rhodopsin phylogenetic trees were calculated in the PHYML program version 2.4.5 (Guindon and Gascuel, 2003; Guindon *et al.*, 2005) under the WAG substitution matrix (Whelan and Goldman, 2001), with a discrete  $\Gamma$  distribution (Yang, 1993) divided into eight rate categories and the proportion of invariable sites estimated. A total of 100 PHYML bootstrap replicates were calculated for each tree.

#### *SSU rRNA phylogenetic analyses*

SSU rRNA sequences representing different lineages of freshwater *Actinobacteria* (Hahn *et al.*, 2003; Warnecke *et al.*, 2004) were used as queries in BLAST searches of unassembled metagenomic data sets of non-marine GOS samples at the CAMERA website (<http://camera.calit2.net/>) (Seshadri *et al.*, 2007). SSU rRNA sequences obtained from BLAST searches and the *Actinobacteria* cultures were aligned in ARB (Ludwig *et al.*, 2004), edited manually and a tree was constructed using MEGA4 (Tamura *et al.*, 2007). Note, the number of nucleotide positions used for the tree construction (745) was restricted by the length of some GOS sequences. Analysis of 121 actinobacterial SSU rRNA sequences from assembled GOS metagenomic data (Seshadri *et al.*, 2007) was performed as follows: sequences were aligned as above, then a Neighbor-joining backbone tree was calculated first with reference sequences representing the phylum *Actinobacteria*, which included longer GOS sequences from assembled data and our own searches

of unassembled data (sequence positions 47–1037, *Escherichia coli* numbering). Second, shorter sequences were added to the tree using a parsimony algorithm (within ARB) and this method almost completely conserved the overall tree topology. Sequences were assigned to previously defined SSU rRNA actinobacterial clusters based on their position in the tree.

#### *Nucleotide sequence accession numbers*

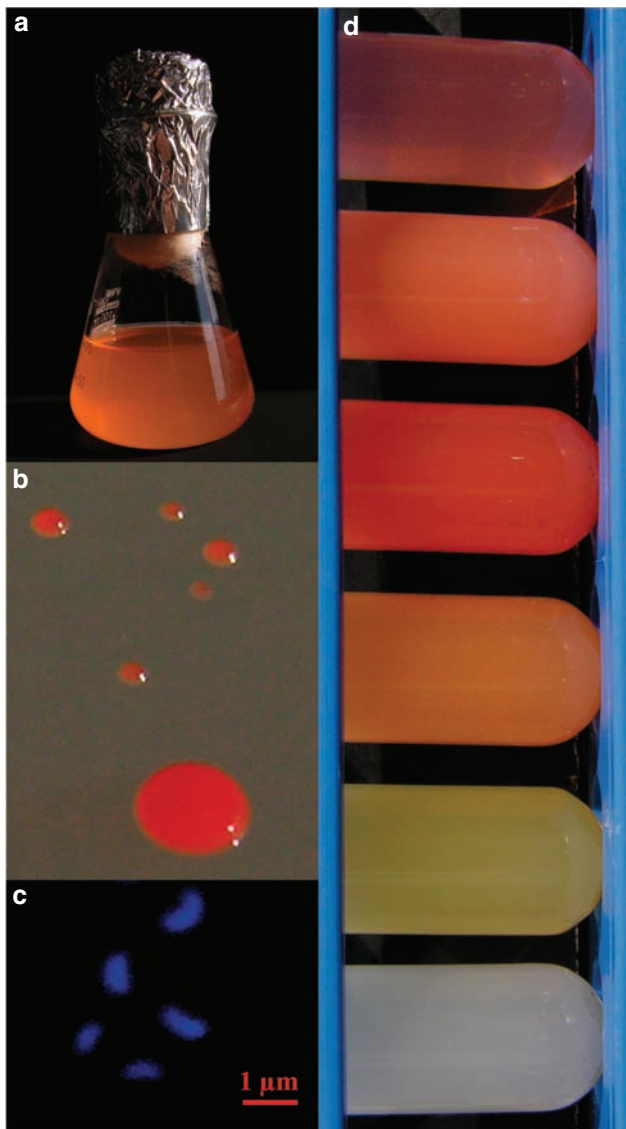
Rhodopsin sequences obtained in this study were deposited in GenBank under the following accession numbers: FJ545138–FJ545171 (Lake Superior), FJ545172–FJ545208 (Lake Erie) and FJ545213–FJ545222 (Luna cultures).

## Results and discussion

To test our hypotheses regarding the environmental and organismal distributions of ActR, we attempted to PCR amplify ActR-encoding gene fragments from both environmental DNA samples and cultivated *Actinobacteria*. Degenerate PCR primers were designed for this study (Table 1) based on the three ActR sequence clades (Figure 1) that we uncovered previously from non-marine GOS samples in Sharma *et al.* (2008). In addition, each primer pair encodes the capability of targeting rhodopsin sequences from additional bacterial phyla (Table 1).

#### *Isolate MWH-Uga1 encodes an LG1-ActR*

In Sharma *et al.* (2008), we used the term 'actinorhodopsin' to designate restricted clades of environmental rhodopsin sequences, which appeared on many GOS-derived metagenomic assemblies (Rusch *et al.*, 2007) linked to genes whose closest relatives were actinobacterial homologues. Although our earlier analysis could not identify the specific organism(s) responsible for ActR sequences from non-marine GOS samples, it predicted that such genes should be present in planktonic *Actinobacteria*. In the current study we investigate a number of cultures containing *Actinobacteria* (Supplementary Table S1) obtained in previous works from a variety of freshwater sources (Hahn *et al.*, 2003, 2004; Hahn and Pöckl, 2005). Our attention was initially drawn to these specific cultures because of the diversity of pigmentation represented by these organisms (Figure 2). Rhodopsin proteins are documented to contribute to similar pigmentation in other rhodopsin-expressing organisms (Bolhuis *et al.*, 2006; Gomez-Consarnau *et al.*, 2007). Alternatively, such coloration is often caused by the presence of carotenoid pigments (Trutko *et al.*, 2005), which increases the probability that these organisms carry rhodopsin genes, because the retinal chromophore of these photoproteins is a by-product of carotenoid biosynthesis (Martinez *et al.*, 2007).



**Figure 2** Phenotypic characteristics of actinobacterial cultures harboring ActR genes. Left column; (a) culture MWH-Dar1 grown in NSY medium, (b) colonies of MWH-Dar1 on NSY agar, (c) microphotograph of MWH-Dar1 cells stained with 4,6-diamidino-2-phenyl indole (DAPI). Right column; (d) 10-fold concentrations of cultures grown in NSY medium. From the top: culture MWH-BeijW16, MWH-EgelM2-3, MWH-Dar1, MWH-Uga1, MWH-Mo1 (actR not detected), *Polynucleobacter* sp. QLW-P1DMWA-1 (*Betaproteobacteria*, rhodopsin-negative based on genome sequence).

Here we validate the prediction that ActR genes are encoded among actinobacterial genomes, through the discovery of a freshwater isolate, strain MWH-Uga1, encoding an ActR gene belonging to the LG1 group outlined in Figure 1. Figure 3 displays the phylogenetic and ecological relationships of LG1 ActRs and shows that the MWH-Uga1 sequence groups among others from the GOS sample of the Punta Cormorant lagoon (Rusch *et al.*, 2007), previously described by us in Sharma *et al.* (2008). Isolate MWH-Uga1 was obtained by Hahn and Pöckl (2005) from a eutrophic pond in Uganda. Phenoty-

pically, this organism is characterized by an extremely small cell size of  $<0.1\ \mu\text{m}^3$  (similar to the cells shown in Figure 2) and orange pigmentation (also see Figure 2). Figure 4 displays the taxonomic placement of this organism as defined by its SSU rRNA gene. Strain MWH-Uga1 is a member of the family Microbacteriaceae and belongs to a group of *Actinobacteria* previously identified from freshwater habitats, named the Luna SSU rRNA cluster. The Luna cluster (Hahn *et al.*, 2003) is named from the Latin translation of moon (luna), which is the name of the lake (Mondsee = moon lake) where the first member of this SSU rRNA cluster was cultivated. For ease of reference, the large monophyletic clade in Figure 4 containing organisms of this SSU rRNA cluster, as well as numerous sequences from cultivated and environmental sources will be defined here as ‘the Luna lineage’.

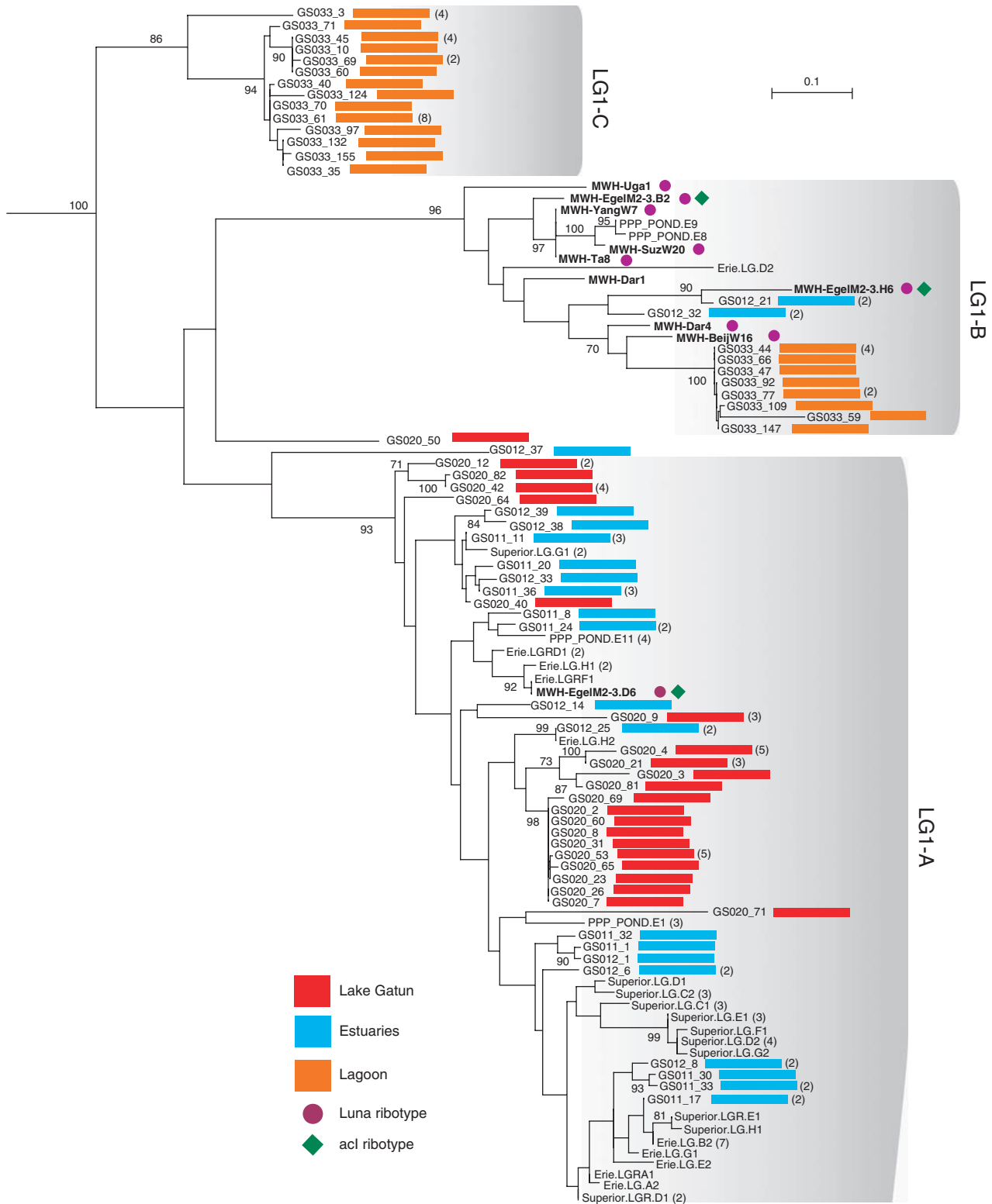
#### LG1 ActRs in other Luna lineage Actinobacteria

LG1 ActR gene fragments were amplified from seven mixed cultures whose primary constituents are also *Actinobacteria* of the Luna lineage (Table 2). Each of these cultures contains a Luna actinobacterium and a non-actinobacterial strain, the latter of which can differ in taxonomic affiliation and proportion of total cell numbers ( $<1$  to  $>20\%$ ) between cultures (Hahn, 2009). The Luna *Actinobacteria* present in mixed cultures share general phenotypic similarities to isolate MWH-Uga1, such as small cell sizes, and pigmentation (Figure 2). Each culture except one (see below) exhibits a single actinobacterial ribotype and a single amplified LG1 ActR, both of which fall into the monophyletic clusters represented by the Luna lineage (Figure 4) and the MWH-Uga1 ActR sequence (Figure 3), respectively. On the basis of these collective observations, we infer that the ActRs recovered from the mixed cultures originate from the Luna strain present in each. These results provide further evidence that ActR genes are encoded in freshwater *Actinobacteria* and suggest that such genes are common among organisms belonging to the Luna lineage. However, of the 20 cultures screened by PCR (Supplementary Table S1) only 8 were positive for ActR genes, even though nearly all contained pigmented Luna *Actinobacteria*. In some instances organisms lacking detectable ActR genes were nearly identical at the SSU rRNA level to rhodopsin-encoding *Actinobacteria* (Figure 4). The absence of gene products for some cultures may result from mismatches to degenerate primers, whereas others may simply lack ActR.

#### ActR gene fragments recovered from environmental DNA samples

To show that ActR genes were present in other freshwater environments, besides those sampled during the GOS by Rusch *et al.* (2007), we used our degenerate ActR primers (Table 1) to amplify





**Figure 3** Best maximum likelihood tree of LG1 ActRs proteins (109 taxa, 96 positions). Only bootstrap values  $\geq 70$  are displayed. The LG1 group is broken down into three subclasses and LG2-type ActRs (outlined in Figure 1) are used as an outgroup. Sequences originally described in Sharma *et al.* (2008), retrieved from the Global Ocean Sampling (GOS) expedition's non-marine metagenomic samples are color coded according to sample origin. ActR sequences amplified in this study from actinobacterial cultures are indicated in bold, and the actinobacterial ribotype(s) associated with each are coded by symbols. Sequences amplified from environmental samples are labeled according to sample origin, and the number of times a particular sequence was detected in our library is indicated in brackets. We also show the number of times a sequence was detected in each non-marine GOS metagenomic sample, by including the results of an analysis from our previous study in Sharma *et al.* (2008), where partial and full-length rhodopsin protein sequences were clustered at a phylogenetic distance of  $\leq 0.1$ . From this analysis, most LG1 sequence clusters formed from each GOS sample have a representative in the tree above.





**Figure 4** Neighbor-joining tree (with Jukes/Cantor correction) of 92 partial actinobacterial small subunit ribosomal RNA (SSU rRNA) sequences (*E. coli* positions 95–840) from the Global Ocean Sampling (GOS) expedition's non-marine metagenomic samples and cultivated freshwater *Actinobacteria*, within the context of selected sequences from the phylum *Actinobacteria*. Bootstrap values (MEGA4) represent percentage support of the nodes based on 1000 resamplings (only values  $\geq 70\%$  are shown). Actinobacterial SSU rRNA sequences from pure and mixed cultures containing ActR genes are indicated in bold. Groups of freshwater *Actinobacteria* defined by past environmental SSU rRNA studies are outlined. Sequences from non-marine GOS samples are color coded according to their sample origin (as in Figure 3) and labeled with CAMERA database accession numbers.

environmental DNA samples obtained from the surface of three freshwater habitats (Table 2). The first was a small (<1Ha) shallow (depth <1m)

pond, named 'Point Pleasant Pond' located in a wooded area in Halifax, Nova Scotia, Canada. The pond was selected as an initial site to test our actR

**Table 2** Freshwater isolates, mixed cultures or metagenomic samples harboring ActR gene(s)

| Sample type<br>(isolate, mixed culture<br>or metagenomic) | Geographical origin           | Habitat | Trophic<br>status  | Climate<br>zone | Conserved acidic aa at<br>positions 97 and 108 <sup>a,b</sup><br>for proton pumping |
|---|-------------------------------|---------|--------------------|-----------------|---|
| Isolate MWH-Uga1  | Jinja, Uganda                 | Pond    | Eutrophic          | Tropical        | Yes (1/1) <sup>c</sup>  |
| Culture MWH-Dar1 <sup>d</sup>                             | Dar es Salaam, Tanzania       | Pond    | Eutrophic          | Tropical        | Yes (1/1) <sup>c</sup>  |
| Culture MWH-Dar4 <sup>e</sup>                             | Dar es Salaam, Tanzania       | Pond    | Eutrophic          | Tropical        | Yes (1/1) <sup>c</sup>  |
| Culture MWH-Ta8 <sup>f</sup>                              | Lake Taihu, China             | Lake    | Hypertrophic       | Subtropical     | Yes (1/1) <sup>c</sup>  |
| Culture MWH-YangW7  | Yangtze River, Nanjing, China | River   | Eutrophic          | Subtropical     | Yes (1/1)   |
| Culture MWH-SuzW20 <sup>f</sup>                           | Suzhou, China                 | Pond    | Eutrophic          | Subtropical     | Yes (1/1)   |
| Culture MWH-BeijW16 <sup>g</sup>                          | Beijing, China                | Pond    | Not determined     | Temperate       | Yes (1/1)   |
| Culture MWH-EgelM2-3 <sup>h</sup>                         | Lake Egelsee, Austria         | Lake    | Eutrophic          | Temperate       | Yes (3/3) <sup>c</sup>  |
| Lake Superior   | North America                 | Lake    | Ultra-oligotrophic | Temperate       | 100% (34/34) <sup>h</sup>   |
| Lake Erie   | North America                 | Lake    | Meso/Oligotrophic  | Temperate       | 100% (37/37) <sup>h</sup>   |
| Point Pleasant Park Pond                                  | Halifax, NS, Canada           | Pond    | Not determined     | Temperate       | 100% (9/9)  |
| Lake Gatun  | Panama Canal                  | Lake    | Meso/Eutrophic     | Tropical        | 97% (75/77) <sup>h</sup>  |
| Delaware Bay  | USA                           | Estuary | Eutrophic          | Temperate       | 95% (41/43) <sup>h</sup>  |
| Chesapeake Bay  | USA                           | Estuary | Supertrophic       | Temperate       | 97% (37/38) <sup>h</sup>  |
| Punta Cormorant lagoon                                    | Galapagos Islands             | Lagoon  | Unknown            | Tropical        | 98% (126/128) <sup>h</sup>  |

<sup>a</sup>Conserved acidic amino acids (D97 and E108) in PR critical for proton pumping function in rhodopsins.

<sup>b</sup>Note that for partial ActR gene sequences amplified in this study, the forward primer encodes the D97 position one codon in from the 3' end (see Table 1).

<sup>c</sup>Cultures from which full-length ActR genes were obtained.

<sup>d</sup>Proposed '*Candidatus* Rhodoluna planktonica' name accepted in Hahn (2009) as are all others shown here.

<sup>e</sup>*Candidatus* Aquiluna rubra.

<sup>f</sup>*Candidatus* Rhodoluna laticola.

<sup>g</sup>One Luna-type cell in this mixed culture associated with an MWH-EgelM2-3 SSU rRNA sequence in Figure 4 named *Candidatus* Limnoluna rubra.

<sup>h</sup>In addition to ActR, percentages include all microbial rhodopsin sequence diversity obtained from these environmental samples that encode the amino-acid positions of interest.

primers and was sampled in late summer 2006, when the water level was severely decreased due to evaporation and was rich in particulate matter. The other two samples originated from the North American Great Lakes. Here we concentrate our efforts on samples from Lakes Superior and Erie previously obtained for diversity studies of cyanobacterial picoplankton (Wilhelm *et al.*, 2006; Ivanikova *et al.*, 2007). Lake Superior represents a unique environment because its waters are extremely phosphate limited, contributing to low rates of primary production similar to non-bloom conditions at the Bermuda Atlantic Time Series Site (RW Sterner, personal communication). Such low rates in turn yield levels of dissolved organic carbon <2 mg l<sup>-1</sup> (Hassler *et al.*, 2009). In contrast, Lake Erie has the highest rates of primary productivity among the Great Lakes, and high growth rates of phytoplankton in the summer lead to seasonal hypoxia in the bottom waters of the shallow (24 m) central basin (Wilhelm *et al.*, 2006).

A total of 52 partial rhodopsin sequences were obtained by the combined use of the LG1 and 2 primer sets shown in Table 1. Of these, 43 were obtained from Lake Superior (3.1 °C in June) and Lake Erie (22 °C in July) and 9 from the initial test site, Point Pleasant Pond. Nearly all sequences recovered with the LG-type primers (51 of 52) encoded proteins that phylogenetically cluster within the LG1 clade of ActRs (Figure 3). Our prior analysis of LG1 sequences from GOS data in Sharma

*et al.* (2008) identified a subclade within this grouping that represented nearly all of the LG1-type ActRs described from the freshwater Lake Gatun sample (44 of 45) and those from Delaware and Chesapeake Bays (32 of 38), which are estuaries that are subject to large inputs from freshwater runoffs. The sequences obtained from freshwater sites sampled in this current study also display a similar pattern (Figure 3), where 48 of 51 LG1-type ActRs fall within that same subclade, which we now define here as LG1-A. Collectively, these environmental sequence data in Figure 3 show that similar ActR genes are present among diverse freshwater habitats. Such observations verify our initial prediction based on our previous findings in Sharma *et al.* (2008) and the knowledge that many environmental SSU rRNA studies show that members of the *Actinobacteria* are widespread in fresh waters (Glockner *et al.*, 2000; Warnecke *et al.*, 2004; Allgaier and Grossart, 2006).

Although the cultures containing Luna-type *Actinobacteria* outlined above were all established from freshwater environments (Table 2), these organisms did not appear to carry genes of the LG1-A subclade that represent most of the ActR sequence diversity recovered from freshwater metagenomic DNA samples (Figure 3). Rather, ActRs from the Luna *Actinobacteria* cluster with sequences recovered from the GOS sample of the hypersaline Punta Cormorant lagoon, in a second subclade we refer to here as LG1-B. These observations suggest that

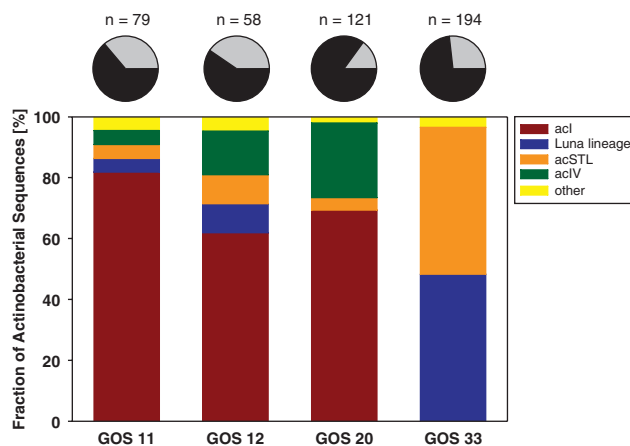
either Luna-type organisms do not belong to the group of *Actinobacteria* responsible for the LG1-A phylotypes or that a paralog representing that clade is undetectable with our primers due to nucleotide mismatches or gene absence. Therefore, in an attempt to understand the taxonomic distribution of *Actinobacteria* present in metagenomic samples containing ActRs, we performed a comparative analysis of SSU rRNA genes.

#### *Analysis of SSU rRNA genes from samples containing ActR*

Assessment of dominant ribotypes in non-marine GOS metagenomic samples by Rusch *et al.* (2007) revealed a large contribution of sequences from *Actinobacteria*, and this observation helped inform our prediction in Sharma *et al.* (2008) that planktonic *Actinobacteria* encode rhodopsins. Our current findings from cultivated *Actinobacteria* substantiate the value of linking strong signals from phylogenetic markers, to inferences based on high-frequency patterns in metagenomic data. Here, signals from SSU rRNA genes are used from metagenomic samples and cultivated *Actinobacteria* to gain insight into the distribution of ActR genes among members of this phylum.

Three SSU rRNA data sets were used for this exercise and the first comes from previous work by Jensen and Lauro (2008), who thoroughly analyzed actinobacterial SSU rRNA sequence diversity from assembled GOS metagenomic data (Rusch *et al.*, 2007) and kindly provided us their data set from non-marine sites. The phylogenetic affinity, as determined by us, for each of the 121 partial and full-length sequences from this data set is summarized in Figure 5. As a second independent assessment, phylogenetic analysis was performed on sequences recovered from BLAST searches of the unassembled GOS metagenomic data (Seshadri *et al.*, 2007) from the same non-marine sample sites using SSU rRNA sequence representatives for each of the known lineages of freshwater *Actinobacteria* as queries (Figure 4). The third data set consists of actinobacterial sequences recovered from a clone library of bacterial SSU rRNA gene fragments amplified from Lake Erie DNA samples (Supplementary Figure S1). Intra and intersite comparisons of ActRs and rRNAs from environmental samples were interpreted in the context of sequences obtained from cultures.

In Sharma *et al.* (2008), we described a set of ActR sequences that appears to originate from Luna-related *Actinobacteria*. Above we outlined the LG1-B clade consisting of Luna-affiliated ActRs and sequences previously described from the hypersaline lagoon GOS sample (Figure 3). Our analysis of GOS lagoon SSU rRNA sequences retrieved from CAMERA (Seshadri *et al.*, 2007) shows two distinct groups that branch within the Luna lineage (Figure 4). It is likely that the



**Figure 5** Group-specific phylogenetic affiliation for 121 actinobacterial small subunit ribosomal RNA (SSU rRNA) sequences identified from assembled metagenomic data from the Global Ocean Sampling (GOS) expedition's non-marine samples. These samples are indicated on the x axis by their GOS identifiers: the estuaries Delaware Bay (GOS11) and Chesapeake Bay (GOS12), the freshwater Lake Gatun (GOS20) and the hypersaline Punta Cormorant Lagoon (GOS33). Only SSU rRNA sequences  $\geq 400$  nucleotides in length were analyzed. Groups of freshwater *Actinobacteria* are color coded according to key and the number of sequences belonging to a specific group in each sample is shown by the proportional representation of a individually colored bar in the graph. Pie charts at the top of each bar depict the total percentage of actinobacterial SSU rRNA sequences (gray) among the total number of SSU rRNA sequences ( $n$ ) contained in the respective metagenomic sample. The sequences used in this analysis were provided by Jensen and Lauro (see text for further details).

organisms represented by these lagoon SSU rRNA sequences are responsible for at least the ActRs that branch in the LG1-B clade with cultivated Luna. In addition, the presence of a few Luna-related ActRs in GOS estuarine samples can also be accounted for by related SSU sequences (Figures 4 and 5).

The lack of Luna lineage rRNAs detected from the GOS Lake Gatun sample (Figure 5) and in the Lake Erie data obtained by us (Supplementary Figure S1) suggests it is unlikely that such organisms account for the LG1-A subclade of sequences in Figure 3 that represents most of ActR gene diversity recovered from freshwater metagenomic DNA samples. Interestingly, SSU rRNA sequences from the uncultured acI lineage comprise over 60% of actinobacterial SSU rRNA genes recovered from the non-marine GOS samples where LG1-A ActRs are found abundant (Figure 5). Organisms belonging to the acI lineage are found widespread and abundant among diverse types of freshwater environments (Allgaier and Grossart, 2006; Newton *et al.*, 2007). The distribution of the acI SSU rRNA genes follows that of the LG1-A ActRs; both are frequently observed in Lake Gatun, Lake Erie (Supplementary Figure S1) and estuarine samples but absent from the Punta Cormorant lagoon (Figure 5). Although these correlations provide a strong lead for future studies, the ability to link acI cell types to LG1-A ActRs is hampered by our current knowledge of this group, which is limited to environmental SSU rRNA

sequences and FISH studies (Warnecke *et al.*, 2005; Newton *et al.*, 2007). In the following section we provide additional evidence to support the hypothesis that acI *Actinobacteria* are the bearers of the LG1-A ActRs in the form of a mixed culture containing an acI cell type.

#### *Culture MWH-EgelM2-3 contains an acI cell type*

MWH-EgelM2-3 was the only mixed culture found to harbor two phylogenetically distinct ActR variants, suggesting that more than one actinobacterial strain was present. This culture, established from a small eutrophic lake in Austria (Table 2), carried ActRs of both the LG1-A and LG-1 B subclades (Figure 3). On the basis of the recovery of LG1-B ActRs from the multiple Luna cultures discussed above, we infer that ActR variants of this type from culture MWH-EgelM2-3 can be accounted for by the presence of Luna organisms (Figure 4). Given the high frequency of acI SSU rRNA genes in metagenomic samples containing LG1-A ActRs (Figure 5), we suspected an acI organism accounted for the LG1-A sequence in MWH-EgelM2-3. Using the acI-specific FISH probe acI-852 (Warnecke *et al.*, 2005), we confirmed that an acI organism was present at an abundance of <0.1% of the total cell population in MWH-EgelM2-3. Subsequently, we obtained the SSU rRNA gene for this organism and demonstrated its affiliation with the acI lineage (Figure 4). Meanwhile, we further analyzed the phylogeny of this acI cell type and proposed to establish the *Candidatus* species '*Candidatus* Planktophilia limnetica' for this strain (Jezbera *et al.*, 2009).

Strikingly, we found that the SSU rRNA sequence of '*Candidatus* Planktophilia limnetica' was identical to a partial acI sequence (Erie.Aughypo31) from Lake Erie (Supplementary Figure S1), which could explain the recovery of identical LG1-A ActR gene fragments obtained from each of these sources (Figure 3). In combination with the high incidence of acI SSU rRNA genes in metagenomic samples harboring LG1-A ActRs, these findings further support the hypothesis that the acI cell type present in this mixed culture is the bearer of an ActR variant absent from all other cultures containing Luna *Actinobacteria*. Although absolute evidence is required to validate this hypothesis, the presence of the acI organism in culture will provide a future means to assess if acI *Actinobacteria* encode ActRs.

#### *Functional homology and sequence characteristics of ActRs*

Of the ActRs we described from non-marine GOS metagenomic data in Sharma *et al.* (2008), nearly all conserved two acidic amino acids, D97 and E108, critical for a light-driven proton pumping function (Table 2). Five full-length ActR genes obtained from actinobacterial cultures here using a walking PCR

technique (Katz *et al.*, 2000) also conserve these residues (Table 2), suggesting that ActRs may function in photoheterotrophy. ActR gene fragments obtained from environmental samples and cultures in this study were amplified with a forward primer that encodes the D97 residue close to the 3' end (Table 1). Therefore, it cannot be stated with certainty that these partial sequences also conserve both acidic amino acids, although each of them conserves the E108 position. Further analysis of ActRs sequence characteristics reveals that genes of the LG1-A and LG-B type exhibit a much lower GC content (~40–50%, Supplementary Figure S2) than commonly perceived for organisms belonging to the phylum Actinobacteria, who are commonly referred to as 'High GC Gram-positive bacteria' (Ventura *et al.*, 2007).

#### *Taxonomic and ecological distributions of rhodopsins in nature*

Together these environmental and genomic sequence data reflect previous observations that planktonic *Actinobacteria* have a global distribution in diverse freshwater environments and suggest that at least some of them use ActRs. In addition to ActR, proteorhodopsin-encoding gene fragments affiliated with proteobacterial and bacteroidetes sequences (Supplementary Table S2) were also amplified from the Great Lakes Superior and Erie. Several recent studies highlight the broad taxonomic and ecological distribution of microbial rhodopsin genes in diverse aquatic environments (Rusch *et al.*, 2007; Atamna-Ismaeel *et al.*, 2008; Sharma *et al.*, 2008), suggesting they are important in the adaptation of microbes to life on the earth's surface. However, exploratory studies such as ours can only provide insight into sequence diversity, generating fundamental questions regarding rhodopsin function and its impact at both cellular and ecosystem levels. A recent analysis by Gonzalez *et al.* (2008) of the genome sequence of a proteorhodopsin-encoding marine bacterium proposed this organism may be capable of both carbon fixation and rhodopsin-based phototrophy. Such ideas highlight the vast potential for novel metabolic strategies involving rhodopsins. Freshwater *Actinobacteria* are an extremely successful group of organisms of unknown metabolic potential that also encode ActRs. The discovery of cultivated actinobacterial representatives encoding these genes represents an exciting opportunity to understand the significance of rhodopsins to their biology and to freshwater ecosystems.

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