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Light-induced transcriptional responses associated with proteorhodopsin-enhanced growth in a marine flavobacterium

Hiroyuki Kimura¹,³, Curtis R. Young¹, Asuncion Martinez¹ and Edward F. DeLong¹,²

¹Department of Civil and Environmental Engineering, Massachusetts Institute of Technology, Cambridge, MA 02139, USA
²Department of Biological Engineering, Massachusetts Institute of Technology, Cambridge, MA 02139, USA
³Department of Geosciences, Faculty of Science, Shizuoka University, Shizuoka, Shizuoka 422-8529, Japan

*Correspondence: Edward F. DeLong
Department of Civil and Environmental Engineering, Massachusetts Institute of Technology,
Parsons Laboratory 48-427, 15 Vassar Street, Cambridge, MA 02139, USA.
E-mail: delong@mit.edu
Tel: (617) 253-0252

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ABSTRACT: Proteorhodopsin (PR) is a photoprotein that functions as a light-driven proton pump in diverse marine Bacteria and Archaea. Recent studies have suggested that PR may enhance both growth rate and yield in some flavobacteria when grown under nutrient limiting conditions in the light. The direct involvement of PR, and the metabolic details enabling light-stimulated growth however, remain uncertain. Here, we surveyed transcriptional and growth responses of a PR-containing marine flavobacterium during carbon-limited growth in the light and the dark. As previously reported (Gómez-Consarnau et al., Nature 445: 210-213, 2007), Dokdonia strain MED134 exhibited light-enhanced growth rates and cell yields under low carbon growth conditions. Inhibition of retinal biosynthesis abolished the light-stimulated growth response, supporting a direct role for retinal-bound PR in light enhanced growth. Among protein-coding transcripts, both PR and retinal biosynthetic enzymes showed significant upregulation in the light. Other light-associated proteins, including bacterial cryptochrome and DNA photolyase, were also expressed at significantly higher levels in the light. Membrane transporters for Na⁺/phosphate and Na⁺/alanine symporters, and the Na⁺-translocating NADH-quinone oxidoreductase (NQR) linked electron transport chain, were also significantly upregulated in the light. Culture experiments using a specific inhibitor of Na⁺-translocating NQR indicated that sodium pumping via NQR is a critical metabolic process in the light-stimulated growth of MED134. In total, the results suggested the importance of both the PR-enabled, light-driven proton gradient, as well as the generation of a Na⁺ ion gradient, as essential components for light-enhanced growth in these flavobacteria.
Introduction

Some prokaryotes possess proteins that interact with light, and convert it into energy for growth or into sensory information. One class of energy-harvesting photoproteins called rhodopsins consist of single, membrane-embedded protein covalently bound to the chromophore retinal (a light-sensitive pigment) (Spudich and Jung, 2005). Ten years ago, prokaryotic rhodopsin, proteorhodopsin (PR), was discovered through metagenomic analyses of marine bacterioplankton genome fragments (reviewed by DeLong and Béjà, 2010). Béjà et al. (2000) found that an uncultivated marine SAR86 clade member in Gammaproteobacteria contained a bacteriorhodopsin-like gene, dubbed PR. Further, the marine SAR86-derived PR functioned as a proton pump, when the recombinant *Escherichia coli* expressing PR is exposed to light. PRs were subsequently detected in many other marine bacteria, some of which appeared to be “tuned” to absorb specific wavelengths of light associated with their habitat of origin; green light in surface waters and blue light in deep waters (Béjà et al., 2001). Additional studies have found PR genes in a diverse array of abundant marine bacterial and archaeal clades (Giovannoni et al., 2005; Frigaard et al., 2006; Brown and Jung, 2006; McCarren and DeLong, 2007). Based on genomic surveys, a large fraction of naturally occurring marine bacterioplankton in oceanic surface seawaters appear to contain the PR gene (de la Torre et al., 2003; Sabehi et al., 2005; Moran and Miller, 2007; DeLong 2009). Interestingly, chromophore biosynthetic genes including a carotenoid biosynthetic gene cluster, and a novel blh gene encoding a 15,15'-β-carotene dioxygenase that cleaves β-carotene to yield retinal, were found linked to the PR gene in some microorganisms (Sabehi et al., 2005). Martinez et al. (2007) demonstrated that the expression of the entire PR photosystem (genetically linked PR and retinal biosynthetic genes) in *E. coli* can result in proton-pumping activity in light, and that the resulting pmf can be used for ATP synthesis via the membrane-embedded ATP synthase. Furthermore, PR in
recombinant *E. coli* can generate a light-driven pmf sufficient to increase the rate of flagellar rotation, providing estimates for energy flux through the photosystem (Walter et al., 2007).

PR-containing marine bacterial isolates have been recently cultured from a variety of marine environments. These isolates include members of SAR11 (Alphaproteobacteria), OM43 (Betaproteobacteria), and SAR 92 (Gammaproteobacteria) clades, as well as members of the Bacteroidetes and Vibrionaceae (Giovannoni et al., 2005; Frigaard et al., 2006; McCarren and DeLong, 2007; Stingl et al., 2007; González et al., 2008). Laboratory experiments examining light-stimulated growth in some of these isolates however have proven equivocal. Some studies could detect no significant light enhancement of either growth rates or cell yields in PR-containing isolates (Giovannoni et al., 2005; Stingl et al., 2007). However, light-enhanced growth rates and cell yields were reported in one PR-containing marine flavobacterium, *Dokdonia* sp. MED134 (Gómez-Consarnau et al., 2007). Additionally, microcosm studies suggested that some of marine flavobacteria and SAR11 populations exhibited enhanced expression of the PR gene in the presence of light (Lami et al., 2009). As well, Gómez-Consarnau et al. (2010) demonstrated the enhanced long-term survival of PR-containing *Vibrio* cells in the light, but not in darkness. Nevertheless, the specific metabolic processes that facilitate light-enhanced growth or survival are not yet well understood.

To better characterize the photophysiology of PR-containing Flavobacteria, we performed transcriptomic analyses targeting total RNA extracted from MED134 exposed to light or in the dark. Transcriptional profiles derived from cultures incubated in the light and dark were analyzed, and these results were used to further direct laboratory experiments using different growth substrates and inhibitors. The effect of light on growth at various carbon concentrations, and the effect of retinal biosynthesis inhibitors on light-enhanced growth, were explored. In addition, the effects of sodium-translocating respiratory chain
inhibitors on light-stimulated on growth were also examined. The combined results from both gene expression studies and physiological experiments were used to develop a model that incorporates some of the important features of photoheterotrophic growth observed in *Dokdonia* strain MED 134.

**Materials and methods**

*Strain and culture conditions*

PR-containing marine flavobacterium, *Dokdonia* sp. MED134, was isolated from surface seawater in Northwest Mediterranean Sea (Gómez-Consarnau et al., 2007). This strain was kindly provided to us by Jarone Pinhassi (University of Kalmar, Sweden). MED134 was grown in artificial seawater (ASW) (35 practical salinity units, prepared from Sea Salts; Sigma) containing low concentration of dissolved organic carbon (DOC) (0.05 mM C). ASW was filter-sterilized through 0.2 μm-pore-size filter system (Nalgene) and autoclaved. Then 250 ml aliquots of ASW (containing a background concentration of 0.05 mM C of DOC), were partially supplemented with full strength medium (FSM; 0.5 g of peptone [Bacto Pepton, BD] and 0.1 g of yeast extract [Bacto Yeast Extract, BD] per 100 ml of ASW), to yield final DOC concentrations of 0.14 and 0.39 mM C, respectively. All media were also supplemented with 225 μM of NH₄Cl and 44.7 μM of Na₂HPO₄·12H₂O, to avoid inorganic nitrogen and phosphate limitation. DOC concentrations were measured using the high temperature combustion method on TOC-V (Shimadzu) with platinized aluminum catalyst. The bacteria were initially grown in ASW enriched to 1.1 mM C, washed in ASW, and then diluted into three different ASW media, each containing a different DOC concentration (0.05, 0.14 and 0.39 mM C). Cultures were incubated at 22°C under continuous white light (approximately 150 μmol of photons m⁻² s⁻¹) or in the darkness.
To determine bacterial cell density, cultures were filtered with pre-blackened Isopore membrane filter (pore size, 0.22 μm; Millipore). Bacterial cells on the filter were stained with SYBR Green I (1:100 dilution; Molecular Probes) for 15 min, and counted under an epifluorescence microscope (Axioskop 2, Zeizz). All culture experiments were performed in triplicate.

**Cultivation for transcriptomic analyses**

MED134 was grown on 900 ml of ASW enriched to 0.14 mM C at 22°C in the darkness for the first 2 days. At this time, 400 ml of culture was filtered onto a pore-size 0.22-μm Durapore membrane filter (25 mm diameter, Millipore), yielding the D2 sample. The remaining culture was split in two 250 ml flasks that were incubated again at 22°C under the continuous white light (approximately 150 μmol of photons m⁻² s⁻¹) or in the darkness. After 2 more days, the cultures were filtered onto Durapore membrane filters (Millipore), yielding samples L2 (light conditions) and D4 (dark conditions), respectively. Filter samples were immediately placed into screw-cap tubes containing 1 ml of RNAlater (Ambion) and stored at -80°C until RNA extraction.

**Total RNA extraction and rRNA subtraction**

Total RNA was extracted from the filter samples using a modification of the mirVana miRNA isolation kit (Ambion) as described previously (Shi et al., 2009; McCarren et al., 2010). Briefly, filter samples were thawed on ice, and the RNAlater surrounding each filter was removed and discarded. The filters were immersed in Lysis/Binding buffer (Ambion) and mixed to lyse attached cells. Total RNA was extracted from the lysate according to the manufacturer's protocol. Remaining genomic DNA in RNA extraction was removed using a TURBO DNA-free kit (Ambion).
Bulk DNA was extracted from MED134 cultured under suitable condition based on a conventional extraction protocol. Cells of strain were lysed with lysozyme and proteinase K solution. Then the genomic DNAs were extracted with phenol-chloroform-isoamyl alcohol and precipitated with ethanol.

16S and 23S rRNAs were removed by the subtractive hybridization described by Stewart et al. (2010). Ribonucleotide probes targeting 16S and 23S rRNA genes were generated from the bulk DNA extracted from MED134. Templates for probe generation were first prepared by PCR using Herculase II Fusion DNA Polymerase (Stratagene) and strain-specific primers flanking nearly the full length of the bacterial 16S and 23S rRNA genes, with reverse primers modified to contain the T7 RNA polymerase promoter sequence (Supplementary Table S1). Biotinylated antisense rRNA probes were generated by in vitro transcription with T7 RNA polymerase, ATP, GTP, CTP, UTP, biotin-11-CTP, biotin-16-UTP (Roche). Biotinylated rRNA probes were hybridized to complimentary rRNA molecules in total RNA sample. Then biotinylated double-stranded rRNA was removed from the sample by hybridization to Streptavidin-coated magnetic beads (New England Biolabs).

The subtraction efficiency was evaluated by monitoring the removal of 16S and 23S peaks from total RNA profiles using a 2100 Bioanalyzer (Agilent).

**RNA amplification, cDNA synthesis, and pyrosequencing**

The rRNA-subtracted RNA (10-15 ng) was amplified using the MessageAmp II-Bacteria kit (Ambion) as described previously (Shi et al., 2009; McCarren et al., 2010). In brief, total RNA were polyadenylated using *Escherichia coli* poly(A) polymerase. Polyadenylated RNA was converted to double-stranded cDNA via reverse transcription primed with an oligo(dT) primer containing a promoter sequence for T7 RNA polymerase and a recognition site for the restriction enzyme BpmI (T7-BpmI-(dT)_16VN) (Supplementary Table S1). cDNA was
transcribed in vitro at 37°C for 12 hr, yielding large quantities (40-60 μg) of single-stranded antisense RNA. The SuperScript double-stranded cDNA synthesis kit (Invitrogen) was used to convert antisense RNA to double-stranded cDNA, which was then digested with BpmI to remove poly(A) tails. Prior to pyrosequencing, poly(A)-removed cDNA was purified by using the AMPure kit (Beckman Coulter Genomics). Purified cDNA was used for the generation of single-stranded DNA libraries and the bead-bound fragments were amplified by emulsion PCR according to established protocols (454 Life Sciences, Roche). The resulting bead-bound single stranded cDNAs were then pyrosequenced on the 454 FLX platform (Roche). All the cDNA sequences generated in this study have been submitted to the GenBank short read archive under accession number SRA029329.

Analyses of pyrosequence data

rRNA and tRNA reads were identified using BLASTN against rRNA and tRNA sequences in MED134 genome data, which are deposited in GenBank under accession no. AAMZ00000000 (Gómez-Consarnau et al., 2007). Reads producing alignments with bit scores greater than 50 were considered as rRNA and tRNA sequences. Protein-encoding cDNAs (from mRNA) were identified using BLASTX against peptide sequences collected from MED134 genome data (bit score ≥50). Small RNAs (sRNAs) were analyzed using the Rfam (version 10.0) website (http://rfam.sanger.ac.uk/). Rfam is a collection of non-coding RNA families, each represented by multiple sequence alignments, consensus secondary structures, and covariance models, including 1,446 families in January 2010 (Gardner et al., 2009). Finally, in order to identify MED134 specific sRNA that might not be represented in Rfam, we assembled a database of intergenic regions (IGRs) in the genome of MED134 longer than 100 bp (total 992 sequences) which might encode putative sRNAs. Reads with matches to the IGRs database (bit score ≥50) were considered sRNA reads.
L2/D4 ratios were calculated based on read number of each cDNA, which was normalized by total number of protein-encoding reads in each sample. The statistic significance of the change observed between cultures in light and dark (L2 and D4) for each cDNA was determined based on false-discovery rate method ($q$-value $\leq 0.05$) (Benjamini and Hochberg, 1995; Storey and Tibshirani, 2003). Clustering analyses of transcriptomics datasets were performed in GenePattern (Reich et al., 2006), using hierarchical clustering (Eisen et al., 1998) by Pearson correlations for both rows and columns, using pairwise complete-linkage.

Culture experiments with specific inhibitors

To confirm an importance of retinal-bound PR for the light-stimulated growth, we performed culture experiments with 2-(4-methylphenoxy)triethylamine hydrochloride (MPTA). MPTA is known to prevent lycopene cyclization in retinal biosynthesis pathway (Cunningham et al., 1994; Armstrong, 1999). First, we cultured MED134 on Marine Agar 2216 (Difco) amended with MPTA at a final concentration of 300 $\mu$M and confirmed the effect of MPTA against strain MED134 based on color of colonies. Next, MED134 was grown in ASW slightly enriched with FSM (0.14 mM C) and amended with MPTA. MPTA was dissolved in methanol and added to ASW at a final concentration of 100 $\mu$M. The same volume of methanol was added to cultures as negative control without MPTA. These cultures were incubated at 22°C under continuous white light (approximately 150 $\mu$mol of photons m$^{-2}$ s$^{-1}$) or in the darkness. The cultures were performed in triplicate. Bacterial cell density was measured every 2 days by the direct counting method with epifluorescence microscope described above. Additionally, colony-forming unit (cfu) was also monitored after spread 100 $\mu$l of cultures on Marine Agar 2216 (Difco) and incubation at 22°C for 48 hr. MPTA was a generous gift of Francis X. (Buddy) Cunningham (University of Maryland, USA).
To determine the importance of sodium pumping in light-driven growth of PR-containing marine flavobacteria, MED134 was grown in ASW with 2-n-heptyl-4-hydroxyquinoline N-oxide (HQNO) (Enzo Life Sciences). HQNO is known to be a specific inhibitor of the electron-transport-linked Na⁺-translocating NQR enzyme complex (Tokuda and Unemoto, 1982; Häse and Mekalanos, 1999). MED134 was incubated in ASW enriched with DL-alanine (0.70 mM C), supplemented with trace element solution (Futamata et al., 2009), and amended with HQNO at a final concentration of 10 μM. DL-alanine was selected as carbon source for the bacterial cultivation, since transcriptomic analyses demonstrated significant over representation of Na⁺/alanine symporters in the presence of light (see Results and discussion). HQNO was prepared in ethanol. The same volume of ethanol was added to culture as negative control without HQNO. These cultures were incubated at 22°C under continuous white light or in the dark. The cultures were performed in triplicate. Bacterial cell densities were measured every 2 days by the direct count and plate count method described above.

Results and discussion

Cultivation in light and darkness

*Dokdonia* sp. MED134 exposed to light reached a maximal abundance of $1.1 \times 10^5$ cell ml⁻¹ in unamended artificial seawater (ASW) (DOC, 0.05 mM C), $1.4 \times 10^6$ cell ml⁻¹ in ASW enriched with 0.14 mM C, and $1.1 \times 10^7$ cell ml⁻¹ in ASW enriched to 0.39 mM C (Figure 1). In contrast, dark-incubated cultures remained below $5.0 \times 10^4$ cells ml⁻¹ in unenriched ASW. In nutrient enriched ASW media, MED134 grew moderately in the dark, but the cell yields were much lower compared to cultures grown in the light. Light/dark ratios of cell yields ranged from 1.6 to 4.6 at the peak of the growth curves. Growth rates in ASW containing
0.14 mM C were 0.69 day⁻¹ in the illuminated culture (logarithmic growth phase, 1.5 to 5
day), and 0.44 day⁻¹ in the dark cultures (logarithmic growth phase, 1.5 to 4 day). Growth
rates in ASW containing 0.39 mM C were 1.17 day⁻¹ in the light, and 1.01 day⁻¹ in the dark
(logarithmic growth phase, 0 to 4 day). These results show the considerable influence of PR
on growth rate at low carbon concentrations, and its lesser influence at higher carbon
concentrations. These findings confirm the previous work of Gómez-Consarnau et al.
(2007), which showed that light has a definite positive impact on the growth of the PR-
containing flavobacteria grown in low carbon conditions.

Transcriptome experiments

For transcriptomic analyses, MED134 was grown again in ASW enriched to 0.14 mM C.
Bacteria grew to 1.0 × 10⁵ cells ml⁻¹ for first 2 days in the dark (D2). After 2 more days,
MED134 reached 3.2 × 10⁵ cells ml⁻¹ in light (L2), whereas bacteria incubated in darkness
remained 1.2 × 10⁵ cells ml⁻¹ (D4).
cDNAs synthesized from the RNA samples were pyrosequenced on the Roche 454
FLX platform, yielding ≈400,000 reads per sample. cDNA derived from intergenic regions
(IGRs) accounted for 21 to 51% of the total cDNA reads (Figure 2). Remarkably, most of
reads derived from IGRs in all samples corresponded to a single gene (399 bp) encoding
transfer-messenger RNA (tmRNA) (Supplementary Table S2). tmRNA is small RNA that
employs both tRNA-like and mRNA-like properties as it rescues stalled ribosomes during
nutrient shortage (Gillet and Felden, 2001; Keiler, 2008; Moore and Sauer, 2007). Since the
proportions of tmRNA further increased with incubation time, the high percentage of tmRNA
is likely to be due to the carbon limiting growth conditions used in this experiment. Protein-
encoding transcripts, identified by comparison with the annotated MED134 genome
(GenBank version in December 2009), represented 38 to 61% of total cDNA reads (Figure
2). Genes with a significant change in L2/D4 ratio ($q$-value $\leq 0.05$) were considered differentially expressed in the light versus the dark. Using this criteria, 601 genes in 2,944 annotated protein-encoding genes were found to be differentially expressed. Specifically, 312 genes were upregulated in the light, whereas 289 genes exhibited downregulation in the light.

PR and retinal biosynthetic enzymes

Previous genomic analysis of MED134 revealed the presence of genes encoding PR, and *crtEBIY* encoding enzymes needed to synthesize β-carotene from farnesyl diphosphate (FPP) (Gómez-Consarnau et al., 2007). Further, a gene (*blh*) encoding an enzyme that converts β-carotene to retinal has been also found next to the PR gene on the genome of MED134. Our transcriptomic survey revealed that the L2/D4 ratios for the PR gene, *crtEBIY* and *blh* were elevated (Figure 3a). In particular, statistical significance tests based on $q$-values showed that the PR, *crtE* and *crtI* genes were significantly upregulated in the culture exposed to light (Table 1). In addition, hierarchical clustering of transcript abundances clustered D2 and D4 together, to the exclusion of L2 (Figure 3b). This clustering pattern reflects the differential response of this PR-containing flavobacterium to light. Gómez-Consarnau et al. (2007) demonstrated by RT-PCR that MED134 had a higher expression of PR gene in the light than in the dark. Lami et al. (2009) also reported that marine flavobacteria and SAR11 in natural costal seawaters displayed significant high expression of PR gene in the presence of light. Our results extend these previous reports and indicate that transcription of the entire PR photosystem is upregulated in the presence of light in this flavobacterium.

The only evidence supporting the role of PR in light-stimulated growth in strain MED134 is the observation that only light corresponding to the wavelengths absorbed by PR elicited growth enhancement (Gómez-Consarnau et al., 2007). To better define the role of
PR in light-stimulated growth in strain MED134, we performed culture experiments with MPTA, a specific inhibitor of lycopene cyclization in the retinal biosynthetic pathway (Cunningham et al., 1994; Armstrong, 1999; see Figure 3a). First, MED134 was grown on agar plates enriched with peptone and yeast extracts with and without MPTA. In the presence of MPTA, MED134 produced light pink colonies, whereas yellow colonies were observed on agar plates without MPTA (Figures 4a and 4b). In general, it is known that bacterial cells accumulating β-carotene display yellow or orange colonies, whereas light pink colonies indicate accumulation of lycopene (Cunningham et al., 1994; Armstrong, 1999). Our results indicate that MPTA effectively prevented β-carotene generation, the precursor for retinal, in strain MED134. MED134 was grown in ASW enriched to 0.14 mM C and amended with MPTA. When strain MED134 was incubated in ASW with MPTA, bacteria grew moderately to $6.3 \times 10^5$ cells ml$^{-1}$ and $4.3 \times 10^5$ cfu ml$^{-1}$ in the presence of light (Figures 4c and 4d). In contrast, MED134 incubated in ASW without MPTA in the light produced significantly higher yields ($1.9 \times 10^6$ cells ml$^{-1}$ and $1.5 \times 10^6$ cfu ml$^{-1}$) than those in ASW with MPTA. In culture experiments in the dark with or without MPTA, bacteria grew similarly to approximately $5.0 \times 10^5$ cells ml$^{-1}$ ($3.0 \times 10^5$ cfu ml$^{-1}$), equivalent to the results with MPTA in the light. The findings suggest that PR bound to retinal plays a critical role in the light-stimulated growth of the PR-containing marine flavobacteria.

**ATP synthetase**

The genome of MED134 harbors genes encoding membrane-embedded ATP synthetase, a multi-unit enzyme consisting of two large complexes. In this study, large numbers of transcripts from ATP synthetase were identified, however, there was no significant difference in ATP synthetase transcript abundance in the light and dark cultures (Supplementary Table S3).
Light sensors

MED134 contains several gene homologs of membrane sensors known to respond to light. González et al. (2008) reported that MED134 contain genes encoding bacterial cryptochrome and several DNA photolyase/cryptochromes that belong to different gene families (DASH family, (6-4) photolyase family, and class I photolyase). Similar genes encoding these light sensor proteins were also identified in the genome sequence of Polaribacter sp. MED152, another marine flavobacterium (González et al., 2008). Further, MED134 has PAS and GAF domains, known to be common components of phytochromes that detect red and far-red light (Taylor and Zhulin, 1999; Anantharaman et al., 2001). PAS domains are able to respond to oxygen levels, redox potential and light, whereas GAF domains work as phototransducers. Another gene associated with light sensing contains the BLUF domain, which specifically responds to blue light (Gomelsky and Klug, 2002). The BLUF domain is also found in the genome of related Polaribacter sp. MED152 (González et al., 2008). In addition to light sensors, several histidine (His) kinases, which might play important roles for secondary transduction and response regulation, are contained in the genome of MED134.

In this study, we found significant upregulation of bacterial cryptochrome and two putative DNA photolyase/cryptochrome genes under light (L2) versus dark (D4) conditions (Table 2). In contrast, genes encoding PAS, GAF, and BLUF domains exhibited no significant difference between light and dark cultures (Supplementary Figure S1). Of the secondary transduction enzymes, one His kinase (MED134_10396) showed very high expression rate (L2/D2 ratio, 18.3) and significant upregulation in light (Table 2). This His kinase may therefore be involved in controlling gene expression in response to light.

Central metabolic pathways
The main energy generating metabolic pathways of MED134 identified by genome analyses were glycolysis, the pentose phosphate cycle, and the TCA cycle (Gómez-Consarnau et al., 2007). Several genes encoding enzymes working in the central metabolic pathways were significantly induced in culture exposed to light (Supplementary Table S4). In particular, genes encoding fructose-bisphosphatase (fbp) in glycolysis, glucose-6-phosphate dehydrogenase (zwf) and phosphogluconate dehydrogenase (gnd) in pentose phosphate cycle, and succinate dehydrogenase (sdhABC) and fumarate hydratase (fumC) in TCA cycle exhibited significant upregulation in light. In addition, hierarchical clustering of transcript abundances clustered D2 and L2 together, to the exclusion of D4 (Supplementary Figure S2). This clustering pattern may reflect the increased levels of carbon available in the D2 culture, and the additional energy source (light) available in the L2 culture, relative to the D4 culture. Hence, the expression pattern of enzymes of the central metabolic pathways may reflect the differential nutrient and energy availability between the different treatments.

MED134 also harbors genes encoding pyruvate carboxylase (pycA) and phosphoenolpyruvate (PEP) carboxylase (ppc) that function in anaplerotic metabolism and that are associated with carbon fixation (Gonzáles et al., 2008). Pyruvate carboxylase generates oxaloacetate from bicarbonate and pyruvate, whereas PEP carboxylase synthesizes oxaloacetate from bicarbonate and PEP (Attwood and Wallace, 2002; Izui et al., 2004; Jitrapakdee et al., 2008). The transcript abundance of pyruvate carboxylase and PEP carboxylase were significantly downregulated in the light, compared to that in the dark (Supplementary Table S5). In contrast, SulP-type bicarbonate transporter and carbonic anhydrase, which is known to interconvert CO₂ and bicarbonate, were not significant different between L2 and D4. Although Polaribacter sp. MED152 has been reported to fix more bicarbonate in the light than in the darkness (Gonzáles et al., 2008), our findings indicate that in MED134 CO₂ incorporation via pyruvate carboxylase and PEP carboxylase
may be equally important for anaplerotic carbon replenishment under both dark and light growth conditions.

Transporters and electron transport chain

Membrane transporters play critical roles in uptake of essential nutrients and minerals. The genome of MED134 contains relatively low numbers of genes encoding membrane transporters, compared to other marine bacteria. Transcriptomic analyses revealed that two predicted Na\(^+\)/alanine symporters (MED134_02355, MED134_14567) were significantly upregulated in the light (Table 1 and Supplementary Figure S3). Further, a Na\(^+\)/phosphate symporter (MED134_11180) also exhibited significant upregulation in the light. Transcriptome analyses also indicated significant upregulation of Na\(^+\)-translocating NQR, succinate dehydrogenase, and cytochrome c oxidase in the light (Table 1 and Supplementary Figure S4). These results suggested the potential importance of the sodium ion gradient for transport fractions in the light, and the potential of indirect light-stimulation of sodium pump activities via the light-driven proton gradient (see Supplementary Figure S5).

To test the importance of sodium ion exchange in light-stimulated physiology of MED134, we performed growth experiments with HQNO, a specific inhibitor of Na\(^+\)-translocating NQR (Tokuda and Unemoto, 1982; Häse and Mekalanos, 1999). MED134 was grown in ASW amended with and without HQNO. For these experiments, DL-alanine was chosen as carbon source, because of the high expression levels of Na\(^+\)/alanine symporter we observed in cultures grown in the light. In cultures without HQNO, MED134 increased to 2.6 × 10\(^5\) cells ml\(^{-1}\) (2.3 × 10\(^5\) cfu ml\(^{-1}\)) in the light and 1.3 × 10\(^5\) cells ml\(^{-1}\) (1.1 × 10\(^5\) cfu ml\(^{-1}\)) in the dark (Figure 5) when grown on DL-alanine. Cell yields in cultures incubated with HQNO in the light were about 3 times less than those grown in the absence of inhibitor. In contrast, for cultures grown in the dark, cell yields decreased by about 1/2 in the presence of
the inhibitor, although the amount of growth on DL alanine was much lower. These findings indicate that the Na⁺-translocating NQR may play a critical role in sodium pumping in light-stimulated growth, and that active PR photophysiology greatly enhances the ability of these flavobacteria to grow on DL alanine.

**Conclusion**

In this study, we characterized gene expression patterns associated with the higher growth rates and cell yields observed in flavobacterium strain MED134 in carbon-limited media, when grown in the light. Among protein-encoding transcripts, a number of genes were upregulated in the light, including PR, retinal biosynthetic enzymes, and several predicted light sensors (Figure 6). Previous studies had suggested the involvement of PR in light enhanced growth, because the action spectra for this response generally matched the PR absorption spectrum. Experiments with MPTA, a specific inhibitor of an enzyme in retinal biosynthetic pathway, confirmed the involvement of PR in the observed light-enhanced growth. MED134 cultures grown with MPTA exhibited much lower cell yields than those without MPTA, when both were grown in the presence of light, while MPTA had no effect on MED134 grown in the dark. Proteins involved in some central metabolic pathways, including fructose-bisphosphatase, glucose-6-phosphate dehydrogenase, phosphogluconate dehydrogenase, succinate dehydrogenase, and fumarate hydratase, also had relatively higher expression in light than dark, suggesting an increased requirement for these enzymes during active cell growth. Transcripts for membrane embedded ATP synthase, and pyruvate carboxylase and PEP carboxylase were well represented in both the light and dark grown cultures. These results likely reflect a central and essential requirement for ATP synthetase and the two carboxylases, in both the light and the dark.
Of the membrane transporters, Na⁺/phosphate symporter and Na⁺/alanine symporter exhibited particularly significant upregulation in cultures exposed to light. The Na⁺-translocating NQR linked electron transport chain also exhibited significantly greater expression levels in the light. The gene expression levels of other sodium pumps however, including the Na⁺/H⁺ antiporter and the Na⁺ efflux pump, did not appear significantly different between the L2 and D4 treatments (Figure 6). These results indicate the potential importance of these symporters, coupled with the central role of NQR in driving energy-requiring nutrient transport via its maintenance of the sodium ion gradient. Culture experiments using a specific inhibitor of Na⁺-translocating NQR, HQNO, also suggested the importance of the sodium ion gradient for MED134 growth, and indicated that sodium pumping via Na⁺-translocating NQR is critical metabolic process for the light-stimulated growth of PR-containing marine flavobacteria, in addition to proton pumping via retinal-bound PR. In total, our findings indicate a direct role for retinal-bound PR in light-enhanced growth in Dokdonia strain MED134. Furthermore, an important role for H⁺/Na⁺ ion exchange, and transport processes that utilize energy derived from the sodium ion gradient, appear particularly important for the photoheterotrophic growth in this flavobacterium strain.
Acknowledgements

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**Figure Legends**

**Figure 1** Growth of MED134 incubated in the light or in the dark. MED134 was grown in unenriched ASW (0.05 mM C) (a), in ASW enriched to 0.14 mM C (b), and in ASW enriched to 0.39 mM C (c). The cultures were incubated under continuous white light (○), or in the darkness (●). Errors bars denote standard deviation for triplicate.

**Figure 2** Inventory of RNAs from cultures in the microbial transcriptomic datasets. MED 134 was first incubated in ASW enriched to 0.14 mM C in the dark for first 2 days (D2). Then culture was split in two flasks, with one incubated in the light (L2), and the other in the dark (D4), for 2 more days. Numbers in the pie charts represent the percentage of total cDNA reads in each transcriptomic dataset. *Subtraction of 16S and 23S rRNAs were performed after total RNA extraction.

**Figure 3** Transcriptomic analyses of proteorhodopsin and retinal biosynthetic genes. (a) Retinal biosynthetic pathway. The colors indicate the L2/D4 ratio of retinal biosynthetic enzymes. The ratio was calculated based on abundance of reads for each specific gene, normalized by the total number of protein-encoding reads for each sample. (b) Cluster analysis of the relative abundance of PR and retinal biosynthetic enzymes. Hierarchical clustering was performed based on the number of cDNA reads, normalized to the total number of protein-encoding cDNAs in each sample, using a Pearson correlation. The heat map shows relative difference of transcript abundance in each sample (red indicates high Pearson correlation; white indicates intermediate; blue indicates low). The numbers of cDNA read are summarized in Table 1. IPP, isopentenyl pyrophosphate; DMAPP, dimethylallyl pyrophosphate; FPP, farnesyl pyrophosphate; GGPP, geranylgeranyl pyrophosphate; MPTA, 2-(4-methylphenoxy)triethylamine hydrochloride.
Figure 4  Colony image and growth of MED134 in culture experiments with a specific
inhibitor of lycopene cyclization, 2-(4-methylphenoxy)triethylamine hydrochloride (MPTA).
(a and b) Colony morphology of MED134 on Marine Agar 2216 (Difco) plate amended with
MPTA (a) and without MPTA (b). (c and d) Microbial cell density in ASW enriched with
peptone and yeast extracts (0.14 mM C) and amended with MPTA in culture exposed to light
(△) and in the dark (▲), and in the ASW amended without MPTA in culture exposed to
light (○) and in the dark (●). Bacterial cells were counted by epifluorescence microscopy (c)
and plate count method (d). cfu, colony-forming unit.

Figure 5  Growth of MED134 in cultures incubated with 2-n-heptyl-4-hydroxyquinoline N-
oxide (HQNO), a specific inhibitor of Na⁺-translocating NADH-quinone oxioreductase
(NQR). MED134 were grown in ASW enriched with DL-alanine (0.7 mM C). Cultures
were incubated in the ASW amended with HQNO in light (△) or dark (▲) and in the ASW
without HQNO in light (○) or dark (●). Bacterial cell densities were determined by
epifluorescence microscopy (a) and plate count method (b). cfu, colony-forming unit.

Figure 6  Model of light-stimulated transcriptional responses in MED134. Proton pumping
processes and retinal biosynthetic pathway are shown in left, whereas central metabolic
pathways, such as glycolysis, pentose phosphate cycle, and TCA cycle, are in right. The
color allows and membrane proteins indicate the value of L2/D4 ratio. The ratio is calculated
based on abundance of reads for each gene normalized by total number of protein-encoding
reads for each sample. PR, proteorhodopsin; PP, pyrophosphatase; NQR, Na⁺-translocating
NADH-quinone oxidoreductase; SDH, succinate dehydrogenase; Cyt, cytochrome oxidase;
NHA, Na⁺/H⁺ antiporter.
Figure 2

Total reads = 300,880

D2

Total reads = 411,697

L2

Total reads = 289,633

D4

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<th>L2</th>
<th>D4</th>
</tr>
</thead>
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<td>0.8%</td>
<td>7.4%</td>
</tr>
<tr>
<td>other small RNA families</td>
<td>2.2%</td>
<td>3.6%</td>
<td>4.7%</td>
</tr>
<tr>
<td>protein-coding RNA</td>
<td>0.7%</td>
<td>1.7%</td>
<td>0.2%</td>
</tr>
<tr>
<td>rRNA*</td>
<td>0.7%</td>
<td>0.2%</td>
<td>0.5%</td>
</tr>
<tr>
<td>tRNA</td>
<td>0.7%</td>
<td>0.2%</td>
<td>0.2%</td>
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<tr>
<td>unknown IGR</td>
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<td>0.2%</td>
</tr>
<tr>
<td>no significant match</td>
<td>0.5%</td>
<td>0.2%</td>
<td>0.2%</td>
</tr>
</tbody>
</table>

protein-coding RNA

tmRNA

other small RNA families

rRNA*

tRNA

unknown IGR

no significant match
Figure 3

(a) Metabolic pathway for carotenoid biosynthesis. 

- IPP \(\rightarrow\) DMAPP \(\rightarrow\) FPP \(\rightarrow\) GGPP
- \(idi\) \(\rightarrow\) \(ispA\) \(\rightarrow\) \(crtE\) \(\rightarrow\) \(crtB\) \(\rightarrow\) Phytolene
- \(crtI\), \(crtY\)
- \(crtB\) (inhibitor of lycopene cyclization)
- \(blih\)

(b) Heat map showing the normalized intensity of various genes involved in carotenoid biosynthesis. The colors indicate the relative expression level with red being high and blue being low.
Figure 5

(a) Cell density (10^5 cells ml^-1) over time (day).

(b) Colony number (10^5 cfu ml^-1) over time (day).
Table 1 Read number and L2/D4 ratio of protein-encoding cDNA that plays a critical role in light-stimulated growth

<table>
<thead>
<tr>
<th>Name</th>
<th>Name</th>
<th>Locus tag</th>
<th>Size (aa)</th>
<th>Read number</th>
<th>L2/D4 ratio</th>
<th>q-value</th>
</tr>
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<tbody>
<tr>
<td>Opsin</td>
<td>Proteorhodopsin</td>
<td>MED134_07119</td>
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<td>1 33 5</td>
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<td>0.0129</td>
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<td>Retinal biosynthetic enzymes</td>
<td>Isopentenyl-diphosphate δ-isomerase (idi)</td>
<td>MED134_07374</td>
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<td>17 36 11</td>
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<td>Farnesyl-diphosphate synthase (ispA)</td>
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<td>Geranylgeranyl pyrophosphate synthase (crtE)</td>
<td>MED134_07466</td>
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<td>53 77 23</td>
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<td>Phytoene synthetase (crtB)</td>
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<td>20 34 8</td>
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<td></td>
<td>Phytoene dehydrogenase (crtI)</td>
<td>MED134_13076</td>
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<td>37 86 20</td>
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<td>Lycopene cyclase (crtY)</td>
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<td>Na+/alanine &amp; glycine symporter</td>
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<td>Na+/phosphate symporter</td>
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<td>Na+-translocating NADH quinone oxidoreductase (nqrC)</td>
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<td>Na+-translocating NADH quinone oxidoreductase (nqrF)</td>
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<td>1.75</td>
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Statistical significance between light and dark cultures was measured based on q-value (false discovery rate method). The features with q-values ≤0.05 are significant (Storey and Tibshirani, 2003), which are underline and in bold.
Table 2  Read number and L2/D4 ratio of cDNA encoding domains and peptides with a role in light absorption and response

<table>
<thead>
<tr>
<th>Name</th>
<th>Locus tag</th>
<th>Size (aa)</th>
<th>Read number</th>
<th>L2/D4 ratio</th>
<th>q-value</th>
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<td>Bacterial cryptochrome, DASH family</td>
<td>MED134_10201</td>
<td>432</td>
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<td>DNA photolyase/cryptochrome, (6-4) photolyase family</td>
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<td>DNA photolyase/cryptochrome, (6-4) photolyase family</td>
<td>MED134_10211</td>
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<td>DNA photolyase/cryptochrome, class I photolyase</td>
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<td>PAS domain</td>
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<td>GAF domain</td>
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<td>BLUF domain</td>
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<td>Two-component system sensor His kinase</td>
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<td>470</td>
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</table>

Statistical significance between light and dark cultures was measured based on q-value (false discovery rate method). The features with q-values ≤0.05 are significant (Storey and Tibshirani, 2003), which are underline and in bold.