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1	Optogenetic control of gut bacterial metabolism to promote longevity
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# 12 Abstract

13 Gut microbial metabolism is associated with host longevity. However, because it requires 14 direct manipulation of microbial metabolism *in situ*, establishing a causal link between these two 15 processes remains challenging. We demonstrate an optogenetic method to control gene 16 expression and metabolite production from bacteria residing in the host gut. We genetically 17 engineer an Escherichia coli strain that synthesizes and secretes colanic acid (CA) under the 18 quantitative control of light. Using this optogenetically-controlled strain to induce CA production 19 directly in the *Caenorhabditis elegans* gut, we reveal the local effect of CA in protecting 20 intestinal mitochondria from stress-induced hyper-fragmentation. We also exploit different 21 intensities of light to determine that the lifespan-extending effect of CA is positively correlated 22 with its levels produced from bacteria. Our results show that optogenetic control offers a rapid, 23 reversible and quantitative way to fine-tune gut bacterial metabolism and uncover its local and 24 systemic effects on host health and aging.

#### 26 Introduction

27 Microbiome studies have identified correlations between bacteria and host aging (Kundu 28 et al., 2017; O'Toole and Jeffery, 2015). For example, 16S rRNA and metagenomic DNA 29 sequencing are used to associate the presence or abundance of specific bacteria to human 30 centenarians (Biagi et al., 2016; Claesson et al., 2012, 2010). However, given the complexity and 31 heterogeneity of the human gut environment, these approaches are unable to elucidate how a 32 specific microbial species contributes to longevity. The nematode *Caenorhabditis elegans* has a 33 short and easily-measured lifespan, features that have revolutionized our understanding of the 34 molecular genetics of aging and longevity (Kenyon, 2010). Studies using C. elegans also provide 35 mechanistic insight into the association between bacterial species and host longevity (Gusarov et 36 al., 2013; Kim, 2013). Importantly, recent studies have revealed that bacterial metabolism can 37 produce specific products to directly influence the aging process in the host C. elegans or 38 modulate the effects of environmental cues on C. elegans lifespan (Cabreiro et al., 2013; Pryor et 39 al., 2019; Virk et al., 2016). These findings highlight the significance of bacterial metabolism in 40 regulating host physiology during the aging process and have inspired interest in directly 41 manipulating bacterial metabolism *in situ* in the host gastrointestinal (GI) tract. 42 In several recent studies, researchers have administered antibiotic- or carbohydrate-based 43 small molecule inducers to modulate gene expression from gut bacteria (Kotula et al., 2014; Lim 44 et al., 2017; Mimee et al., 2015). While this approach has enabled the *in situ* analysis of a gut 45 bacterial-host interaction (Lim et al., 2017), chemical effectors may have unwanted side-effects

46 on host or microbial physiology and subject to slow and poorly-controlled transport and
47 degradation processes that ultimately limit their precision.

Optogenetics combines light and genetically-engineered photoreceptors to achieve 48 49 unrivaled control of biological processes (Olson and Tabor, 2014). Previously, we and others 50 have engineered bacterial photoreceptors that activate or repress gene expression in response to 51 specific wavelengths of light (Levskaya et al., 2005; Li et al., 2020; Ohlendorf et al., 2012; Ong 52 and Tabor, 2018; Ong et al., 2017; Ramakrishnan and Tabor, 2016; Ryu and Gomelsky, 2014; 53 Schmidl et al., 2014). These photoreceptors have been used to achieve precise quantitative 54 (Olson et al., 2017, 2014), temporal (Chait et al., 2017; Milias-Argeitis et al., 2016; Olson et al., 55 2017, 2014), and spatial (Chait et al., 2017; Levskaya et al., 2005; Ohlendorf et al., 2012; Tabor 56 et al., 2010) control of bacterial gene expression in culture conditions. They have also been used 57 to characterize and control transcriptional regulatory circuits (Chait et al., 2017; Olson et al., 58 2014; Tabor et al., 2009) and bacterial metabolic pathways (Fernandez-Rodriguez et al., 2017; 59 Tandar et al., 2019) in vitro. Here, we hypothesized that optogenetic control of bacterial gene 60 expression might provide a new way to manipulate bacterial metabolism in vivo in the host GI 61 tract, with high temporal and spatial precision and no unwanted side-effects.

62 We address this possibility using the *E. coli-C. elegans* interaction model, a testable 63 system with known mechanistic links between bacterial metabolism and host longevity and with 64 complete optical transparency. In particular, CA is an exopolysaccharide synthesized and 65 secreted from E. coli, which can extend the lifespan of the host C. elegans through modulating 66 mitochondrial dynamics (Han et al., 2017). We thus have genetically engineered an E. coli strain 67 to put its biosynthesis of CA under a switchable control between green and red lights, and then 68 utilized green light to induce CA production from this strain in the gut of the host *C. elegans*. We 69 discovered that light-induced CA from bacteria residing in the host is sufficient to modulate 70 mitochondrial dynamics and lifespan, which gives even more potent effects than dietary

supplementation of CA. Furthermore, this optogenetic manipulation allowed us to investigate the local effect of CA on intestinal cells in a time-controlled manner and its systemic effect on organisms in a quantitative way, which are not possible with purified CA or CA-overproducing genetic mutants. This work paves the road for future application of bacterial optogenetics in understanding bacteria-host interaction with temporal, spatial and quantitative controls and minimal chemical interference.

77

### 78 **Results**

79 To demonstrate optogenetic control over gut bacterial gene expression, we first 80 engineered E. coli strain LH01, wherein our previous light-switchable two-component system 81 CcaSR (Schmidl et al., 2014) controls expression of superfolder green fluorescent protein (*sfgfp*), 82 and *mcherry* is expressed constitutively to facilitate identification of the bacteria (**Fig. 1a**, 83 Supplementary Fig. 1, Supplementary Tables 1-3). In LH01, green light exposure switches 84 CcaS to an active state in the presence of chromophore phycocyanobilin (PCB), wherein it 85 phosphorylates the response regulator CcaR. Phosphorylated CcaR then activates transcription of sfgfp from the P<sub>cpcG2-172</sub> output promoter. Red light reverts active CcaS to the inactive form, de-86 87 activating *sfgfp* expression (**Fig. 1a**).

We then reared two groups of *C. elegans* from the larval to the adult stage on plates of LH01 under red or green light, respectively (**Fig. 1b**). Next, we washed away external bacteria, applied the paralyzing agent levamisole to prevent expulsion of gut contents, and transferred the worms to agar pads. Finally, we switched the light color from red to green, or green to red, and used epi-fluorescence microscopy to image the resulting changes in fluorescence in the gut lumen over time (**Fig. 1c**). In the red-to-green (step ON) experiment, we observed that sfGFP fluorescence in the worm gut lumen starts low, begins to increase within 2 hours, and reaches a saturated high level at 6 hours (**Fig. 1d**). In contrast, in the green-to-red (step OFF) experiment, sfGFP fluorescence begins high, and decreases exponentially between hours 1-7 (**Fig. 1d**). This light response is abolished when the PCB biosynthetic operon is removed ( $\Delta$ PCB) (**Fig. 1d**), demonstrating that sfGFP levels are specifically controlled by CcaSR.

99 Next, we used flow cytometry to achieve high-throughput single-cell resolution analysis 100 of this *in situ* bacterial light response. Specifically, we reared worms in red and green light as 101 before, but then washed, paralyzed, and placed them into microtubes prior to light switching 102 (Fig. 1b). At several time points over the course of 8 hours, we homogenized the animals, 103 harvested the gut contents, and sorted bacterial cells and measured fluorescence via cytometry. 104 This experiment revealed that bacteria in the host gut remain intact (Supplementary Fig. 2) and 105 respond to light in a unimodal fashion (Fig. 1e-g). Furthermore, the temporal dynamics of the 106 gene expression response and dependence on PCB recapitulate our microscopy results 107 (Supplementary Fig. 3). We also confirmed that residual bacteria on the exterior of worms do 108 not contribute to the flow cytometry measurements (Supplementary Fig. 4). Together, these 109 experiments demonstrate that we can use optogenetics to rapidly, reversibly induce gene 110 expression of *E. coli* residing in the *C. elegans* gut.

111 Next, we sought to utilize our optogenetic method to modulate the production of specific 112 metabolites in bacteria residing in the gut of live hosts. It is well-known that bacterial genes 113 involved in the same metabolic process are often clustered into operons and co-regulated at the 114 transcriptional level. We took advantage of this coordinated mode of regulation and chose the 115 *cps* operon and its transcription activator RcsA for testing optogenetic control of bacterial 116 metabolism. The *cps* operon in *E. coli* consists of 19 genes that encode enzymes required for the

117 biosynthesis and secretion of CA (Torres-Cabassa and Gottesman, 1987), and CA-overproducing 118 bacterial mutants  $\Delta lon$  and  $\Delta hns$  promote longevity in the host C. elegans (Han et al., 2017). To 119 place CA biosynthesis under optogenetic control, we engineered an E. coli strain (MVK29) 120 lacking genomic rcsA and expressing a heterologous copy of rcsA under the control of CcaSR 121 (Fig. 2a, Supplementary Tables 1-3). We first examined whether MVK29 could respond to 122 green light and induce CA production and secretion. To this end, we grew the strain in batch 123 culture under red or green light and quantified supernatant CA levels. In red light, MVK29 124 secretes CA to concentrations below the limit of detection of the assay, similar to the  $\Delta rcsA$ 125 mutant (Fig. 2b). Green light, on the other hand, induces MVK29 to secrete high levels of CA, 126 and removal of the PCB biosynthetic operon abolishes this response (Fig. 2b). Moreover, 127 mutation of the CcaS catalytic histidine to a non-functional alanine (H534A), or the CcaR 128 phosphorylation site from an aspartic acid to a non-functional asparagine (D51N) abolishes 129 detectable CA production (Fig. 2b). Importantly, the level of secreted CA increases sigmoidally 130 with green light intensity, similar to the response of CcaSR itself (Fig. 2c) (Schmidl et al., 2014). 131 We conclude that we have placed CA production under the control of the CcaSR system, and 132 that we can use light to tune the production of bacterial metabolites.

We then used this approach to study a gut bacterial metabolite-host interaction pathway *in vivo*. We first reared worms expressing mitochondrially-localized GFP (mito-GFP) (**Supplementary Table 3**) on MVK29 with red light. We then continued this red-light exposure for one group, and switched a second to green, for an additional 6 hours, and immediately imaged intestinal cell mitochondrial morphology using confocal microscopy (**Fig. 3b**). We found that mitochondrial fragmentation increases in worms exposed to the bacteria with light-induced 139 CA secretion (Fig. 3c). This result recapitulates the phenotype observed in worms supplemented
140 with purified CA (Han et al., 2017).

141 Next, we induced CA production directly from bacteria residing within the host gut. To 142 this end, we paralyzed the worms carrying mito-GFP, split them into two groups, and treated one 143 with red and the second with green for 6 hours (Fig. 3a). We then analyzed mitochondrial 144 morphology in intestinal cells using confocal microscopy. We found that the 6-hour levamisole 145 treatment does not kill worms but leads to mitochondrial hyper-fragmentation (Fig. 3d), which 146 might be due to the inhibitory effect of levamisole on mitochondrial NADH-oxidizing enzymes 147 (Köhler and Bachmann, 1978). This stress-induced effect resembles mitochondrial decay related 148 to aging and age-related neurodegenerative diseases (Cho et al., 2009; Exner et al., 2007; 149 Sebastián et al., 2017). Interestingly, we found that green light exposure counteracts this hyper-150 fragmentation in paralyzed worms bearing MVK29 in the gut (Fig. 3d). Importantly, we found 151 no such effects in worms bearing the  $\Delta PCB$ , CcaS(H534A), CcaR(D51N) or  $\Delta rcsA$  mutant 152 strains (Fig. 3d), suggesting that this protective effective is a result of light-induced CA 153 overproduction in the gut. These results not only show that optogenetics can be utilized to induce 154 CA secretion from gut-borne E. coli in vivo, but also reveal a local protective effect of CA on 155 intestinal cells.

Finally, we took the advantage of the quantitative control afforded by optogenetics to investigate how the lifespan-extending effect of CA relates to CA levels. Beginning at the day-1 adult stage, we exposed worms bearing MVK29 to red, or two green light intensities resulting in intermediate or high CA secretion, and measured their lifespans. We found that the lifespan extension increases proportionally with green-light intensity (**Fig. 4a**), revealing the prolongevity effect of CA is dose dependent. We also noticed that the extent of lifespan extension

162 by light is much stronger than that caused by dietary supplementation of purified CA (Han et al., 163 2017), suggesting the high efficacy of optogenetic induction to modulate host physiology. As a 164 control, we repeated the experiment with worms bearing the CA-overproducing  $\Delta lon$  mutant and 165 showed that the lifespan extension caused by  $\Delta lon$  is independent of light exposure (Fig. 4b). 166 The extent of lifespan extension is similar to the MVK29 intermediate green light condition, but 167 less than the MVK29 high green light condition (**Fig. 4a**, **b**), suggesting that MVK29 is capable 168 of producing higher levels of CA than  $\Delta lon$ . In addition, the lifespan of worms bearing the  $\Delta rcsA$ 169 mutant is also light-independent and is comparable to that of MVK29 worms under red light 170 (Fig. 4a, c). These results suggest that optogenetic control is sufficient to induce bacterial 171 production of pro-longevity compounds and improve host health, and can exert stronger 172 beneficial effects than administration of a bacterial mutant or supplementing purified 173 compounds. Importantly, unlike the traditional approach of introducing a bacterial mutant, 174 optogenetic control of bacterial metabolism can modulate a host-level phenotype in a 175 quantitative manner.

176

### 177 Discussion

Our method has broad applications for studying microbe-host interactions *in situ*. For example, we have identified about two dozen additional *E. coli* genes that are unrelated to CA biosynthesis and that enhance worm longevity when knocked out (Han et al., 2017), though the mechanisms by which they act remain largely unclear. By using light to induce their expression in the gut and measuring acute host responses such as changes in mitochondrial dynamics, the role of these genes in gut microbe-host interactions could be further explored. In another example, the quorum-sensing peptide CSF and nitric oxide, both of which are produced by *Bacillus subtilis* during biofilm formation, have been found to extend worm lifespan through downregulation of the insulin-like signaling pathway (Donato et al., 2017). We have recently ported CcaSR into *B. subtilis* and demonstrated that it enables rapid and precise control of gene expression dynamics (Castillo-Hair et al., 2019). The method we report here should enable *in situ* studies of how gene expression and metabolite production from this important Gram-positive model bacterium impact longevity as well.

191 Multiple photoreceptors could also be combined to study more complex microbe-host 192 interaction pathways. Specifically, we and others have co-expressed CcaSR with independently-193 controllable blue/dark and red/far-red reversible light sensors in order to achieve simultaneous 194 and independent control of the expression of up to three genes in the same bacterial cell 195 (Fernandez-Rodriguez et al., 2017; Olson et al., 2017; Tabor et al., 2010). Such optogenetic 196 multiplexing could be performed in situ and used to study potential synergistic, antagonistic, or 197 other higher-order effects of multiple bacterial genes or pathways. A large number of eukaryotic 198 photoreceptors have also been developed, enabling optical control of many cell- and 199 neurobiological processes (Deisseroth, 2015; Gautier et al., 2014; Goglia and Toettcher, 2018; 200 Leopold et al., 2018). Bacterial and eukaryotic photoreceptors could be combined to enable 201 simultaneous optical manipulation of bacterial and host pathways in order to interrogate whether 202 or how they interact. Optogenetics could also be used to manipulate bacterial and/or host 203 pathways at specific locations within the gut to examine location- or tissue-dependent 204 phenomena.

Finally, our method could be extended to other bacteria or hosts. In particular, it should be possible to port CcaSR or other bacterial photoreceptors into native *C. elegans* symbionts (Zhang et al., 2017) or pathogens(Couillault and Ewbank, 2002). Because these strains stably

colonize the host, the use of these bacteria could eliminate the need for paralysis, and facilitate longer-term experiments. It is likely that light can also be used to control gut bacterial gene expression in other model hosts such as flies, zebrafish, or mammals. Red-shifted wavelengths and corresponding optogenetic tools(Ong et al., 2017; Ryu and Gomelsky, 2014) may prove superior for less optically transparent or larger animals. Overall, by enabling precision control of bacterial gene expression and metabolism *in situ*, we believe that optogenetics will greatly improve our understanding of a wide range of microbe-host interactions.

215

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225

#### 226 Author Contributions

JJT and MW conceived of the study. LAH and MVK designed experiments. MVK and LAH

- 228 constructed plasmids and strains. LAH, MVK, BH, CJL, LG, and MW performed experiments.
- 229 CJL and LG scored single-blinded mitochondrial confocal micrographs. LAH, MVK, and MW
- analyzed and interpreted results. LAH, MW, and JJT wrote the manuscript.

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# **Declaration of Interests**

233 The authors declare no competing interests.

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235 Figures and legends



237 Figure 1. Optogenetic control of C. elegans gut bacterial gene expression. (a) Strain LH01. 238 (b) Microscopy and cytometry workflows. (c) Fluorescence microscopy images 0 and 8 h after 239 green light exposure in the step ON experiment. Scale bar: 10 µm. (d) Response dynamics in the 240 step ON (green) and step OFF (red) microscopy experiments. Black:  $\Delta PCB$  strain (step ON 241 experiment). Individual- (light lines) and multi-worm average (dark lines) data are shown. n = 7, 242 4, 6 worms for green, red, black data sets (measured over 2, 3, 1 days, respectively). Error bars: 243 SEM. (e-g) Flow cytometry histograms for response dynamics experiments. MEFL: molecules of 244 equivalent fluorescein.

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246

Figure 2. Optogenetic control of colanic acid biosynthesis. (a) Strain MVK29. (b) CA

secretion levels for MVK29 and control strains exposed to red and green light. JW1935-1 is the

249 E. coli rcsA background strain used in this study. N.D.: below assay limit of detection. (c)

250 Green light intensity versus CA secretion level for MVK29. Data points represent 3 biological

251 replicates collected on a single day. Dashed line: limit of detection. Error bars indicate standard

252 deviation of the three biological replicates.

253



255

Figure 3. Light-regulated CA secretion modulates C. elegans mitochondrial dynamics. (a) 256 257 Schematic of experiment for activating CA biosynthesis in situ. (b) Representative images of the 258 mitochondrial network of anterior intestinal cells immediately distal to the pharynx are scored as 259 fragmented, intermediate, or tubular, as previously(Han et al., 2017). Scale bars: 10 µm. (c) 260 Mitochondrial fragmentation profiles of un-paralyzed worms fed MVK29 while exposed to red 261 or green light for 6 h. (d) Fragmentation profiles for worms fed the indicated strain, then 262 paralyzed for 6 h while exposed to red or green light. The number of worms included in each 263 condition is indicated below each bar. The Chi-Squared Test of Homogeneity was used to 264 calculate *p*-values between conditions.

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Figure 4. Optogenetically-regulated CA biosynthesis extends worm lifespan. (a) When exposed to green light, worms grown on MVK29 live longer than those exposed to red light, and the magnitude of lifespan extension is proportional to green light intensity (p < 0.0001 green vs. red, log-rank test). (b-c) The lifespans of worms grown on the  $\Delta lon$  (b) or the  $\Delta rcsA$  (c) controls are not affected by light condition (p > 0.1 green vs. red, log-rank test).

273

#### 275 Methods

#### 276 <u>E. coli plasmids, strains, and media</u>

277 Plasmids used in this study are described in **Supplementary Table 1**. Genbank accession 278 numbers are given in Supplementary Table 2. All plasmids constructed in this study were 279 assembled via Golden Gate cloning(Engler et al., 2009). Primers were ordered from IDT 280 (Coralville, IA). Assembled plasmids were transformed into E. coli NEB10β (New England 281 Biolabs) for amplification and screening. All plasmid sequences were confirmed by Sanger 282 sequencing (Genewiz; S. Plainfield, NJ). To construct pLH401 and pLH405, pSR58.6(Schmidl et al., 2014) was modified by inserting an mCherry expression cassette composed of a 283 284 (J23114; http://parts.igem.org/Promoters/Catalog/Anderson), constitutive promoter RBS 285 (BBa\_B0034; http://parts.igem.org/Part:BBa\_B0034), mCherry, and a synthetic transcriptional 286 terminator (L3S1P52 (Chen et al., 2013)). To construct pLH405, pLH401 was further modified 287 by exchanging the superfolder GFP gene (sfgfp) for gfpmut3\*. pMVK201.2 was built by 288 modifying pSR58.6 to control expression of *rcsA*.

All *E. coli* strains are described in **Supplementary Table 3.**  $\Delta rcsA$  (JW1935-1) was obtained from the Coli Genetic Stock Center.  $\Delta lon$  (JW0429-1) was obtained from the Keio *E. coli* knockout library(Baba et al., 2006), a gift from the Herman lab. All *E. coli* strains were maintained in LB media supplemented with appropriate antibiotics (chloramphenicol 34 µg/mL, spectinomycin 100 µg/mL, kanamycin 100 µg/mL) in a shaking incubator at 37 °C and 250 rpm unless otherwise noted.

#### 296 <u>C. elegans strains and media</u>

297 All C. elegans strains (Supplementary Table 3) were provided by the Caenorhabditis 298 Genetics Center (University of Minnesota), which is funded by the NIH office of Research 299 Infrastructure Programs (P40 OD010440). Worms were grown at 20°C on 1.7% NGM-agar 300 plates in 60 mm Petri dishes inoculated with a lawn of E. coli (CGSC str. BW28357), as 301 described in the CGC WormBook (wormbook.org), unless otherwise specified. The common 302 strain E. coli OP50 was not used for worm feeding, as it produces CA during normal growth(Han 303 et al., 2017). M9 buffer for C. elegans (abbreviated M9Ce to distinguish from E. coli M9 media) 304 was composed of 3 g KH<sub>2</sub>PO<sub>4</sub>, 6 g Na<sub>2</sub>HPO<sub>4</sub>, 5 g NaCl, 1 mL 1 M MgSO<sub>4</sub>, H<sub>2</sub>O to 1 L, and 305 sterilized by autoclaving (wormbook.org).

306

### 307 Optogenetic control of CA production

308 3 mL starter cultures of appropriate E. coli strains were inoculated from a -80°C freezer and grown 12 h at 37 °C. These starters were diluted to  $OD_{600} = 1 \times 10^{-2}$  in M9 minimal media (1x 309 310 M9 salts, 0.4% w/v glucose, 0.2% w/v casamino acids, 2 mM MgSO<sub>4</sub>, 100 µM CaCl<sub>2</sub>) 311 supplemented with appropriate antibiotics. The M9/cell mixtures were then distributed into 3 mL 312 aliquots in 15 mL clear polystyrene culture tubes and grown at 37 °C in a shaking incubator at 313 250 rpm while illuminated with the appropriate light wavelength and intensity, using the Light 314 Tube Array (LTA) device (Gerhardt et al., 2016). After 22 h, cultures were removed and iced to 315 halt growth and the OD<sub>600</sub> was measured. Culture samples were collected for CA quantification.

# 317 <u>CA quantification</u>

318 We adapted a previous CA quantification protocol(DISCHE, 1947; DISCHE and 319 SHETTLES, 1948) that takes advantage of the fact that it is the only exopolysaccharide 320 produced in our E. coli strains that incorporates fucose. In particular, we quantified the amount 321 of fucose in cell-derived exopolysaccharides (EPS), and used that value as a proxy CA levels. 322 First, EPS was liberated from cells by boiling 2 mL of culture for 15 min. in a 15 mL conical 323 tube. The sample was then centrifuged in 1.5 mL Eppendorf tubes for 15 min. at 21,000 x g. 324 Then, 0.7 mL of supernatant was dialyzed against water for at least 12 h using Pur-A-Lyzer Midi 325 3500 dialysis mini-tubes (Sigma-Aldrich, PURD35100-1KT) to remove monomeric fucose from 326 the sample.

327 Fucose monomers were then liberated from the EPS polymers by hydrolyzing 0.2 mL of 328 dialyzed media with 0.9 mL of  $H_2SO_4$  solution (6:1 v/v acid:water). This mixture was boiled in a 329 15 mL conical for 20 min and then cooled to room temperature. The absorbance at 396 nm and 330 427 nm was measured. Next, 25 µL of 1 M L-cysteine HCl was added and mixed thoroughly by 331 pipetting. The absorbance at 396 nm and 427 nm was measured again. Simultaneously, 332 absorbance measurements of L-fucose standards pre- and post-L-cysteine addition were also 333 recorded. Absorbance change, given by D in the formula below, were used to compare the L-334 fucose standard samples to the dialyzed culture samples and estimate the L-fucose concentration 335 in the dialyzed product.

$$D = \left( \left( A_{post}^{396} - A_{pre}^{396} \right) - \left( A_{post}^{427} - A_{pre}^{427} \right) \right)$$

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336

#### 337 Preparation of NGM-agar plates for worm feeding

338 3 mL E. coli starter cultures were inoculated from -80°C freezer stocks and grown for 12 h at 37 °C. These starters were then diluted to  $OD_{600} = 1 \times 10^{-6}$  in M9 minimal media 339 340 supplemented with appropriate antibiotics. The M9/cell mixture was then distributed into 3mL 341 aliquots in 15 mL clear polystyrene culture tubes and grown at 37 °C in a shaking incubator at 342 250 rpm while illuminated with the appropriate light in the LTA. Once cultures reached  $OD_{600} =$ 343 0.1-0.4, tubes were iced for 10 min and subsequently concentrated to  $OD_{600} \sim 20$  by 344 centrifugation (4°C, 4000 rpm, 10 min) and resuspension in fresh M9 media. 400-600 µL of 345 dense bacterial culture was then applied to sterile NGM-agar plates and allowed to dry in a dark 346 room, or a room with green overhead safety lights if cultures were preconditioned in green light. 347 Plates were wrapped in foil and refrigerated at 4 °C for no more than 5 days until needed.

348

### 349 <u>Time-lapse microscopy</u>

350 To obtain age-synchronized worm cultures, axenized C. elegans (strain glo-1) eggs were 351 isolated and allowed to arrest in L1 by starvation in M9 buffer (distinct from M9 media: 3 g 352 KH<sub>2</sub>PO<sub>4</sub>, 6 g Na<sub>2</sub>HPO<sub>4</sub>, 5 g NaCl, 1 mL 1M MgSO<sub>4</sub>, and water to 1 L, sterilized by autoclaving 353 at 121°C for 20 min) at room temperature for 12-18 h. 10-100 larvae were transferred to a 354 previously prepared NGM-agar plate containing a lawn of the appropriate bacterial strain. The 355 plate was then placed in a 20°C incubator and illuminated with appropriate optogenetic light 356 provided by a single LED positioned 1cm above the Petri dish. Adult worms were transferred to 357 a fresh plate as necessary to maintain only a single generation.

358 Individual worms aged 1-3 days were removed from the dish and prepared for time-lapse 359 epifluorescence imaging. A 1.5% agar pad was prepared using M9 buffer as previously 360 described<sup>45</sup>, and punched into  $\frac{1}{2}$ " circles with a hollow punch. A 4  $\mu$ L droplet of 2 mM 361 levamisole was deposited on a pad and 5 adult worms were transferred from the NGM plate to 362 the droplet. An additional 4 µL of levamisole solution was added and the worms were gently 363 washed to remove external bacteria. Worms were then transferred to a fresh pad with a 4 µL 364 droplet of levamisole solution, which was allowed to dry, thereby co-localizing and aligning the 365 worms on the pad. The pad was then inverted and placed into a 13 mm disposable microscopy 366 petri dish with a #1.5 coverslip on the bottom (Cell E&G; Houston, TX). Another coverslip was 367 placed on the top of the pad in the dish to curtail evaporation.

368 The dish was then placed on the stage of a Nikon Eclipse Ti-E inverted epifluorescence 369 microscope (Nikon Instruments, Inc; Melville, NY). Complete paralysis was induced by 370 incubating the dish at room temperature ( $\sim 23 \circ C$ ) for 30 min. Meanwhile, worms were exposed 371 to appropriate preconditioning light supplied by a circular array of 8 LEDs (4 x 660 nm, 4 x 525 372 nm) mounted to the microscope condenser ring, about 2 cm above the Petri dish. Light was then 373 switched from the preconditioning to the experimental wavelength, and worms were imaged 374 periodically using 10x, 40x, and 60x objectives. For each time point, the LEDs were turned off 375 and images acquired in the brightfield (DIC) and fluorescent channels. Afterwards, the LEDs 376 were turned on again to maintain optogenetic control.

377

## 378 Epifluorescence image analysis

379 All epifluorescence images were analyzed using the Nikon Elements software package380 (Nikon Instruments, Inc; Melville, NY). The mCherry signal was used as a marker for the gut

381 lumen, and only cells in this region were included in the analysis. Image ROIs were created by 382 thresholding the sfGFP signal to identify the boundaries of cell populations. Out of focus regions 383 were eliminated from analysis. The average sfGFP pixel intensity inside the ROIs was calculated 384 and recorded for each time point.

385

### 386 <u>Flow cytometry</u>

387 1-3 day old *glo-1* worms were prepared for flow cytometry of the microbiome 388 constituents by washing, using a protocol adapted from previous work(Portal-Celhay et al., 389 2012). Groups of 5 worms were washed 2x in a 5 µL droplet of lytic solution: C. elegans M9 390 buffer containing 2 mM levamisole, 1% Triton X-100, and 100 mg/mL ampicillin. The worms 391 were then washed 2x in 5  $\mu$ L droplets of M9 buffer containing 2 mM levamisole only. Finally, 392 the worms were transferred to clear 0.5 mL Eppendorf tubes containing 50  $\mu$ L of M9 buffer + 2 393 mM levamisole, ensuring that 5 worms were deposited in the liquid contained in each tube. Each 394 tube was then exposed to light by placing it within one well of a 24-well plate (AWLS-303008, ArcticWhite LLC) atop a Light Plate Apparatus (LPA) containing green and red LEDs<sup>47</sup> for 8 h 395 396 at room temperature. In separate control experiments, we demonstrated that any stray bacteria 397 that may escape the worms over this period, or which were inadvertently added to the 50  $\mu$ L of 398 M9 buffer, are incapable of responding to optogenetic light (Supplementary Fig. 4).

At the conclusion of the experiment, tubes were removed from the plate and immediately chilled in an ice slurry for 10 min in the dark. Worms were homogenized using an anodized steel probe sterilized between samples via 70% ethanol treatment and flame before being cooled.

402 Next, we used our previous antibiotic-based fluorescent protein maturation 403 protocol(Olson et al., 2014) to allow unfolded proteins to mature while preventing the production 404 of new protein. In particular, 250  $\mu$ L PBS containing 500 mg/mL Rifampicin was added to the 405 50  $\mu$ L homogenized worm samples and transferred to cytometry tubes. These tubes were 406 incubated in a 37 °C water bath for precisely 1 h, then transferred back to an ice slurry.

407 These samples were measured on a BD FACScan flow cytometer. For gating, an 408 FSC/SSC polygon gate was first created using non-fluorescent bacteria grown *in vitro* at 37 °C 409 (Supplementary Fig. 2). Events outside this region were excluded as non-bacterial material. To 410 isolate the engineered gut bacteria, only events with a high mCherry signal (FL3 > 1200 a.u., 411 FL3 gain: 999) were included (Supplementary Fig. 2). Samples were measured until 20,000 412 events were recorded or the sample was exhausted.

413

### 414 <u>Flow cytometry data analysis</u>

415 All flow cytometry data (FCS format) were analyzed using FlowCal(Castillo-Hair et al., 416 2016) and Python 2.7. We wrote a standard cytometry analysis workflow that truncated the 417 initial and final 10 events to prevent cross-sample contamination, removed events from saturated 418 detector bins at the ends of the detection range, and added 2D density gate on SSC/FSC retaining 419 the densest 75% of events (Supplementary Fig. 2a). GFPmut3\* fluorescence units were 420 converted into standardized units of molecules of equivalent fluorescein (MEFL) using a 421 fluorescent bead standard (Rainbow calibration standard; cat. no. RCP-30-20A, Spherotech, Inc.) 422 as described previously(Castillo-Hair et al., 2016). Finally, to eliminate events associated with C. *elegans* autofluorescence (Supplementary Fig. 2b), any events in the region  $FL1 \le 1200$  MEFL 423 424 were discarded.

#### 426 <u>Mitochondrial Fragmentation Assays</u>

Synchronized L1 worms (strain *ges-1*) were applied to NGM agar plates containing bacterial strain BW25113 and allowed to develop until adulthood. This parental bacterial strain is used to allow all worms to develop at the same rate, avoiding any developmental/growth effects the experimental strains may exert on the worms. All experimental bacterial strains were preconditioned in red light with optogenetic light provided by a single LED positioned 1 cm above the Petri dish.

433 After 3-5 days (between days 1-3 of adulthood), worms allocated for the experiment were 434 transferred to experimental strains for approximately 60-90 minutes to thoroughly inoculate the 435 GI tract. In the case of the unparalyzed worms, red or green light was then applied for an 436 additional 6 hours. For the paralyzed worm experiments, 1.5% low-melt agar pads were prepared 437 as described above and placed on individual slides. About 15 adult worms were transferred from 438 the experimental strain Petri dish to an agar pad containing 10  $\mu$ L of C. elegans M9 buffer + 2 439 mM levamisole (M9Ce+Lev), where worms were gently washed before being transferred to a 440 fresh pad also containing 10 µL of M9Ce+Lev. The majority of M9Ce+Lev on the pad was 441 allowed to evaporate, which causes the worms to align longitudinally before a cover slip was 442 applied. Slides were then exposed to either red or green light by placing them under a single 443 LED positioned 1 cm above the Petri dish for 6 h. Afterward, the anterior intestinal cells were 444 imaged using confocal microscopy (Olympus Fluoview 3000) in the brightfield and GFP 445 channels.

#### 447 <u>Confocal Microscopy Image Analysis</u>

448 All confocal images for an experiment were manually cropped to display only the 449 anterior intestinal cells of a single worm (in the GFP channel). These cropped images were then 450 anonymized, randomized and the mitochondrial networks of each were blindly classified by two 451 researchers independently as either tubular, fragmented, or intermediate. Tubular samples were 452 marked by a high degree of network connectivity throughout. Fragmented samples were 453 composed almost exclusively of isolated clusters of fluorescence with high circularity. 454 Intermediate samples contained regions of both types. Scores were then de-randomized and 455 aggregated. For each experimental strain, the red and green light conditions were compared for 456 statistical significance using the chi-squared test of homogeneity.

457

#### 458 <u>Lifespan experiments</u>

459 3 mL starter cultures of  $\Delta lon$ ,  $\Delta rcsA$  or MVK29 were inoculated from -80°C freezer 460 stocks into LB supplemented with appropriate antibiotics and grown shaking for 12 h at 37 °C at 250 rpm. These cultures were diluted to  $OD_{600} = 1 \times 10^{-6}$  in 27 mL M9 media supplemented with 461 462 appropriate antibiotics. 1.5 mL of each M9/cell mixture was added to each of 18 wells on three 463 24-well plates and grown in 3 LPA devices under the appropriate light conditions at 37 °C and 464 250 rpm. Once cultures reached  $OD_{600} = 0.1-0.4$ , all tubes were iced for 10 min and subsequently 465 concentrated 10 times by centrifugation (4°C, 4000 rpm, 10 min). Approximately 50 µL of this 466 dense bacterial culture was then applied to sterile NGM-agar plates with no antibiotics and 467 allowed to dry in a dark room. The plates were then illuminated with the appropriate light 468 wavelength and intensity for 16 h at room temperature, and immediately used for lifespan assays.

During the longitudinal lifespan assay, exposure to white light is limited to the minimal 469 470 level. To reach this goal, the sqt-3(e2117) temperature sensitive mutant (Supplementary Table 471 3) was used to perform longitudinal analyses at 25 °C, which avoids time-consuming animal 472 transfer without interrupting normal reproduction. sqt-3(e2117) is a collagen mutant of C. 473 elegans that reproduces normally but is embryonically lethal at 26 °C, and has been used 474 previously in longitudinal studies(Han et al., 2017; Wang et al., 2014). Worms were age-475 synchronized by bleach-based egg isolation followed by starvation in M9 buffer at the L1 stage 476 for 36 hours. Synchronized L1 worms were grown on BW25113 E. coli at 15 °C until the L4 477 stage, when worms were transferred to 24-well plates (~15 worms/well) with  $\Delta lon$ ,  $\Delta rcsA$  or 478 MVK29 (Supplementary Table 3). The plates were placed in LPA. The LPA LEDs were programmed to illuminate wells with constant red (10 µmol/m<sup>2</sup>/s), low-intensity green (0.25 479 480  $\mu$ mol/m<sup>2</sup>/s), or high-intensity green light (10  $\mu$ mol/m<sup>2</sup>/s). The apparatus was then transferred to a 481 26 °C incubator. The number of living worms remaining in each well was counted every day. 482 Death was indicated by total cessation of movement in response to gentle mechanical 483 stimulation. Statistical analyses were performed with SPSS (IBM Software) using Kaplan-Meier 484 survival analysis and the log-rank test (Supplementary Table 4).

485

### 486 Supplemental Information Titles and Legends

Supplementary Figure 1. *In vitro* characterization of GFP reporter strains used in this study. All optogenetic strains carry our previously published CcaSR v2.0 system, which is encoded on plasmids (a) pSR43.6 and (b) pSR58.6. (c) pLH401 (used for microscopy) and (d) pLH405 (used for cytometry) genetic device schematics. We replaced *sfgfp* with *gfpmut3*\* in pLH405 as we hypothesized that the latter may be less stable and thus result in faster response 492 dynamics. However, we observed no difference in dynamics. (e) Batch culture light responses of 493 CcaSR v2.0 and all GFP reporter strains used in this study. GFP fluorescence was measured by 494 flow cytometry. The dynamic range (ratio of GFP fluorescence in green versus red light) is 495 shown above each data set. We note that CcaSR v2.0 exhibits 77-fold dynamic range in the 496 reference strain BW29655 ( $\Delta envZ$ ,  $\Delta ompR$ ), which is similar to our previous measurement of 497 this strain at 120-fold (Schmidl et al., 2014). The calculated fold-change is sensitive to 498 fluctuations in the measured *E. coli* autofluorescence and red-light expression level, which likely 499 explains the slight discrepancy. CcaSR v2.0 dynamic range increases slightly to 84.2-fold in 500  $\Delta rcsA$  (JW1935-1), which is used throughout this work, due to higher sfGFP expression in green 501 light. The mCherry cassette in pLH401 decreases dynamic range to 38.3-fold due to higher leaky 502 sfGFP expression in red light. In worms, the fold-change in response to green light decreases 503 further to 5.52  $\pm$  2.4-fold (Fig. 1d). The use of *gfpmut3*\* in pLH405 further decreases dynamic 504 range to 13.5  $\pm$  0.0-fold for reasons that are not clear, while in worms the fold-change is 8.63  $\pm$ 505 3.6-fold (Fig. 1e). Data represent the mean of three independent, autofluorescence-subtracted, 506 biological replicates acquired on 3 separate days. Error bars: standard deviation.

507

**Supplementary Figure 2. Flow cytometry gating.** (a) Strain LH05 (**Supplementary Table 3**), which expresses only mCherry, was fed to worms, isolated, and analyzed by flow cytometry to quantify *E. coli* autofluorescence through the FL1 (GFP) channel. To eliminate events corresponding to cytometer noise and autofluorescence of homogenized worm samples, three gates were then applied: (1) a density gate for the most homogeneous 75% of samples in forward scatter (FSC) vs side-scatter (SSC) (the area within the bold black line in the plots in Column 1), (2) a threshold gate on the FL1 (GFP) channel (marked by a red dashed line; **Methods**), and (3) a threshold gate for events exhibiting high FL3 (mCherry, red dashed line; **Methods**). (b) Applying these gates to samples from the step ON experiment (**Fig. 1e**) reveals robust isolation of bacteria with a FSC/SSC profile consistent with our previous *in vitro* experiments<sup>24,47</sup> and an increase in expression of GFPmut3\* in response to green light. Ungated data are shown for comparison to gated data in all plots.

520

521 Supplementary Figure 3. Gene expression dynamics from flow cytometry experiments. (a) 522 LH03 and (b) LH04 (ΔPCB) step ON results. (c) LH03 and (d) LH04 step OFF results. Each 523 individual trajectory (faint lines) corresponds to a single biological replicate. A biological 524 replicate comprises the homogenized contents of five worms collected on a single day. Each 525 biological replicate on a given plot was run on a different day. Population medians taken across 526 all trajectories are shown in bold (**Methods**). Data are composed of six and eight trajectories in 527 the step ON and step OFF experiments, respectively. Error bars: SEM.

528

529 Supplementary Figure 4. Bacteria on the exterior of worms do not contribute to the 530 measured light response in flow cytometry experiments. Worms were suspended in clear 531 tubes containing M9Ce+Lev media and levamisole in the flow cytometry experiments. To 532 demonstrate that bacterial cells outside the worm gut (e.g. on the exterior of the worm) do not 533 respond to light in these conditions, samples of pre-conditioned bacteria from the NGM plates 534 were suspended in the M9Ce+Lev or E. coli M9 media supplemented with casamino acids and 535 glucose (M9Ec). These samples were then exposed to either green or red light for 8 h, then 536 measured via flow cytometry after the fluorophore maturation protocol (Methods). As expected,

- 537 the bacterial populations are only responsive to light when grown in M9Ec and when the PCB
- 538 operon is intact (strain JW1935-1/pLH405/pSR43.6, aka LH03). We conclude that the responses
- shown in data in Fig. 1e-f and Supplementary Fig. 3 are not due to bacteria outside the worm,
- 540 nor to bacteria that escape the worm over the course of the experiment, as such cells do not
- respond to light in the experimental buffer (M9Ce). Data represent 7, 6, 3, and 5 replicates (left);
- and 8, 3, 8, and 3 replicates (right) over 8 and 10 days, respectively; error bars: SEM. Individual
- 543 data points for each condition are overlaid as white markers.
- 544 Supplementary Table 1. Plasmids used in this study
- 545 Supplementary Table 2. Genbank Accession Numbers
- 546 Supplementary Table 3. Bacterial and worm strains used in this study
- 547 Supplementary Table 4. Statistical analysis of worm lifespan experiments
- 548
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