1	An Escherichia coli nitrogen starvation response					
2	is important for mutualistic coexistence with Rhodopseudomonas palustris					
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19 Abstract

20 Microbial mutualistic cross-feeding interactions are ubiquitous and can drive important community 21 functions. Engaging in cross-feeding undoubtedly affects the physiology and metabolism of individual 22 species involved. However, the nature in which an individual's physiology is influenced by cross-feeding 23 and the importance of those physiological changes for the mutualism have received little attention. We 24 previously developed a genetically tractable coculture to study bacterial mutualisms. The coculture 25 consists of fermentative Escherichia coli and phototrophic Rhodopseudomonas palustris. In this 26 coculture, E. coli anaerobically ferments sugars into excreted organic acids as a carbon source for R. 27 *palustris*. In return, a genetically-engineered *R. palustris* constitutively converts N_2 into NH_4^+ , providing 28 E. coli with essential nitrogen. Using RNA-seq and proteomics, we identified transcript and protein levels 29 that differ in each partner when grown in coculture versus monoculture. When in coculture with R. 30 palustris, E. coli gene-expression changes resembled a nitrogen starvation response under the control of 31 the transcriptional regulator NtrC. By genetically disrupting E. coli NtrC, we determined that a nitrogen 32 starvation response is important for a stable coexistence, especially at low *R. palustris* NH₄⁺ excretion 33 levels. Destabilization of the nitrogen starvation regulatory network resulted in variable growth trends and 34 in some cases, extinction. Our results highlight that alternative physiological states can be important for 35 survival within cooperative cross-feeding relationships.

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37 Importance

Mutualistic cross-feeding between microbes within multispecies communities is widespread. Studying how mutualistic interactions influence the physiology of each species involved is important for understanding how mutualisms function and persist in both natural and applied settings. Using a bacterial mutualism consisting of *Rhodopseudomonas palustris* and *Escherichia coli* growing cooperatively through bidirectional nutrient exchange, we determined that an *E. coli* nitrogen starvation response is important for maintaining a stable coexistence. The lack of an *E. coli* nitrogen starvation response ultimately destabilized the mutualism and, in some cases, led to community collapse after serial transfers. Our findings thus inform on the potential necessity of an alternative physiological state for mutualistic
coexistence with another species compared to the physiology of species grown in isolation.

47

48 Introduction

49 Within diverse microbial communities, species engage in nutrient cross-feeding with reciprocating 50 partners as a survival strategy (1). In cases where species are not obligate mutualists, transitioning from a 51 free-living lifestyle to one based on cross-feeding can change the physiological state of the cells involved, 52 the extent to which depends on the nature of the cross-feeding relationship. For example, cross-feeding 53 can promote physiological changes that increase virulence (2, 3) or drastically alter cellular metabolism 54 (4), in some cases allowing for lifestyles that are only possible during mutualistic growth with a partner 55 (4–7). Aside from these examples, relatively little is known about how cell physiology is influenced by 56 mutualistic cross-feeding, despite the prevalence of cross-feeding in microbial communities.

57 Synthetic communities, or cocultures, are ideally suited for studying the physiological responses 58 to cooperative cross-feeding given their tractability (8, 9). We previously developed a bacterial coculture 59 that consists of fermentative Escherichia coli and the N₂-fixing photoheterotroph Rhodopseudomonas 60 palustris (Fig. 1) (10). In this coculture, E. coli anaerobically ferments glucose into organic acids, 61 providing R. palustris with essential carbon. In return, a genetically engineered R. palustris strain (Nx) 62 constitutively fixes N_2 gas, resulting in NH_4^+ excretion that provides *E. coli* with essential nitrogen. The 63 result is an obligate mutualism that maintains a stable coexistence and reproducible growth trends (10) as 64 long as bidirectional nutrient cross-feeding levels are maintained within a defined range (11, 12).

Here we determined how nutrient cross-feeding between *E. coli* and *R. palustris* Nx alters the physiological state of each partner population. Using RNA-seq and proteomic analyses, we identified genes in both species that were differentially expressed in coculture compared to monoculture, with *E. coli* exhibiting more overall changes in gene expression than *R. palustris* Nx. Specifically, *E. coli* geneexpression patterns resembled that of nitrogen-deprived cells, as many upregulated genes were within the nitrogen-starvation response regulon, controlled by the master transcriptional regulator NtrC. Genetic disruption of *E. coli ntrC* resulted in variable growth trends at low *R. palustris* NH_4^+ excretion levels and prevented long-term mutualistic coexistence with *R. palustris* across serial transfers. Our results highlight the fact that cross-feeding relationships can stimulate alternative physiological states for at least one of the partners involved and that adjusting cell physiology to these alternative states can be critical for maintaining coexistence.

76

77 **Results**

78 Engaging in an obligate mutualism alters the physiology of cooperating partners. In our coculture, E. 79 coli and R. palustris Nx carry out complementary anaerobic metabolic processes whose products serve as 80 essential nutrients for the respective partner. Specifically, E. coli ferments glucose into acetate, lactate, 81 and succinate, which serve as carbon sources for R. palustris Nx, while other fermentation products such 82 as formate and ethanol accumulate; in return R. palustris Nx fixes N₂ and excretes NH_4^+ as the nitrogen 83 source for E. coli (Fig. 1). We previously demonstrated that our coculture supports a stable coexistence 84 and exhibits reproducible growth and metabolic trends when started from a wide range of starting species 85 ratios, including single colonies (10). However, we hypothesized that coculture conditions would affect 86 the physiology of each species, particularly E. coli, based on the following observations. First, as growth 87 is coupled in our coculture, E. coli is forced to grow 4.6-times slower in coculture with R. palustris Nx than it does in monoculture with abundant NH_4^+ due to slow NH_4^+ cross-feeding from *R. palustris* Nx 88 89 (10). In contrast, R. palustris Nx grows at a rate in coculture that is comparable to that in monoculture 90 (12), consuming a mixed pool of excreted organic acids from E. coli. Second, coculturing pulls E. coli 91 fermentation forward due to removal of inhibitory end products. For example, we observed higher yields 92 of formate, an E. coli fermentation product that R. palustris does not consume, in cocultures compared to 93 E. coli monocultures (10).

To determine changes in gene-expression patterns imposed by coculturing, we performed RNAseq and comparative proteomic analyses (13) on exponential phase cocultures and monocultures of *E. coli* and *R. palustris* Nx. To make direct comparisons, all cultures were grown in the same basal anaerobic

97 minimal medium, and monocultures were supplemented with the required carbon or nitrogen sources to 98 permit growth for each species. Cocultures and E. coli monocultures were provided glucose as a sole 99 carbon source, whereas a mixture of organic acids and bicarbonate was provided to R. palustris Nx 100 monocultures, as R. palustris does not consume glucose. For a nitrogen source, all cultures were grown 101 under a N_2 headspace, and E. coli monocultures were further supplemented with NH₄Cl, as E. coli is 102 incapable of using N_2 . We identified several differentially expressed genes between monoculture and 103 coculture conditions in both species with more differences observed in E. coli compared to R. palustris 104 Nx, in agreement with our initial hypothesis (Fig. 2). For E. coli, out of 4377 ORFs, 55 were upregulated 105 and 68 were downregulated (Table 1) (log2 value cutoff=2). Out of 4836 ORFs in R. palustris Nx, 14 106 were upregulated and 20 were downregulated (Table 1) (log2 value cutoff=2). We also considered that 107 due to lower E. coli abundance in coculture, the apparently larger E. coli gene response may be partly due 108 to decreased resolution and thus increased error variance. Reassuringly, many of the genes identified as 109 being differentially expressed by RNA-seq were in agreement with the proteomic results (Table 2). Both 110 RNA-seq and proteomic analyses identified the E. coli ammonium transporter AmtB as an important, 111 upregulated gene in coculture, corroborating our previous findings that E. coli AmtB activity is important 112 for stable coexistence with R. palustris (12). Many E. coli genes involved in amino acid and purine 113 biosynthesis were downregulated in coculture (Table 1, Table 2), consistent with the lower observed 114 growth rate. Additionally, many E. coli flagellar and chemotaxis proteins were downregulated in 115 coculture (Table 1, Table 2), perhaps suggesting that motility is not important for coculture growth. 116 Alternatively, lower flagellar and chemotaxis transcript levels could be part of a general stress response 117 (14), perhaps associated with nitrogen limitation in cocultures. Whereas many of the differentially 118 expressed E. coli genes have been characterized in the literature, the R. palustris genes showing the 119 were uncharacterized genes encoding largest differential expression upregulated putative 120 alcohol/aldehyde dehydrogenases and a downregulated putative TonB-dependent receptor/siderophore 121 (Table 1, Table 2). Together, these datasets provide insight on how engaging in obligate cross-feeding 122 changes the lifestyle of each partner.

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124 An E. coli nitrogen starvation response is important for mutualistic growth with R. palustris. We 125 chose to further examine differential gene expression patterns in E. coli as its growth rate and 126 fermentation profile are drastically affected by coculturing, whereas the R. palustris Nx growth rate is 127 similar to that in monoculture. We identified several E. coli genes and proteins that were upregulated in 128 coculture with R. palustris Nx compared to monoculture growth (Table 1, Table 2). We hypothesized that 129 the deletion of highly upregulated *E. coli* genes would negatively affect its growth in coculture. We made 130 deletions in E. coli genes that were identified in both RNA-seq and proteome datasets as well as the 131 highest upregulated E. coli transcript (rutA). We did not examine the effect of deleting amtB in this case 132 as we previously determined it to be important for coculture growth (12). These selected E. coli genes 133 were all involved in metabolism of alternative nitrogen sources such as D-ala-D-ala dipeptides (ddpX, 134 ddpA) (15), pyrimidines (*rutA*) (16), amino acids (*argT*) (17), and polyamines (*patA*, *potF*) (18). In 135 monocultures with 15mM NH₄Cl, there were negligible differences in growth or fermentation profiles 136 between WT E. coli and any of the single deletion mutants (Fig. S1). These results are consistent with 137 findings by others, as these genes are only important when scavenging alternative nitrogen sources that 138 are not present in our defined medium. We next tested these E. coli mutants in coculture with R. palustris 139 Nx to determine if these genes were important when NH_4^+ is slowly cross-fed from R. palustris Nx. All 140 cocultures using the E. coli mutants paired with R. palustris Nx exhibited similar growth and population 141 trends to cocultures with WT E. coli (Fig. 3). Additionally, there were no significant differences in the 142 growth rates, growth yields, or product yields from cocultures containing the E. coli mutants (Fig. S2). 143 These data suggest that none of these highly expressed E. coli genes are solely important for coculture 144 growth. While it is possible that synergistic expression of these genes is important for E. coli's lifestyle in 145 coculture, the actual nitrogen sources accessed by expression of these genes are absent in the defined 146 medium. Thus, unless E. coli gains access to alternative nitrogen sources that we are unaware of in 147 coculture with *R. palustris* Nx, synergistic expression of these genes likely provides little to no benefit.

148 Even though individual deletions of the *E. coli* genes showing high expression in coculture had 149 no effect on coculture trends, we noted that they were all involved in nitrogen scavenging and fell within 150 the regulon of the transcription factor, NtrC, which controls the nitrogen starvation response (19). During 151 nitrogen limitation, the sensor kinase NtrB phosphorylates the response regulator NtrC (19). 152 Phosphorylated NtrC then binds to DNA and activates expression of ~45 genes (20), including those we 153 tested genetically above and *amtB*, which we previously determined to be important for coculture growth 154 (12). To examine the importance of the E. coli nitrogen starvation response in coculture, we deleted ntrC. 155 We first checked for any general defects of the resulting $\Delta NtrC$ mutant in monoculture with 15 mM 156 NH₄Cl and found that it exhibited similar growth and metabolic trends to WT E. coli (Fig. S3). We then 157 paired E. coli Δ NtrC with R. palustris Nx in coculture. Compared to cocultures using WT E. coli, 158 cocultures with *E. coli* Δ NtrC exhibited slower growth rates, longer lag periods (Fig. 4A), and lower final 159 E. coli cell densities (Fig. 4D). The long lag phase was less prominent in cocultures inoculated from 160 single colonies (Fig. S4A) compared to cocultures inoculated with a 1% dilution of stationary cocultures 161 (Fig. 4A). This result suggests that starting E. coli Δ NtrC cocultures from single colonies stimulated early 162 growth, perhaps by increasing the E. coli frequency to be similar to that of R. palustris when started with 163 colonies of similar sizes rather than a dilution of stationary cocultures wherein the E. coli frequency was 164 low (~0.1%; Fig. 4D). A higher initial *E. coli* frequency might help *E. coli* acquire excreted NH_4^+ before 165 it is taken back up by *R. palustris* cells and thereby promote reciprocal cross-feeding, similar to what we 166 observed previously in cocultures with *E. coli* Δ AmtB mutants that were defective for NH₄⁺ uptake (12).

167 The overall coculture metabolism was also altered when *E. coli* Δ NtrC was paired with *R.* 168 *palustris* Nx. In cocultures pairing WT *E. coli* with *R. palustris* Nx, glucose is typically fully consumed 169 within 5 days coinciding with the accumulation of formate and ethanol (10). Cocultures pairing *E. coli* 170 Δ NtrC with *R. palustris* Nx differed in this regard, leaving ~40% of the glucose unconsumed after 10 171 days and exhibiting little to no formate and ethanol accumulation (Fig. S4B). Even despite the lower 172 glucose consumption, the final *R. palustris* cell density of cocultures pairing *R. palustris* Nx with *E. coli* 173 Δ NtrC was similar to those with WT *E. coli*. This unexpectedly high cell density could be explained by 174 consumption of formate and ethanol by *R. palustris* Nx, though we have never observed consumption of 175 formate by *R. palustris* Nx in monoculture. Alternatively, a lack of formate and/or ethanol production by 176 *E. coli* could explain the high cell density if the fermentation profile were shifted towards organic acids 177 that *R. palustris* normally consumes, namely acetate, lactate and succinate. Together, these data indicate 178 that misregulation of the nitrogen starvation response affected coculture growth and metabolism.

179 As noted above, the low E. coli Δ NtrC population and decreased coculture growth rate when 180 paired with *R. palustris* Nx resembled trends from cocultures that contained *E. coli* Δ AmtB mutants (12). 181 We previously found that the *E. coli* NH_4^+ transporter, AmtB, was required for coexistence with *R*. 182 palustris Nx across serial transfers as the transporter gives E. coli a competitive advantage in acquiring 183 the transiently available NH_4^+ before it can be reclaimed by the *R. palustris* population (12). To determine 184 if E. coli Δ NtrC was capable of maintaining a stable coexistence in coculture, we inoculated cocultures of 185 E. coli Δ NtrC paired with R. palustris Nx at equivalent CFUs and performed serial transfers every 10 186 days. While average final *E. coli* frequencies were consistently between 0.6 - 2.8 % (Fig. 5A), the values 187 became variable over serial transfers, as did coculture growth rates, lag periods, and net changes in both 188 E. coli and R. palustris cell densities (Fig. 5). This variability was due to 2 of the 4 lineages exhibiting 189 improved coculture growth over successive transfers (Fig. 5B, C), perhaps due to the emergence of 190 compensatory mutations, while the other two lineages showed declining growth trends (Fig. 5D, E). 191 Indeed, by transfers 5 and 6 there was little to no coculture growth in the slower-growing lineages (Fig 192 4D, E). The heterogeneity in growth trends through serial transfers of cocultures with E. coli Δ NtrC is in 193 stark contrast to the stability of cocultures with WT E. coli, which we have serially transferred over 100 194 times with no extinction events (McKinlay, unpublished data). The nitrogen starvation response thus 195 appears to be important for long-term survival of the mutualism.

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197 Increased NH_4^+ cross-feeding levels can compensate for the absence of a nitrogen starvation 198 response. The NtrC regulon is critical during periods of nitrogen starvation, activating a wide variety of 199 genes that are important for scavenging diverse nitrogen sources (20). We hypothesized that higher *R*. 200 *palustris* NH₄⁺ cross-feeding levels could mitigate the poor growth of E. coli Δ NtrC in coculture by 201 making the nitrogen starvation response less important for survival. Previously, we engineered an R. 202 *palustris* Nx strain that excretes 3-times more NH_4^+ by deleting *R. palustris* NH_4^+ transporters encoded by 203 *amtB1* and *amtB2* (Nx Δ AmtB) (10). N₂-fixing bacteria use AmtB to reacquire NH₄⁺ that leaks outside the 204 cell, and \triangle AmtB mutants thus accumulate NH₄⁺ into the supernatant (10, 12, 21). In agreement with our 205 hypothesis, cocultures with R. palustris Nx Δ AmtB exhibited similar growth trends regardless of the E. 206 *coli* strain used (Fig. 4B, D). As *R. palustris* Nx Δ AmtB excretes more NH₄⁺ than *R. palustris* Nx, it was 207 previously shown to result in faster WT E. coli growth and subsequent fermentation rates in coculture, 208 ultimately leading to the accumulation of consumable organic acids (Fig. S4B) and acidification of the 209 medium, inhibiting R. palustris growth (10). Cocultures pairing R. palustris Nx Δ AmtB and E. coli Δ NtrC 210 similarly exhibited growth (Fig. 4B,D), and fermentation profile trends (Fig. S4B) that were 211 indistinguishable from cocultures pairing R. palustris NxAAmtB with WT E. coli. These similar trends 212 indicate that high *R. palustris* NH_4^+ excretion can eliminate the trends observed when the *E. coli* nitrogen 213 starvation response is compromised due to a Δ NtrC mutation.

214 One possibility for why high NH_4^+ cross-feeding levels eliminate the need for *E. coli ntrC* is that 215 the free NH_4^+ levels might be sufficiently high enough to prevent activation of the *E. coli* NtrC regulon. 216 However, comparative RNA-seq and proteomic analyses revealed that the same *E. coli* genes within the 217 NtrC regulon that were highly upregulated in cocultures pairing WT *E. coli* with *R. palustris* Nx were 218 also upregulated in cocultures with *R. palustris* Nx Δ AmtB (Table 1, Table 2). Thus, even though the *E. 219 coli* nitrogen-starvation response is activated when cocultured with *R. palustris* Nx Δ AmtB, this response 220 is likely dispensable if there is sufficiently high NH_4^+ cross-feeding.

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E. coli NtrC is required for adequate AmtB expression to access cross-fed NH_4^+ in coculture. While a high level of *R. palustris* NH_4^+ excretion can compensate for an improper *E. coli* nitrogen-starvation response, less NH_4^+ excretion could potentially exaggerate problems emerging from the absence of NtrC. We previously constructed an *R. palustris* Δ AmtB strain that excreted $1/3^{rd}$ of the NH_4^+ than *R. palustris* 226 Nx in monoculture and which could not coexist in coculture with E. coli Δ AmtB (12). The reason for this 227 lack of coexistence was due to R. palustris Δ AmtB outcompeting E. coli Δ AmtB for the lower level of 228 transiently available NH_4^+ , thus limiting E. coli growth and thereby the reciprocal supply of fermentation 229 products to R. palustris (12). Expression of E. coli amtB is thus important in coculture in order to 230 maintain coexistence. Indeed, RNA-seq and proteomic analyses revealed that E. coli AmtB transcript and 231 protein levels were upregulated in all cocultures pairing WT E. coli with any of the three R. palustris 232 strains (Nx, Nx Δ AmtB, Δ AmtB) (Table 1, Table 2). We thus wondered whether *E. coli* Δ NtrC would 233 coexist with the low NH₄⁺-excreting strain R. palustris Δ AmtB in coculture, as E. coli amtB expression is 234 transcriptionally activated by NtrC. Consistent with our previous findings, R. palustris Δ AmtB supported 235 a high relative WT E. coli population in coculture (Fig. 4D) (12). When cocultured with WT E. coli, R. 236 *palustris* Δ AmtB responds to NH₄⁺ loss to *E. coli* by upregulating nitrogenase activity since it has a wildtype copy of NifA (12). As a result, R. palustris Δ AmtB cross-feeds enough NH₄⁺ to stimulate a high WT 237 238 E. coli frequency and subsequent accumulation of consumable organic acids, similar to cocultures with R. 239 palustris Nx Δ AmtB (Fig 3D, Fig. S4B) (12). In contrast, when we paired E. coli Δ NtrC with R. palustris 240 Δ AmtB, little to no coculture growth was observed (Fig. 4C), similar to previous observations in 241 cocultures pairing E. coli \triangle AmtB with R. palustris \triangle AmtB (12). Cocultures inoculated with single 242 colonies of each species in this pairing grew to low cell densities (Fig. S4A), and cocultures inoculated 243 from these cocultures resulted in little to no growth, even after prolonged incubation (Fig. 4C).

As AmtB is under the control of NtrC (20), we hypothesized that cocultures pairing *E. coli* Δ NtrC with *R. palustris* Δ AmtB resulted in insufficient *E. coli amtB* expression, leading to a decreased ability to capture NH₄⁺, which *R. palustris* will reaquire if given the chance (12). We thus predicted that increased expression of *amtB* in *E. coli* Δ NtrC would result in increased net growth of both species, as *E. coli* Δ NtrC would be more competitive for essential NH₄⁺ and be able to grow and produce more organic acids for *R. palustris* Δ AmtB. To test this prediction, we obtained a plasmid harboring an IPTG-inducible copy of *amtB* (pamtB) for use in *E. coli* Δ NtrC. AmtB is typically tightly regulated and only expressed when 251 NH_4^+ concentrations are below 20 μ M, as cells acquire sufficient NH_4^+ through passive diffusion of NH_3 252 across the membrane at higher concentrations (22). Additionally, excessive NH_4^+ uptake through AmtB 253 transporters that exceeds the rate of assimilation can result in a futile cycle, as excess NH_3 inevitably 254 diffuses outside the cell (19). We first tested the effect of pamtB in WT E. coli monocultures with 15 mM 255 NH₄Cl. Induction with 1 mM IPTG prevented growth whereas 0.1 mM IPTG permitted growth albeit at a 256 decreased growth rate (Fig. S5). We thus decided to use 0.1 mM IPTG to induce *amtB* expression in all 257 cocultures described below. In cocultures pairing E. coli Δ NtrC pamtB with R. palustris Δ AmtB, more 258 growth was observed than in cocultures with E. coli Δ NtrC harboring an empty vector (pEV) (Fig. 6A). In 259 cocultures with *E. coli* Δ NtrC pEV, the *R. palustris* Δ AmtB cell density increased whereas the *E. coli* cell 260 density did not (Fig. 6B). The *R. palustris* growth was likely due to growth-independent cross-feeding of 261 fermentation products from E. coli maintenance metabolism, a phenomenon we described previously 262 (11). In contrast, cell densities of both species increased in cocultures pairing R. palustris Δ AmtB with E. 263 *coli* Δ NtrC pamtB (Fig. 6C), in agreement with our hypothesis that poor E. coli amtB expression 264 contributed to the lack of growth in this coculture pairing. While E. coli amtB expression in this coculture 265 pairing was sufficient to restore growth of both species, there are likely other genes within the NtrC 266 regulon that contribute to E. coli growth in coculture. For example, the E. coli NtrC-regulated 267 serine/threonine kinase yeaG has been shown to play a role in survival during nitrogen starvation by 268 promoting metabolic heterogeneity (23). Indeed, E. coli yeaG and its associated protein of unknown 269 function yeaH are both highly upregulated in coculture (Table 1). Thus, while we cannot rule out that 270 other genes within the E. coli ntrC regulon are not important for coculture growth, the necessity of NtrC 271 to upregulate *amtB* is clearly important.

272

273 Discussion

In this study, we found that reciprocal nutrient cross-feeding between *E. coli* and *R. palustris* resulted in significant changes in gene expression in both species compared to monocultures. Based on the RNA-seq and proteomic analyses, we determined that *E. coli* alters its physiology to adopt a nitrogen-starved state 277 in response to low NH_4^+ cross-feeding levels from *R. palustris*. We subsequently determined that this 278 nitrogen-starved state is important for coexistence as genetic elimination of the master transcriptional 279 regulator, NtrC, resulted in variable population outcomes. Mutualistic nutrient cross-feeding has also been 280 shown to change the lifestyle of interacting partners in other systems. In natural communities, nutrient 281 cross-feeding can alter gene-expression patterns to adapt each species to a syntrophic lifestyle (24–27). In 282 some cases, the lifestyles exhibited within a mutualism might not even be possible during growth in 283 isolation. For example, in synthetic communities that pair the sulfate-reducer Desulfovibiro vulgaris with 284 the methanogen Methanococcus maripaludis, the methanogen consumes H₂, which maintains low partial 285 pressures that permit the sulfate reducer to adopt a fermentative lifestyle that would otherwise be 286 thermodynamically infeasible (5). Similarly, in an experimental Geobacter coculture, direct electron 287 transfer from Geobacter metallireducens to Geobacter sulfurreducens makes ethanol fermentation by G. 288 *metallireducens* thermodynamically possible (7).

289 Similar to our mutualistic system, the mutualism between D. vulgaris and M. maripaludis 290 represents a facultative mutualism, at least in the short term prior to evolutionary erosion of independent 291 lifestyles (28). For mutualistic relationships to persist between partners that are conditionally capable of a 292 free-living lifestyle, the relationship must exhibit resilience, or the ability to recover its function after a 293 disturbance (29). One important resilience factor is the activation of regulatory networks that allow for 294 microbes to quickly respond to environmental perturbations. Whereas flexible gene expression is useful 295 for an individual microbe's survival, excessive flexibility can sometimes lead to community collapse 296 between mutualists in a fluctuating environment (30, 31). In the coculture of D. vulgaris and M. 297 maripaludis, alternating between coculture and monoculture conditions, which require different metabolic 298 lifestyles, resulted in community collapse (30, 31). Surprisingly, community collapse could be avoided by 299 mutations that disrupted the D. vulgaris regulatory response needed to adapt cells for optimal growth rates 300 in monoculture (30). Disruption of this regulatory response resulted in a heterogeneous D. vulgaris 301 population, ensuring that a subpopulation would be primed for immediate mutualistic growth upon 302 transition between growth conditions (31). In our system, the E. coli nitrogen starvation regulatory

303 network was specifically activated by coculturing with *R. palustris* and was important for coculture 304 stability. It is currently unclear if transitioning *E. coli* between monoculture and coculture conditions 305 would result in similar community collapse or whether the NtrC-regulated network would adjust rapidly 306 enough to meet the demands of each condition.

307 Nutrient starvation and other stress responses are widely conserved in diverse microbes and are 308 primarily regarded as necessary for an individual's survival in nutrient-limited environments (32-35). 309 Many microbial communities are composed of primarily slow-growing or even non-growing 310 subpopulations (36–38). However, lack of microbial growth in these communities does not imply 311 cessation of cross-feeding, as bacteria often carry out growth-independent maintenance processes at slow 312 rates (39), and such activities can be coupled to cross-feeding (11). Our findings suggest that nutrient 313 starvation and perhaps other stress responses can help stabilize microbial cross-feeding interactions, 314 especially at low nutrient cross-feeding levels. The extent to which specific starvation or stress responses 315 are active in diverse mutualistic relationships remains unclear, yet likely depends on the environmental 316 context. Together our results highlight the important role that alternate physiological states, including 317 stress responses, can play in establishing and maintaining mutualistic cross-feeding relationships.

318

319 Materials and Methods

320 Strains and growth conditions. Strains, plasmids, and primers are listed in Table S1. All R. palustris 321 strains contained $\Delta uppE$ and $\Delta hupS$ mutations to facilitate accurate colony forming unit (CFU) 322 measurements by preventing cell aggregation (40) and to prevent H_2 uptake, respectively. E. coli was 323 cultivated on Luria-Burtani (LB) agar and R. palustris on defined mineral (PM) (41) agar with 10 mM 324 succinate. (NH₄)₂SO₄ was omitted from PM agar for determining R. palustris CFUs. Monocultures and 325 cocultures were grown in 10 mL of defined M9-derived coculture medium (MDC) (10) in 27-mL 326 anaerobic test tubes under 100% N₂ as described (10). For harvesting RNA and protein, 100-mL cultures 327 were grown in 260-mL serum vials. In both cases, MDC was supplemented with cation solution (1 % v/v; 328 100 mM MgSO₄ and 10 mM CaCl₂) and glucose (25 mM), unless indicated otherwise. R. palustris 329 monocultures were further supplemented with 15 mM sodium bicarbonate, 7.8 mM sodium acetate, 8.7 330 mM disodium succinate, 1.5 mM sodium lactate, 0.3 mM sodium formate, and 6.7mM ethanol. E. coli 331 monocultures were further supplemented with 2.5 mM NH₄Cl. Kanamycin was added to a final 332 concentration of 30 µg/ml for E. coli where appropriate. Chloramphenicol was added to a final 333 concentration of 5 µg/ml for both R. palustris and E. coli where appropriate. All cultures were grown at 334 30°C laying horizontally under a 60 W incandescent bulb with shaking at 150 rpm. Starter cocultures 335 were inoculated with 200 µL MDC containing a suspension of a single colony of each species. Test 336 cocultures and serial transfers were inoculated using a 1% dilution from starter cocultures. For 337 experiments requiring a starting species ratio of 1:1, E. coli and R. palustris starter monocultures were 338 grown to equivalent cell densities, and inoculated at equal volumes.

Generation of *E. coli* mutants. P1 transduction (42) was used to introduce deletions from Keio
collection strains into MG1655. The genotype of kanamycin-resistant colonies was confirmed by PCR
and sequencing.

Analytical procedures. Cell density was assayed by optical density at 660 nm (OD_{660}) using a Genesys 20 visible spectrophotometer (Thermo-Fisher, Waltham, MA, USA). Growth curve readings were taken in culture tubes without sampling (i.e., tube OD_{660}). Specific growth rates were determined using readings between 0.1-1.0 OD_{660} where there is linear correlation between cell density and OD_{660} . Final OD_{660} measurements were taken in cuvettes and samples were diluted into the linear range as necessary. Glucose, organic acids, formate and ethanol were quantified using a Shimadzu high-performance liquid chromatograph (HPLC) as described (43).

Sample collection for transcriptomics and proteomics. Monocultures and cocultures were grown in 100-mL volumes to late exponential phase and chilled in an ice-water bath. A 1-mL sample was collected for protein quantification using a Pierce BCA Protein Assay Kit as per the manufacturer's protocol. A 5ml sample was removed for RNA extraction and 90 ml was used for proteomic analysis. All samples were centrifuged at 4°C, supernatants discarded, and cell pellets frozen in liquid N₂ and stored at -80°C. 354 **RNA-seq.** Total RNA was isolated from cell pellets using the RNeasy kit (Qiagen, Valencia, CA, USA) 355 as per the manufacturer's protocol. In order to calculate baseline expression levels, RNA sequencing 356 reads resulting from monoculture were mapped to their corresponding reference genome (E. coli str. K-12 357 substr. MG1655 (44), NCBI RefSeq: NC_000913.3; R. palustris CGA0009 (45), NCBI RefSeq: 358 NC 005296.1) using the Tuxedo protocol for RNA expression analysis (46) (Workflow deposited 359 at https://github.com/behrimg/Task3/RNASeq). Specifically, split-reads were aligned to the reference 360 genome with Tophat2 (v.2.1.0) (47) and Bowtie2 (v.2.1.0) (48). Following mapping, transcripts were 361 assembled with cufflinks (v.2.2.0) (49), and differential expression was identified with the cufflinks tool, 362 cuffdiff (v.2.2.0). To assure that crossmapping of homologous sequencing reads would not complicate 363 expression analysis from the co-culture experiments, monoculture reads were additionally mapped as 364 described to the opposing genome. As all potential crossmapping was confined to residual rRNA reads, 365 these regions were excluded from the analysis and the co-culture RNA-seq reads where analyzed by 366 mapping the sequenced reads to both reference genomes with no further correction.

367 Preparation of protein samples for MS. Cell pellets were resuspended in 1 mL total protein buffer

368 (TPB; 20mM HEPES-NaOH pH7.4, 150mM NaCl, 2mM EDTA, 0.2mM DTT, 1:100 PMSF, 1:100

369 protease inhibitors cocktail IV) and sonicated at 20% intensity (7 seconds on, 7 seconds off) for 5 min in

an ice bath. Then 1/10 volume of 20% SDS was added. Samples were vortexed, boiled for 5 min, and

immediately placed on ice. Debris was cleared by centrifuging for 30 s at 10,000 x g at 4°C and the

372 supernatant was collected. Protein content of different lysates was analyzed by Coomassie staining

373 following SDS-PAGE and sample aliquots containing 200 µg protein were subjected to

374 chloroform:methanol protein extraction as described (50).

Analysis by LC-MS/MS. Mass spectrometry was performed at the Mass Spectrometry and Proteomics Research Laboratory (MSPRL), FAS Division of Science, at Harvard University. Samples were individually labeled with tandem mass tag (TMT) 10-plex reagents according to the manufacturer's protocol (ThermoFisher Scientific) and mixed. The mixed sample was dried in a speedvac and re-diluted with Buffer A (0.1 % formic acid in water) for injection for HPLC runs. The sample was submitted for a 380 single liquid chromatography coupled to tandem mass spectrometry (LC-MS/MS) experiment which was 381 performed on a LTQ Orbitrap Elite (ThermoFisher Scientific) equipped with Waters (Milford, MA) 382 NanoAcquity HPLC pump Peptides were separated onto a 100 µm inner diameter microcapillary trapping 383 column packed first with approximately 5 cm of C18 Reprosil resin (5 µm, 100 Å, Dr. Maisch GmbH, 384 Germany) followed by analytical column ~20 cm of Reprosil resin (1.8 µm, 200 Å, Dr. Maisch GmbH, 385 Germany). Separation was achieved through applying a gradient from 5–27% ACN in 0.1% formic acid 386 over 90 min at 200 nl min-1. Electrospray ionization was enabled through applying a voltage of 1.8 kV 387 using a home-made electrode junction at the end of the microcapillary column and spraved from fused 388 silica pico tips (New Objective, MA). The LTQ Orbitrap Elite was operated in data-dependent mode for 389 the mass spectrometry methods. The mass spectrometry survey scan was performed in the Orbitrap in the 390 range of 395 –1.800 m/z at a resolution of 6×10^4 , followed by the selection of the twenty most intense 391 ions (TOP20) for CID-MS2 fragmentation in the ion trap using a precursor isolation width window of 2 392 m/z, AGC setting of 10,000, and a maximum ion accumulation of 200 ms. Singly charged ion species 393 were not subjected to CID fragmentation. Normalized collision energy was set to 35 V and an activation 394 time of 10 ms. Ions in a 10 ppm m/z window around ions selected for MS2 were excluded from further 395 selection for fragmentation for 60 s. The same TOP20 ions were subjected to HCD MS2 event in Orbitrap 396 part of the instrument. The fragment ion isolation width was set to 0.7 m/z, AGC was set to 50,000, the 397 maximum ion time was 200 ms, normalized collision energy was set to 27V and an activation time of 1 398 ms for each HCD MS2 scan.

Mass spectrometry data analysis. Raw data were submitted for analysis in MaxQuant 1.5.6.5 (13). Assignment of MS/MS spectra was performed by searching the data against a protein sequence database including all entries from the *E. coli* MG1655 proteome (51), the *R. palustris* CGA009 proteome (45), and other known contaminants such as human keratins and common lab contaminants. MaxQuant searches were performed using a 20 ppm precursor ion tolerance with a requirement that each peptide had N termini consistent with trypsin protease cleavage, allowing up to two missed cleavage sites. 10-plex TMT tags on peptide amino termini and lysine residues were set as static modifications while methionine 406 oxidation and deamidation of asparagine and glutamine residues were set as variable modifications. MS2 407 spectra were assigned with a false discovery rate (FDR) of 1% at the protein level by target-decoy 408 database search. Per-peptide reporter ion intensities were exported from MaxQuant (evidence.txt). Only 409 peptides with a parent ion fraction greater than or equal to 0.5 were used for subsequent analysis (6063 of 410 9987 peptides). Intensities were calculated as the sum of peptide intensities. Ratios between conditions 411 were computed at the peptide level, and the protein ratio was computed as the mean of peptide ratios. All 412 ratios were normalized by dividing by the median value for proteins from the same species. Ratio 413 significance for coculture conditions at an FDR of 1% was computed by determining the ratio r at which 414 99% of genes have ratio less than r when comparing biological replicate monocultures.

415 **Expression of** *E. coli amtB* in coculture. The ASKA collection (52) plasmid harboring an IPTG-416 inducible copy of *amtB* (pCA24N *amtB*) was purified from strain JW0441-AM and introduced by 417 electroporation into WT *E. coli* and *E. coli* Δ NtrC. Cocultures were inoculated with either single colonies 418 of each species or at a 1:1 starting species ratio, as indicated in the figure legends. IPTG and 5 µg/ml 419 chloramphenicol were supplemented to cocultures to induce *E. coli amtB* expression in cocultures and 420 maintain the plasmid, respectively.

421

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567 **Figure Legends**

568 Table 1. Selected differentially expressed transcripts in cocultures of E. coli and R. palustris compared to monocultures

			Dr. Nive I	E . WT	Dr. Nr. A A mat		Dr. A AmtD	
			<i>Rp</i> Nx +	FDR	<i>Rp</i> NxAAmt	$\frac{\mathbf{D} + Ec \text{ W I}}{\mathbf{FDR}}$	<i>Rp</i> AmtB	$\frac{\mathbf{FDR}}{\mathbf{FDR}}$
	Gene		Fold	adjusted	Fold	adjusted	Fold	adjusted
Species	symbol	Gene description	change ^c	P-value	change	P-value	change	P-value
E. coli	rutA ^b	Pyrimidine monooxygenase	114.5 ± 0.0	0.09	108.0 ± 0.0	0.09	118.0 ± 0.1	0.09
<i>L. con</i>	rutC ^b	Aminoacrylate peracid reductase	60.7 ± 0.1	0.01	58.0 ± 0.1	0.01	60.9 ± 0.1	0.01
	ddpX ^{ab}	D-ala dipeptidase	58.3 ± 0.1	0.01	59.9 ± 0.1	0.01	50.1 ± 0.0	0.01
	rutD ^b	Aminoacrylate hydrolase	56.9 ± 0.0	0.01	52.9 ± 0.1	0.01	56.6 ± 0.1	0.01
	rutE ^b	Malonic semialdehyde	48.8 ± 0.1	0.01	44.4 ± 0.1	0.01	48.2 ± 0.1	0.01
	rutF ^b	FMN reductase	45.2 ± 0.1	0.01	40.3 ± 0.1	0.01	45.5 ± 0.1	0.01
	patA ^{ab}	Putrescine aminotransferase	36.3 ± 0.1	0.01	33.6 ± 0.1	0.01	34.4 ± 0.0	0.01
	argT ^{ab}	Lysine/arginine/ornithine binding protein	35.1 ± 0.3	0.01	38.9 ± 0.3	0.01	35.3 ± 0.3	0.01
	rutG ^b	FMN reductase	28.5 ± 0.0	0.01	26.9 ± 0.0	0.01	29.0 ± 0.1	0.01
		Probably dipeptide binding periplasmid						
	ddpA ^{ab}	protein	23.7 ± 0.0	0.01	26.8 ± 0.0	0.01	21.0 ± 0.0	0.01
	amtB ^{ab}	Ammonium transporter	21.3 ± 0.2	0.02	25.0 ± 0.2	0.01	24.1 ± 0.2	0.01
		-						
	yeaG ^b	Eukaryotic-like serine/threonine kinase	13.6 ± 0.0	0.08	15.2 ± 0.0	0.04	14.5 ± 0.1	0.06
	yeaH ^b	Unknown	12.8 ± 0.0	0.06	14.2 ± 0.1	0.05	14.0 ± 0.1	0.06
	metE	Methionine biosynthesis	-16.2 ± 0.1	0.03	23.6 ± 0.6	0.03	22.8 ± 0.5	0.02
	fimF	Fimbriae regulatory protein	-16.3 ± 0.0	0.01	18.4 ± 0.0	0.01	20.3 ± 0.1	0.01
	tar	Methyl-accepting chemotaxis protein II	-16.3 ± 0.2	0.01	15.8 ± 0.2	0.02	15.4 ± 0.2	0.01
	purL ^a	Purine biosynthesis	-16.8 ± 0.0	0.03	20.4 ± 0.1	0.02	18.8 ± 0.0	0.02
	flgD	Flagellar basal body rod modification protein	-17.1 ± 0.1	0.02	16.9 ± 0.0	0.01	17.4 ± 0.1	0.01
	ilvL ^a	Isoleucine biosynthesis	-17.4 ± 0.7	0.02	14.9 ± 0.4	0.02	14.2 ± 0.5	0.02
	pgaB	Glucosamine deacetylase	-17.9 ± 0.0	0.02	18.8 ± 0.0	0.03	17.3 ± 0.0	0.04
	ilvC ^a	Isoleucine biosynthesis	-18.0 ± 0.2	0.03	17.1 ± 0.2	0.04	17.6 ± 0.2	0.03
	metK	Methionine biosynthesis	-19.2 ± 0.1	0.03	17.5 ± 0.1	0.03	17.4 ± 0.1	0.04
	tap	Methyl-accepting chemotaxis protein IV	-19.7 ± 0.3	0.01	22.0 ± 0.2	0.01	22.1 ± 0.2	0.01
	flgC	Flagellar basal body	-20.1 ± 0.1	0.05				
	-						$21.02 \pm$	
	purK ^a	Purine biosynthesis	-20.7 ± 0.1	0.03	25.1 ± 0.1	0.01	0.05	0.03
	metA	Methionine biosynthesis	-21.0 ± 0.1	0.02	20.6 ± 0.1	0.02	20.8 ± 0.2	0.02
		-					$22.14 \pm$	
	ilvG ^a	Isoleucine biosynthesis	-22.1 ± 0.1	0.01	19.3 ± 0.1	0.03	0.07	0.01

	metF	Methionine biosynthesis	-23.3 ± 0.1	0.01	22.5 ± 0.1	0.01	17.62 ± 0.38 23.74 ±	0.03
	nadB	Aspartate oxidase	-24.3 ± 0.0	0.08	29.1 ± 0.1	0.05	0.01	0.07
R. palustris	RPA1206 ^a	Aldehyde dehydrogenase	36.0 ± 0.9	0.02			62.4 ± 0.4	0.01
-	RPA1205 ^a	Putative alcohol dehydrogenase	32.8 ± 0.5	0.02			28.6 ± 0.4	0.01
	RPA0538	Putative porin	31.6 ± 2.3	0.03				
	RPA1009 ^a	Possible cytochrome P450	10.4 ± 0.8	0.03				
	RPA3101 ^a	Unknown	9.4 ± 0.3	0.03			10.3 ± 0.3	0.04
	RPA4045 ^a	Putative aa ABC transport	8.8 ± 0.4	0.02				
	RPA3100	Unknown	7.8 ± 0.2	0.02				
	RPA1010	Beta-lactamase-like	7.7 ± 0.4	0.04				
	RPA4020 ^a	Putative aa ABC transport permease	7.7 ± 0.2	0.02				
	RPA1204	Unknown	7.4 ± 0.1	0.02			7.4 ± 0.1	0.03
	RPA2376	Unknown	-6.9 ± 0.1	0.04	15.4 ± 0.2	0.04	9.0 ± 0.2	0.03
	RPA2142	Putative fatty acid CoA ligase	-7.3 ± 0.1	0.03				
	RPA2377	Unknown	-8.4 ± 0.2	0.02	16.4 ± 0.6	0.05	7.3 ± 0.1	0.02
	RPA2379	Probable acetyltransferase	-8.5 ± 0.3	0.02				
	RPA2390	Possible Rhizobactin siderophore biosynthesis	-9.6 ± 0.2	0.06	22.8 ± 0.2	0.05	16.8 ± 0.5	0.03
	RPA1260 ^a	Universal stress protein	-10.5 ± 0.0	0.02			7.2 ± 0.0	0.07
	RPA2380	Possible tonB dep iron siderophore	-11.4 ± 0.6	0.03	17.1 ± 0.1	0.06	18.4 ± 0.2	0.01
	RPA1259	Putative cation-transporting P-type ATPase	-11.6 ± 0.4	0.02			10.6 ± 0.0	0.06
	RPA2378 ^a	Putative TonB-dep receptor	-13.1 ± 0.1	0.03	24.1 ± 0.3	0.06	17.5 ± 0.3	0.02

569

570 Genes shown in table were directly or indirectly mentioned in the text. For a full list of differentially-expressed genes, see Supplementary Data.

- ^a Genes were also identified as differentially expressed proteins in coculture (Table 2).
- ^b Gene is transcriptionally activated by *E. coli* NtrC during nitrogen limitation.
- ^c Fold-change values represent mean ± SD. Positive values indicate gene was upregulated in coculture. Negative values indicate gene was
- 574 downregulated in coculture. Initial cutoff was set to a log₂ value of 2 in at least 2 of 3 biological replicates. For a complete list of all differentially
- 575 regulated transcripts, refer to supplementary data. Differential expression was determined with the Cufflinks tool cuffdiff (v.2.2.0) (46)

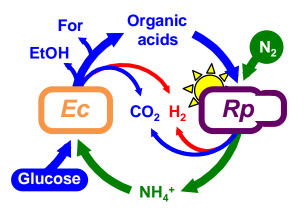
Species	Gene Symbol	Gene Description	Rp Nx + Ec WT Normalized Relative Protein Intensity ^c	Rp NxAAmtB + Ec WT Normalized Relative Protein Intensity ^d	Rp ∆AmtB + Ec WT Normalized Relative Protein Intensity ^d
E. coli	argT ^{ab}	Lysine/arginine/ornithine binding protein	10.9	11.1	10.3
	ddpA ^{ab}	D-ala dipeptide permease	5.8	7.2	6.8
	gss	Bifunctional glutathionylspermidine synthetase/amidase	4.5	4.7	4.0
	tktB	Transketolase	4.1	5.5	3.8
	potF ^{ab}	Putrescine-binding periplasmic protein	3.8	4.2	4.1
	modA	Molybdate-binding periplasmic protein	3.8	4.0	4.9
	gabD ^{ab}	Succinate-semialdehyde dehydrogenase	3.7	4.8	3.5
	dapB	4-hydroxy-tetrahydrodipicolinate reductase	3.6	2.8	2.7
	talA	Transaldolase A	3.6	4.2	4.4
	$\operatorname{amtB}^{\operatorname{ab}}$	NH4+ Transporter	3.5	3.5	3.5
	asnS	Asparagine biosynthesis	-2.1	-1.9	-1.9
	serA	Serine biosynthesis	-2.1	-2.5	-2.3
	secE	Protein translocase subunit	-2.1	-1.8	-2.3
	glf	LPS biosynthesis	-2.1	-1.9	-1.9
	yjiM	Putative dehydratase	-2.2	-1.9	-1.8
	sstT	Serine/threonine transporter	-2.2	-2.4	-2.4
	rmlA1	Carbohydrate biosynthesis	-2.3	-2.1	-2.4
	ompF	Outer membrane protein	-2.3	-2.3	-2.6
	ribĒ	Riboflavin biosynthesis	-2.3	-1.7	-1.9
	secY	Protein translocase subunit	-2.6	-2.0	-2.0
	glyA	Glycine biosynthesis	-3.2	-3.0	-3.4
	purE ^a	Purine biosynthesis	-3.3	-3.6	-3.5
	yqjI	Transcriptional regulator	-3.6	-3.0	-3.4
	asnA	Aspartate-ammonia ligase	-6.4	-3.8	-3.7
R. palustris	RPA1206 ^a	Aldehyde dehydrogenase	10.0		3.3
_	RPA1205 ^a	Putative alcohol dehydrogenase	7.8	1.2	3.9
	RPA3101 ^a	Unknown	7.1	1.5	2.6
	RPA3093	ABC transporter urea/short-chain binding protein	4.8	1.6	3.4
	RPA3297	ABC transporter urea/short-chain binding protein	4.7	1.5	3.2
	RPA4019	Putative aa ABC transporter system substrate-binding protein	3.9	1.4	2.5
	RPA4045 ^a	Putative aa ABC transport	3.3	1.4	2.1
	RPA1009 ^a	Possible cytochrome P450	3.2	1.3	2.1
	RPA1748	Putative branched-chain amino acid transport system	-2.1	-1.4	-2.5

576 Table 2. Selected differentially expressed proteins in cocultures of *E. coli* and *R. palustris* compared to monocultures

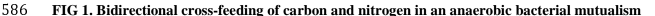
	substrate-binding protein			
RPA2378 ^a	Putative tonB-dependent receptor protein	-2.1	-1.2	-1.1
RPA2124	TonB dependent iron siderophore receptor	-2.3	-1.5	-1.7
RPA1260 ^a	Universal stress protein	-2.5	-1.5	-1.8
RPA2050	Unknown	-2.7	-1.6	-2.6
	Putative ABC transporter periplasmic solute-binding protein			
RPA3669	precursor	-2.8	-1.1	-1.3
RPA2120	Periplasmic binding protein	-6.0	-1.6	-1.7

- 577
- 578 Proteins shown in table were directly or indirectly mentioned in the text. For a full list of differentially-expressed proteins, see Supplementary
- 579 Data.
- ^a Genes were also identified as differentially expressed transcripts in coculture (Table 1)
- ^b Gene is transcriptionally activated by *E. coli* NtrC
- 582 Values represent mean normalized relative protein intensity for either two^c or one^d biological replicate. Positive values indicate gene was
- 583 upregulated in coculture. Negative values indicate gene was downregulated in coculture.

584





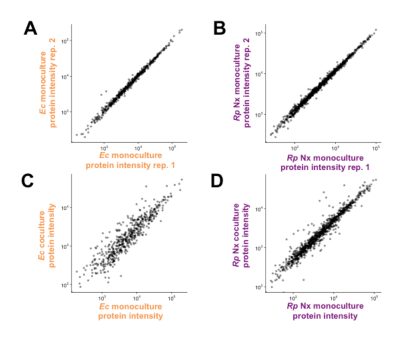


587 between fermentative *Escherichia coli* (*Ec*) and phototrophic *Rhodopseudomonas palustris* (*Rp*). *E*.

588 *coli* anaerobically ferments glucose into excreted organic acids that *R. palustris* Nx consumes (acetate,

10.589 lactate and succinate) and other products that *R. palustris* Nx does not consume (formate (For) and

- thanol (EtOH)). In return, *R. palustris* Nx constitutively fixes N_2 gas and excretes NH_4^+ , supplying *E*.
- 591 *coli* with essential nitrogen. *R. palustris* Nx grows photoheterotrophically wherein organic compounds are
- used for carbon and electrons, and light is used for energy.

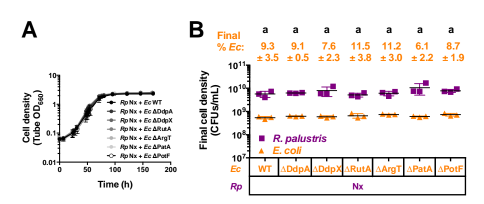


593

594 FIG 2. Coculture conditions result in altered protein expression patterns in both species, with more

- 595 differences in WT E. coli compared to R. palustris Nx. Protein expression (estimated by LC-
- 596 MS/MS intensity) of wild-type E. coli (left, A,C) and R. palustris Nx (right, B, D) comparing
- 597 protein expression patterns between monoculture biological replicates (rep. 1 versus rep. 2, **A**, **B**)
- and monoculture (average over monoculture replicates) versus coculture (**C**, **D**).





601

602 FIG 3. Single deletions of upregulated *E. coli* genes do not impair mutualistic growth with *R*.

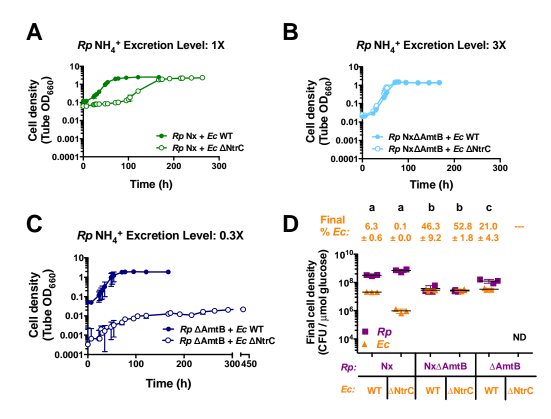
603 *palustris* Nx. Growth curves (A) and final cell densities (B) from cocultures pairing *E. coli* (*Ec*) mutants

604 with deletions in highly upregulated genes with *R. palustris* (*Rp*) Nx. Final cell densities (**B**) were taken at

605 the final time point in (A). Cocultures were started with a 1% inoculum of stationary starter cocultures

606 grown from single colonies. Error bars indicate SD, n=3. Different letters indicate statistical differences, p

607 < 0.05, determined by one-way ANOVA with Tukey's multiple comparisons posttest.





609 **FIG 4.** *R. palustris* **NH**₄⁺ excretion level affects growth and population trends in cocultures with *E.*

610 coli NtrC. Growth curves (A, B, C) and final cell densities normalized to glucose consumption (D) from 611 cocultures pairing WT E. coli (Ec) (filled circles) or Δ NtrC (open circles) with R. palustris (Rp) strains 612 with different NH_4^+ excretion levels. Final cell densities (**D**) were taken at the final time point in the 613 respective growth curve (A, B, C), except for cocultures pairing R. palustris \triangle AmtB with E. coli \triangle NtrC 614 which were sampled at 260 h. Cell densities were normalized to glucose consumed to account for 615 incomplete glucose consumption in cocultures containing E. coli Δ NtrC. Cocultures were started with a 616 1% inoculum of stationary starter cocultures grown from single colonies. Error bars indicate SD, n=3. 617 Different letters indicate statistical differences, p < 0.05, determined by one-way ANOVA with Tukey's 618 multiple comparisons posttest. ND, not determined.

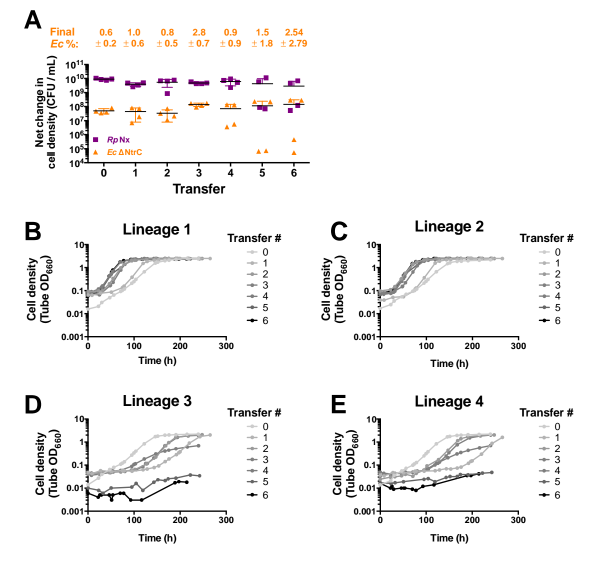
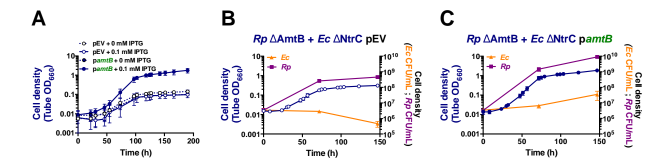


FIG 5. Lack of *E. coli* NtrC results in variable coculture growth trends across serial transfers. Net changes in cell densities (A) and replicate growth curves (B-E) of cocultures pairing *E. coli* (*Ec*) Δ NtrC with *R. palustris* (*Rp*) Nx across serial transfers. Cocultures were initially inoculated (Transfer 0) at a 1:1 starting species ratios based on CFUs/mL from *R. palustris* and *E. coli* monocultures. A 1% inoculum was used for each serial transfer. Transfers were performed every 10 d. Error bars indicate SD, n=4.

619





626 FIG 6. Ectopic expression of *amtB* in *E. coli* ΔNtrC permits mutualistic growth with *R. palustris*

627 ΔAmtB. Growth curves (A-C) and cell densities for each species (B, C) from cocultures pairing *R*.

628 *palustris* (*Rp*) ΔAmtB with *E. coli* (*Ec*) ΔNtrC harboring a plasmid encoding an IPTG-inducible copy of

629 *amtB* (pamtB, filled circles) or an empty vector (pEV, open circles). To maintain plasmids, all cocultures

630 were supplemented with 5 µg/ml chloramphenicol, which is otherwise lethal to *E. coli* but not to *R*.

631 *palustris* (Fig. S6). Cocultures were inoculated with a single colony of each species (A) or at a 1:1

632 starting species ratio based on equivalent CFUs/mL from starter *R. palustris* and *E. coli* monocultures (**B**,

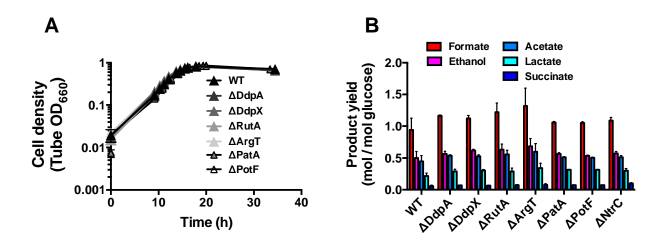
633 C). 0.1 mM IPTG was added to the cocultures at the initial time point. Error bars indicate SD, n=3.

634 Supplemental

635 **Table S1. Strains and plasmids**

Strain or plasmid	Description or Sequence (5'-3'); <u>Designation in this study</u>	Source or Purpose
R. palustris strains		-
CGA009	Wild-type strain; spontaneous Cm ^R derivative of CGA001	(45)
CGA4005	$CGA009 \Delta hupS \Delta uppE nifA*; Nx$	(10)
CGA4021	CGA4005 \Delta amtB1 \Delta amtB2; Nx\Delta AmtB	(10)
CGA4026	CGA009 $\Delta hupS \Delta uppE \Delta amtB1 \Delta amtB2; \Delta AmtB$	(12)
E. coli strains		
MG1655	Wild-type K12 strain; WT	(53)
K-12 JW1483	Keio collection $\Delta ddpX::Km$	(54)
K-12 JW5240	Keio collection $\Delta ddpA::Km$	(54)
K-12 JW0997	Keio collection $\Delta rut A::Km$	(54)
K-12 JW2307	Keio collection $\Delta argT::Km$	(54)
K-12 JW5510	Keio collection $\Delta patA::Km$	(54)
K-12 JW0838	Keio collection $\Delta potF::Km$	(54)
K-12 JW3840	Keio collection $\Delta ntrC::Km$	(54)
K-12 pCA24N (pASKA)	ASKA collection pCA24N	(52)
MG1655 pCA24N -GFP	ASKA collection pCA24N with gfp removed using NotI digest	This study
K-12 JW0441-AM pASKAamtB	ASKA collection pCA24N-N-His-amtB (gfp minus)	(52)
MG1655\DdpX	MG1655 $\Delta ddpX::Km; \Delta DdpX$	This study
MG1655∆DdpA	MG1655 Δ <i>ddp</i> A:: <i>Km</i> ; <u>ΔDdpA</u>	This study
MG1655∆RutA	MG1655 $\Delta rutA::Km; \Delta RutA$	This study
MG1655∆ArgT	MG1655 Δ <i>argT::Km</i> ; <u>ΔArgT</u>	This study
MG1655∆PatA	MG1655 Δ <i>patA::Km</i> ; <u>ΔPatA</u>	This study
MG1655∆PotF	MG1655 Δ <i>potF::Km</i> ; <u>ΔPotF</u>	This study
MG1655ΔNtrC	MG1655 Δ <i>ntrC::Km</i> ; <u>ΔNtrC</u>	This study
MG1655 pEV	MG1655 pCA24N; <u>WT pEV</u>	This study
MG1655∆NtrC pEC	MG1655 Δ <i>ntrC::Km</i> pCA24N; ΔNtrC pEV	This study
MG1655 pamtB	MG1655 pCA24N-N-His-amtB+; WT pamtB	This study
MG1655ANtrC pamtB	MG1655 ΔntrC::Km pCA24N-N-His-amtB+; ΔNtrC pamtB	This study
Plasmids		
pCA24N	Cm ^R ; ASKA collection empty vector with IPTG-inducible promoter	(52)
pCA24N-amtB+	Cm ^R ; ASKA collection vector with IPTG-inducible promoter in front of N-terminal His-tagged <i>amtB</i> gene	(52)
Primers		

ALM47	cggaaagcgcagcaatttttgt	<i>ddpX</i> upstream flanking region (<i>E. coli</i>)
ALM48	gagcaatgtgggacgaaacg	<i>ddpX</i> downstream flanking region (<i>E. coli</i>)
ALM45	atateceetggcacacage	<i>ddpA</i> upstream flanking region (<i>E. coli</i>)
ALM46	ccagcagcgttggcgtaaaata	<i>ddpX</i> downstream flanking region (<i>E. coli</i>)
ALM10 ALM51	ccgctttgcaaacaagcc	<i>rutA</i> upstream flanking region (<i>E. coli</i>)
ALM52	atcagcgcactttgctgc	<i>rutA</i> downstream flanking region (<i>E. coli</i>)
ALM49	gcaaacacacacaacacaacacaac	argT upstream flanking region (<i>E. coli</i>)
ALM50	ccatcaggtacagcttccca	argT downstream flanking region (E. coli)
ALM53	tgaaagcgtgctgttaacgc	<i>patA</i> upstream flanking region (<i>E. coli</i>)
ALM54	atcccgattttcgcgatcg	patA downstream flanking region (E. coli)
ALM55	ctggccgggagaaagttct	<i>potF</i> upstream flanking region (<i>E. coli</i>)
ALM56	ttacgggttttcgcctgc	<i>potF</i> downstream flanking region (<i>E. coli</i>)
MO 7	caatctttacacacaagctgtgaatc	<i>ntrC</i> upstream flanking region (<i>E. coli</i>)
MO 8	cctgcctatcaggaaataaagg	ntrC downstream flanking region (E. coli)
		ASKA pCA24N upstream into IPTG-
pCA24N.for	gataacaatttcacacagaattcattaaagag	inducible promoter upstream of cloned gene
		ASKA pCA24N downstream into IPTG-
pCA24N.rev	cccattaacatcaccatctaattcaac	inducible promoter upstream of cloned gene



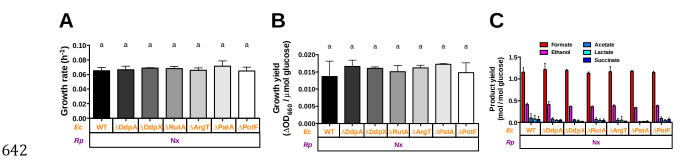
637

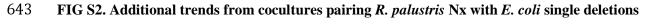
638 **FIG S1. Single deletions of** *E. coli* **genes that were upregulated in coculture no effect in monoculture**

639 with 15 mM NH₄⁺. Growth curves (A) and product yields (B) from *E. coli* monocultures grown with 15

640 mM NH₄Cl. Product yields were taken in stationary phase. Error bars indicate SD, n=3.

641





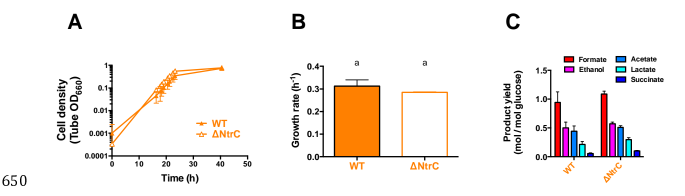
644 mutants. Growth rates (A), growth yields (B), and product yields (C) after a one-week culturing period

from cocultures pairing *E. coli* mutants with deletions in highly upregulated genes with *R. palustris* Nx.

646 Growth and product yields were taken at the final time point indicated in Fig. 3A. Cocultures were started

647 with a 1% inoculum of stationary starter cocultures grown from single colonies. Error bars indicate SD,

- 648 n=3. Different letters indicate statistical differences, p < 0.05, determined by one-way ANOVA with
- 649 Tukey's multiple comparisons posttest.



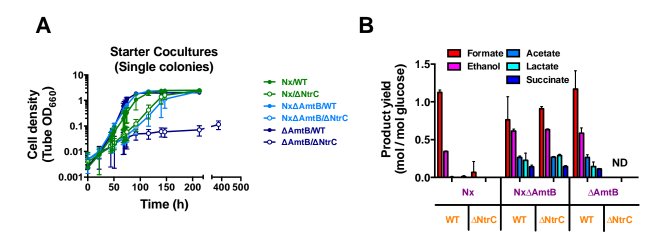
651 FIG S3. E. coli ΔNtrC growth and metabolic trends are similar to those of WT E. coli in

652 monoculture with 15 mM NH_4^+ . Growth curves (A), growth rate (B) and product yields (C) from WT E.

653 *coli* (filled) or ΔNtrC (open) monocultures grown with 15 mM NH₄Cl. Product yields were taken in

654 stationary phase. Error bars indicate SD, n=3.

655



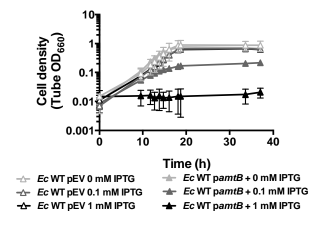
656

657 FIG S4. Additional trends from cocultures of *E. coli* ΔNtrC paired with different *R. palustris*

658 **partners.** Growth curves of starter cocultures inoculated with single colonies of each species (A) and

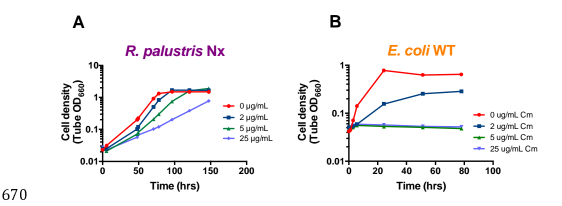
product yields from test cocultures (**B**). Product yields (**B**) were taken at the final time point indicated in

- the respective growth curve in Fig. 4. Test cocultures were started with a 1% inoculum of stationary
- 661 starter cocultures. Error bars indicate SD, n=3. ND, not determined.



662

FIG S5. Increased *amtB* expression is harmful to *E. coli* in monocultures with 15 mM NH_4^+ . Growth curves of WT *E. coli* monocultures harboring a plasmid encoding an IPTG-inducible copy of *amtB* (p*amtB*, filled) or empty vector (pEV, open) and grown at different IPTG concentrations. All monocultures were supplemented with 15 mM NH_4Cl and 5 µg/ml chloramphenicol to maintain the plasmid. Cultures were inoculated with a 1% inoculum from stationary monocultures grown in 0 mM IPTG. After inoculation, IPTG was added to the indicated final concentration. Error bars indicate SD, n=3. ND, not determined.



671 FIG S6. Determination of a chloramphenicol concentration to maintain pamtB in E. coli without

harming *R. palustris*. Representative growth curves of *R. palustris* Nx (A) and WT *E. coli* (B) at

673 different concentrations of chloramphenicol. All cultures were grown anaerobically in MDC with a 1%

674 inoculum from stationary monocultures. *R. palustris* Nx was provided 20 mM sodium acetate as a carbon

source with a 100% N₂ headspace for nitrogen. WT *E. coli* was provided 25 mM glucose, 10 mM cation

676 solution, and 15 mM NH₄Cl.