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Re-defining how mRNA degradation is coordinated with transcription and translation in bacteria

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15 **Abstract**

16

17 In eukaryotic cells, transcription, translation, and mRNA degradation occur in distinct
18 subcellular regions. How these mRNA processes are organized in bacteria, without
19 employing membrane-bound compartments, remains unclear. Here, we present
20 generalizable principles underlying coordination between these processes in bacteria. In
21 *Escherichia coli*, we found that co-transcriptional degradation is rare for mRNAs except
22 for those encoding inner membrane proteins, due to membrane localization of the main
23 ribonuclease, RNase E. We further found, by varying ribosome binding sequences, that
24 translation affects mRNA stability not because ribosomes protect mRNA from degradation,
25 but because low translation leads to premature transcription termination in the absence
26 of transcription-translation coupling. Extending our analyses to *Bacillus subtilis* and
27 *Caulobacter crescentus*, we established subcellular localization of RNase E (or its
28 homolog) and premature transcription termination in the absence of transcription-
29 translation coupling as key determinants that explain differences in transcriptional and
30 translational coupling to mRNA degradation across genes and species.

31

32 **Keywords**

33 mRNA degradation, RNase E, transcription-translation coupling, premature transcription
34 termination, transertion, co-transcriptional regulatcanion

35 Introduction

36 Unlike eukaryotic cells, bacterial cells do not have a nucleus, and the transfer of genetic
37 information from DNA to protein takes place within a common space, the cytoplasm,
38 permitting concurrence of translation and even mRNA degradation while mRNA is
39 transcribed^{1,2}. How transcription, translation, and mRNA degradation are coordinated
40 during the life cycle of an mRNA, in the absence of membrane-bound compartments, is
41 a fundamental question that underpins gene expression regulation in bacterial cells.
42 Addressing this question will enable understanding of how protein expression levels are
43 regulated by the cell in response to different environments³⁻⁵ and also inform the design
44 of synthetic gene expression systems, where precise manipulation of gene expression is
45 essential⁶.

46 Decades of research in a model bacterium, *E. coli*, has provided strong evidence that
47 transcription is coupled to translation⁷⁻¹¹, that mRNA degradation can start during
48 transcription¹²⁻¹⁵, and that translation affects mRNA degradation^{16,17}. However, whether
49 this picture is broadly applicable across all genes and different bacterial species remains
50 unclear. Identifying the molecular and sequence variables affecting coupling between
51 transcription, translation, and mRNA degradation will lead to a generalizable model for
52 understanding the regulation of bacterial gene expression. In this work, we investigated
53 when (during the life cycle of mRNA) and where (within a cell) mRNAs are degraded in
54 coordination with transcription and translation in bacterial cells and identified key factors
55 that contribute to commonalities and differences in co-transcriptional and post-
56 transcriptional control of gene expression in *E. coli*, *B. subtilis*, and *C. crescentus*.

57 The possibility of co-transcriptional mRNA degradation has been discussed since
58 early studies of long operons in *E. coli*. In *lac* and *trp* operons, mRNA sequences from
59 the promoter-proximal gene were shown to decay before the promoter distal genes were
60 transcribed¹²⁻¹⁴. A genome-wide measurement of mRNA lifetimes in *E. coli* compared
61 transcription elongation time and mRNA lifetime and suggested that many long genes
62 that exhibit transcription elongation times longer than mRNA lifetimes may experience co-
63 transcriptional mRNA degradation¹⁵. Co-transcriptional mRNA degradation can have a
64 significant impact by reducing the number of proteins made per transcript, which can be
65 beneficial when a quick stop in protein synthesis is needed to respond to changing cellular

66 needs. However, whether co-transcriptional degradation is indeed possible in *E. coli*
67 remains in question because the main ribonuclease controlling mRNA degradation,
68 RNase E, is found on the inner membrane of the cell, away from the nucleoid¹⁸⁻²¹. Co-
69 transcriptional mRNA degradation would therefore need to invoke the dynamic
70 relocalization of a gene locus to the membrane, which has been observed for certain
71 genes^{22,23}. Furthermore, unlike in *E. coli*, RNase E is localized in the cytoplasm in *C.*
72 *crescentus*²⁴⁻²⁶, raising a question about how mRNA degradation is differentially
73 controlled in *C. crescentus* cells in comparison to *E. coli* cells.

74 In contrast to the lack of clarity how mRNA degradation can be coupled to transcription
75 in bacterial cells, several studies have supported the coupling of mRNA degradation to
76 translation, such that mRNAs with a strong ribosome binding sequence have long
77 lifetimes^{16,17,27-31}. This trend has been explained by the notion that ribosomes protect
78 mRNA from degradation. However, what aspect of ribosome activity—for example,
79 whether it is the rate of loading at the 5' end of the mRNA or whether it is ribosomal
80 density across the mRNA—is responsible for the protective role remains unclear^{16,17,32}.
81 Understanding the exact mechanism of translation that affects mRNA lifetime would help
82 make quantitative predictions for expression output for different genes.

83 In this work, we used *lacZ* as a model gene to study how transcription, translation,
84 and mRNA degradation are coordinated in bacterial cells. The *lac* operon in *E. coli* is a
85 paradigm of bacterial gene regulation, and our current understanding of transcription-
86 translation coupling in bacteria and dependency between translation and mRNA stability
87 has been established by seminal studies that used *lacZ* as a model gene^{9,28,33,34}. Its
88 regulatory mechanisms are well characterized, allowing us to manipulate parameters for
89 *lacZ* gene expression and test hypotheses toward a generalizable model. For example,
90 we introduced the effect of transertion to *lacZ* to emulate what happens to genes encoding
91 inner membrane proteins²², we varied the 5' untranslated region (5'-UTR) sequence of
92 *lacZ* to test variable translation efficiencies across the genome^{3,35}, and we perturbed the
93 subcellular localization of RNase E to capture differences across bacterial species^{36,37}.
94 From this approach, we identified spatial and genetic design principles that bacteria have
95 evolved to differentially regulate transcriptional and translational coupling to mRNA
96 degradation across various genes and species.

97

98 **Results**

99 ***lacZ* mRNA is degraded post-transcriptionally, uncoupled from transcription**

100 While the possibility of nascent mRNA degradation during transcription has been
101 discussed^{12-15,27,38,39}, the actual rate of co-transcriptional mRNA degradation has never
102 been reported. We used *lacZ* gene under the *lac* promoter in *E. coli* as a model to
103 measure the rates of co-transcriptional and post-transcriptional mRNA degradation (k_{d1}
104 and k_{d2} , respectively; **Fig. 1A**). Earlier studies discussed co-transcriptional degradation
105 of *lacZ* based on the observation that *lacZ* decays before the synthesis of *lacY* and *lacA*
106 in the original *lac* operon. However, this result can be explained by co-transcriptional
107 mRNA processing at the intergenic region between *lacZ* and *lacY*⁴⁰⁻⁴², instead of real co-
108 transcriptional degradation of *lacZ*. Therefore, we deleted *lacY* and *lacA* genes from the
109 original *lac* operon in the chromosome of wild-type, MG1655 to yield a monocistronic *lacZ*
110 (strain SK98). The intrinsic terminator sequence after *lacA* follows the coding sequence
111 of *lacZ* to ensure the dissociation of RNA polymerase (RNAP) from DNA after finishing
112 the transcription of *lacZ* (**Fig. 1B**). To follow the degradation kinetics of *lacZ* mRNA after
113 the stoppage of transcription initiation, transcription of *lacZ* was induced with membrane-
114 permeable inducer isopropyl b-D-1-thiogalactopyranoside (IPTG) and re-repressed with
115 glucose 75 seconds (s) after addition of IPTG (**Fig. 1B**). Importantly, glucose was added
116 before the first RNAPs finished transcription, so that co-transcriptional mRNA degradation
117 can be observed. During this time course, a population of cells was acquired every 20-30
118 s, from which 5' and 3' *lacZ* mRNA levels were quantified by quantitative real-time PCR
119 (qRT-PCR) using probes denoted as Z5 and Z3, respectively. A key feature of this time
120 course experiment is the temporal separation of *lacZ* mRNA status between nascent and
121 released (**Fig. 1B**). Until the first RNAPs finish transcription of *lacZ* (T_3), all *lacZ* mRNAs
122 are expected to be nascent (time window i). After the last RNAPs finish transcription of
123 *lacZ* at t_3 , all *lacZ* mRNAs are released (time window iii). In between, nascent and
124 released *lacZ* mRNAs co-exist (time window ii). Hence, we can measure the rates of co-
125 transcriptional and post-transcriptional mRNA degradation by fitting Z5 level changes with
126 an exponential decay function in the time windows i and iii, respectively.

127 If the co-transcriptional degradation of *lacZ* mRNA takes place ($k_{d1} > 0$), Z5 level will
128 decrease in the time window *i*, as demonstrated by a mathematical model (**Fig. 1C** and
129 **S1A**). However, our data shows that Z5 level stays constant in the time window *i*,
130 suggesting that k_{d1} is close to zero (**Fig. 1D**). From biological replicates, we obtained k_{d1}
131 = $0.042 \pm 0.0598 \text{ min}^{-1}$ and $k_{d2} = 0.43 \pm 0.067 \text{ min}^{-1}$ (**Fig. S1B**). Essentially, the mean
132 lifetime of nascent *lacZ* mRNA ($1/k_{d1}$) is 24 min, much longer than transcription elongation
133 time (~ 3.5 min), suggesting that *lacZ* mRNA is unlikely to experience degradation during
134 transcription elongation.

135

136 **Membrane localization of RNase E accounts for uncoupling of transcription and** 137 **degradation of *lacZ* mRNA**

138 Among various ribonucleases in *E. coli*, the endoribonuclease RNase E has been
139 considered the main enzyme to initiate mRNA degradation^{21,36,43-45}, including *lacZ*
140 mRNA^{27,46,47}. To confirm that the observed k_{d1} and k_{d2} of *lacZ* mRNA are controlled by
141 RNase E, we repeated the experiment in a strain carrying a temperature-sensitive RNase
142 E allele (*rne3071*, strain SK519), in which RNase E can be inactivated by a 10-min shift
143 to 43.5°C⁴⁸. We performed IPTG induction of *lacZ* transcription at 43.5°C after 10 min of
144 the temperature shift. Because transcription elongation is faster at this high temperature
145 ($T_3' = 100$ s), glucose was added at 50 s after IPTG induction, so that we can still capture
146 the time window *i* to measure k_{d1} . When RNase E was inactivated, k_{d1} and k_{d2} were about
147 7 times smaller than those measured in wild-type RNase E at 43.5°C and about 2-3 times
148 smaller than those measured at 30°C (**Fig. S2A**), confirming that RNase E controls k_{d1}
149 and k_{d2} of *lacZ* mRNA. In *E. coli* cells, RNase E is associated with the inner membrane
150 via the membrane targeting sequence (MTS)²⁰. Therefore, the lack of co-transcriptional
151 degradation of *lacZ* mRNA (very low k_{d1}) could be accounted for by the membrane
152 localization of RNase E, away from the nucleoid (or transcription site).

153 *E. coli* cells are viable even when the MTS sequence of RNase E is removed and
154 RNase E is localized to the cytoplasm^{20,49} (RNase E Δ MTS, **Fig. 2A**). When RNase E is
155 in the cytoplasm, instead of anchored to the membrane, it can interact with nascent and
156 released mRNAs more frequently and likely affect k_{d1} and k_{d2} of *lacZ* mRNA. To check
157 this possibility, we measured k_{d1} and k_{d2} of *lacZ* mRNA in the strain expressing RNase E

158 Δ MTS. We found that cytoplasmic RNase E increases both k_{d1} and k_{d2} ; especially, k_{d1}
159 increases about 7 fold, to $0.31 \pm 0.084 \text{ min}^{-1}$, in comparison to the wild-type RNase E
160 strain (**Fig. 2B**).

161 We tested another cytoplasmic RNase E mutant, RNase E (1-529) (**Fig. 2A**). This
162 mutant lacks the MTS as well as the C-terminal domain, which provides binding sites for
163 other RNA degradosome components⁵⁰ (**Fig. S2B**). To interpret the effect of RNase E
164 localization in the absence of the C-terminal domain, we compared k_{d1} and k_{d2} of *lacZ*
165 mRNA from cells expressing the RNase E (1-529) mutant with those from cells expressing
166 a RNase E (1-592) mutant, which also lacks the C-terminal domain but is localized to the
167 membrane via MTS. We found that k_{d1} and k_{d2} of *lacZ* mRNA are higher in the cytoplasmic
168 RNase E (1-529) than in the membrane-bound RNase E (1-592) (**Fig. 2B**), supporting
169 that mRNA degradation is faster when RNase E is localized in the cytoplasm. We note
170 that the absence of C-terminal domain in RNase E (1-592) results in lower k_{d1} and k_{d2} of
171 *lacZ* mRNA in comparison to those in the wild-type RNase E, suggesting the importance
172 of having the C-terminal domain for the catalytic activity of RNase E happening at the N-
173 terminal domain^{51,52} (See **Fig. S2C** for additional data). Altogether, our results show that
174 the membrane localization of RNase E slows down the degradation of *lacZ* mRNA,
175 especially during transcription, giving rise to the uncoupling of transcription and mRNA
176 degradation.

177

178 **Proximity of nascent mRNAs to the membrane alone does not affect their** 179 **degradation rates**

180 Since slow co-transcriptional mRNA degradation is likely due to the spatial separation
181 between membrane-localized RNase E and nascent mRNAs, we considered a scenario
182 where nascent mRNAs are positioned close to the membrane. When mRNAs coding for
183 a transmembrane protein are transcribed, co-transcriptional translation may be
184 accompanied by membrane insertion of the nascent protein, a process known as
185 transertion⁵³⁻⁵⁵. A previous study showed that expression of *lacY* (encoding the lactose
186 permease localized in the inner membrane) brings the *lacY* locus and nearby DNA region
187 (~90 kb) close to the membrane²³. This suggests that even a gene encoding a
188 cytoplasmic protein (such as *lacZ*) can be localized close to the membrane if it is adjacent

189 to an actively transcribed *lacY* gene locus on the chromosome. Therefore, we inserted a
190 constitutively expressed *lacY* gene downstream of *lacZ* (strain SK435; **Fig. 3A**) to test if
191 the transertion of *lacY* can bring *lacZ* closer to the inner membrane and increase k_{d1} . As
192 a control, we made a strain where *lacY* is replaced with *aadA*, a gene encoding a
193 cytoplasmic protein, spectinomycin adenylyltransferase, that does not undergo
194 transertion (strain SK390).

195 To test the effect of transertion on the localization of nascent *lacZ* mRNA, we
196 performed fluorescence in situ hybridization (FISH) using Cy3B-labeled probes binding
197 to the first 1-kilobase region of *lacZ* mRNA ($Z5_{\text{FISH}}$; **Fig. 3A**)⁵⁶. Transcription of *lacZ* was
198 induced with IPTG and re-repressed with glucose as in the qRT-PCR experiment (**Fig.**
199 **1B**). Cells were sampled every 1 min interval and fixed immediately. The $Z5_{\text{FISH}}$ signal
200 appeared as diffraction-limited foci (**Fig. 3A** and **S3A**). Their centroid coordinates along
201 the short and long axes of the cell were normalized to cell width and length, respectively,
202 and combined into a 2D histogram (**Fig. 3B-3C**). Notably, until T_3 (or the end of the time
203 window i , $t = 210$ s), most of the $Z5_{\text{FISH}}$ signals are expected to be from nascent mRNAs
204 tethered to gene loci (**Fig. 1B**). Hence, the location of $Z5_{\text{FISH}}$ at $t = 1, 2,$ and 3 min after
205 induction allows us to examine the subcellular localization of the nascent mRNAs (and
206 their gene loci) exclusively. We observed that already at $t = 1$ min, $Z5_{\text{FISH}}$ in SK435 (*lacZ*
207 followed by constitutively transcribed *lacY*) were localized off the center long axis, while
208 those in SK390 (*lacZ* followed by constitutively transcribed *aadA*) were close to the center
209 long axis of the cell (**Fig. 3B-3C**). As time progresses to $t = 2$ and 3 min, $Z5_{\text{FISH}}$ in both
210 strains localized away from the center long axis, likely due to the *lacZ* gene locus moving
211 to the periphery of the nucleoid upon induction—an effect previously observed in the *lacZ*
212 locus in *E. coli*⁵⁷. In all three time points, $Z5_{\text{FISH}}$ in SK435 were localized closer to the
213 membrane than those in SK390 (**Fig. 3B-3C**), suggesting that the transertion of *lacY*,
214 which does not occur with *aadA*, results in the neighboring *lacZ* gene transcription taking
215 place close to the inner membrane.

216 Next, we measured k_{d1} and k_{d2} of *lacZ* mRNA in SK435 and SK390 by qRT-PCR to
217 check if the proximity to the membrane allows the nascent and released *lacZ* mRNAs to
218 be degraded faster. We found that k_{d1} and k_{d2} of *lacZ* mRNAs were invariable between

219 the two strains and almost identical to the original *lacZ*-only strain without *lacY* or *aadA*
220 (**Fig. 3D**).

221 We wondered if nascent *lacZ* mRNAs in SK435 were not close enough to the
222 membrane to facilitate their degradation. To bring nascent *lacZ* mRNAs even closer to
223 the membrane, we constructed a translational fusion of *lacZ* with the first two
224 transmembrane segments of *lacY* (*lacY2*), so that *lacZ* is directly linked to the transertion
225 element (**Fig. 3E**). We also fused the *venus* gene at the 3' end of *lacZ* sequence to verify
226 the membrane localization of LacZ proteins by fluorescence imaging (strain SK575; **Fig.**
227 **S3B**). FISH imaging of 5' *lacZ* mRNA expressed from the *lacY2-lacZ-venus* fusion at t =
228 1, 2, 3 min after induction showed that as soon as two transmembrane segments (*lacY2*
229 of mRNA length 222 nt) are transcribed, the *lacZ* sequence is strongly enriched near the
230 membrane in comparison to the original *lacZ* strain without the *lacY2* fusion (**Fig. 3F-3G**
231 and more information in **Fig. S3C**). This result suggests that transertion takes place
232 immediately after induction and brings nascent transcripts to the membrane. Also, the
233 direct translational fusion of *lacY2* element to *lacZ* placed the nascent *lacZ* mRNAs closer
234 to the membrane in comparison to the previous *lacZ-lacY* context where the transertion
235 effect came from the neighboring *lacY* gene locus (strain SK435; **Fig. 3B**).

236 While nascent *lacZ* mRNAs were closer to the membrane, their k_{d1} was not larger than
237 that of the original *lacZ*-only strain (strain SK98; **Fig. 3H**). This result further supports the
238 notion that proximity to the inner membrane (where RNase E is localized) is not sufficient
239 to increase the rate of co-transcriptional *lacZ* mRNA degradation.

240 In the *lacY2-lacZ-venus* fusion strain, we can also measure the degradation kinetics
241 of the *lacY2* region of the transcripts using a set of qRT-PCR primers amplifying that
242 region. Remarkably, we found that *lacY2* exhibits fast co-transcriptional mRNA
243 degradation with $k_{d1} = 0.34 \pm 0.041 \text{ min}^{-1}$ (**Fig. 3I** and **S3D**). This likely represents the
244 characteristics of the original *lacY* transcript, as we measured a similar rate of co-
245 transcriptional mRNA degradation from the full-length *lacY* gene (**Fig. S3E**). This finding
246 indicates that coupling between transcription and mRNA degradation is possible for
247 transcripts encoding inner-membrane proteins. Considering that the proximity of nascent
248 mRNAs to the membrane alone does not affect the co-transcriptional degradation rate of

249 *lacZ* mRNA, the fast co-transcriptional degradation observed with *lacY* mRNA is likely
250 due to additional factors other than its proximity to the membrane (see **Discussion**).

251
252 **Another cytoplasmic protein-coding gene, *araB*, exhibits high k_{d1} due to its RBS**
253 **sequence**

254 To determine if the result we observed for *lacZ* is generalizable to other genes in *E. coli*,
255 we examined k_{d1} and k_{d2} of another gene encoding a cytoplasmic protein, *araB*, which is
256 under the control of the arabinose-inducible promoter (P_{ara}) of the *araBAD* operon on the
257 chromosome of *E. coli* strain MG1655 (**Fig. 4A**). We deleted *araA* and *araD* genes to
258 make *araB* a monocistronic gene (P_{ara} -*araB*; strain SK472). Transcription from the P_{ara}
259 promoter was induced with arabinose and re-repressed by glucose 50 s afterward. In
260 contrast to very slow co-transcriptional degradation observed in *lacZ* mRNA, 5' *araB*
261 mRNAs were degraded before 3' *araB* mRNAs were transcribed (time window indicated
262 as a blue box in **Fig. 4B**), resulting in $k_{d1} = 0.55 \pm 0.134 \text{ min}^{-1}$ (**Fig. 4C**). This is quite
263 striking because in *E. coli*, co-transcriptional degradation does not seem to occur for
264 genes encoding cytoplasmic proteins due to the membrane localization of RNase E (**Fig.**
265 **2B**).

266 We found that this high k_{d1} is not due to any aspects of the *araB* sequence, because
267 replacing *araB*'s coding region with that of *lacZ* (P_{ara} -*lacZ*; SK477) resulted in similarly
268 high k_{d1} of $0.56 \pm 0.146 \text{ min}^{-1}$ of *lacZ* mRNA, in contrast to the low k_{d1} observed at the
269 native *lac* locus (SK98; **Fig. 4C**). The high k_{d1} of *lacZ* mRNA produced from P_{ara} was not
270 due to the chromosomal position either, because bringing the *lacI-lacZ* region from the
271 native *lac* locus to the *ara* locus (P_{lac} -*lacZ*; SK499) did not change the original (low) k_{d1} of
272 *lacZ* mRNA (SK98; **Fig. 4C**). The high k_{d1} likely originates from what P_{ara} -*araB* (SK472)
273 and P_{ara} -*lacZ* (SK477) have in common: the sequence in 5'-UTR (**Fig. 4A**), which is
274 different from 5'-UTR of native *lacZ* (SK98 and SK499). Therefore, the high k_{d1} of P_{ara}
275 may originate from a certain feature of the 5'-UTR sequence.

276 5'-UTR of an mRNA contains ribosome binding site (RBS), including Shine-Dalgarno
277 (SD) sequence, which governs translation initiation and protein expression level⁵⁸⁻⁶⁰.
278 Henceforth, we refer to the SD sequence and its surrounding sequence as the RBS. RBS
279 sequences are known to affect the energetics of ribosome binding and translation

280 initiation, such that one can quantitatively predict the RBS strength, or protein expression
281 outcome from the sequence⁶¹⁻⁶³. However, weakening RBS strength by changing its
282 sequence has also been known to destabilize the mRNA²⁷⁻³¹, thus reducing the overall
283 protein expression by reducing both translation initiation rate and mRNA lifetime.

284 Conforming to this expectation, *lacZ* transcripts with the native RBS of *araB* (P_{ara} -*lacZ*
285 in SK477) produced 30-fold lower LacZ protein expression than P_{lac} -*lacZ* at the same
286 chromosome location (**Fig. 4D**). This result came from measuring LacZ protein
287 expression by Miller assay. To corroborate this finding, we replaced the RBS sequence
288 in P_{ara} -*lacZ* (SK477) with a strong RBS sequence designed using an RBS calculator⁶³
289 (SK613 in **Fig. 4A**). The synthetic RBS sequence yielded increased LacZ protein
290 expression, higher than that from native *lacZ* RBS (SK499; **Fig. 4D**). Also, *lacZ* mRNA
291 with this strong synthetic RBS sequence exhibited low k_{d1} as observed in the native *lacZ*
292 RBS (SK98 or SK499; **Fig. 4C**). These results support that the hypothesis that the weak
293 RBS sequence in P_{ara} -*araB* is responsible for the high k_{d1} of *araB* mRNA.

294

295 **RBS strength affects *lacZ* mRNA localization due to premature transcription** 296 **termination**

297 We have shown that P_{lac} and P_{ara} , two inducible promoters widely used in gene
298 expression studies⁶⁴⁻⁶⁶, have vastly different RBS strengths. Indeed, RBS sequences and
299 their expected strengths vary widely among genes in the *E. coli* genome^{3,35}. While
300 mRNAs with a weaker RBS are expected to have shorter lifetime^{16,17}, how RBS
301 sequences affect co-transcriptional and post-transcriptional mRNA degradation rates has
302 not been studied. To address this question, we compared the original *lacZ* strain (with the
303 native RBS; SK98) with a weak RBS mutant, which was created by changing five bases
304 in the original SD sequence (**Fig. 5A** and **S4A** for LacZ protein expression). We found
305 that mutating the RBS sequence increases k_{d1} by 15 fold to $0.65 \pm 0.171 \text{ min}^{-1}$ without
306 affecting k_{d2} (**Fig. 5B**). We confirmed that the high k_{d1} is largely controlled by RNase E
307 because the temperature-sensitive RNase E allele (*rne3071*) showed much lower k_{d1} for
308 this weak RBS at the non-permissive temperature in comparison to the wild-type RNase
309 E at the same temperature (**Fig. S4B-S4C**). This brings us to the next question: How does

310 membrane-bound RNase E carry out co-transcriptional degradation of mRNAs with a
311 weak RBS?

312 To test the possibility that nascent mRNAs are localized differently depending on the
313 RBS strength, we visualized 5' *lacZ* mRNAs by FISH (Z5_{FISH}). We reasoned that nascent
314 mRNAs with a weak RBS sequence would be difficult to detect by FISH because they are
315 quickly degraded (**Fig. 5B**). Therefore, we performed FISH in strains harboring the
316 *rne3071* allele to inactivate RNase E (strain SK519 and SK591 for the native and weak
317 RBS sequences, respectively). At the non-permissive temperature, *lacZ* expression was
318 induced with IPTG and re-repressed with glucose at 50 s after the induction. qRT-PCR
319 analysis of RNA samples from this time-course experiment showed that Z3, probing the
320 3' end of the mRNA, appears above the basal level at t = 100 s after induction (**Fig. S4C**),
321 indicating that before t = 100 s, all 5' *lacZ* mRNAs would be nascent and visualized as
322 diffraction-limited foci originating from the gene loci that they are tethered to. Surprisingly,
323 the 2D histogram of the relative positions of Z5_{FISH} in this time window showed different
324 mRNA localization patterns between native RBS and weak RBS strains (**Fig. 5C-5D**).
325 While Z5_{FISH} signals from the native RBS were localized at a specific location with a high
326 probability (red bins in the histogram) as seen earlier in WT RNase E (**Fig. 3G**), Z5_{FISH}
327 signals from the weak RBS were localized at random places throughout the cytoplasm,
328 such that the dense region (red color) did not show up in the histogram (**Fig. 5D**).
329 Additionally, before t = 100 s, the weak RBS strain contained a higher number of Z5_{FISH}
330 spots per cell that have weaker fluorescence intensity in comparison to those in the native
331 RBS strain (**Fig. S4D-S4E**). For example, at t = 60 s, there are up to two *lacZ* gene loci
332 per cell (**Fig. S4F**), but the weak RBS strain had 16% of cells with 3 or more Z5_{FISH} spots
333 per cell, in contrast to 4% observed in the native RBS strain (**Fig. 5E**). These results are
334 consistent with a scenario, in which 5' *lacZ* mRNAs with the weak RBS become physically
335 separated from gene loci even when all of them are expected to be tethered to the gene
336 loci and form only one or two diffraction-limited fluorescence spots per cell (**Fig. S4F**).

337 The spatial dispersion of mRNAs with the weak RBS in the time window i is
338 reminiscent of premature transcription termination, previously shown to follow
339 transcription-translation uncoupling due to nonsense mutation, antibiotic treatment, and
340 amino acid starvation^{33,67-69}. To check the possibility of premature RNAP termination in

341 our weak RBS construct, we examined Z5 and Z3 levels at steady state after induction.
342 In the native RBS strain (SK98), Z5 and Z3 levels were equal at the steady state (**Fig.**
343 **5F**). Considering that Z5 and Z3 have equal lifetimes (**Fig. S1B**), the equal steady state
344 level means that 100% of RNAPs that passed the Z5 probe region reached the Z3 probe
345 region at the end of the gene, i.e., 0% premature transcription termination. However, in
346 the weak RBS *lacZ* strain (SK421), we observed that the steady state level of Z3 is about
347 half of that of Z5, indicating significant premature transcription termination (**Fig. 5G**). We
348 confirmed that this premature transcription termination is controlled by the rho factor
349 because treatment with bicyclomycin (BCM) rescued the Z5 and Z3 difference, bringing
350 the steady-state Z5 and Z3 levels to equal in the weak RBS strain (**Fig. 5H**).

351 These results are consistent with the hypothesis that transcription-translation coupling
352 requires a strong RBS, which allows the loading of the first ribosome to the RBS as soon
353 as the RBS sequence is transcribed by an RNAP. In the case of a weak RBS, in which
354 the first ribosome loading event is delayed, an RNAP might not be coupled with a
355 ribosome and experience premature termination by the rho factor^{68,70} (**Fig. 5I**). Then, the
356 prematurely released (short) transcripts may diffuse to the membrane and get degraded
357 by RNase E on the inner membrane.

358 359 **Translation affects mRNA degradation via premature transcription termination, not** 360 **via ribosome protection**

361 A notable lesson from the weak RBS strain is that there are prematurely released
362 transcripts in the time window t_i , in which we measured k_{d1} assuming all transcripts are
363 nascent. Hence, the high k_{d1} observed in weak RBS strains (including the ones observed
364 with *araB*'s RBS in **Fig. 4C**) likely includes the degradation of prematurely released
365 mRNAs and is not a true rate of co-transcriptional mRNA degradation. To address this
366 problem, we modeled k_{d1} as a weighted average of real co-transcriptional degradation
367 rate of nascent mRNAs (k_{d1^*}) and post-release degradation rate of prematurely terminated
368 mRNAs (k_{dPT}):

$$k_{d1} = k_{d1^*} \cdot (1 - PT) + k_{dPT} \cdot PT \quad (1)$$

369 where PT is the probability of premature termination during transcription.

370 If premature transcription termination leads to high k_{d1} in a weak RBS strain, we expect
371 to see a good correlation between k_{d1} and PT . To test this prediction, we used nine strains
372 harboring *lacZ* with varying RBS sequences at the *ara* and *lac* loci (see **Table S4**). We
373 measured k_{d1} and k_{d2} of *lacZ* mRNAs from re-repression (with glucose addition; **Table S5**)
374 and calculated PT from steady-state levels of Z5 and Z3 after induction (without glucose
375 addition). Because the ratio between steady-state levels of Z5 and Z3 (e.g. **Fig. 5F-5H**) is
376 related to PT as well as k_{dPT} (**Fig. S5A**), we performed iterative fitting of k_{d1} and estimated
377 PT using equation (1) and obtained best PT value for each strain and k_{dPT} common among
378 nine strains (**Fig. 5J**; see **method details**). The optimal fitting of equation (1) gave k_{d1}^* of
379 $0.025 \pm 0.0372 \text{ min}^{-1}$ and k_{dPT} of $0.80 \pm 0.0587 \text{ min}^{-1}$ ($R^2 = 0.93$). We note that k_{d1}^* value
380 is very similar to k_{d1} of strong RBS cases (where premature termination is 0%; e.g. SK98),
381 and k_{dPT} value is larger than most of k_{d2} we have observed for transcripts released after
382 transcription is completed. Possibly, prematurely released transcripts are degraded faster
383 because they diffuse faster than longer, full-length mRNAs and/or because they lack
384 certain features at the 3' end that full-length mRNAs have, such as a stem-loop structure,
385 making them more easily degraded, by 3'-to-5' exonuclease, PNPase.

386 One of the models explaining the RBS effect on mRNA lifetime is based on the notion
387 that ribosomes protect mRNA from the attack of RNase E^{16,17}. According to this model,
388 transcripts with a weak RBS sequence, or those showing high probability of premature
389 transcription termination (PT), would undergo fast degradation because there are fewer
390 ribosomes on the mRNAs. To test this model across different RBS sequences, we
391 examined k_{d2} , the decay rate of Z5 after $t_{3'}$ (last RNAP passes the end of *lacZ* gene) in
392 time window iii. k_{d2} is largely determined by the degradation rate of full-length transcripts
393 that have the 3' sequence and not affected by prematurely released transcripts, which
394 are degraded rather quickly (k_{dPT}) and minimally contribute to the Z5 signal in this time
395 window. Nine strains of varying RBS sequences showed that k_{d2} is independent of PT
396 (**Fig. 5K**) with very little correlation ($P = -0.078$). This result is in contrast to what would
397 be expected from the ribosome protection model, which would expect higher k_{d2} in
398 transcripts with weaker RBS, or higher probability of premature transcription termination,
399 because the transcripts carry fewer ribosomes on average. Therefore, our results suggest

400 that translation affects mRNA lifetime mainly by affecting the percentage of prematurely
401 released transcripts (**Fig. 5J**).

402
403 **Premature transcription termination and subcellular localization of RNase E (or its**
404 **homolog) affect the degradation of *lacZ* mRNA in other bacterial species**

405 Our results so far imply that in *E. coli*, the fate of mRNA is determined by the RBS
406 sequence because of its effect on transcription-translation coupling. Next, we examined
407 if this conclusion can be generalized to other bacterial species. For example, in *B. subtilis*,
408 RNAP was shown to translocate faster than the ribosome during expression of *lacZ*,
409 preventing the ribosome from coupling to RNAPs^{34,71}. We tested if the transcription-
410 translation uncoupling in *B. subtilis* results in premature transcription termination and
411 potentially a high k_{d1} . First, we repeated the experiment done by previous papers
412 measuring the transcription and translation times of *lacZ* by qRT-PCR and Miller assay in
413 *B. subtilis* (strain GLB503; **Fig. 6A**), respectively. The transcription time was acquired
414 from the initial increase of Z3 signal from the baseline after induction with IPTG, indicating
415 the moment first RNAPs reach the end of the gene. The translation time was acquired
416 from the initial increase of LacZ protein levels from the baseline after induction, indicating
417 the moment first ribosomes reach the end of *lacZ* mRNA. In a slow growth condition³⁴,
418 we observed that the translation time was 2.6 ± 0.054 min, much longer than the
419 transcription time of 1.3 ± 0.56 min (**Fig. S6A-S6C**). The steady-state levels of Z5 and Z3
420 were similar (**Fig. 6B**), implying that premature transcription termination does not take
421 place even if transcription and translation are uncoupled in *B. subtilis*.

422 Next, we measured k_{d1} and k_{d2} of *lacZ* mRNAs by re-repressing transcription by
423 adding rifampicin, a drug that stops transcription initiation⁷²⁻⁷⁴, at $t = 30$ s after induction
424 (**Fig. 6C**). We obtained k_{d1} of 0.025 ± 0.0036 min⁻¹ and k_{d2} of 0.14 ± 0.026 min⁻¹ (**Fig.**
425 **S6D**). Since premature transcription termination is not observed (**Fig. 6B**), k_{d1} can be
426 attributed to co-transcription degradation, and the lifetime of nascent mRNA can be
427 estimated as 40 min ($1/k_{d1}$), much longer than the transcription time of 1.3 min. Hence,
428 co-transcriptional degradation of *lacZ* mRNAs is likely very rare in *B. subtilis*, like in *E.*
429 *coli*. The lack of co-transcriptional degradation can be explained by the membrane
430 localization of the main endoribonuclease performing mRNA degradation, RNase Y and

431 RNase E, in *B. subtilis* and *E. coli*, respectively^{18,75,76}. We note that k_{d2} in *B. subtilis* is low
432 relative to *E. coli* (see **Fig. 5K**), and *lacZ* mRNA levels do not return to the basal level
433 within 10 min (**Fig. 6C**). This is quite surprising because the amount of LacZ proteins
434 expressed from the 30-s induction was minimal according to the Miller assay using a
435 sensitive fluorogenic LacZ substrate (**Fig. S6E**), suggesting that the remaining *lacZ*
436 mRNAs do not support protein synthesis. Possibly, translation initiation is slow in this
437 strain, and/or functional inactivation of mRNAs precedes the chemical degradation of
438 mRNAs in *B. subtilis*.

439 In contrast to *E. coli* and *B. subtilis*, *C. crescentus* is known to have cytoplasmic RNase
440 E²⁴⁻²⁶. Hence, we investigated the possibility of co-transcriptional degradation of *lacZ*
441 mRNA in *C. crescentus* using a strain where *lacZ* is placed under the xylose-inducible
442 promoter in the chromosome (strain LS2370⁷⁷; **Fig. 6D**). First, we measured the
443 transcription and translation times of *lacZ* by qRT-PCR and by Miller assay after induction
444 with xylose. The translation time was 2.5 ± 0.21 min (**Fig. 6E** and **S6F**), similar to the
445 transcription time of 2.3 ± 0.27 min (**Fig. 6F**), suggesting that transcription and translation
446 are coupled. To measure k_{d1} and k_{d2} of *lacZ* mRNAs, transcription was re-repressed with
447 rifampicin at $t = 50$ s after addition of xylose. qRT-PCR data show that Z5 decays very
448 quickly after rifampicin addition (**Fig. 6G**). Strikingly, Z3 does not increase above the basal
449 level, such that the time windows i and iii cannot be defined for fitting Z5 for k_{d1} and k_{d2} .
450 If we take $T_{3'}$ of 2.3 min from the induction-only experiment (**Fig. 6F**) to estimate the time
451 window i (blue box in **Fig. 6G**), we obtain k_{d1} of 1.3 ± 0.13 min⁻¹. The absence of 3' *lacZ*
452 mRNA signals in the re-repression experiment (also minimal LacZ protein expression;
453 **Fig. S6G**) indicates significant premature transcription termination, which contributes to
454 high k_{d1} .

455 Indeed, when we blocked the rho factor activity with BCM, the steady-state levels of
456 Z5 and Z3 increased, suggesting that rho-dependent premature termination affected *lacZ*
457 mRNA levels in non-treated cells (**Fig. S6H** vs. **6F**). Repeating k_{d1} measurement in BCM-
458 treated cells allowed us to obtain the true rate of co-transcriptional degradation, k_{d1}^* of
459 0.71 ± 0.093 min⁻¹ (**Fig. S6I**). Through mathematical modeling, we estimated that
460 prematurely terminated mRNAs are degraded at k_{dPT} of 3.4 ± 0.61 min⁻¹ and the
461 probability of premature transcription termination (*PT*) to be $69 \pm 4.4\%$ (see **method**

462 **details**). The fast mRNA degradation likely originates from the cytoplasmic distribution of
463 RNase E in *C. crescentus* cells. Interestingly, RNase E in *C. crescentus* has been shown
464 to interact with the rho factor⁷⁸. We speculate that the cooperation between the rho factor
465 and RNase E results in the high k_{d1} and high probability of premature transcription
466 termination we observed in *C. crescentus* (**Fig. 6G**).

467 The high probability of premature transcription termination (~69%) agrees with the
468 absence of 3' *lacZ* mRNA signal when *lacZ* transcription was induced only for 50 s (**Fig.**
469 **6G**). However, it seems incompatible with transcription-translation coupling concluded
470 based on the synchronized transcription and translation times (**Fig. 6E-6F**). We note that
471 the transcription and translation times were determined by the first (fastest) RNAPs and
472 ribosomes arriving at the 3' end, and they can be the same even though only a small
473 fraction of RNAPs are coupled to a ribosome. Hence, it is likely that a significant fraction
474 of RNAPs is uncoupled with a ribosome during the transcription of *lacZ* in *C. crescentus*
475 and experiences premature transcription termination.

476

477 Discussion

478 Our findings have implications for gene regulation based on when and where mRNAs are
479 degraded within a bacterial cell. In bacteria, such as *E. coli* and *B. subtilis*, where major
480 ribonuclease and RNA degradosome are localized to the membrane, co-transcriptional
481 mRNA degradation is likely negligible for most genes, and mRNA degradation takes place
482 exclusively on the membrane once mRNAs are released from the gene loci (**Fig. 7A-7B**).
483 The lack of co-transcriptional degradation would be advantageous when more proteins
484 need to be made per transcripts.

485 Our data showing co-transcriptional degradation of *lacY* mRNA suggests an exception
486 to this rule for genes encoding inner membrane proteins (**Fig. 7C**). We note that the high
487 k_{d1} of 5' *lacY* mRNA measured in *lacY2-lacZ-venus* (SK575; **Fig. 3I**) and in full *lacY* mRNA
488 (SK564; **Fig. S3E**) likely reflects genuine co-transcriptional degradation, without
489 premature transcription termination because (1) the native (strong) *lacZ* RBS was used
490 and (2) full *lacY* transcript (SK564) showed 0% premature transcription termination (**Fig.**
491 **S3F**). In terms of the mechanism, membrane localization of nascent mRNAs may not be
492 the only reason that *lacY* mRNA is degraded co-transcriptionally. The *lacZ* sequence

493 within *lacY2-lacZ-venus* mRNA had similar k_{d1} and k_{d2} as those of the *lacZ*-alone case
494 (SK98) even though their localization (or the proximity to the membrane) were vastly
495 different (**Fig. 3H**). Hence, we speculate that there are additional features in the *lacY2*
496 sequence (i.e. the first two transmembrane segments) that promote its co-transcriptional
497 mRNA degradation. For example, the signal recognition particle (SRP) and SecYEG,
498 proteins involved in transertion of LacY^{79,80}, might interact with RNase E to promote the
499 degradation of *lacY2* sequence. Although this idea remains to be tested, such a
500 mechanism can also explain earlier results that translational fusion of a SRP signal
501 peptide to a random gene decreases the transcript's lifetime⁸¹ and that fast degradation
502 of *ptsG* mRNA encoding transmembrane glucose transporter (IIBC^{glc}) requires the
503 transmembrane segment of its protein⁸². Also, this hypothesis predicts that the rate of co-
504 transcriptional mRNA degradation of *lacY* would decrease when RNase E is localized in
505 the cytoplasm. Indeed, previous genome-wide characterization of mRNA lifetimes in the
506 RNase E Δ MTS strain showed that many genes encoding inner membrane proteins are
507 preferentially stabilized in this cytoplasmic RNase E mutant^{81,83}. These results suggest
508 that membrane localization of RNase E is important for differential regulation of
509 membrane protein expression in comparison to cytoplasmic proteins. Since membrane
510 surface area is limited (more than the cytoplasmic volume)⁸⁴ and since membrane
511 channel proteins (such as LacY) have a higher activity cost when expressed^{84,85}, tight
512 regulation of membrane protein expression, by employing co-transcriptional degradation
513 mechanism, is likely beneficial for cellular fitness.

514 Another important determinant of mRNA degradation is the timing that transcripts are
515 released from the gene. This timing can vary depending on the gene length and RNAP
516 speed. Also, in the case of polycistronic genes, mRNA processing in the intergenic
517 region⁴⁰⁻⁴² can release the promoter proximal gene first while promoter distal gene is
518 being transcribed. Adding to this list, our work highlights the important role played by RBS
519 sequences in permitting premature release of incomplete transcripts (**Fig. 5J**).

520 Based on our model (**Fig. 1A**), the mean mRNA lifetime in the steady state is affected
521 by the degradation rates of nascent (k_{d1^*}), fully-transcribed (k_{d2}), and prematurely-
522 released (k_{dPT}) transcripts because these three types of mRNA can have distinct
523 degradation rates. If ribosomes indeed protect mRNA from degradation^{16,17}, each of these

524 rates may increase with lower RBS strength. However, our data suggests that these rates
525 do not vary much among the RBS mutant strains we examined; instead, the portion of
526 prematurely released transcripts varies significantly (**Fig. 5J-5K**), eventually yielding
527 different protein output for different RBS sequences (**Fig. S5B-S5E**). Considering that
528 premature transcription termination is a hallmark event in the absence of transcription-
529 translation coupling^{86,87} (**Fig. 5I**), we identified RBS sequences as the key genetic feature
530 that can modulate the probability of transcription-translation coupling and subsequently
531 mean mRNA lifetimes across the genome.

532 The RBS sequences we tested cover a wide range of translation initiation strengths
533 observed in the genome⁸⁸ (**Fig. S5G-S5H**). If we compare the maximum translation
534 initiation strength we observed non-zero percentage of premature transcription
535 termination (strain SK420 and SK518 in **Fig. S5G**) with endogenous translation initiation
536 strengths across the *E. coli* genome⁸⁸, we estimate that at least 58% of all genes may
537 experience some percentage of premature transcription termination due to compromised
538 transcription-translation coupling (**Fig. 5I** and **S5H**). This estimation is consistent with the
539 high percentage of 3'-end mRNAs detected at the 5' UTRs and inside of genes in a recent
540 *E. coli* transcriptome analysis⁸⁹. Collectively, these results support that transcription-
541 translation uncoupling arising from low translation initiation rate and the resulting
542 premature transcription termination are likely common across the *E. coli* genome.

543 T7 transcription systems in *E. coli*, often used for bioengineering and synthetic biology
544 field⁹⁰, are known to experience transcription-translation uncoupling because T7 RNAP
545 outpaces the host ribosome (8-fold speed difference⁹¹), yet T7 RNAP does not
546 prematurely terminate⁹². Based on our model of mRNA degradation in *E. coli*, we predict
547 that transcripts made by T7 RNAPs are degraded once transcription is completed, as
548 opposed to experiencing co-transcriptional degradation as proposed previously⁴⁶.

549 We found that gene expression in *B. subtilis* is analogous to the T7 system, such that
550 premature transcription termination is negligible even though RNAP and ribosome are
551 uncoupled (**Fig. 6B**). We note that a recent study showed that *B. subtilis* RNAPs can
552 prematurely dissociate from DNA during transcription of *lacZ* in a rho-independent
553 manner, especially when their speed is slow⁷¹. Hence, premature transcription
554 termination may be possible under certain conditions and in certain genes that are under

555 the control of riboswitches and attenuators⁹³ and help down-regulate protein expression
556 in *B. subtilis*.

557 In bacteria, such as *C. crescentus*, where major ribonuclease and RNA degradosome
558 are located in the cytoplasm, mRNA degradation may start during transcription (**Fig. 7D**).
559 We observed in *C. crescentus*, high rate of co-transcriptional degradation rate (k_{d1^*}) for
560 *lacZ* mRNA and significant premature transcription termination ($PT = 69\%$). This high
561 premature transcription termination suggests that many RNAPs transcribing *lacZ* were
562 not coupled to a ribosome and points out that the equality between transcription and
563 translation times (**Fig. 6E-6F**) may not be a good indicator for the percentage of
564 transcription-translation coupling. Together with the fact that the rho factor physically
565 interacts with RNase E in *C. crescentus*⁷⁸, our results imply that transcription, premature
566 transcription termination, and mRNA degradation are highly coupled in the cytoplasm of
567 *C. crescentus*.

568 In conclusion, our work overall identifies subcellular localization of RNase E (or its
569 homologue) and premature transcription termination in the absence of transcription-
570 translation coupling (arising from weak RBS sequences) as spatial and genetic design
571 principles by which bacteria have evolved to differentially regulate transcriptional and
572 translational coupling to mRNA degradation across genes and species. These principles
573 will serve the basis for quantitative modeling of protein expression levels across the
574 genome^{24,81,94} and for comprehending the subcellular localization patterns of mRNAs
575 found for different genes and in different bacteria species^{24,81,94}. In the future, it would be
576 interesting to investigate whether our findings are relevant to the coordination of
577 transcription, translation, and mRNA degradation in other contexts where there is a lack
578 of membrane-bound microcompartments, such as archaea⁹⁵, chloroplast³⁶, and
579 mitochondria⁹⁶.

580

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591

592 **Author contributions**

593 Conceptualization, Se.K. and Sa.K.; Methodology and investigation, Se.K., Y.W., A.H.,
594 and Sa.K.; Writing and visualization, Se.K., Y.W., A.H., and Sa.K.; Supervision and
595 funding acquisition, Sa.K.

596

597 **Figure Legends**

598 **Figure 1** Two-phase mRNA degradation in bacteria. **(A)** Definition. mRNA degradation
599 can occur during transcription (on nascent transcripts) and after transcription (on released
600 transcripts) with different rates k_{d1} and k_{d2} , respectively. **(B)** Schematics of the time-
601 course assay, in which *lacZ* transcription is pulse-induced and its transcripts are
602 quantified with qRT-PCR primers amplifying the 530-660 nucleotide (nt) region (Z5) and
603 2732-2890 nt region (Z3) of *lacZ* (gene length = 3072 nt). The first and last RNAPs pass
604 the Z5 probe site at T_5' and t_5' , respectively, and they pass the Z3 probe site at T_3' and t_3' ,
605 respectively. Blue and yellow shaded boxes indicate the time when k_{d1} and k_{d2} can be
606 measured by an exponential decay fit, respectively. **(C)** Anticipated result when mRNA
607 degradation takes place with $k_{d1} = 0.18 \text{ min}^{-1}$ and $k_{d2} = 0.42 \text{ min}^{-1}$. **(D)** Time course data
608 of 5' and 3' *lacZ* mRNA (Z5 and Z3) after induction with 0.2 mM IPTG at $t = 0$ and re-
609 repression with 500 mM glucose at $t = 75 \text{ s}$ (strain SK98, grown in M9 glycerol at 30°C).
610 Error bars represent the standard deviation from three biological replicates.

611

612 **Figure 2** Effect of RNase E localization and its C-terminal domain on k_{d1} and k_{d2} of *lacZ*
613 mRNA. **(A)** Schematic description of wild-type RNase E and mutants forming different
614 RNA degradosome complexes. The wild-type RNase E interacts with PNPase (green),
615 Enolase (yellow), and RhlB (purple) to form the RNA degradosome. Not drawn to scale.
616 **(B)** k_{d1} and k_{d2} of *lacZ* mRNA in RNase E localization mutant strains (strain SK98, SK339,

617 SK370, and SK369). Transcription of *lacZ* was induced with 0.2 mM IPTG at $t = 0$ and re-
618 repressed with 500 mM glucose at $t = 75$ s. Error bars represent the standard deviation
619 from three biological replicates. ** and * indicate $p < 0.01$ and $p < 0.05$, respectively (two-
620 sample t test).

621
622 **Figure 3** Effect of transertion on k_{d1} and k_{d2} of *lacZ* mRNA. **(A)** Localization of 5' *lacZ*
623 mRNA in the presence of active transcription of *lacY* or *aadA* from a constitutive promoter
624 P_{con} . An example FISH image of 5' *lacZ* mRNA when transcription was induced with 0.2
625 mM IPTG and re-repressed with 500 mM glucose at 75 s after induction. The example
626 image is from cells taken at $t = 1$ min. Scale bar = 1 μ m. **(B-C)** 2D histogram of $Z5_{FISH}$
627 localization at different time points along the time-course assay. Colors denote the
628 probability of finding $Z5_{FISH}$ in a certain bin location. To minimize noise, the normalized
629 positions of foci along the cell long and short axes were calculated in the first quartile and
630 extended to the other three quartiles using mirror symmetry. The bin size is 70-80 nm.
631 The white ovals are cell outlines, and the white lines are the axes of symmetry. For each
632 histogram, over 5,000 foci were analyzed. The same color scale was used for both
633 histograms. **(D)** k_{d1} and k_{d2} of *lacZ* mRNA with different downstream genes: *lacY* (SK390),
634 *aadA* (SK435), or none (SK98). **(E)** A pair of strains to study the direct transertion effect
635 on *lacZ* mRNA degradation kinetics. **(F-G)** 2D histogram of $Z5_{FISH}$ localization in *lacY2*-
636 *lacZ-venus* (SK575, **F**) and *lacZ* (SK98, **G**). Transcription of *lacZ* was induced and re-
637 repressed the same way as described in panel A. For each histogram, over 2,000 mRNA
638 foci were analyzed. Except only 564 foci were used for SK98 $t = 2$ min. **(H)** k_{d1} and k_{d2} of
639 *lacZ* mRNA in SK575 and SK98. **(I)** Relative mRNA level of the *lacY2* sequence in SK575
640 measured by qRT-PCR during the time course assay described in panel A. *lacY2* is
641 probed by primers amplifying 80-222 nt region of *lacY2* sequence. Blue and yellow
642 shaded boxes indicate the time windows i and iii for k_{d1} and k_{d2} of *lacY2*, respectively. In
643 all panels, error bars represent the standard deviation from three biological replicates.
644 Also, ns indicates a statistically nonsignificant difference (two-sample t test).

645
646 **Figure 4** Degradation kinetics of *araB* mRNA and the effect of RBS sequences on k_{d1} .
647 **(A)** Design of strains used in this figure. 5'-UTR sequences from the first base (+1) of the

648 transcript to the start codon (atg) are shown. SD elements estimated by an RBS
649 calculator⁶³ are underlined. We note that *lacZ* in the original *lac* locus was deleted when
650 *lacZ* was placed in the *ara* locus. **(B)** *araB* mRNA level change from induction with 0.2%
651 arabinose at $t = 0$ and re-repression with 500 mM glucose at $t = 50$ s. 5' *araB* and 3' *araB*
652 were probed by qRT-PCR primers amplifying 33-210 nt and 1536-1616 nt regions of *araB*.
653 Blue and yellow boxes denote the time windows where k_{d1} and k_{d2} are measured. **(C)** k_{d1}
654 measured in strains shown in panel A. P_{ara} and P^*_{ara} were induced with 0.2% arabinose,
655 and P_{lac} was induced with 0.2 mM IPTG. 500 mM glucose was added at $t = 50$ s (for *araB*)
656 or 75 s (for *lacZ*) to turn off the promoter. **(D)** LacZ protein expression measured by Miller
657 assay. LacZ expression was induced and re-repressed the same way as in the qRT-PCR
658 experiment (panel C), and the total LacZ protein produced from the pulsed induction were
659 calculated from each strain. In all panels, error bars indicate the standard deviations from
660 three or more biological replicates (except D, from two replicates). ***, **, and * indicate
661 $p < 0.001$, 0.01, and 0.05, respectively, and ns indicates a statistically nonsignificant
662 difference (two-sample t test).

663
664 **Figure 5** Origin of fast k_{d1} observed in *lacZ* mRNA with a weak RBS. **(A)** 5' UTR
665 sequences of native *lacZ* and a weak RBS mutant. mRNA sequences from the first base
666 of the transcript to the start codon (atg) are shown. SD sequences estimated by an RBS
667 calculator⁶³ are underlined. **(B)** k_{d1} and k_{d2} of *lacZ* mRNA measured by induction with 0.2
668 mM IPTG at $t = 0$ and re-repression with 500 mM glucose at $t = 75$ s. Error bars represent
669 the standard deviation from three biological replicates. *** denotes $p < 0.001$, and ns
670 indicates a statistically nonsignificant difference (two-sample t test). **(C-D)** 2D histogram
671 of $Z5_{FISH}$ localization depending on the RBS sequence. After shifting the temperature to
672 43.5°C for 10 min, *lacZ* expression was induced with 0.2 mM IPTG at $t = 0$ and re-
673 repressed with 500 mM glucose at $t = 50$ s. In each case, over 25,000 mRNA foci were
674 analyzed. **(E)** Number of fluorescent $Z5_{FISH}$ spots detected per cell at $t = 60$ s during the
675 time-course experiment described in panel C-D. Error bars represent the standard error
676 from bootstrapping. **(F-H)** $Z5$ and $Z3$ levels after *lacZ* transcription was induced with 0.2
677 mM IPTG at $t = 0$. In (H), 100 $\mu\text{g}/\text{mL}$ BCM was added 5 min before IPTG addition. Error
678 bars represent the standard deviation from two biological replicates. **(I)** Effect of RBS

679 strength on the fate of mRNA. **(J-K)** Relationship between k_{d1} or k_{d2} and the probability
680 of premature transcription termination in various RBS-*lacZ* mRNAs and $P_{ara-araB}$ mRNA.
681 See **Table S4** for the list of strains used. The line fit is based on the equation (1). Error
682 bars for k_{d1} or k_{d2} represent the standard deviation from three replicates and those for *PT*
683 were calculated from the steady-state ratio of Z5 and Z3 in two replicates.

684
685 **Figure 6** Degradation kinetics of *lacZ* mRNA in *B. subtilis* and *C. crescentus*. **(A)** IPTG-
686 inducible *lacZ* in the chromosome of *B. subtilis*. For qRT-PCR, we used the same Z5 and
687 Z3 primers used in *E. coli lacZ*. **(B-C)** Z5 and Z3 levels after induction with 5 mM IPTG at
688 $t = 0$, probed by qRT-PCR. To measure *lacZ* mRNA degradation rates in **(C)**, transcription
689 was re-repressed with 200 $\mu\text{g}/\text{mL}$ rifampicin at $t = 30$ s. The time windows used for k_{d1}
690 and k_{d2} fitting are indicated as blue and yellow boxes. *B. subtilis* cells were grown in
691 MOPS media supplemented with maltose at 30°C. Error bars represent the standard
692 deviation from two (B) or three (C) biological replicates. **(D)** Xylose-inducible *lacZ* in *C.*
693 *crescentus*. For qRT-PCR, we used the same Z5 and Z3 primers used in *E. coli lacZ*. **(E)**
694 Translation kinetics of LacZ protein expression in *C. crescentus* after adding 0.3% xylose,
695 probed by Miller assay using MUG (3-methylumbelliferyl-beta-D-galactopyranoside) as a
696 sensitive LacZ substrate. Error bars represent the standard deviation from three biological
697 replicates. **(F-G)** Z5 and Z3 levels after induction with 0.3% xylose at $t = 0$, probed by
698 qRT-PCR. To measure *lacZ* mRNA degradation rates in **(G)**, transcription was re-
699 repressed with 200 $\mu\text{g}/\text{mL}$ rifampicin at $t = 50$ s. The time window used for on k_{d1} fitting
700 for Z5 is indicated as blue box. *C. crescentus* cells were grown in M2G at 28°C. Error bars
701 represent the standard deviation from five (F) or three (G) biological replicates.

702
703 **Figure 7** Generalizable model of mRNA degradation in bacteria. **(A-C)** Scenarios in *E.*
704 *coli* (and possibly other bacterial species having the main ribonuclease on the membrane)
705 for genes encoding cytoplasmic proteins with strong RBS **(A)** and weak RBS **(B)** and for
706 genes encoding inner membrane proteins **(C)**. **(D)** A scenario in *C. crescentus* and
707 possibly other bacterial species having the main ribonuclease in the cytoplasm. The
708 cartoon is drawn to reflect that nucleoid takes a large area of the cytoplasm in *C.*
709 *crescentus*⁹⁷.

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981

Figure 1

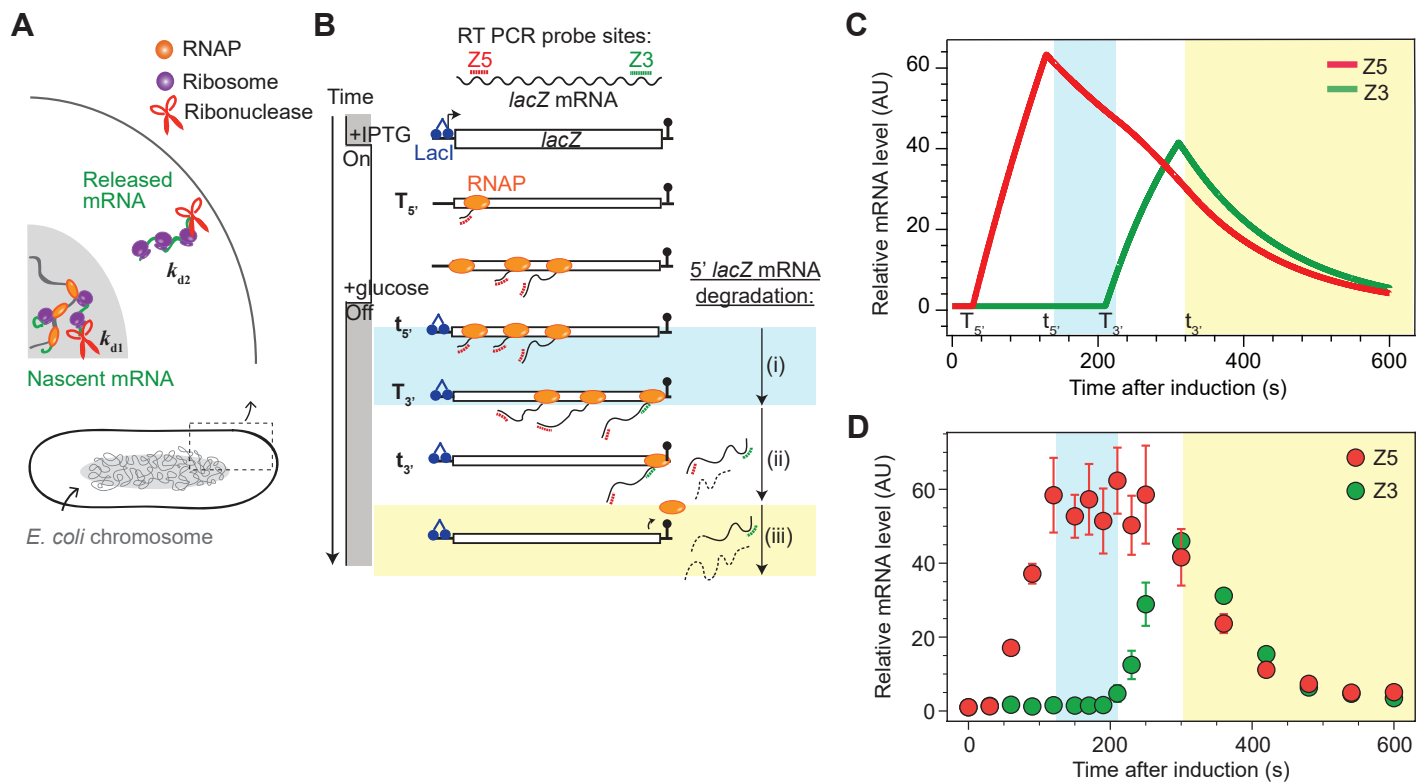


Figure 2

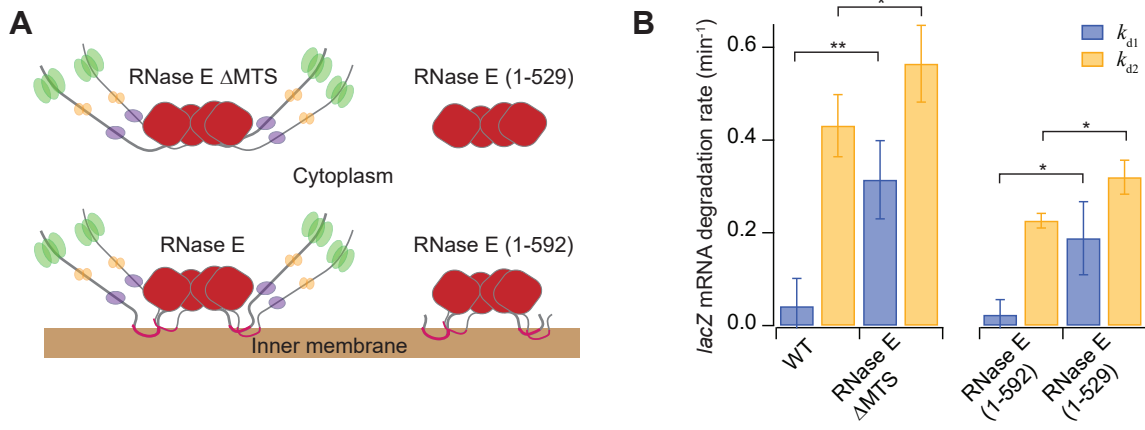


Figure 3

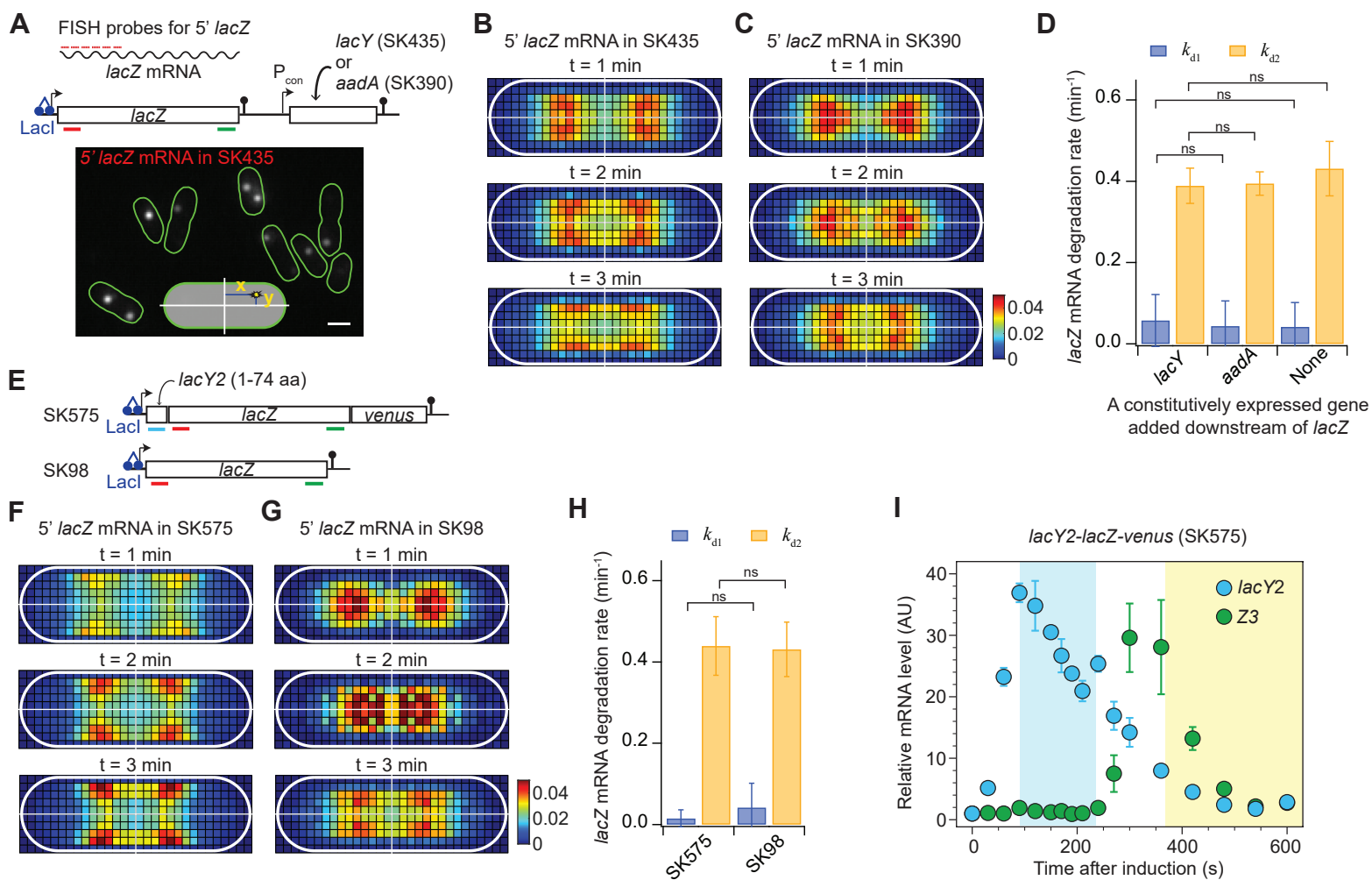


Figure 4

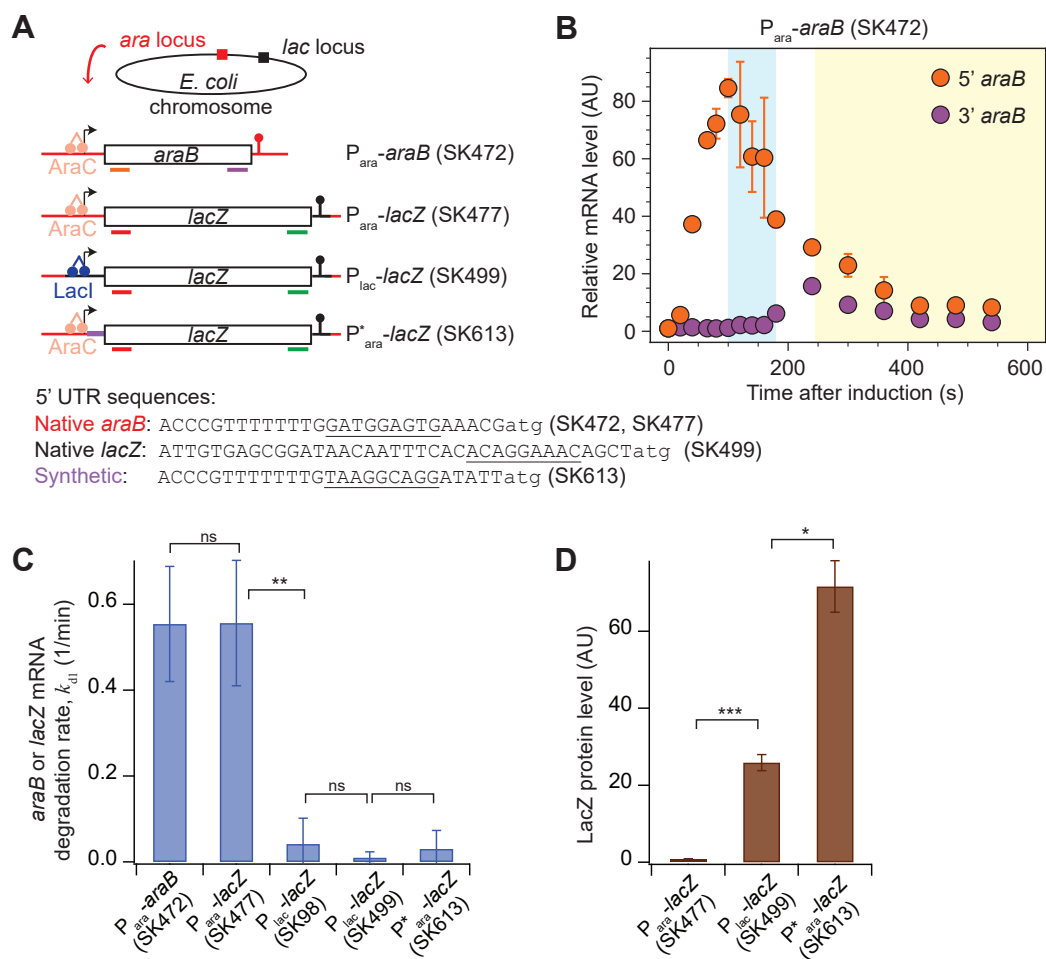


Figure 5

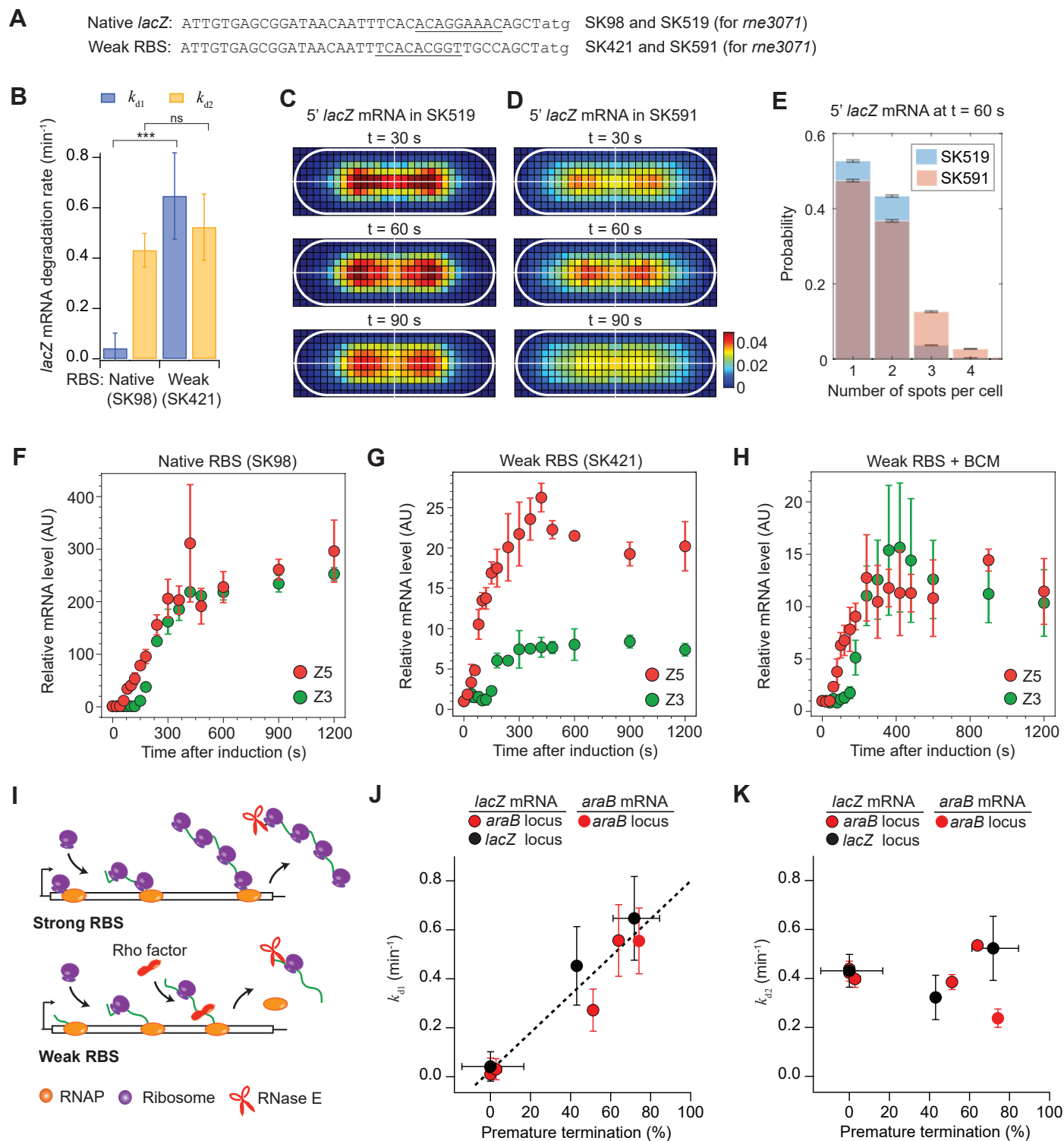


Figure 6

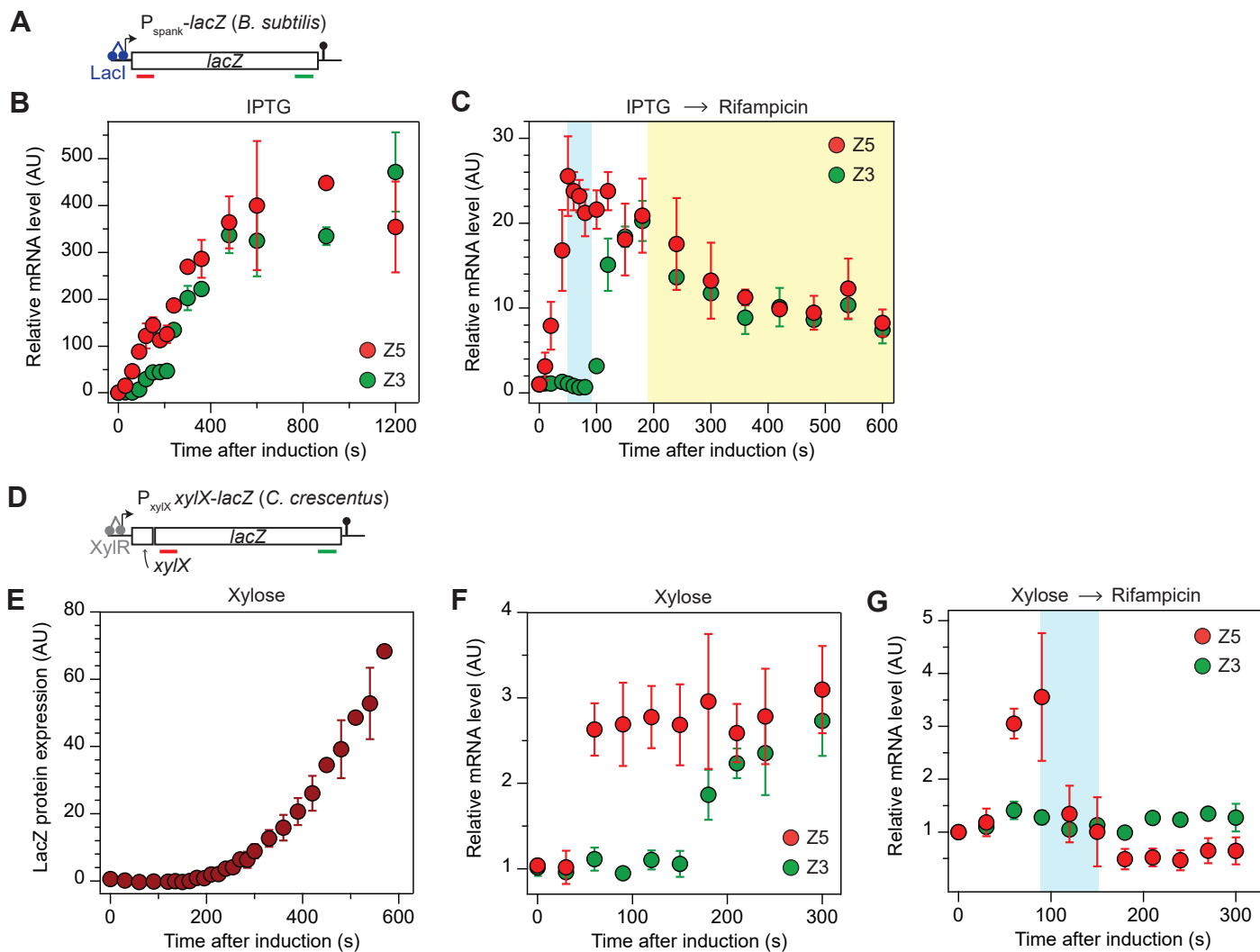


Figure 7

