

1 Systematic identification of novel regulatory interactions controlling biofilm formation in the bacterium
2 *Escherichia coli*

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27 **Running Title:** Regulatory network for biofilm formation in bacteria

28 **Keywords:** Biofilm formation in bacteria, regulatory network, global regulation

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50 **ABSTRACT**

51 Here, we investigated novel interactions of three global regulators of the network that controls biofilm formation in the
52 model bacterium *Escherichia coli* using computational network analysis, an *in vivo* reporter assay and physiological
53 validation experiments. We were able to map critical nodes that govern planktonic to biofilm transition and identify 8
54 new regulatory interactions for CRP, IHF or Fis responsible for the control of the promoters of *rpoS*, *rpoE*, *flhD*, *fliA*,
55 *csgD* and *yeaJ*. Additionally, an *in vivo* promoter reporter assay and motility analysis revealed a key role for IHF as a
56 repressor of cell motility through the control of FliA sigma factor expression. This investigation of first stage and
57 mature biofilm formation indicates that biofilm structure is strongly affected by IHF and Fis, while CRP seems to
58 provide a fine-tuning mechanism. Taken together, the analysis presented here shows the utility of combining
59 computational and experimental approaches to generate a deeper understanding of the biofilm formation process in
60 bacteria.

61

62 **INTRODUCTION**

63 Bacteria shift from their free-swimming lifestyle to adopt a community structure to benefit from the micro-environment
64 created in a biofilm ¹, gaining protection against hazardous substances, and leading in some cases, to antibiotic
65 resistance ^{1,2,3}. It is well-understood that when differentiating from the planktonic to a biofilm structure bacteria transit
66 through specific stages, with each functional stage accompanied by changes to expression of specific genes of the
67 flagella-biofilm regulatory network ^{1,2,3}. In *Escherichia coli*, the complex transcriptional regulatory network of flagella
68 function and curli fimbriae production, the principal biofilm structure indicator, has been investigated in various
69 reports ^{4,5,6,7}. In this organism, the process is controlled by the RpoS sigma factor and FlhDC regulator. These
70 master regulators receive major regulatory inputs from c-di-GMP, cAMP and ppGpp, which are modulated by a series
71 of environmental and physiological stresses ^{8,9,10}. In this sense, the *flhDC* genes are expressed at post-exponential
72 phase and their products control more than 60 genes involved in flagella synthesis and related functions, such as
73 chemotaxis ¹¹. Also at stationary phase, the general stress response master regulator RpoS is produced and controls

74 over 500 genes in a highly complex regulatory network ⁷. Therefore, the interplay between those master regulators
75 and downstream-activated genes modulates the complex transition between planktonic and biofilm stages.

76 In the case of planktonic bacteria, mainly found in the exponential phase, flagella maintenance is due to the
77 activation of effector molecules such as FliA sigma factor. Whilst, at stationary phase, biofilm formation is initiated by
78 increasing levels in the secondary messenger c-di-GMP, as well as expression and activation of *rpoS*, which in turn
79 leads to the activation of *csgD* ^{5,12}. Additionally, there are alternative mechanisms that involve different key genes of
80 the biofilm pathway, also leading to *csgD* activation and scale-up of curli fiber production; such as CpxR and ClpX
81 which play a complex dual role during bacterium development to inhibit or activate both programs ^{8, 10, 13}.
82 Undoubtedly, the interplay between these molecules acts to modulate the proper execution of the flagella-biofilm
83 program ^{14,15}. Experimentally, the transition between flagella and biofilm formation is evaluated using a “batch cells”
84 assay, in which cells grow and attach to the chemically-inert surface of a microliter dish under static conditions,
85 generating a biofilm in an “aquatic environment”. Additionally, the “macro-colonies” assay allows a bacterial
86 population to grow over extended periods of time on agar plates, leading to striking morphological patterns,
87 resembling biofilms growing in biological materials.

88 While hundreds of reports have dissected the molecular mechanisms leading to biofilm formation in several bacterial
89 groups, a full understanding the dynamics of the complex regulatory network controlling planktonic to biofilm
90 transition still lies out of reach. This is particularly true since most studies analyzed the effect of particular regulators
91 on the expression of specific sets of genes, without addressing the complexity of the network using a more
92 systematic approach. In particular, since a few global regulators (GRs) are able to control most of the genes in
93 genome of *E. coli* ¹⁶, it is anticipated that the role of these transcriptional factors in the biofilm formation network is
94 rather underestimated. GRs are expressed or activated differentially during growth or under specific input signals ¹⁷.
95 Specifically, CRP is related to the control of metabolic processes in bacteria and it is differentially activated by
96 substrates ¹⁸, while IHF (growth rate dependent) and Fis (expressed at early exponential phase) are dual nucleoid
97 associated proteins (NAPs) that can work both as activators and repressors ^{19,20}. During the transition from motile
98 cells to biofilm structure, a mixture of different developmental stages occur. In this sense, small molecules and GRs

99 direct the activation of signaling proteins that are sensed by neighboring cells, coordinating the construction of this
100 complex community structure^{14,15}. Therefore, mapping hidden interactions in the flagella-biofilm regulatory network
101 would provide a better understanding of this developmental-like program and provide new targets for intervention
102 and/or engineering^{21,22}.

103 In this work, we used computational tools and molecular biology, together with microbiology approaches to
104 investigate the effect of the GRs CRP, IHF and Fis on the *E. coli* flagella-biofilm network. First, to analyze regulatory
105 interactions already reported, we reconstructed the flagella-biofilm transcriptional regulatory network *in silico* using
106 previous data and we analyzed its structure. Next, by using promoter expression assays, we mapped novel
107 regulatory interactions between the three GRs and key genes controlling biofilm formation. Finally, novel regulatory
108 interactions mapped using this approach were added to the original network to generate an enhanced appreciation of
109 biofilm formation. The general strategy used here is depicted in **Fig. 1**. These results suggest that IHF and Fis are
110 important modulators of the process while CRP seems to exert a fine-tuning effect. Additionally, using motility,
111 adherence and biofilm formation assays, we validated key regulatory effects observed using transcriptional fusions.
112 All together, the systematic investigation presented here adds new understanding of the complex regulatory program
113 that controls biofilm formation in *E. coli*, revealing a novel player in this network.

114 **RESULTS AND DISCUSSION**

115 ***In silico* analysis of the regulatory network controlling planktonic to biofilm transition in *E. coli***

116 To reconstruct the flagella-biofilm regulatory network, information regarding the transcriptional regulation of *E. coli*
117 was gathered from scientific articles and compiled with that from RegulonDB²³ and KEGG DB²⁴. The resulting
118 network consisted of 37 nodes representing the connections between different transcription factors (TFs), GRs and
119 small molecules (**Fig. 2**). The degree analysis (which shows the most connected nodes) of the flagella-biofilm
120 transcriptional regulatory network identified three major hubs, *rpoS*, *flhD* and *csgD* (**Fig. S1A**). The betweenness and
121 edge betweenness analysis which provides information about the most important nodes and paths in the network,
122 shows that from 37 nodes, 8 seem to be the principal effectors that modulate the proper gene expression of the

123 network (*rpoS*, *csgD*, c-di-GMP, *rpoE*, *cpxR*, *clpX*, *flhD*, *fliA*). Interestingly, *rpoS* is the main effector in the network
124 and it is modulated by different inputs. From those, c-di-GMP presents high betweenness and edge connection
125 values (**Fig. 2**). From the 37 nodes, *rpoD* and CRP were the main nodes, which exert more out-regulatory signals to
126 other nodes, while all others possessed fewer out-connections (**Fig. S1B**). Additionally, this analysis shows three
127 paths that seem to be important to maintain balance in the network. The first critical connection that maintains
128 flagella-biofilm program equilibrium lies between *rpoS* and *rpoE*, activating both *fliA* for motility and its repressor
129 *matA*, to eventually develop adherence. Secondly, *rpoS* with the TF *cpxR*, which modulates *rpoE*, but also has an
130 important connection with the TF *nsrR*, which represses *fliA* expression. Thirdly, the activation of *clpX* ATPase by
131 *rpoS* that exerts *flhD* inhibition at post-transcriptional and/or post-translational levels ²⁵. From the analysis presented
132 here, we selected ten genes in order to perform a systematic investigation of the effect of CRP, IHF and Fis, which
133 are among the main GRs of the regulatory network of *E. coli* ²⁶. We expected that this approach would reveal some
134 hidden interactions controlling the critical nodes of the flagella-biofilm network. The selected genes of interest were
135 *rpoS*, *rpoE*, *csgD*, *cpxR*, *flhD*, *fliA*, *matA*, *ompR*, *adrA* and *yeaJ*. In order to analyze the promoter activity of the
136 selected genes, the promoter regions of the genes were cloned upstream of a GFP_{lva} reporter system ²⁷ and used to
137 transform wild-type and Δihf and Δfis mutant strains of *E. coli*. As control, an empty reporter plasmid and a strong
138 synthetic promoter named BBa_J23100 (http://parts.igem.org/Part:BBa_J23100, here referred as *Pj100*) were also
139 used to transform the same strains (**Fig. S2**). The 36 reporter strains were then analyzed as presented in the next
140 section.

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142 **CRP, IHF and Fis modulate promoters of *rpoS*, *rpoE* and *flhD* master regulators**

143 We started our investigation by considering the three putative master regulators of the transition from planktonic to
144 biofilm phases. The sigma factor RpoS has been proposed as a main regulator of the flagella-biofilm network ^{7,8}, in
145 agreement with our network analysis. Thus, we were interested to understand the effect of these three GRs on the
146 promoter region of the *rpoS* gene. For this, overnight cultures of *E. coli* wild-type strain carrying the transcriptional
147 *PrpoS::GFP_{lva}* fusion were diluted 1:100 in fresh M9 minimal medium supplemented with glycerol as sole carbon

148 source, and GFP fluorescence was measured at 20 minute time intervals over 8 hours at 37°C. As can be seen in
149 **Fig. 3A**, the analyzed promoter presented a growth-phase-dependent activity with increased activity toward the end
150 of the growth curve. When the same reporter system was analyzed in *E. coli* Δihf and Δfis mutant strains, we
151 observed a significant increase in promoter activity in the absence of IHF and a strong activity in the absence of Fis
152 (**Fig. 3A left panel**, red and green lines, respectively). These data strongly suggest that IHF and Fis GRs are acting
153 to repress the *rpoS* promoter. To understand the role of CRP over the *rpoS* promoter region, we employed the
154 glucose inducible CCR system to down-modulate the activity of CRP by adding glucose in the same experimental
155 conditions as described above ²⁸. When the three reporter strains were analyzed in the presence of 0.4% glucose,
156 we observed the same general expression profile as in **Fig. 3A left panel** but a remarkably reduced promoter activity
157 (**Fig. 3A right panel**), confirming the previously reported positive role of CRP on the *rpoS* promoter ²⁹. Additionally,
158 since the synthetic constitutive promoter (*Pj100*) was not significantly affected in the strains and conditions used (**Fig.**
159 **S2**), we concluded that the effects observed for *rpoS* promoter are true regulatory events taking place at this
160 element.

161 When we analyzed the activity of the *rpoE* promoter, which controls the expression of a sigma factor linking RpoS
162 activity with flagella genes (**Fig. 2**), we observed a strong increase in promoter activity in the Δihf and Δfis strains of
163 *E. coli* when compared to the wild-type strain (**Fig. 3B left panel**), indicating that IHF and Fis have a negative effect
164 on the activity of this promoter. However, contrary to the observation for *rpoS*, the addition of glucose (**Fig. 3b right**
165 **panel**) to the media (which inhibits CRP activity) did not result in any significant change in *rpoE* promoter activity.

166 After investigating the activity of *rpoS* and *rpoE* promoters, we analyzed the promoter of *flhD* gene, which encodes a
167 master regulator of flagellar genes and of the flagella-related *fliA* sigma factor ³⁰. Using the same experimental
168 conditions as described above, we first observed that *flhD* promoter activity has a strong growth-dependence, with
169 activity increasing during the growth curve when assayed in the wild-type strain of *E. coli* (**Fig. 3C left panel**). When,
170 we assayed the promoter activity in Δihf and Δfis mutant strains, we observed a similar expression profile as in the
171 wild-type, with the exception that in both strains the initial promoter activity was significantly higher than in the wild-
172 type. When we assayed promoter response in the three strains in the presence of glucose (to trigger CRP

173 inactivation), activity was strongly impaired in all strains (**Fig. 3C right panel**), confirming the reported role of CRP as
174 activator of *flhD* expression ³¹.

175 **Novel regulatory effects for CRP, IHF and Fis exerted on *csgD*, *fliA* and *yeaJ* promoters**

176 Once we had investigated the effect of the GRs of flagella to biofilm transition, we decided to analyze the promoters
177 of genes related to biofilm formation and an additional lower flagella regulator (*fliA*). CsgD is a central protein related
178 to, and an indicator of, biofilm formation ^{12, 32}. When we analyzed the activity of *csgD* promoter in the wild-type strain,
179 we observed a peak of expression at about 200 min of growth with a reduced activity later in the growth phase (**Fig.**
180 **4A left panel**). Investigating this promoter in Δihf and Δfis mutants, we observed a strong decay in promoter activity,
181 indicating that both proteins positively modulate the activity of the promoter. Moreover, when we added glucose to
182 the growth media (**Fig. 4A right panel**), we observed a generalized decay in promoter activity in all three strains,
183 indicating a positive role of CRP for this promoter. Reflecting on these results, it is interesting to note that, while CRP
184 and IHF have been reported as positive regulators of the *csgD* promoter ^{33, 34}, the positive effect of Fis has never
185 been demonstrated before.

186 In the case of the *fliA* gene that codes a sigma factor specifically related to flagella genes, analysis of promoter
187 activity in both wild-type and Δfis mutant strains revealed a very low level of activity throughout the growth curve,
188 both in the presence and absence of glucose (**Fig. 4B**). However, in the Δihf mutant strain, this promoter displays a
189 strong increase (30-fold) in activity in the absence of glucose (left panel), while this level was lower when glucose
190 was added (right panel). Since *fliA* expression is dependent on FlhD ³⁰, the observed decay of *fliA* promoter activity
191 during growth in the presence of glucose could be the result of a cascade process, with the apparent IHF repression
192 of this promoter suggesting a previously unreported regulatory interaction. To test the direct effect of IHF on FliA, we
193 identified two putative binding sites for this global regulator at *fliA* promoter **Fig. S3A**. We then constructed a mutant
194 version of this promoter with 11 point mutations that fully abolish both sites and tested this new variant as before. As
195 shown in **Fig. S3B**, expression of the mutated version of *fliA* promoter was low in the wild-type and Δfis mutant

196 strains at the same level as the original promoter, and displayed a higher activity in the *Δihf* mutant, both in the
197 absence and presence of glucose. These data suggest that the effect of IHF on *fliA* promoter is indirect.

198 We next analyzed the effect of GRs on the expression of *yeaJ*, a diguanylate cyclase coding gene, which modulates
199 the levels of c-di-GMP in the cell. In *E. coli*, deletion of *yeaJ* results in strains with reduced motility at 37 °C⁸,
200 however no regulatory proteins have been demonstrated to play a role in the expression of this gene. The expression
201 profile of the *yeaJ* promoter in wild-type and *Δihf* mutants strains showed a strong peak of activity at 200 min of
202 experiment (**Fig. 4C**). However, when this promoter was assayed in the *Δfis* mutant, we observed a significant
203 decrease in promoter activity, indicating that the Fis regulator could play a positive role in its expression.
204 Furthermore, addition of glucose to the growth media (**Fig. 4C right panel**) resulted in an increase in the promoter
205 activity in the first minutes of growth, suggesting the existence of additional, potentially CRP-dependent, regulatory
206 mechanisms at this promoter. Finally, the additional three promoters selected for investigation (from *adrA*, *cpxR* and
207 *ompR* genes, **Fig. S4**) displayed maximal activity very similar to that of the negative control (the empty reporter
208 vector, pMR1 in **Fig. S2**), which precludes unequivocal conclusions concerning their regulation.

209 **Reconstruction and analysis of the regulatory network with novel interactions**

210 In order to generate fresh insight into the regulatory network controlling biofilm formation in *E. coli*, we added the new
211 regulatory interactions identified in the previous sections to the network from **Fig. 2**. For this, the effect of CRP, IHF
212 and Fis over the promoter regions obtained by our promoter activity assay was transformed into activation or
213 inhibition links. From the 27 interactions tested, we suggested 8 new interactions, and were able to confirm 5
214 described interactions (**Table S1**). We integrated these new interactions into the network and re-performed the
215 centrality measurements. Using this approach, we observed that the integration of the experimental data changed the
216 topology of the network, while maintaining the three major hubs identified (**Fig. S5A**). Yet, the out-degree analysis
217 showed that *rpoD*, CRP, IHF, and Fis in minor degree are the main nodes exerting the most out-connections to all
218 other nodes (**Fig. S5B**), while the edge betweenness analysis indicates that, upon the addition of the new interactions,
219 the critical nodes of the network remain the same (**Fig. 5**). Taken together, these data suggest the existence of

220 previously unidentified regulatory interactions in the biofilm regulatory network that play a critical role in the planktonic
221 to biofilm transition.

222 **Phenotypic consequences of loss of GRs for the planktonic and biofilm stages of *E. coli***

223 Flagella play an important function during the different developmental phases of biofilm formation such as attachment
224 at a surface, as well as bacterial cell cohesion within the biofilm ^{2,30}. Therefore, we were interested in understanding
225 the effect of the GRs CRP, IHF and Fis on the flagella function. As it can be seen in **Fig. 6**, after 24h of incubation in
226 a low agar plate, *E. coli* BW25113 wild-type strain showed reduced motility in the absence of glucose. However,
227 analysis of the Δihf strain in similar conditions revealed a strong motile phenotype, with this effect more apparent
228 after 24h of incubation with full spreading over the plate observed (**Fig. S6**). Finally, under similar conditions, the Δfis
229 strain presented reduced motility, which is more evident at 24h. It is important to notice that addition of glucose to the
230 media generated a general decrease in motility of all strains (**Figs. 6 and S6**, right panels). Interestingly, the
231 observed strong phenotypic effect of Δihf on the motility of *E. coli* can be traced directly to the strong increase of *fliA*
232 promoter activity in this mutant strain, since FliA sigma factor controls several genes related to flagella formation in
233 this organism ³⁵.

234 We next analyzed the capability of the different strains to attach to a solid surface, which is indicative of early biofilm
235 formation, using the protocol described by O'Toole *et al.*, 2011. For this, we performed the experiments at 30 °C and
236 37 °C, since different reports have used different temperatures ^{36,37}. In general, *E. coli* cells adhered to the wall of the
237 96-well plates in all conditions. At 37°C, two- or three-fold increases in biofilm yields were observed when compared
238 with *E. coli* cells cultured at 30°C, as can be seen by comparing the biofilm indexes in **Fig. 7A**. In general terms, the
239 Δihf mutant strain of *E. coli* displayed decreased adhesion at both temperatures, which is in agreement with the
240 increased motility observed for this strain in **Fig. 6**. By the same token, adhesion of the Δfis mutant strain at 37 °C
241 was significantly increased when compared to the wild-type, which could be due to the reduced motility observed
242 before. In most of the cases, the general adhesion observed for the strains was decreased in the presence of
243 glucose, with the exception of the adhesion level of Δihf mutant at 30 °C. These data suggest that, at 37°C, CRP and

244 IHF are important effectors to activate the adherence program. They also suggest that Fis is acting as a repressor of
245 the early biofilm program, since in its absence, motility is higher at 37 °C. Altogether, adherence yields are better
246 when induced at 37°C than at 30°C, and the three GRs CRP, IHF and Fis participate differentially to regulate the
247 adherence capabilities of *E. coli*.

248 As depicted in **Fig. S7**, all the strains were capable of developing biofilm structures at 30 °C and 37 °C in the
249 presence or absence of glucose, as previously reported. In general, *E. coli* cells grown at 30 °C produced a biofilm
250 structure resembling that reported for 28 °C, while those grown at 37 °C conditions showed a more visually complex
251 biofilm structure with three morphologically distinct zones, as reported by Serra *et al.*, 2013 (**Fig. 7B**). More explicitly,
252 in wild-type cells a concentric ring delineates zone 3, which presents an intense red color suggesting curli formation
253 by the different layers of stationary phase cells. Zone 2 represents the intrinsic capability of wild-type community to
254 produce wrinkles and curli. Finally, zone 1 presents weak red color, an indicative of bacteria at exponential phase
255 related with colony expansion but not with curli production ⁷. In the presence of glucose, the biofilm structure formed
256 by the wild-type strain presented slight differences. When we analyzed the Δihf mutant strain, we observed a
257 significant reduction in the colony size and degree of pigmentation (that could be related to lower curli production)
258 when compared to the two other strains (**Fig. 7B**). This size reduction can be evidenced by the systematic decrease
259 in the three zones of the colony in this strain. Yet, analysis of biofilm formation in this strain in the presence of
260 glucose showed a recovery in colony size generated by a high expansion of zone 1, while pigmentation was
261 apparently not affected. Finally, the analysis of Δfis mutant strain revealed a colony where the zone 1 (expansion
262 zone) could not be detected. Additionally, this strain presented a darker zone 3, which could be indicative of higher
263 production of curli fimbriae than wild-type strain, with addition of glucose to the growth media only generated small
264 changes in the colony such as an apparent reduction in size.

265 Altogether, morphology assays have shown that IHF and Fis GRs are important contributors to proper biofilm
266 development, as could be indicated by drastic changes in the three well-characterized zones in these mutants. In
267 contrast, the role of CRP protein (which was indirectly assayed by the addition of glucose to the media) during
268 mature biofilm formation could not be dependably inferred, since the change in the nutritional state of the colonies

269 could have a strong impact in the process. Taken together, these data indicate that these GRs could have an
270 important role in the final biofilm structure in *E. coli* that could be the result of the detected change in gene expression
271 of key regulatory elements in the planktonic to biofilm regulatory network.

272 **Conclusions**

273 The work presented here describes the systematic investigation of the regulatory interaction mediated by GRs during
274 the transition from planktonic to biofilm in the bacterium *E. coli*. While several works have addressed this issue
275 before, data generated has been fragmented and few targets have been investigated at each time. The systematic
276 investigation used here allowed the identification of novel interactions mediated by CRP, Fis and IHF. More
277 importantly, the modulation of some critical nodes, such as *fliA* by IHF, could explain the strong phenotypic effects
278 observed for motility and attachment assays, as this regulator has also been found to be a modulator of genes
279 involved in the biofilm program as *rpoS*, *matA* and *csgD*^{33, 38, 39, 40}. The molecular mechanism involved in the different
280 phases of biofilm formation by IHF remains to be determined. However, in this work, we present evidence that IHF is
281 a key element to biofilm formation by modulating gene expression level of the flagella-biofilm program, suggesting
282 that IHF could be a good candidate to disrupt/engineer bacterial biofilm structure. Here, we also demonstrated that
283 Fis plays a more major role in the biofilm formation network than anticipated, both using promoter assays as well as
284 physiological tests. The evidence in this study indicates that, CRP, IHF and Fis are important effectors that modulate
285 the flagella-biofilm network. Interestingly, no condition in this study shows absence of motility or biofilm formation,
286 suggesting that, while these GRs are important modulators of these processes, they are not essential genes to
287 suppress completely the flagella-biofilm program. Additionally, it has been demonstrated that bacteria without the
288 main effectors *rpoS* or *csgD* (also *rpoE*, *csgB*, *flhDC* or *fliA*) are still capable of producing a biofilm^{7, 41}. Altogether,
289 the high plasticity observed in the flagella-biofilm program clearly shows that the network has evolved the robustness
290 to compensate for the loss of critical nodes, indicating the importance of this program for bacterial survival. Finally,
291 the systematic identification of novel connections for a specific network will allow prediction of the behavior of the
292 biological system in the absence of different TFs. It opens the possibility of using genetic engineering to achieve a
293 balanced connectivity between synthetic circuits and the entire bacterial network when performing a specific task. In

294 other words, that rational interactive synthetic circuits can operate in accordance with the natural network of a
295 specific organism, thereby allowing enhanced performance of a desired biological task.

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297

298 **EXPERIMENTAL PROCEDURES**

299 **Bacterial strains, media and growth conditions**

300 The list of bacterial strains, plasmids and primers is presented in **Table1**. *Escherichia coli* wild-type (BW25113) or
301 strains genetically depleted of IHF or Fis GRs ⁴² were used as the host of all plasmids. *E. coli* DH5 α or DH10b were
302 used as cloning strains. DNA of *E. coli* BW25113 and MG1655 were used as templates. *E. coli* cells were grown in
303 Luria Broth (LB) or M9 minimal media (6.4g/L Na₂HPO₄·7H₂O, 1.5g/L KH₂PO₄, 0.25g/L NaCl, 0.5g/L NH₄Cl)
304 supplemented with 2mM MgSO₄, 0.1mM CaCl₂, 0.1mM casamino acids, and 1% glycerol or 0.4% glucose as the
305 carbon source. Chloramphenicol was added (34 μ g/mL) to ensure plasmid maintenance. Cells were grown at 37°C
306 with constant shaking at 220 rpm overnight.

307 **Plasmid construction**

308 Plasmids and primers used in this work are listed on Table 2 and 3 respectively. As a control of all experiments we
309 transformed *E. coli* BW25113 wild-type and mutant strains with an empty pMR1 vector ²⁷ and pMR1-Pj100. To
310 analyze the signal integration of GRs at the studied genes, we amplified the promoter regions by PCR and cloned
311 into pMR1 vector using standard protocols ⁴³. DNA from *E. coli* BW25113 wild-type was used as a template for all
312 constructions except for *flhD* promoter, for which *E. coli* MG1655 was used. PCR was performed using 50 ng of DNA
313 template with 50pmol of each primer (purchased from Sigma and extended) and 1U Phusion® High-Fidelity DNA
314 Polymerase (New England Biolabs M0530L) in a 50 μ l reaction volume. Initial denaturation was at 98°C for five
315 minutes; subsequently, amplification was done over 30 cycles at 98°C, 60°C and 72°C, each with duration of 30
316 seconds. The final extension step was performed at 72°C for ten minute . All PCR products were gel purified and

317 cloned as EcoRI/BamHI fragments in pMR1, previously digested with the same enzymes. The resulting plasmids
318 were DNA sequence verified and correct plasmids were used to transform *E. coli* BW25113 (wild-type), *E. coli*
319 JW1702 (Δihf) and *E. coli* JW3229 (Δfis).

320

321 **Promoter activity assay**

322 For promoter activity measurements, single colonies of wild-type and mutant *E. coli* strains harboring the different
323 plasmids were picked from fresh plates. Each strain was inoculated and grown overnight at 37 °C in 2 mL of M9
324 medium with glycerol or glucose containing chloramphenicol with shaking at 225 rpm. Stationary phase cultures were
325 diluted to a final optical density (OD) of 0.05 in 200 μ L of M9 medium containing either glycerol or glucose as
326 required, and supplemented with chloramphenicol. Cell growth and GFP fluorescence was quantified every 20
327 minutes over an 8 hr incubation at 37 °C using Victor X3 plate reader (PerkinElmer, Waltham, Massachusetts, USA).
328 Under these conditions, all strains presented similar growth rates (**Fig. S8**). Data were analyzed from at least three
329 biological samples, arbitrary units were calculated by dividing GFP-corrected by OD600-corrected, subsequently, the
330 mean and standard error for each promoter was calculated and graphed using Microsoft Excel and R software.

331 **Bacteria motility assay**

332 The effect of GRs on motility was evaluated as following. *E. coli* wild-type or mutant strains harboring pMR1 plasmid
333 were grown overnight at 37°C on LB plates. Inoculation of a single colony onto motility plates (tryptone 1%, NaCl
334 0.25%, agar 0.3% and when indicated glucose was added) was done by using a toothpick ⁴⁴. The motility halos were
335 measured at 18h and 24h. Each strain was evaluated in triplicate from independent plates.

336 **Biofilm formation in liquid media**

337 Biofilm formation was assayed using the Microtiter Dish Biofilm Formation Assay ⁴⁵. Single colonies of *E. coli* wild-
338 type or mutant strains harboring all the constructions were grown overnight at 37°C in M9 supplemented with glycerol
339 or glucose. The cultures were diluted 1:100 in 200 μ l of fresh M9 media with either glycerol or glucose. A 96-well plate

340 was incubated at 37°C for 72h, after which cells were gently washed four times with distilled water and left at room
341 temperature for 15 minutes with 200µL of crystal violet 0.1% w/v solution. Subsequently, vigorous washing was
342 performed with distilled water (four times) and the plates left to dry at room temperature for 24h. Finally, 200µL acetic
343 acid 30% w/v was added and after 15 minutes, the solution was transferred to a new plate and Optical Density at
344 550nm was measured in FLUOstar Omega (BMG Labtech) plate reader. All the culture dilutions, staining and
345 quantification steps were performed by using procedures developed for a robotic platform called Edwin ⁴⁶. Data from
346 biological triplicates were analyzed by Microsoft Excel software. Average values and standard deviation was
347 determined.

348 **Biofilm formation in solid media**

349 Biofilm morphology was evaluated by Congo red assay ⁴⁷. Chloramphenicol was added during all steps to ensure
350 plasmid maintenance. Wild-type and mutant *E. coli* strains with pMR1 plasmid were grown overnight at 37°C on LB
351 agar plates. Single colonies were picked and grown for 14h at 37°C on 1ml LB at 220rpm. The cultures were washed
352 twice with MgSO₄ and resuspended in M9 media with either glycerol or glucose as required, to an OD at 600nm (~
353 0.5). 5 µL drop of each culture was added into YESCA-CR plates ⁴⁸. YESCA media (1 g/L yeast extract, 20 g/L agar)
354 was complemented with Congo red 50 µg/ml diluted in KPi buffer (50 mM potassium phosphate buffer, pH 7.2, 28.9
355 mM KH₂PO₄, 21.1 mM K₂HPO₄ in water), where indicated glucose was added at 0.4% final concentration. YESCA-
356 CR seeded plates were allowed to dry in a sterile environment and, left to grow at 37°C. Biofilm development was
357 followed (documented) for 6 days using a fluorescence stereoscope (Leica MZ16). Biological triplicate images were
358 analyzed by ImageJ and prisma software was used to determine the statistical significance of the experiments by
359 using one-way ANOVA.

360 **Network Construction**

361 To generate the transcriptional regulatory network that will define our genes of interest, we gathered information from
362 research articles that contained information on genes that modulate the transition between planktonic to biofilm
363 formation. Eventually, the data were compared with information available at Regulon DB, EcoCyc DB and Kegg

364 pathway DB. Cytoscape 3.4.1 was used to generate the transcriptional network with slight modifications ⁴⁹. Briefly,
365 we transformed our mined data into activation or inhibition statements for each of the partner molecules. Secondly, to
366 construct the network graph, we used Cytoscape 3.4.1 in which different connections were added to each node on
367 the collected data. Next, networks were analyzed using different algorithms, such as degree, out-degree,
368 betweenness centrality by using Cytoscape 3.4.1 software. Finally, we transformed our GFP dynamics gene
369 expression experimental results into functional interactions (inhibits or activates). Results with same function as
370 reported were maintained without changes in the rewired network, while those with different activity were changed in
371 the network. During the data collection from the literature, some TFs were reported with dual effect (activation and
372 repression) for the same target gene. For instance, *rpoS* expression was reported to be activated but also inactivated
373 by the CRP global regulator ^{50, 51, 52}. Additionally, most of the genes that were reported as being regulated by different
374 TFs, but in the web-databases the same gene did not present a *cis*-element to any TFs. To overcome those
375 constrains we considered for the network that any TF, master regulator, global regulator, small molecule or RNAs
376 could be a node. This last, despite being gathered, were not included because RNA-node analysis of regulatory gene
377 expression significantly increases the complexity of the network ^{12, 53} and because of the complex requirements of
378 molecular tools to evaluate this topic. For edges direction analysis, we considered the activation or inhibition most
379 reported, or both when divergent reports were found. The data were plotted in a power-law graph using an organic
380 algorithm to give cluster visualization generating the transcriptional regulatory network of both planktonic and biofilm
381 structure. Furthermore, we performed degree analysis in order to present a general view of the most connected
382 nodes, followed by edge and node betweenness analysis ⁵⁴ in order to disclose the main effectors and the logic
383 pathway that could describe the network. Finally, we also performed the out degree analysis at the integrated
384 network to identify the potential talkative nodes.

385

386 **ACKNOWLEDGMENTS**

387 The authors express their thanks to members of the Silva-Rocha & Elfick labs for insightful discussion and
388 suggestions in developing this work. This work was supported by a FAPESP Young Research Award (grant number
389 2012/22921-8) and by a Royal Society Newton International Exchange award (NI140137). GRA was supported by a
390 CAPES PhD fellowship while ASM was supported by FAPESP Scientific Initiation Award (2015/22386-3). AdIH was
391 supported by an EPSRC Programme Grant (EP/J02175X) to the Flowers Consortium.

392

393 **AUTHOR CONTRIBUTIONS**

394 G.R.A. and R.S.R. conceived the work. G.R.A., A.dH. and A.S.M. performed the experiments. G.R.A., A.dH. and
395 A.M.S. analyzed the data. G.R.A. and R.S.R. drafted the manuscript. A.dH. and A.E. corrected the manuscript. A.E.
396 and R.S.R. provided funding for the execution of the work. All authors read and approved the manuscript.

397 **COMPETING FINANCIAL INTEREST**

398 The authors declare no competing financial interest.

399 **MATERIAL & CORRESPONDENCE**

400 Correspondence and material request should be addressed to Rafael Silva Rocha.

401 **ADDITIONAL INFORMATION**

402 Supplementary Information accompanies this paper online.

403

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Tables

Table 1. Bacterial strains, plasmids and primers used in this study.

Strain	Description	Reference
<i>E. coli</i> DH5α	F ⁻ <i>endA1 glnV44 thi-1 recA1 relA1 gyrA96 deoR nupG purB20</i> φ80 <i>dlacZ</i> ΔM15 Δ(<i>lacZYA-argF</i>)U169, <i>hsdR17</i> (<i>r_Km_K</i> ⁺), λ ⁻ .	55
<i>E. coli</i> BW25113	<i>lacI^rrmB_{T14} ΔlacZ_{WJ16} hsdR514 ΔaraBAD_{AH33} ΔrhaBAD_{LD78} rph-1</i> Δ(<i>araB-D</i>)567 Δ(<i>rhaD-B</i>)568 Δ <i>lacZ4787</i> (:: <i>rmB-3</i>) <i>hsdR514 rph-1</i> .	56
<i>E. coli</i> JW1702	<i>E. coli</i> BW25113 Δ <i>ihfA</i> mutant strain	56
<i>E. coli</i> JW3229	<i>E. coli</i> BW25113 Δ <i>fis</i> mutant strain	56
<i>E. coli</i> MG1655	K-12 F ⁻ λ- <i>ilvG-</i> <i>rfb-50 rph-1</i>	57
Plasmids		
pMR1	CmR, <i>oriP15a</i> ; GFP <i>lva</i> promoter probe vector	27
pMR1-Pj100	CmR, <i>oriP15a</i> ; pMR with synthetic constitutive promoter BBa_J23100	This study
pMR1-PrpoS	CmR, <i>oriP15a</i> ; pMR1- <i>rpoS</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-PrpoE	CmR, <i>oriP15a</i> ; pMR1- <i>rpoE</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-PcsgD	CmR, <i>oriP15a</i> ; pMR1- <i>csgD</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-PflhD	CmR, <i>oriP15a</i> ; pMR1- <i>flhD</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-PfliA	CmR, <i>oriP15a</i> ; pMR1- <i>fliA</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-PyeaJ	CmR, <i>oriP15a</i> ; pMR1- <i>yeaJ</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-AdrA	CmR, <i>oriP15a</i> ; pMR1- <i>adrA</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-PcpxR	CmR, <i>oriP15a</i> ; pMR1- <i>cpxR</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-PmatA	CmR, <i>oriP15a</i> ; pMR1- <i>matA</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-PompR	CmR, <i>oriP15a</i> ; pMR1- <i>ompR</i> -GFP <i>lva</i> transcriptional fusion	This study
pMR1-PfliA/IHFmut	CmR, <i>oriP15a</i> ; pMR1- <i>fliA</i> /IHFmut-GFP <i>lva</i> transcriptional fusion	This study
pMR1-PyeaJ/Fismut	CmR, <i>oriP15a</i> ; pMR1- <i>yeaJ</i> /Fismut-GFP <i>lva</i> transcriptional fusion	This study
Primers		
Sequence		
5 <i>adrA</i> EcoRI	5'-GCGCGAATTCGAAAAAAGTTTGACGCCAC-3'	This study
3 <i>adrA</i> BamHI	5'-GCGCGGATCCCAATTTTCCCAAATTATAGAGACGG-3'	This study
3 <i>adrA</i> SphI	5'-GCGCGCATGCGCACGTTTACGCCATTAC-3'	This study
5 <i>cpxR</i> EcoRI	5'-GCGCGAATTCGCGTGGCTTAATGAACTGAC-3'	This study
3 <i>cpxR</i> BamHI	5'-GCGCGGATCCTGTTTAAATACCTCCGAGGCA-3'	This study
3 <i>cpxR</i> SphI	5'-GCGCGCATGCATGAAGCAGAAACCATCAGATAG-3'	This study
5 <i>matA</i> EcoRI	5'-GCGCGAATTCCTTTTCACTCAAACCTGTTAAGATG-3'	This study
3 <i>matA</i> BamHI	5'-GCGCGGATCCCCGGAAGTAAATAAGATACGT-3'	This study
3 <i>matA</i> SphI	5'-GCGCGCATGCACCAATAATTTGCTAAGGCC-3'	This study
5 <i>ompR</i> EcoRI	5'-GCGCGAATTCCTCGTTGATTCCCTTTGTCT-3'	This study
3 <i>ompR</i> BamHI	5'-GCGCGGATCCGCAACAATTTGTAAGCGTGT-3'	This study
3 <i>ompR</i> SphI	5'-GCGCGCATGCCACCAGGTAACATTAATCCAG-3'	This study
5 <i>fliA</i> - IHFmut overlap	5'-CCC GGGTTGCACATTC CCGGGG CCGGATAAGGCGT-3'	This study
3 <i>fliA</i> - IHFmut overlap	5'-CCCCGGGAATGTGCAACCCGGGTAAATTGCAATTCAACTGTAGGC-3'	This study
5 <i>yeaJ</i> -Fismut EcoRI	5'-GCGCGAATTCGAAGCGAAAAGCGAGGG-3'	This study

585 **Figure legends**

586
587 **Figure 1. General strategy to define novel regulatory interactions controlling planktonic/biofilm transition.**

588 The flagella-biofilm network was constructed and analyzed based on available data from the literature. Subsequently,

589 analysis of the promoter architecture regions of the principal nodes effectors was performed to confirm the
590 interactions reported. Next, cloning of the natural promoter regions in the pMR1-reporter system were developed.
591 Promoter activity was determined for each promoter in different conditions as established in material and methods.
592 Finally, GFP activity was transform into connectivity data and loaded into flagella-biofilm network to gain a better
593 understanding of this program.

594 **Figure 2. Flagella-biofilm transcriptional regulatory network.** The principal nodes and paths, which drive the
595 flagella-biofilm network were analyzed using Cytoscape 3.4.1. The network was analyzed by using betweenness
596 centrality and edge betweenness centrality algorithms⁵⁴. Size of the nodes (circles) indicates betweenness analysis
597 and width of the Edges (lines connections the circles) indicates the edge betweenness analysis. Scale colors from
598 red to bright to dark indicate the high to low values.

599 **Figure 3. Effect of CRP, IHF and Fis GRs over the promoter activity of *rpoS*, *rpoE* and *flhD*.** Promoter activity
600 assay of (A) pMR1-*PrpoS*, (B) pMR1-*PrpoE*, and (C) pMR1-*PflhD* were evaluated in *E. coli* BW25113 *wild type* (blue
601 line), Δihf (red line) and Δfis (green line) in 96well plate as described in methods in the absence (left panel) or
602 presence (right panel) of 0.4% of glucose. GFP fluorescence was measured each 20 minutes at 37 °C during 8 hours
603 in static conditions and normalized by OD600. Solid lines represents the average from three independent
604 experiments while dashed lines are the upper and lower limits of standard error of the mean (S.E.M).

605 **Figure 4. Effect of CRP, IHF and Fis GRs on the promoter activity of *csgD*, *fliA* and *yeaJ*.** GFP promoter activity
606 assay of (A) pMR1-*PcsgD*, (B) pMR1-*PfliA*, and (C) pMR1-*PyeaJ* were evaluated in *E. coli* BW25113 *wt* (blue line),
607 Δihf (red line) and Δfis (green line) in 96well plate as described in methods in the absence (left panel) or presence
608 (right panel) of 0.4% of glucose. GFP fluorescence was measured each 20 minutes at 37 °C during 8 hours in static
609 conditions and normalized by OD600. Solid lines represents the average from three independent experiments while
610 dashed lines are the upper and lower limits of standard error of the mean (S.E.M).

611 **Figure 5. Experimental data integration to the Flagella-biofilm transcriptional regulatory network.** Promoter
612 activity values were transformed into activation or repression connections and were loaded into Cytoscape 3.4.1. An
613 organic algorithm was used to the properly cluster visualization. The network was analyzed by use the betweenness
614 centrality and edge betweenness centrality measurements. Size of the nodes (circles) indicates betweenness
615 analysis and width of the Edges (lines connections the circles) indicates the edge betweenness analysis. Scale
616 colors from red to bright to dark indicate the high to low values.

617 **Figure 6. Effect of GRs in the motility program at 24h.** Motility phenotype of *E. coli* BW25113 wild-type and
618 mutant strains were evaluated by cell motility assay at 24h in the presence or absence of glucose as depicted.

619 Divergent motility capability is observed between the different conditions, proving the effect of the GRs CRP, IHF and
620 Cis to modulate the motility program. The results are representative of 3 independent experiments.

621 **Figure 7. Capability of *E. coli* and mutant strains to develop adherence and mature biofilm.** A) Adherence
622 capability of *E. coli* BW25113 *wild-type*, Δihf and Δfis were evaluated in 96-well plate using violet crystal method.
623 Comparisons of adherence capability of BW25113 wt and mutant strains at 30 and 37°C are shown. Vertical bars are
624 standard deviations calculated from three independent experiments. B) Mature biofilm formation of *E. coli* BW25113
625 *wt*, Δihf and Δfis strains were perform using congo red plate assay. Comparisons of the mature biofilm morphological
626 characteristics of wild type and mutant strains at 37°C in the presence or absence of glucose is shown. Dashed line,
627 Zone III; white line, zone II; black line, zone I; arrows represent wrinkles and clefts structures. The results are
628 representative of 3 independent experiments.

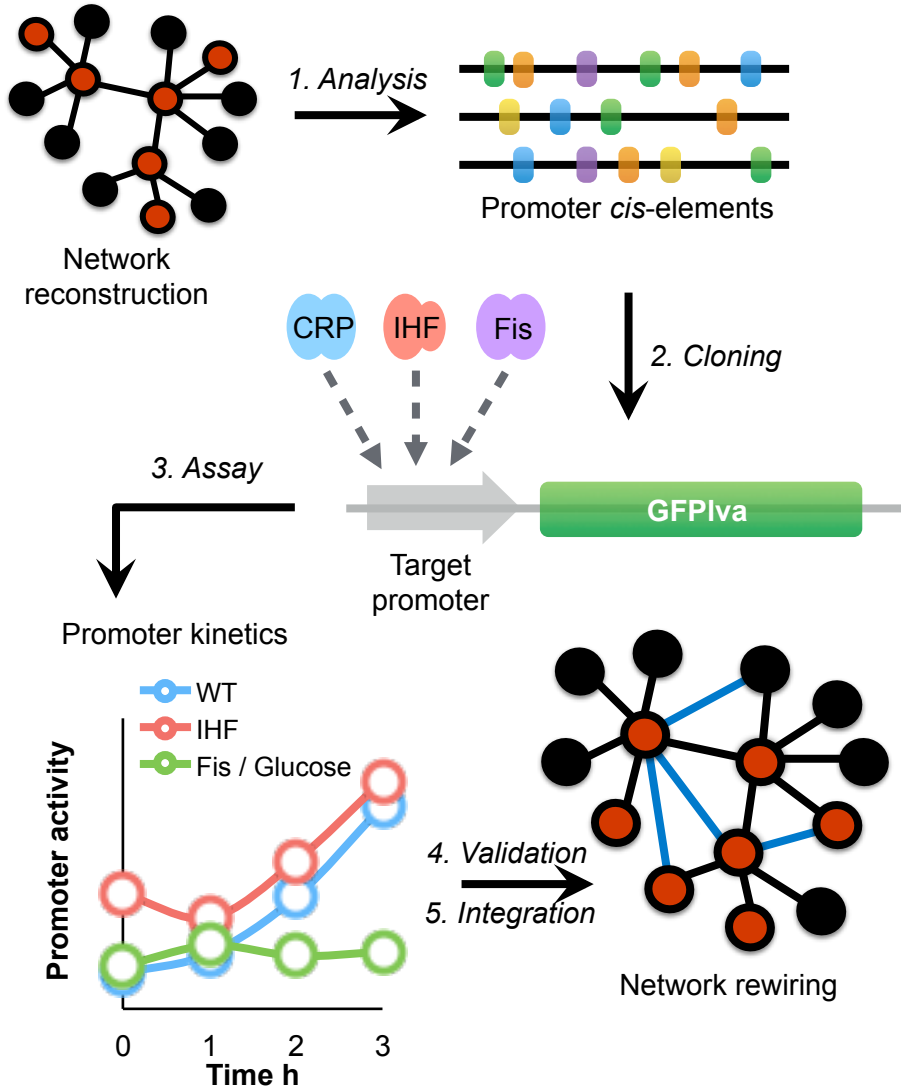


Figure 1

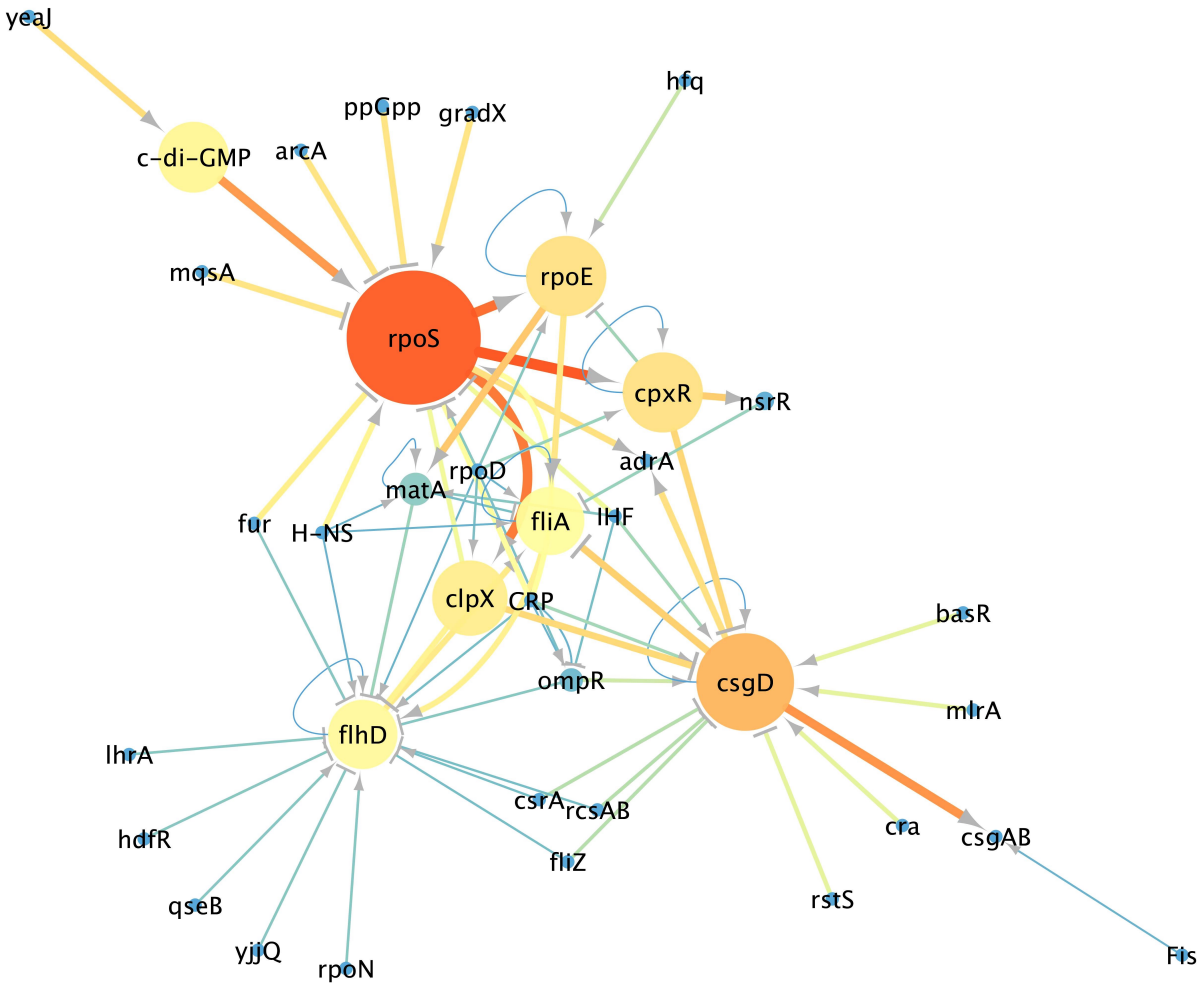


Figure 2

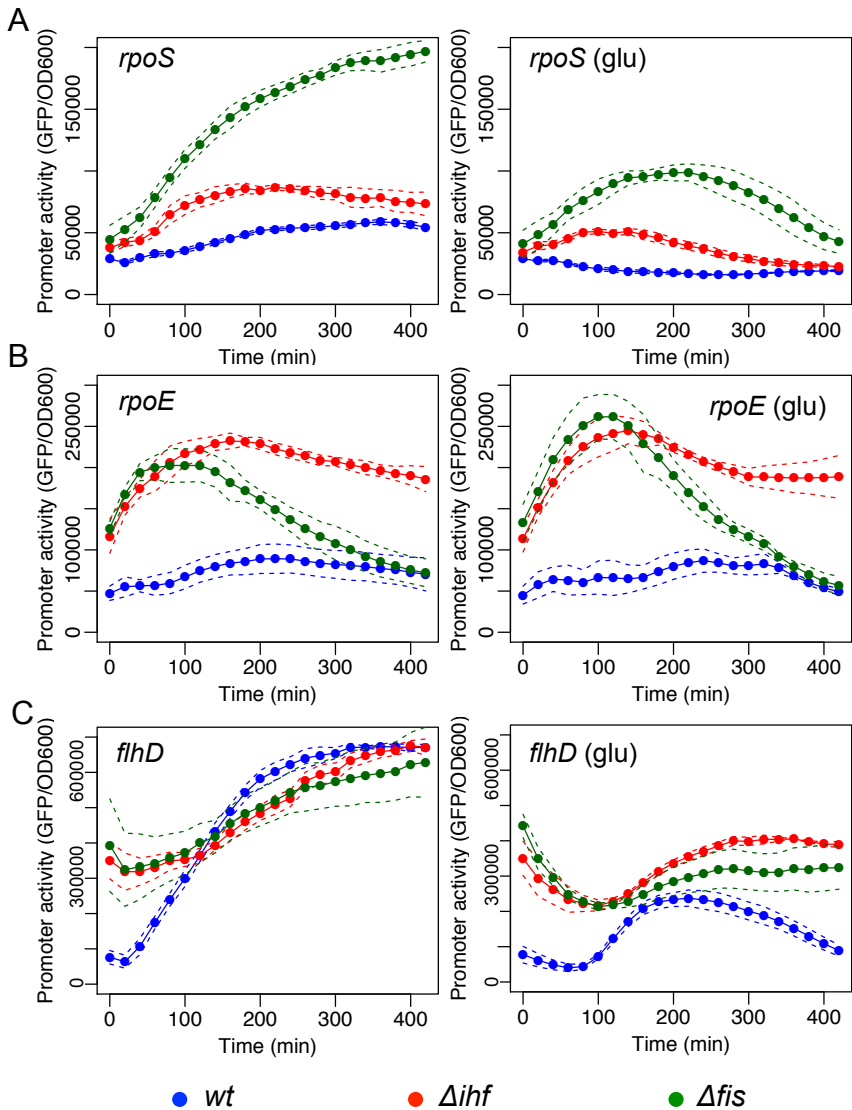


Figure 3

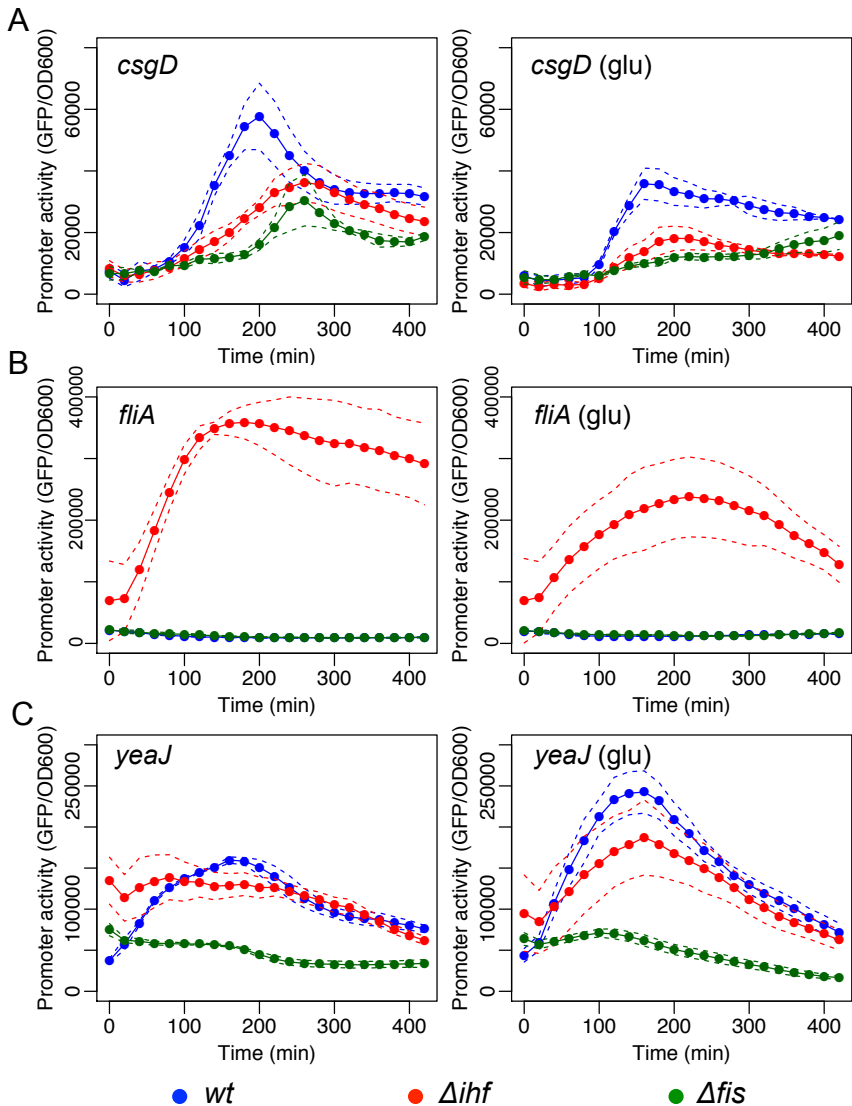


Figure 4

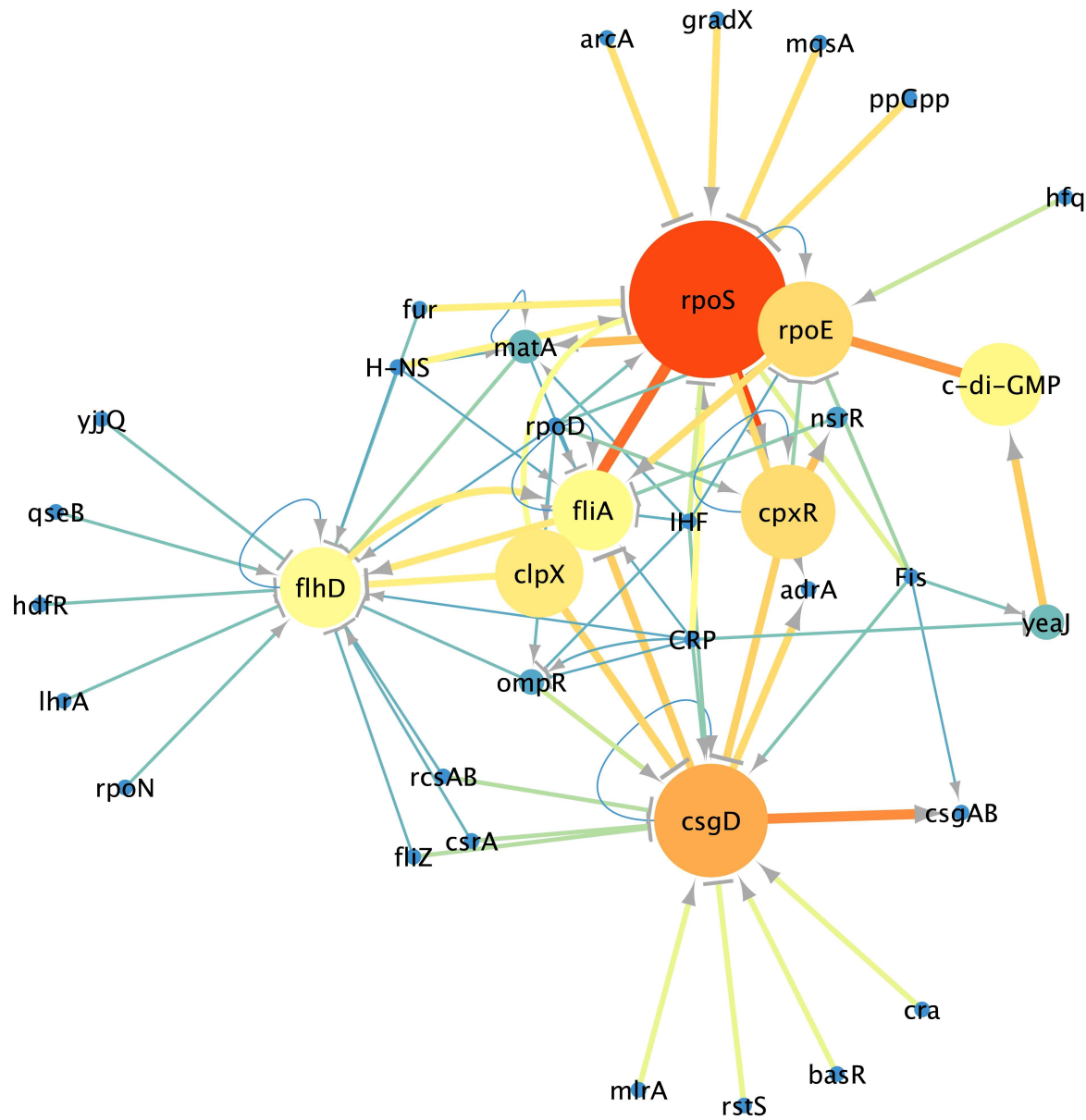
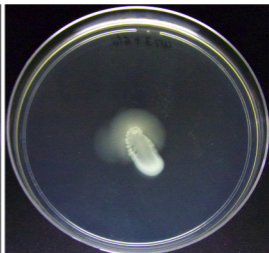
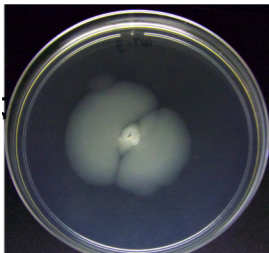


Figure 5

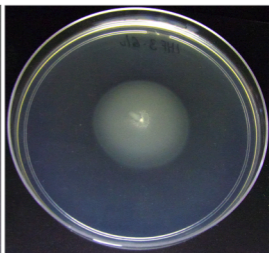
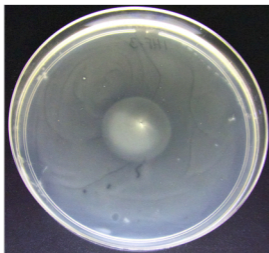
Control

Glucose

Wild-type



Δ lih1



Δ fis1

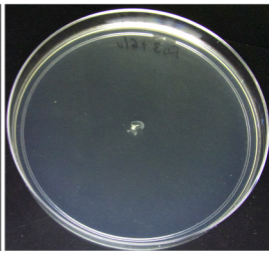
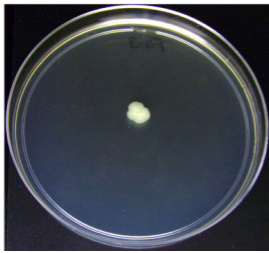


Figure 6

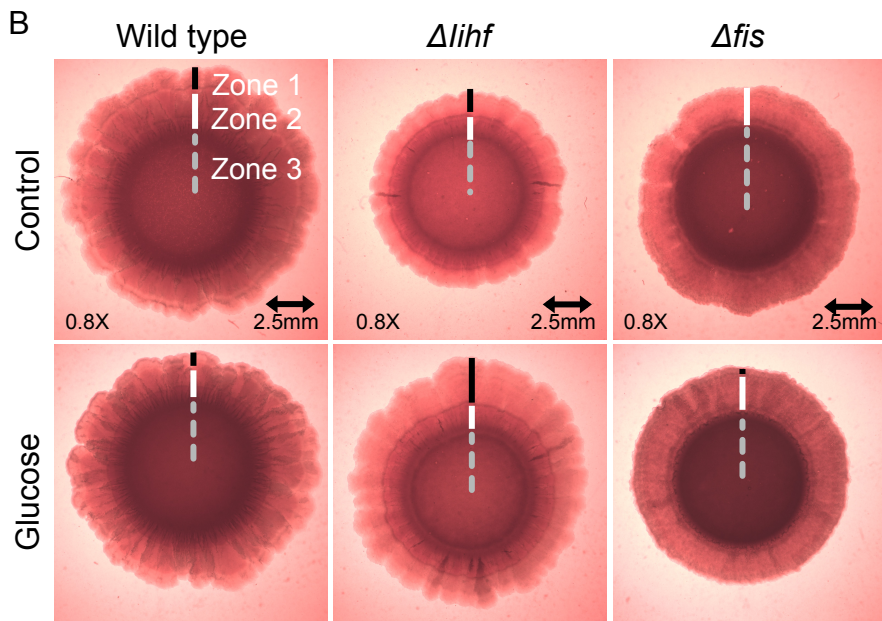
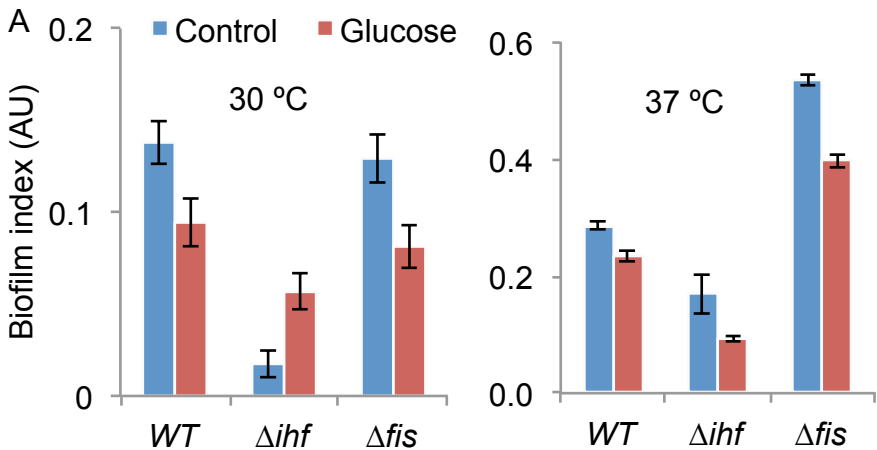


Figure 7