



Multichromatic Control of Gene Expression in *Escherichia coli*

Jeffrey J. Tabor¹, Anselm Levskaya² and Christopher A. Voigt^{1*}

¹Department of Pharmaceutical Chemistry, School of Pharmacy, University of California San Francisco, MC 2540, Room 408C, 1700 4th Street, San Francisco, CA 94158, USA

²Graduate Program in Biophysics, University of California San Francisco, San Francisco, CA 94158, USA

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Light is a powerful tool for manipulating living cells because it can be applied with high resolution across space and over time. We previously constructed a red light-sensitive *Escherichia coli* transcription system based on a chimera between the red/far-red switchable cyanobacterial phytochrome Cph1 and the *E. coli* EnvZ/OmpR two-component signaling pathway. Here, we report the development of a green light-inducible transcription system in *E. coli* based on a recently discovered green/red photoswitchable two-component system from cyanobacteria. We demonstrate that the transcriptional output is proportional to the intensity of green light applied and that the green sensor is orthogonal to the red sensor at intensities of 532-nm light less than 0.01 W/m². Expression of both sensors in a single cell allows two-color optical control of transcription both in batch culture and in patterns across a lawn of engineered cells. Because each sensor functions as a photoreversible switch, this system should allow the spatial and temporal control of the expression of multiple genes through different combinations of light wavelengths. This feature aids precision single-cell and population-level studies in systems and synthetic biology.

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Introduction

Several genetically encoded tools have been developed for the optical regulation of molecular interactions inside of living cells. These include light-regulated transcriptional regulatory systems in bacteria^{1,2} and yeast³; light-dependent metabolic,⁴ signaling,⁵ and protein-splicing⁶ enzymes; a light switchable protein dimerization system⁷; and light-regulated neuronal ion channels⁸ and adrenergic receptors.⁹ These molecular genetic tools are unique in that they allow exquisite spatial and in some cases temporal control of cell states with minimal invasiveness.

Many biological and biotechnological applications require external control of cellular gene expression. To this point, all of the engineered light-regulated gene expression systems^{1–3} have been monochromatic: transcription from a given promoter is regulated (reversibly or irreversibly) by one set of light wavelengths. The development of multichromatic gene regulatory systems, where different light wavelengths regulate the expression of different genes, will allow more advanced control of synthetic and natural gene regulatory networks.

Phytochromes, a ubiquitous family of proteins that switch between active and inactive signaling states in response to red and far-red light,¹⁰ have previously been used for synthetic control of living cells. In the first reported example, a phytochrome/phytochrome-interacting factor (PIF) pair was adapted to a classical two-hybrid system to construct a light-regulated promoter in yeast.³ In another study, we fused the phytochrome Cph1 from *Synechocystis* PCC6803 to the *Escherichia coli*

*Corresponding author. E-mail address:

cavoigt@picasso.ucsf.edu.

Current address: J. J. Tabor, Department of Bioengineering, Rice University, MS142, 6100 Main Street, Houston, TX 77005, USA.

histidine kinase EnvZ to engineer a red light-regulated transcription system in *E. coli*.¹ More recently, we used a phytochrome/PIF pair from *Arabidopsis* to engineer rapidly photoswitchable protein dimerization (seconds time scale) in mammalian cells.⁷ Other approaches have employed the blue light-responsive LOV (light, oxygen, or voltage) domain to control gene expression and signal transduction.^{2,5} In contrast to the phytochrome-based tools, however, LOV-based systems respond unidirectionally to light exposure with dark-dependent relaxation of signaling occurring on the order of minutes to hours.¹¹

Recently, a cyanobacterial two-component system has been shown to induce the expression of a phycobilisome-related gene in response to green light.¹² The two-component system consists of the membrane-associated histidine kinase CcaS and its response regulator CcaR. CcaS is a member of the cyanobacteriochrome family of proteins, a cyanobacteria-specific relative of the phytochromes with blue-shifted absorption spectra.¹³ As in phytochromes, a bilin chromophore (in this case phycocyanobilin) binds at a conserved cysteine within an N-terminal GAF (cyclic GMP phosphodiesterase, adenyl cyclase, FhlA) domain and imparts reversible photoactivation of signaling activity with maximal responses to 535-nm (green) and 672-nm (red) light. Absorption of green light increases the rate of CcaS autophosphorylation, phosphotransfer to CcaR, and transcription from the promoter of the phycobilisome linker protein *cpcG2*, while absorption of red light reverses this process.¹²

Because they share a common chromophore and light absorption mechanism but have different chromatic specificities and transcriptional outputs, we hypothesized that CcaS/R may be able to function alongside our previously constructed red sensor (Cph8) for multichromatic control of gene expression in *E. coli*. Because CcaS is inactivated in the red band to which the Cph1/EnvZ chimera Cph8 responds, green and red light could be differentially applied to specifically induce transcription from each system. Moreover, because both sensors are photoreversible, such a system would allow multiplexed spatiotemporal control of gene expression.

Results

Cloning of the *ccaS/ccaR* cluster

To investigate whether the green light-inducible two-component system could function in *E. coli*, a plasmid expressing CcaS and CcaR and carrying a *lacZ* reporter fused to the P_{cpcG2} promoter (pJT118; Supplementary Fig. S2) was constructed. First, the

ccaS/ccaR/cpcG2 cassette was amplified from the genome of *Synechocystis* PCC6803 and cloned into a multicopy vector, generating plasmid pJT116 (Supplementary Fig. S2). The open reading frame of the output gene *cpcG2* was then seamlessly replaced with that of *lacZ* (Materials and Methods). The product of *lacZ*, β -galactosidase, was chosen as a reporter because it has previously proven tractable in both batch culture and plate-based light regulation experiments.^{1,14}

The plasmid pPLPCB(S) (Materials and Methods) was used to produce the chromophore PCB for green light sensor experiments. pPLPCB(S) carries the *Synechocystis* PCC6803 genes *ho1* and *pcyA*, which convert heme into PCB via a two-step oxidation/reduction process. pJT118 and pPLPCB(S) were cotransformed into *E. coli* strain JT2, a derivative of the strain previously used for red light sensor experiments (RU1012)¹⁵ from which a genomic fusion between the *ompC* promoter and *lacZ* was deleted (Materials and Methods).

Green light-induced gene expression in *E. coli*

Green light-induced transcription from P_{cpcG2} was assayed by growing *E. coli* expressing CcaS/R in liquid medium for 10 cell divisions in the dark or under 0.080 W/m² 532-nm light as described previously.¹⁴ Miller assays were conducted to determine the abundance of β -galactosidase per cell under each condition. Dark-exposed bacteria produced 24.7 ± 1.3 Miller units (M.U.), while those exposed to green light produced 50.7 ± 3.1 M.U. (Fig. 2a, $n=4$).

To determine whether the *E. coli* green light sensor functions as previously demonstrated *in vitro*,¹² cells were then exposed to inactivating red light. Exposure to 0.080 W/m² 650-nm light resulted in a slight reduction in β -galactosidase levels as compared to dark-grown cells (Fig. 2a). To determine whether the green light-dependent increase in gene expression is a specific effect of light absorption by the CcaS chromophore, the experiments were repeated in a strain lacking PCB. This strain showed no response to green or red light (Fig. 2a). These gene expression data agree with *in vitro* assays¹² indicating that in *E. coli*, the bilin-ligated (holo) form of CcaS is produced in the inactive green light-absorbing state (P_g) and is activated by green light and repressed by red light in a PCB-dependent manner.

Solid-phase light exposure experiments were then conducted to determine if the green light response could be visualized as patterns of gene expression across a lawn of cells. In agreement with the data in Fig. 2a, expression of β -galactosidase was induced only in areas of green light (Fig. 2b). Because CcaS adopts the inactive ground state in the dark, the rates of phosphotransfer to CcaR and resulting transcription of *lacZ* were low in dark-exposed

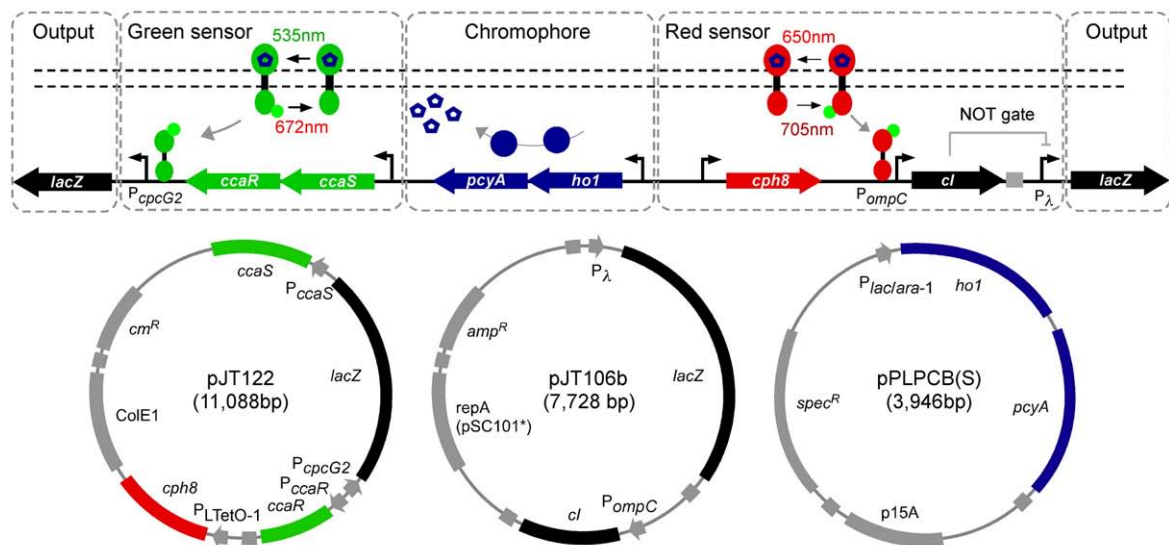


Fig. 1. Engineered two-color transcription induction system in *E. coli*. The green sensor and chromophore biosynthetic pathways are as described in the main text. The red light-sensing protein Cph8 is expressed from the $P_{LTetO-1}$ promoter in the phosphorylated ground state. It is switched to the unphosphorylated state by 650-nm light and back to the phosphorylated state by 705-nm light.³⁸ When phosphorylated, Cph8 passes a phosphoryl group to OmpR, which then binds to and activates transcription from P_{ompC} . Because it is inactivated by red light, Cph8 can be considered a NOT red sensor. A genetic inverter or NOT gate is used to invert the response of the NOT red sensor to that of a red light sensor. (Bottom) Plasmid maps of the green + red sensor plasmid pJT122, the red light inverter plasmid pJT106b, and pPLPCB(S), a variant of pPLPCB³⁸ in which the kanamycin resistance cassette has been replaced by a spectinomycin resistance cassette (Materials and Methods). Note that the true configuration of the DNA encoding this system is represented by the plasmid maps, while the version shown atop this figure is simplified for clarity.

areas of the plate. However, in green light-exposed areas, CcaS kinase activity increased, increasing the abundance of β -galactosidase and the cleavage of its chromogenic black substrate in the medium (Materials and Methods). Because the output signal is black, this results in a negative print of the projected image on the bacterial plate. Plate-based bacterial films expressing the green sensor do not respond to red images and are dependent on PCB (Fig. 2b).

Construction of a red light-activated genetic circuit

Transcription from the output promoter of the previously constructed red light sensor (P_{ompC}) is inversely proportional to the intensity of red light.¹ For many applications, including an initial demonstration of two-color optical gene regulation, a sensor that is activated by red light (analogous to the green sensor) is desirable. For this purpose, a genetic inverter¹⁶ was placed between the red light sensor and *lacZ*. Similar to our previously reported inverted red sensor,¹⁴ the CI repressor from phage λ is expressed as the output of P_{ompC} , and *lacZ* is expressed under the control of a CI repressible promoter (Fig. 1). Dark exposure therefore results in high-level production of CI repressor and repression

of *lacZ* transcription, while exposure to red light relieves this repression.

The performance of the red sensor was examined using Miller assays. Cells were grown for 10 generations in the dark or under 0.080 W/m^2 650-nm light (Materials and Methods). Dark-exposed cells produced 0.58 ± 0.01 M.U., while red light-exposed cells generated 1.41 ± 0.03 M.U (Fig. 2a). This 2.4-fold induction is similar to the green light response and is dependent on PCB (Fig. 2a). Unlike the green sensor, which remains inactivated in red light, the red sensor shows a minor response to high levels (0.080 W/m^2) of green light (532 nm; Fig. 2a). Lawns of bacteria expressing the red sensor print images of red light as negatives but do not respond significantly to images of green light (Fig. 2b).

Characterization of spectral transfer functions

The transfer function describes the quantitative relationship between the input and output of a genetic circuit.^{17–20} In the case of the light sensors, the input can be light wavelength or light intensity. The spectral transfer functions of the green and red sensors were determined by measuring transcriptional output relative to dark-exposed cells at different wavelengths of light between 430 nm and

730 nm (Fig. 2c). For each wavelength in Fig. 2c, high levels (0.080 W/m^2) of the respective wavelength were applied. In agreement with *in vitro* measurements of the absorbance of the CcaS holoprotein,¹² the green light sensor shows strongest transcriptional activation between 490 and 570 nm, with a maximum response near 535 nm. There is very little

induction in 610 nm (orange) and the sensor is inactive in 650 nm (red) light. By contrast, the red sensor is strongly induced in the 610–650-nm range. As expected, the red sensor is inactive in the far-red. The red sensor also has a long tail into the blue regions of the spectrum, although the magnitude of the response decreases significantly below 610 nm

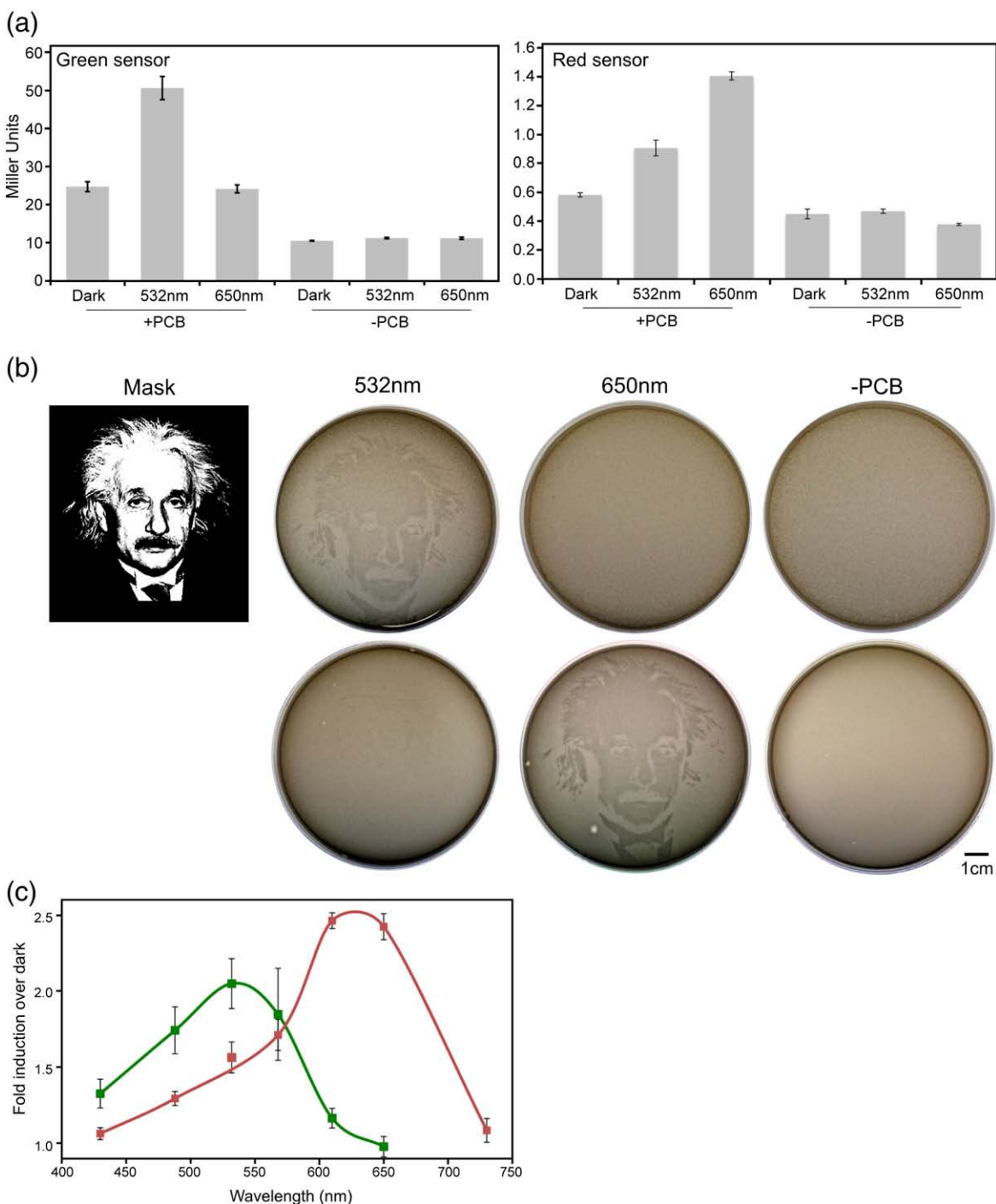


Fig. 2 (legend on next page)

(Fig. 2c). Despite the slight overlap, we determined that the separation in the action spectra of the two sensors could allow them to be combined for multiwavelength optical control of gene expression in a single cell.

Two-color optical control of gene expression

To investigate whether the two light sensors can function simultaneously in a single cell, a plasmid expressing both the green and red sensors (pJT122) was constructed (Fig. 1 and Materials and Methods). The light intensity transfer functions of the three sensor combinations (green only, red only, and both) were then determined for 532-nm and 650-nm inputs. Cells expressing the green sensor show sharp activation between dark and 0.01 W/m^2 532-nm light, above which point the response saturates (Fig. 3a). By contrast, bacteria expressing the red sensor show a small linear response to 532-nm light between dark and 0.080 W/m^2 . Cells expressing both sensors have a transfer function very similar to that of cells expressing the green sensor alone, although the total Miller unit output is slightly lower (Fig. 3a).

E. coli expressing only the red sensor are continuously induced by 650-nm light between dark and 0.01 W/m^2 , after which point the response largely saturates (Fig. 3a). The shape of the red sensor transfer function to 650-nm light is similar to that of the green sensor to 532-nm light, and both sensors respond over similar light intensities. While cells expressing only the green sensor are slightly repressed by 650-nm light, the addition of the red sensor causes the cells to be induced with a transfer function similar to cells expressing the red sensor alone (Fig. 3a).

The transfer functions of the red and green sensors are nonadditive when combined in a single cell. For

example, the decrease in Miller unit output of the green sensor in 650-nm light would be expected to offset the increase in Miller output from the red sensor, but this is not observed in the data. In fact, the presence of the green sensor leads to a greater increase in Miller output by the red sensor in response to 650-nm light (Fig. 3a). There are numerous direct or indirect interactions that could cause such nonadditivity. For example, the red sensor could more effectively compete for the chromophore, diminishing the response of the green sensor when both are present in a single cell. Alternatively, the kinase domain of the red sensor could dephosphorylate CcaR, the response regulator of the green sensor pathway, reducing signaling through the green pathway. Nonspecific effects such as competition for ribosomes²¹ or protein degradation machinery²² could also affect the expression level of a given sensor when the other is overexpressed. Follow-up investigations of these effects could inform future efforts in phytochrome engineering and the construction of synthetic signaling pathways in bacteria in general.

The data in Fig. 3a demonstrate that appropriate dosing of light wavelengths and intensities allows independent control of the sensors in a single cell. This was then demonstrated by projecting a composite green–red image onto agarose-embedded films of engineered bacteria. The intensity of projected green light was set at 0.02 W/m^2 , just above the saturation point of the green sensor, so as not to trigger unwanted induction of the red light sensor (Fig. 3b). When a strain expressing only the green sensor is exposed to this two-color image, β -galactosidase abundance increases sharply in the green areas and within regions of white light, but not in the red areas. Conversely, a strain expressing the red sensor is induced for β -galactosidase in red

Fig. 2. Transcriptional response of green and red sensors to different light conditions. (a) *E. coli* cultures were grown in the dark under 0.080 W/m^2 532-nm light or 0.080 W/m^2 650-nm light. +PCB, strain JT2 carrying the green (pJT118) or red sensor (pCph8 + pJT106b3) plasmids and pPLPCB(S). –PCB, JT2 carrying only the green or red sensor plasmids. Each data point represents the average of four separate cultures grown and measured in parallel on a single day. Data taken under different light conditions were collected on different days. Miller assays were conducted as reported previously.¹⁴ Error bars represent ± 1 SD. (b) Plate-based assays of green and red sensors. The mask shown was used to project an image of 532-nm or 650-nm filtered light onto an agarose-embedded film of bacteria expressing the green (top) or red (bottom) sensors. The chromogenic substrate S-gal (Sigma) and ferric ammonium citrate are added to agarose medium such that the product of *lacZ*, β -galactosidase, produces a visible black pigment when expressed. For all trials, 0.030 W/m^2 532-nm and 0.080 W/m^2 650-nm red light were projected through the mask. A slightly lower 532-nm intensity was used because the red sensor shows a minor response to 0.080 W/m^2 532-nm light [panel (a) and Fig. 3a]. The green sensor strain is the same as in panel (a). The red-sensor strain is JT2 carrying pCph8, pPLPCB(S), and pJT106b (a variant of pJT106b3 with a stronger ribosome binding site upstream of *lacZ*) for higher pigment production on plates. The –PCB condition indicates a given strain lacking pPLPCB(S) exposed to its cognate light wavelength. After 21 h, the bacterial plates produce images that can easily be seen by the naked eye with no further image enhancement. (c) Spectral transfer functions. *E. coli* carrying the green or red sensor [strains as in panel (a)] were exposed to saturating levels of a given light wavelength, and Miller assays were conducted as described in Materials and Methods. Data are reported as fold induction over dark-exposed cells. This is calculated by dividing the Miller unit value of the light-exposed cells by the Miller unit value of the same strain grown in the dark. Each data point represents the average of four separate cultures grown and measured in parallel on a single day. Data at different light wavelengths (or dark) were collected on different days. Error bars represent ± 1 SD. Miller assays were conducted as reported previously.¹⁴

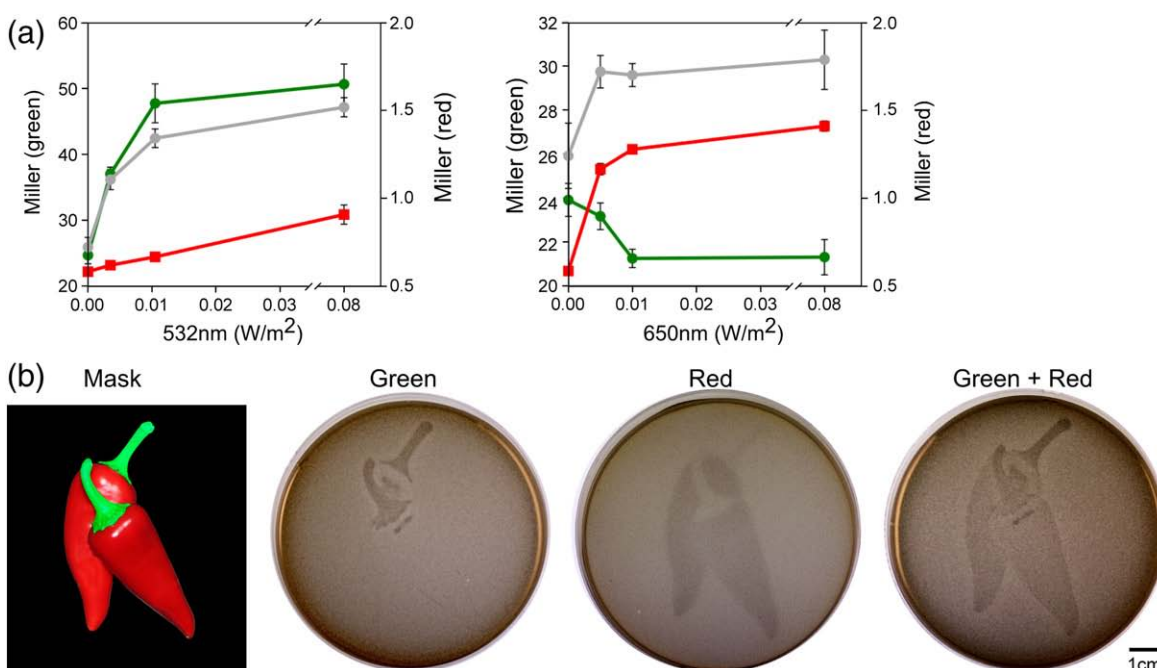


Fig. 3. Two-color optical control of gene expression in *E. coli*. (a) Light intensity transfer functions of strains carrying each sensor alone or both sensors. Strains expressing the green sensor only (green circles), red sensor only (red squares), or both (gray circles) were exposed to varying intensities of 532-nm or 650-nm light, and Miller assays were conducted as described in **Materials and Methods**. The green and green+red data (circles) correspond to the left axis, while the red sensor data (squares) correspond to the right axis. Two axes were used because the absolute Miller unit output of the RBS-weakened red sensor is low compared to the green sensor. Error bars represent ± 1 SD. (b) Two-color bacterial photography. A two-color mask was generated from a color-enhanced photograph of chili peppers. Green light passing through the stem regions of the image was set at 0.02 W/m^2 , slightly above the saturation point of the green sensor. At these illumination levels, the mask transmits $0.02\text{--}0.025 \text{ W/m}^2$ 650-nm light, above the saturation point of the red sensor. The same light intensities were used for all three plates. Green sensor- and red sensor-only strains are as described in the legend to Fig. 2a. Green+red strain is JT2 carrying plasmids pJT122, pJT106b3, and pPLPCB(S).

areas but only very slightly in green regions. Finally, in a strain expressing both sensors, β -galactosidase expression is induced by both colors of light (Fig. 3b).

The ribosome binding site upstream of *lacZ* in the red sensor was engineered to be weak (Supplementary Information), resulting in ~ 30 -fold lower β -galactosidase output from the red sensor as compared to the green sensor (Figs. 2a and 3a). On plates, this causes green light-exposed areas to appear darker than red light-exposed areas (Fig. 3b). The translation of color information to differences in monochrome intensity results in grayscale effects, which compensate for the lack of visually distinct (color) outputs. Despite the slight nonadditivities that occur when the sensors are combined, the data in Fig. 3 demonstrate that the expression of a second sensor does not significantly change the response of a given sensor to its cognate light wavelength, and that coexpression of the two sensors allows two-color optical control of gene expression in a single cell.

Discussion

Several light-regulated transcriptional regulatory systems have previously been constructed.^{1–3} By combining an *E. coli* red light sensor with a recently discovered green light sensor from *Synechocystis*, we have engineered a multichromatic gene regulatory system where different promoters are controlled by different wavelengths in a single cell. This system has several unique properties. Because most gene regulatory systems rely on the addition of chemicals to the growth medium, modulation of gene expression is often unidirectional, with reversal depending on the decay or dilution of the effector compound. By contrast, both sensors reported here function as switches that can be toggled between states by different light wavelengths,^{12,23,24} a feature that allows more precise temporal control of gene expression.

The reversible behavior of the green and red light sensors begins at the phycocyanobilin (PCB) chromophore. After PCB ligation, the holoprotein adopts

a stable green- or red-absorbing ground state (P_g or P_r). Picoseconds after absorption of the activating photon, isomerization of PCB drives a conformational rearrangement of the surrounding protein, which occurs on the order of milliseconds to seconds.²⁵ Structural changes in the light-sensing domains are then transmitted to the kinase domains, activating phosphosignaling. Phosphotransfer from the histidine kinase to its response regulator then occurs in milliseconds.²⁶ When phosphorylated, the response regulator binds its cognate promoter and induces transcription. This occurs in minutes but can take on the order of 1 h to reach steady state.²⁷ The light sensors should therefore allow reversible control of transcription on the minutes time scale. Because of the relatively slow nature of gene expression and protein decay, however, time periods on the order of hours will be required to switch between on and off steady states. This would also be the case for chemically regulated transcription systems, although these systems do not have the benefit of reversibility.

Modern optical methods such as two-photon excitation or digital micromirror devices allow the projection of light patterns at subcellular resolution. If combined with such optics⁷ and fluorescent or luminescent reporter genes, the two-color system described here should allow real-time control and observation of the expression of multiple genes in individual cells within a larger population. This would enable facile external patterning of phenotypes and studies of time-dependent multicellular phenomena such as biofilm formation.

The action spectra for the two light sensors in this study partially overlap (Fig. 2c). The maximum inactivating wavelength of the green sensor is effectively the same as the maximum activating wavelength of the red sensor. The result is that there are only three “cognate” control wavelengths for four possible states. The red sensor also has a long blue tail, showing induction in response to wavelengths as low as 490 nm (Fig. 2c). Despite these overlaps, there are a variety of strategies for achieving independent control of the four sensor states. For example, intensities of 532-nm light less than ~ 0.01 W/m² activate the green sensor while leaving the red sensor inactive (Fig. 3a). Far-red light (730 nm) can also be applied concomitantly with 532 nm to inhibit the red sensor while activating the green sensor. To activate both sensors simultaneously, high levels (0.08 W/m² or greater) of an intermediate wavelength such as 575 nm can be used (Fig. 2c).

To improve performance, the action spectra of the light sensors themselves could also be engineered. A number of mutations in the light-sensing domains of phytochrome-related proteins have been generated and shown to alter the absorbance spectra.^{28–30} As expected, mutations in the chromophore binding pocket affect absorption, but other more dramatic mutations in domain architecture likely play a role

as well. Indeed, the green-sensing cyanobacteriochrome protein used here has several binding pocket mutations in conserved residues as well as domain organization differences as compared to the red sensor.¹² Because we have linked the sensors to gene expression outputs in *E. coli*, standard laboratory evolution methods targeting critical amino acids in the chromophore binding domain or even the domain shuffling methods altering the overall architecture of the phytochrome could potentially be used to rapidly generate new light sensors or to narrow the spectral sensitivities of existing sensors.

Synthetic gene circuits could be also used to filter the responses of the existing light sensors. For example, a bistable genetic switch³¹ could be placed between the light sensors and the output genes. Bistable circuits cannot rest in intermediate output states but switch digitally from low to high output in response to continuous changes in input signal.³² Because the responses of the two light sensors decrease symmetrically with distance from the maximal inducing wavelength (Fig. 2c), a bistable switch could cut off responses below a certain threshold, effectively narrowing the action spectra of the sensors.

The output of the two light sensors reported here changes continuously with input (Fig. 3a). Gradients of light can therefore be used to set different transcription levels across space in solid-phase experiments (Supplementary Fig. S1). The light sensors could be connected to genetic circuits, and one- or two-dimensional light gradients could be applied to determine circuit transfer functions over a continuous range of inputs in a single experiment. The ability to measure two-dimensional transfer functions in a single step could prove very useful in both systems³³ and synthetic biology studies.^{34,35}

The system reported here represents the first engineered multichromatic gene regulatory system, whereby the expression of different genes can be controlled by different wavelengths of light. Multichannel optical regulation of neuronal membrane potentials stands to revolutionize neurobiology by allowing unprecedented temporal control of neuronal activity *in vivo*.³⁶ The multiplexed optical control of gene expression should find broad utility in scientific, engineering, and industrial applications.

Materials and Methods

Plasmid construction

Construction of pJT116

The fragment of the *Synechocystis* PCC6803 genome bearing the *ccaS-ccaR* cluster (chromosomal position 3399457–3405249) was amplified with the primers TACTAGACTAGATCAGAGTACGATCAGTCATGACTAGACGATCGGACGTCCTAAGCTCGAGGCAAATGG

and TGTCATGTATCGTCAATGGTACTGACTCTACTC-AATACGTTCTAGATCTTCTAGACTAGTTTTTCCC-TTGGCAC from purified genomic DNA and cloned into the pProTet.E333 backbone (Clontech, Mountain View, CA) at the AatII and XbaI sites (underlined). The use of these sites removes the $P_{LTetO-1}$ promoter, ribosome binding site, 6 \times His tag, and MCS while leaving the downstream transcription terminator. The endogenous *Synechocystis* promoters are therefore responsible for the expression of CcaS and CcaR in this plasmid. pJT116 was maintained with 34 μ g/mL chloramphenicol.

Construction of pJT118

The green light-inducible *cpcG2* open reading frame¹² in pJT116 was seamlessly replaced with *lacZ* using the MEGAWHOP protocol.³⁷ *lacZ* was amplified from pEX-*PlacZ* (Invitrogen, Carlsbad CA) with the primers GATA-TAACAGTATAGATTTTTGTCAGCCTTCAGCTTGGCTT-TACCGTCAAAAAAATTAGACTCGAGCGGCCG and CACA TACCAGTTATTGGCTGGACATTTAAA-CAACTTTTAAGTTAATTACTA AACTTTATCTATGATAGATCCCGTCGTTTACAACG to generate the green light-responsive reporter plasmid pJT118 (*lacZ* binding regions are underlined). pJT118 was maintained with 34 μ g/mL chloramphenicol.

Construction of pJT122

An expression cassette for the red light-responsive *oph1/envZ* chimera *cph8*¹ was added to the green light reporter plasmid pJT118 to generate plasmid pJT122. The $P_{LTetO-1}$ promoter, ribosome binding site, and *cph8* open reading frame¹ were amplified using the primers GCCCTAGACCTAGGGCGTTCGGCTGCGGC-GAGCGGTATCACCTTTCGTTCTTACCTCGAG and GTTCTTTCCTGCGTTATCCCCTGATTCTGTGGATTAACCGTATTACCGCCTTTGAGTGAGCT-TACCTTCTTTGTATGCCCC (promoter and *cph8* binding sequences are underlined, respectively). This PCR product was then used as a megaprimer in a MEGAWHOP reaction to clone the cassette downstream of the *ccaR* transcription terminator in pJT118. pJT122 was maintained with 34 μ g/mL chloramphenicol.

Construction of pJT106b

pJT106b encodes a red light inverter circuit driving a *lacZ* reporter gene. pJT106b is derived from pJT106, which carries the P_{ompC} promoter BBa_R0082 driving the *cl* gene, the product of which represses the LuxR+3OC₆HSL-activated, CI-repressed output promoter BBa_R0065.¹⁴ In pJT106b, R0065 is replaced by a LuxR+3OC₆HSL-independent CI repressible promoter, BBa_J64067. To make J64067 LuxR+3OC₆HSL independent (and increase the overall transcription rate), the weak -35 site of R0065 (TTTACG) was replaced with a consensus TTGACA site, the suboptimal 16-bp spacer between the -35 and -10 sites was replaced with a 17-bp spacer, and the -16 nucleotide was swapped from T to G. A megaprimer encoding these four mutations was generated by amplifying the R0065 region of pJT106b with the primers CGTACAGGTTGACAAACAAGAAAATGGTGTGTTATAGTCG and CAT-TAAATGTGAGCGAGTAACAACCCG (mutations are

underlined). This megaprimer was then used to extend pJT106 in a MEGAWHOP to generate pJT106b. pJT106b was maintained with 50 μ g/mL ampicillin.

Construction of pPLPCB(S)

Because strain JT2 bears native kanamycin resistance, pJT118 and pJT122 bear chloramphenicol resistance, and pJT106b bears ampicillin resistance, a variant of plasmid pPLPCB³⁸ carrying a spectinomycin resistance marker was constructed. To this end, the spectinomycin resistance cassette (including promoter, ribosome binding site, and *spec^R* gene) were amplified from plasmid pKD13³⁹ using the primers AGAGCCTAGACCATAGACATAGAATA-TACGTACGGGCCCCAGCAAGCGAACC GGAAATTGCC and TATATTGACTCTAGCTCTAACTCTATGGGCTC-TAGAGCTCTTATTGCGGACTACCTTGG (primer binding sites are underlined) and cloned into pPLPCB using ApaI and SacI, which remove the kanamycin resistance cassette. pPLPCB(S) was maintained with 100 μ g/mL spectinomycin.

Bacterial strains

Strain JT2 (RU1012 $\Delta P_{ompC-lacZ}$) was used for all experiments. JT2 was constructed using the Datsenko–Wanner method³⁹ to knock out the region of the RU1012 genome within which the *lacZ* gene was fused to the *ompC* open reading frame.¹⁵ The entire knocked out region contains, in order, the *ompC* promoter followed by the first 789 bp of the *ompC* gene, a translational fusion between the first 177 bp of the *E. coli* tryptophan synthase α subunit and *lacZ*, *lacY*, a truncated *lacA*, and a second copy of the *ompC* promoter driving a second copy of the *ompC* gene, which is internally disrupted by a Tn5 transposon carrying a kanamycin resistance marker. Although the embedded kanamycin resistance marker used to make this fusion was deleted in this step, the strain maintained resistance to kanamycin at 50 μ g/mL. This suggests that at least one additional, unannotated kanamycin resistance marker is present in the genome of RU1012. Strain JT2 was grown in the presence of 50 μ g/mL kanamycin for all experiments. The primers used to generate the knockout PCR fragment were

GAATTATTATTGCTTGATGTTAGGTGCT-TATTTCCGCAATCCGCAATAATCTTAAAAAGTGTG-TAGGCTGGAGCTGCTTC

and

TTGTACGCTGAAAACAATGAAAAAGGGCCCG-CAGGCCCTTTGTTTCGATATCAATCGA-GAATCCCGGGGATCCGTCGACC,

which bear homology to the region immediately upstream of the *ompC* promoter and immediately downstream of the end of the *ompC* ORF.

Miller assays

Overnight cultures were grown in 3 mL of unbuffered LB broth (Lennox formulation) + appropriate antibiotics to OD₆₀₀ ~3–4. These cultures were diluted to OD₆₀₀ = 0.001

in 1 mL of fresh LB (Lennox)+0.1 M Hepes (pH 6.6)+ appropriate antibiotics, grown for 10 cell divisions (to $OD_{600}=1.0$) and subjected to Miller assays as described previously.¹⁴ Light was projected onto the growing cultures as described before¹⁴ using the following band-pass filters (Edmund Optics, Barrington NJ): 430 nm NT43-160, 488 nm NT43-168, 532 nm NT43-174, 568 nm NT43-179, 610 nm NT43-183, 650 nm NT43-189, and 730 nm NT43-195. Replicates were grown in parallel on a single day, while data for different light intensities and wavelengths were collected on different days.

Determination of light intensity

The intensity of light was measured in power units of watts per square meter using a EPP2000 UVN-SR calibrated spectroradiometer (Stellarnet, Tampa, FL) with a collection window ± 30 nm from the reported (peak) wavelength. The bandpass filters used in these experiments have 10-nm transmission windows centered on the peak emission wavelength.

Plate assays

Plate assays were conducted as described previously¹⁴ except that starter cultures were grown overnight in unbuffered LB broth+appropriate antibiotics. The light exposure step was carried out for 21 h, except in the case of red sensor-only cells carrying the weak *lacZ* ribosome binding site (plasmid pJT106b3), in which case, light exposure was carried out for 48 h to allow the accumulation of more black pigment.

The two-color mask used in Fig. 3b was generated by taking a photograph of chili peppers (Whole Foods, San Francisco, CA) with a Canon EOS Rebel SLR camera with a macro lens and hood. The background was made black and the RGB characteristics of the chilis were then enhanced using Adobe Photoshop. Although the color composition of the chili regions of the image was greater than 90% red, the stems contained significant red, green, and blue components. To remove the red and blue components, the stems were isolated using the magic wand function and their color balance was minimized away from red and blue (toward cyan and yellow) and toward green (away from magenta). A color-enhanced tiff file was then used as the template to fabricate a 35-mm slide (Oscar's Photo Lab, San Francisco, CA), which was used to mask a white light projector as reported previously.¹⁴ Light transmission through the stem and chili regions of the mask was verified to be almost exclusively green and red, respectively, using a spectroradiometer as described earlier.

Imaging of agarose plates

Agarose plates were placed face up on a white fluorescent light box, and photographs were taken in a darkened room with a Canon EOS Rebel SLR camera with a macro lens and hood. Image levels, tone, contrast, and shadowing were adjusted using Adobe Photoshop (Adobe Systems Inc., San Francisco, CA) to more accurately represent the appearance of the agarose plates to the naked eye.

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Supplementary Data

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.jmb.2010.10.038](https://doi.org/10.1016/j.jmb.2010.10.038)

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