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Notes:
Light-powering *Escherichia coli* with proteorhodopsin

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Proteorhodopsin (PR) is a light-powered proton pump identified by community sequencing of ocean samples. Previous studies have established the ecological distribution and enzymatic activity of PR, but its role in powering cells and participation in ocean energy fluxes remains unclear. Here, we show that when cellular respiration is inhibited by depleting oxygen or by the respiratory poison azide, *Escherichia coli* cells expressing PR become light-powered. Illumination of these cells with light coinciding with PR’s absorption spectrum creates a proton motive force (pmf) that turns the flagellar motor, yielding cells that swim when illuminated with green light. By measuring the pmf of individual illuminated cells, we quantify the coupling between light-driven and respiratory proton currents, estimate the Michaelis–Menten constant ($K_m$) of PR (10² photons per second/nm²), and show that light-driven pumping by PR can fully replace respiration as a cellular energy source in some environmental conditions. Moreover, sunlight-illuminated PR⁺ cells are less sensitive to azide than PR⁻ cells, consistent with PR⁺ cells possessing an alternative means of maintaining cellular pmf and, thus, viability. Proteorhodopsin allows *Escherichia coli* cells to withstand environmental respiration challenges by harvesting light energy.

Results and Discussion

We tracked swimming PR⁺ *E. coli* in two dimensions by using dark-field microscopy, periodically illuminating the cells with green light at PR’s absorption maximum. We observed single cells to characterize rapid responses of the cellular pmf to light. No detectable increase in cell swimming velocity occurred upon illumination with green light. We surmised that light-driven proton pumping may benefit the cell only under certain environmental conditions, as suggested by Giovannoni et al. (14).

To test the possibility that light-driven proton pumping is most beneficial to aerobically grown cells when their ability to respire is suddenly impaired, we energy-depleted the cells. Because *E. coli* is difficult to energy-deplete by nutrient limitation because of its endogenous energy stores (20, 21), we additionally used the respiratory poison azide, which has multiple cellular effects (22, 23) but primarily inhibits cytochrome oxidase and, thus, proton extrusion by the respiratory chain, stopping the flagellar motor (18).

Strikingly, with respiration inhibited by azide, PR⁺ cells responded to green light. PR⁺ cells in 30 mM azide swam slowly in red illumination, but they showed a marked velocity increase with green illumination (Fig. 1b). Upon removal of the green light, they slowed to their previous velocity. To increase the accuracy of our flagellar rotation measurements, we subsequently used a tethered cell geometry (Fig. 2a), permitting extended observation of the same bacterium in different illumination conditions. PR⁺ cells were allowed to stick to the coverslip via a flagellum, and we then monitored the angular rotation rate of cells (Fig. 2a). A typical tethered cell rotated at a mean rate of 0.2–1 Hz, depending on its length and the position of the stuck flagellum along its body. To facilitate data interpretation, we deleted the *cheY* gene (24, 25), yielding smooth-swimming mutants whose flagellar motors do not reverse.

As expected, there was no effect of green light on the cell’s rotation rate in the absence of azide (Fig. 2b). However, as we inhibited respiration by adding azide, the cells again became light-responsive. PR⁺ cells sped up upon illumination with green light [Fig. 2a and c, see also supporting information (SI) Movie 1]. The PR⁺ cells were converting light energy into an electro-

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**Author contributions:** J.M.W. designed research; J.M.W. performed research; J.M.W., D.G., and J.L. wrote the paper; D.G. analyzed data; D.G. performed model simulations; and C.B. and J.L. conceived the experiment.

The authors declare no conflict of interest.

Abbreviations: pmf, proton motive force; PR, proteorhodopsin.

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increased rapidly with intensity up to 10 mW/cm² (15 mM azide) or 20 mW/cm² (60 mM azide) (Fig. 3).

To determine the extent to which light can replace respiration as an energy source, we varied the azide concentration. Increasing the azide concentration from 0 to 110 mM caused cellular pmf levels, and therefore average cellular angular velocity, to decrease (SI Fig. 4a). The pmf dropped to 50% of its original value at an azide concentration of 55 mM, in agreement with inhibition studies of isolated bo cytochrome (26). When the cells were illuminated with intense green light, their rotation rate was restored to the speed of cells with an unimpaired respiratory system. As proton extrusion by the respiratory system dropped with increasing azide, PR provided an ever larger fraction of the pmf. As protonophore carbonyl cyanide 3-chlorophenylhydrazone (100 µM) to low oxygen cultures eliminated all bacterial motility, and movement was not restored with 50 seconds of green illumination (Fig. 2c; P ≤ 0.005, one-sided t test). Addition of the protonophore carbonyl cyanide 3-chlorophenylhydrazone (100 µM) to low oxygen cultures eliminated all bacterial motility, and movement was not restored with 50 seconds of green illumination (n = 300), demonstrating that cells unable to maintain a pmf cannot be revived.

To clarify the relationships between pmf, azide, and light, we constructed a highly simplified model of E. coli membrane fluxes. Our cells have multiple proton pumps that can contribute to the pmf (28), including PR, the respiratory chain, and the ATPase (Fig. 3c). Pmf is consumed by the flagellar motor and numerous transporters. In addition, the bacterial membrane has a basal permeability to protons (29). In a simpler system, Rotterdam et al. (30) showed that lipid vesicles with only one kind of ion pump reach a steady-state pmf whose magnitude is well approximated by a simple RC circuit.

Hypothesizing that an analogous circuit might be able to capture the functional relationship between pmf, azide, and chemical potential used to do mechanical work. PR⁺ cells stopped moving or slowed considerably when the green illumination light was removed. For example, at low concentrations of azide (5–15 mM), angular velocity dropped by one-fourth when the green light was removed. At higher azide concentrations (80–110 mM), angular velocity dropped further (50–60%) because more of the cells’ cytochromes were bound by azide (26).

To account the absorption cross-section of PR and the solar irradiation spectrum (detailed information is published as SI Text).

It was now essential to confirm the postulated mechanism of action. Reduced proton pumping by the respiratory system causes the pmf to drop. Light-based proton pumping by proteorhodopsin can then increase the pmf. Removal of oxygen from the cell culture also should lead to light-responsive bacteria with none of the possible confounding effects of azide. We needed to reduce O₂ levels substantially, because at 0.3 ppm, the E. coli respiratory system still generates ~50% of the normal pmf (27). We gently bubbled nitrogen through the cell culture for 15 min and then used a nitrogen-filled glovebox to prepare sealed imaging chambers containing 2 µl of cells. Just as we had seen upon addition of azide, PR⁺ cells became light-responsive upon oxygen depletion. Illumination increased the cells’ angular velocity by 45 ± 25% (Fig. 2c; P < 0.005, one-sided t test). Addition of the protonophore carbonyl cyanide 3-chlorophenylhydrazone (100 µM) to low oxygen cultures eliminated all bacterial motility, and movement was not restored with 50 seconds of green illumination (n = 300), demonstrating that cells unable to maintain a pmf cannot be revived.

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light, we constructed the model shown in Fig. 3d (see also SI Fig. 5) and parameterized it by fitting to the data shown in Fig. 3a and b and in SI Fig. 4. This model describes in vivo time-dependent dynamics between light-driven proton pumping and respiration and is described in detail in SI Text. Despite the model’s simplicity, it suggests why no effect of PR on growth rates has been reported. The model indicates that the maximum potential PR can generate by using the free energy from photon absorption ($V_{PR}$) is similar to the potential generated by *E. coli* respiration. Thus, in *E. coli* grown at neutral pH in rich or minimal media, or in *E. coli* respiring aerobically by using endogenous energy stores, PR cannot pump protons. Only when the pmf falls below the maximum potential ($V_{PR}$) during respiratory stress does PR begin to pump, and the proton flux through PR increases as the pmf falls. PR is able to maintain *E. coli* cellular pmf near this maximum potential ($V_{PR} \approx -0.2V$) with sufficiently bright illumination ($K_M \approx 60mW/cm^2$).

The cell motility studies together with the results of the pmf modeling raised the possibility that PR could pump sufficient protons to increase cell viability in addition to powering the

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*Fig. 2.* Flagellar response to green light measured in PR$^+$ and PR$^-$ bacteria observed at different levels of respiratory inhibition. (a) Movie frames showing a tethered cell in red light and green light (full movie is available at SI Movie 1), a plot of the angular position versus time of a cell in 60 mM azide, and a schematic of tethered cell geometry. In red light, the cell rotationally diffuses about its attachment point. Green light (shaded area) leads to counter clockwise rotation of the cell. (b) Cells without (b) and with (c) PR. Solid, dotted, and dashed lines show three representative cells per condition. Note the considerable cell-to-cell variation in absolute rotation rates due to variation of cell length and tethering geometry. With or without azide, cells lacking PR show no response to green light. Removal of oxygen also leads to light-responsiveness. At high azide concentrations (c Top), angular velocity is nearly zero until cells are illuminated with green light.
flagellar motor. To clarify a possible relationship between viability and proton pumping by PR, we plated cell cultures after their exposure to 30 mM azide for 30 min in sunlight. The cells lacking PR but having retinal were slightly more azide resistant; 1% of cells survived (SI Table 1). The cells with both PR and retinal were significantly more azide resistant than in all three other conditions (11% of cells survived; \( P \leq 0.005 \), Mann–Whitney \( U \) test). Consistent with the motility studies and the pmf model, PR is able to sustain cellular pmf at a level that increases viability. These findings directly illustrate the survival benefit of PR-based proton pumping under natural illumination conditions.

Our data demonstrate that during respiratory challenges, light-driven proton pumping by PR can augment the cellular pmf to the extent that it powers cell motility and increases cell survival, yielding cells that can withstand respiratory poisons and oxygen depletion via light harvesting. Conservation of the basic features of energy metabolism in proteobacteria such as SAR86 and SAR11 makes it likely that PR expression will confer similar benefits to other members of this class and, perhaps, even to more distantly related Gram-negative bacteria. Therefore, it may be possible to synthesize a diverse array of light-powered bacteria for a variety of purposes via PR expression combined with modulation of the cell’s natural pmf-generating mechanisms.

**Materials and Methods**

**Cell Culture.** We expressed the SAR86 \( \gamma \)-proteobacterial PR-variant (Geneart, Toronto, ON, Canada) in *E. coli* cells (RP437 DE3 \( \Delta \)cheY Cm\(^R\)) by using a T7-based expression system (pET200, Kan\(^R\), Invitrogen, Carlsbad, CA). Cells were grown in T broth (1% tryptone/0.5% NaCl) supplemented with kanamycin (25 \( \mu \)g/ml). In mid-log phase, PR expression was induced with 1 mM isopropyl \( \beta \)-d-thiogalactoside and the medium was supplemented with 10 \( \mu \)M ethanolic \textit{all-trans}-retinal. Cells were collected in late log phase by gentle centrifugation (4,500 \( \times \)g for 5min) and carefully resuspended in motility medium (1 mM PBS (Ambion, Foster City, CA)/0.1 mM EDTA, pH 7.4). Throughout the article, PR\(^+\) cells express proteorhodopsin, and PR\(^−\) denotes cells (RP437 DE3 \( \Delta \)cheY Cm\(^R\)) without the PR plasmid. PR\(^−\), unless otherwise noted, were also induced with \textit{all-trans}-retinal and grown in T broth with chloramphenicol (25 \( \mu \)g/ml). Unless otherwise noted, all experiments were done in glucose-free motility medium at room temperature.

**Instrumentation.** Throughout the article, power density values for “green light” refer to the power density passed by a D540/25\( \times \) filter (Chroma, Rockingham, VT) and originating at a 175 W Xenon bulb (Lambda light source; Sutter Instruments, Novato, CA), or, for “red light,” to the power density passed by a HQ620/60\( \times \) filter and originating at a 100 W Quartz Halogen bulb (Nikon). To visualize the cells, the sample chamber was illuminated continuously with faint red light (0.09 mW/cm\(^2\)) at the tail of PR’s absorption spectrum (4) (Fig. 1a). We periodically illuminated the sample chamber with bright green light (160 mW/cm\(^2\)) coinciding with the maximum of PR’s absorption spectrum, \(~=\) 525 nm (4). Cells were imaged at a framerate of 5

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**Fig. 3.** The response of PR\(^+\) bacteria to green light increases with respiratory inhibition and light intensity. (a) Benefit of illumination versus degree of inhibition of the respiratory system. The difference between angular velocity in green light and red light (\( \omega_{g} - \omega_{r} \)) becomes pronounced in PR\(^+\) bacteria (filled circles) as respiration is inhibited by low oxygen or sodium azide. PR\(^+\) cells (open circles) show no change between red and green illumination. To facilitate comparison between cells, the angular velocities are normalized by each bacterium’s maximum velocity, \( n = \frac{\omega_{g} - \omega_{r}}{\omega_{\max}} \). Green line, fit to model described in d. (b) The rotation speed of PR\(^+\) cells depends on the intensity of green illumination. Individual PR\(^+\) spinner cells were exposed to six intensities of green light. The mean angular velocity at each intensity is plotted \( (n = 5-6 \text{cells for each intensity}) \), normalized by the velocity at maximum illumination. Dashed lines, fits to model described in d. (c) Overview of transmembrane fluxes and proton pumping in PR\(^+\) *E. coli*. Sources of proton motive force include respiration and PR. Sinks include rotation of the flagellar motor and ATP synthesis. (d) Model including sources of pmf (respiration and proteorhodopsin), sinks (such as the flagellar motor), and the membrane capacitance. The variable resistors \( R_b \) and \( R_m \) model the effect of azide and light on proton extrusion by respiration and PR, respectively. The voltmeter (top-most circuit element) measures the potential difference across the membrane (equivalent to the pmf).
Hz by using an Andor (South Windsor, CT) iXon camera mounted to a Nikon TE2000 microscope. The Nikon microscope was modified for dark-field work by attaching a Zeiss 1.2–1.4 N.A. oil immersion dark-field condenser to the Nikon condenser turret using a custom adapter. Custom software written in C++ was used to control the Andor camera, and Matlab was used to process the images. All errors and error bars represent standard errors of the mean.

**Cell Viability Experiments.** PR+ and PR− cells were grown in LB plus Kanamycin (PR+) or LB plus Chloramphenicol (PR−) to an OD600 of 0.25–0.3, induced with 1 mM isopropyl β-D-thiogalactoside and supplemented with 10 μM all-trans-retinal (if retinal+), grown to an OD600 of 0.5–0.6, spun down (4,500 × g for 5 min), resuspended in motility buffer (1 mM PBS/0.1 mM EDTA) to an OD600 of 0.1–0.2. Colonies were counted manually after overnight incubation. The one-sided Mann–Whitney U test (31) was used to test the null hypothesis that the ability to pump protons does not increase cell survival.

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