The Small RNA CyaR Activates Translation of the Outer Membrane Haem Receptor chuA in Enterohemorrhagic Escherichia coli

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To sense the transition from environment to host, bacteria use a range of environmental cues to control expression of virulence genes. Iron is tightly sequestered in host tissues and in the human pathogen enterohemorrhagic Escherichia coli (EHEC) iron-limitation induces transcription of the outer membrane haem transporter encoded by chuAS. ChuA expression is post-transcriptionally activated at 37°C by a FourU RNA thermometer ensuring that the haem receptor is only expressed under low iron, high temperature conditions that indicate the host. Here we demonstrate that expression of chuA is also independently regulated by the cAMP-responsive small RNA (sRNA) CyaR and transcriptional terminator Rho. These results indicate that chuAS expression is regulated at the transcription initiation, transcript elongation, and translational level.

We speculate that additional sensing of the gluconeogenic environment allows further precision in determining when EHEC is at the gastrointestinal epithelium of the host. With previous studies, it appears that the chuAS transcript is controlled by eight regulatory inputs that control expression through six different transcriptional and post-transcriptional mechanisms. The results highlight the ability of regulatory sRNAs to integrate multiple environmental signals into a layered hierarchy of signal input.

Keywords: small RNA, Hfq, heme, iron, post-transcriptional, non-coding RNA, EHEC

INTRODUCTION

During infection, nutrients required by pathogenic bacteria are withheld by the host in a process known as nutritional immunity (Weinberg, 1975; Hood and Skaar, 2012). An example can be found in iron, a transition metal that is essential for pathogenic bacteria due to its role in essential physiological processes, such as respiration (Skaar, 2010; Evstatiev and Gasche, 2012). To deprive pathogens of this vital nutrient, the host sequesters iron in the high affinity iron binding molecules haem, ferritin and transferrin (Barber and Elde, 2014; Cornelissen, 2018). These ensure that the concentration of extracellular free iron in the host is too low to support bacterial growth and infection. To overcome this nutritional deficit, pathogens such as
Haemophilus influenzae, Campylobacter jejuni, Pseudomonas aeruginosa, and uropathogenic Escherichia coli have evolved multiple systems that allow for the scavenging of iron, these include siderophores and haem uptake systems (Rossi et al., 2001; Torres et al., 2001; Ong et al., 2006; Ridley et al., 2006; Hagan and Mobley, 2009; Skaar, 2010; Fournier et al., 2011; Garcia et al., 2011; Morgenthau et al., 2013; Zambolin et al., 2016).

Enterohemorrhagic E. coli (EHEC) is an enteric pathogen that causes hemorrhagic colitis which can progress into potentially fatal hemorrhagic uremic syndrome. The primary reservoir of this pathogen are ruminants and transmission mainly occurs via the consumption of contaminated foodstuffs (Kaper et al., 2004; Gyles, 2007; Karpman and Ståhl, 2014). EHEC can express an outer membrane haem receptor from the iron regulated chu haem uptake operon (Torres and Payne, 1997). The chu locus is homologous to the shu locus found in Shigella, and contains genes that allow for haem uptake, transport, utilization and degradation. The locus consists of a bicistronic operon (chuAS) as well as two polycistronic operons (chuTWXY and chuUV). ChuA is a TonB-dependent outer membrane haem receptor that imports haem into the periplasm (Celia et al., 2016). The periplasmic binding protein ChuT binds to haem, and via the ABC transporter ChuUV, is shuttled into the cytoplasm. Haem is processed by the haem oxidase ChuS that converts haem into biliverdin, carbon monoxide and free iron. This breakdown also prevents haem toxicity (Anzaldi and Skaar, 2010). Under anaerobic conditions, this role is taken up by the SAM-methyltransferase ChuW and the anaerobilin reductase ChuY (LaMattina et al., 2016, 2017).

The haem uptake operon is part of a larger regulatory network that allows EHEC to achieve iron homeostasis. Transcription of chuA is repressed by the iron-dependent transcriptional regulator Fur. In iron-rich conditions, Fur binds to iron which increases the affinity of Fur for its DNA-binding site ~1000-fold (Bags and Neilands, 1987). Fur also regulates a Hfq-dependent small RNA (sRNA) RyhB that is central to maintaining iron homeostasis (Massé and Gottesman, 2002; Massé et al., 2007; Waters and Storz, 2009; Wagner and Romby, 2015). RyhB prevents the expression of non-essential proteins that require Fe as a cofactor such as the TCA cycle and respiration genes sdhC and fumAC, and iron storage genes such as bacterioferritin. Genes that contribute to iron uptake, such as the shikimate permease shiA and the colicin I receptor cirA, are positively regulated by this sRNA (Massé and Gottesman, 2002; Ongley-Sherrouse and Murphy, 2013; Chargey and Mandin, 2018).

Enterohemorrhagic Escherichia coli utilizes the Chu system when it is in an iron-poor, haem-rich host. For that reason, as well as the toxicity that accompanies haem and iron over-accumulation, it is necessary to precisely control expression of this operon. FourU RNA-thermometers are used in different bacterial pathogens, such as Shigella dysenteriae, Yersinia pseudotuberculosis, Vibrio cholerae, and Salmonella typhimurium, to regulate expression of their virulence factors in a temperature-dependent manner (Waldminghaus et al., 2007; Kouse et al., 2013; Weber et al., 2014; Righetti et al., 2016). A FourU RNA-thermometer was found in the chuA 5'UTR that blocks the ribosomal binding site (RBS) and is formed at temperatures below 37°C, indicating that the pathogen is outside of the host (Kouse et al., 2013). This limits chuA expression outside of the host where haem is unlikely to be available.

Identification of binding sites for the small RNA chaperone Hfq in EHEC identified the phage-encoded sRNA Assr that positively regulates the haem oxygenase chuS by sponging interactions with the negatively regulating sRNA, FnrR (Tree et al., 2014). Further, analysis of the EHEC sRNA interactome revealed that the outer membrane haem receptor chuA was also negatively regulated by the EHEC-specific Hfq-binding sRNA Esr41 (Waters et al., 2017). Here, we show that translation of chuA is activated by the Crp-cAMP regulated sRNA CyaR in a temperature- and Rho-termination-independent manner. Our results suggest that EHEC employs an AND-OR logic gate that integrates information on iron availability, temperature and carbon availability to sense its location within the host for appropriate expression of chuA.

RESULTS

The Outer Membrane Haem Receptor chuA Is Regulated by CyaR and ChiX

In previous work the binding sites of the small RNA chaperone Hfq were mapped using UV-crosslinking, denaturing purification and sequencing of Hfq-crosslinked RNA (termed CRAC) (Tree et al., 2014). Deletions introduced at the site of protein-RNA crosslinking during reverse transcription of UV-crosslinked RNAs can be used to identify the site of direct Hfq-RNA contact. Analysis of our previous Hfq-CRAC dataset reveals that Hfq binds strongly to the 5'UTR of the outer membrane haem receptor chuA (Figures 1A,B). Canonically, gene regulation through sRNAs occurs through direct base-pairing of the sRNA to the RBS within the 5'UTR of mRNA targets. Hfq-CRAC sequencing reads and contact-dependent deletions peaked at the RBS consistent with RyhB and Esr41 negative regulation at this site (Waters et al., 2017; Banerjee et al., 2020). A further Hfq-binding peak was located at position +160 to +200 suggesting that chuA may be subject to additional regulatory inputs within the 5'UTR (Figures 1A,B). The chuA transcription start site is predicted to be ~300 nt upstream of the chuA start codon (Nagy et al., 2001). To precisely map the transcription start site of chuA in EHEC str. Sakai, we analyzed our previously published differential RNA-seq (dRNA-seq) data, which enriches for primary transcripts using terminator exonuclease (TEX) (Sharma et al., 2010; Sy et al., 2020). This showed that the +1 site for chuA is at position 4,391,862, and that the 5'UTR is 291 nucleotides long, which concurs with previous predictions (Figure 1B; Nagy et al., 2001).

In previous analyses we found that sRNA-mRNA seed sites are often positioned within Hfq-bound reads recovered by Hfq-CRAC, and that the Hfq binding site can be used to restrict the sequence space for predicting sRNA-mRNA interactions (Tree et al., 2014; Wang et al., 2018). To identify sRNAs that may regulate chuA, IntaRNA (Busch et al., 2008; Mann et al., 2017) was used to predict interactions between the chuA 5'UTR and
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FIGURE 1 | The chuA 5′UTR is regulated by Hfq-dependent small RNAs. (A) RNA-sequencing data mapping to the chuAS operon. From top to bottom: total RNA-seq (dark blue; RPM), differential RNA-seq in samples untreated (light green) and treated (dark green) with TEX, Term-seq (red; raw 3′ end reads) (BioProject PRJNA601151), Hfq-binding sites from UV-crosslinking datasets (BioProject PRJNA197291). Sequencing deletions which indicate direct contact with Hfq are indicated in black. (B) Same as (A), but focusing on the 5′UTR of chuA. (C) Fluorescence measurements of the EHEC chuA-sfGFP fusion in the presence of predicted binding sRNAs. (D) Fluorescence measurements of the UPEC chuA-sfGFP fusion in the presence of CyaR or ChiX (**p < 0.01, ***p < 0.001).

44 known Hfq-dependent sRNAs (Li et al., 2013). Small RNAs that were predicted to interact within Hfq read peaks were retained. Through this analysis, we predicted that the sRNAs CyaR, SgrS, SdsR, ChiX, FnrS, RyhB, DicF, SroC, and MicC may have regulatory effects on chuA.

To confirm these predicted interactions the 5′UTR of chuA, starting from the +1 site up until the 15th codon of the coding sequence was fused to sfGFP by cloning into the plasmid pXG10SF (Corcoran et al., 2012). The fluorescence of this translational fusion was monitored in E. coli strain DH5α in the presence and absence of the candidate sRNAs expressed from constitutive expression plasmids (Figure 1C). This revealed that the Hfq-dependent sRNAs CyaR and ChiX have an activating and repressive effect on chuA, respectively.

CyaR Regulation of chuA Is Conserved in Uropathogenic Escherichia coli

Earlier work from the Murphy Lab indicated that chuA 5′UTR from EHEC and Shigella are more closely related than
the 5′UTR from EHEC and uropathogenic *E. coli* (UPEC) (Kouse et al., 2013). We next examined whether CyaR and ChiX regulation are conserved in the divergent EHEC and UPEC 5′UTRs. In UPEC, haem acquisition using *chuA* is required for maximal colonization of the kidneys (Hagan and Mobley, 2009). The 5′UTRs of EHEC and UPEC (*chuA*EHEC and *chuA*UPEC) *chuA* share 83.4% identity (Nagy et al., 2001). The UPEC 5′UTR sequence has only 67.4% identity within the Hfq-binding site suggesting that *chuA*UPEC may not be regulated by Hfq-dependent sRNAs. To test whether *chuA*UPEC is post-transcriptionally regulated by CyaR and ChiX we constructed a translational fusion of *chuA*UPEC with sfGFP and monitored fluorescence in the presence or absence of CyaR and ChiX overexpression plasmids. The *chuA*UPEC fusion was activated by CyaR consistent with our results for *chuA*EHEC but was not significantly repressed by ChiX (Figure 1D). Notably, the *chuA*UPEC fusion was 55% more fluorescent than the *chuA*EHEC fusion. This suggests that while *chuA* in both pathotypes is subject to regulation by CyaR, the divergent 5′UTR sequence has de-repressed *chuA* expression in UPEC.

**CyaR Activates *chuA* Translation Through Direct Base-Pairing**

Both CyaR and ChiX are class II sRNAs that are known to modulate sRNA regulation by displacing sRNAs from Hfq when overexpressed (Moon and Gottesman, 2011; Santiago-Frangos et al., 2016). To determine whether CyaR and ChiX control *chuA* through direct RNA–RNA interactions with the *chuA* 5′UTR or indirectly through titration of Hfq, point mutations were made in both *chuA* and the sRNAs CyaR and ChiX to disrupt the predicted interaction sites. A predicted 3 nt compensatory mutation was made in the top scoring *chuA*–ChiX interaction as predicted by IntaRNA. The M1 mutation made in ChiX resulted in the de-repression of *chuA*-sfGFP expression. However, this effect was not observed when the compensatory *chuA* M1 mutation was introduced. This suggested that while the nucleotides mutated in ChiX contributed to *chuA* regulation, they did not interact with the predicted *chuA* interaction site. A second set of compensatory point mutants (M2) were made for the next highest scoring interaction that used the same ChiX seed region that was previously tested. Mutations in either *chuA* or ChiX alone resulted in the disruption of the *chuA*–ChiX interaction, but testing the mutants together did not restore the repression of *chuA* (Supplementary Figure 1). These results indicated that while ChiX affects translation of *chuA*, it is not due to a direct interaction with the *chuA* M1 or M2 site and may occur indirectly.

CyaR is predicted to bind approximately 15 nt upstream of the *chuA* Shine–Dalgarno sequence. The interaction between CyaR and *chuA* results in a 1.3-fold activation of translation. Mutating CyaR at the M1 site reduced CyaR activation by a modest 8.1%, while the *chuA* point mutant completely abolished the interaction (Figure 2). Providing the CyaR-M1 mutant partially restore translational activation of *chuA*-M1, supporting a direct interaction between CyaR and the 5′UTR of *chuA* that activates translation.

**CyaR Activates *chuA* Translation Independently of Temperature**

Regulation of *chuA* occurs on both the transcriptional and translational level. Transcription of *chuA* is inhibited by Fur when cells are grown in iron-rich conditions (Torres and Payne, 1997), while the level of translation is dependent on the environmental temperature. At 25°C, translation is inhibited by the formation of a FourU RNA thermometer that occludes the RBS (Kouse et al., 2013). This inhibitory structure lies 15 nt downstream of the CyaR binding site. To determine whether CyaR-mediated activation of *chuA* translation occurs by inhibiting the formation of this RNA-thermometer, a U273A point mutation known to disrupt the formation of the FourU hairpin loop was made in the 5′UTR of the *chuA*-GFP translational fusion (Figure 3A; Kouse et al., 2013). Expression of wild-type and mutant *chuA*-sfGFP translational fusions were monitored in the presence and absence of CyaR overexpression plasmids at 25°C. Disruption of the RNA thermometer via the U273A point mutation resulted in a 2.6-fold increase in fluorescence compared to the wild-type, confirming the FourU temperature-dependent regulation of *chuA* translation. Disruption of this secondary structure however did not affect CyaR-mediated activation of *chuA*, as a 1.6-fold increase in fluorescence was observed when CyaR was provided with the U273A mutation in *chuA* (Figure 3B). This demonstrated that CyaR-mediated activation of *chuA* acts independently of post-transcriptional inhibition through the FourU RNA thermometer structure.

**The *chuA* 5′UTR Contains a Sequence That Inhibits Its Expression**

While the activation of *chuA* translation by CyaR is not due to disruption of the RNA thermometer, *in silico* folding predictions on the 5′UTR of *chuA* indicated potential secondary structure throughout the 5′UTR, including some that occluded sequences downstream of the RBS (Supplementary Figure 2). To understand whether the activating *chuA*–CyaR interaction requires structured sequences within the *chuA* 5′UTR, we progressively removed sequence from the 5′ end in four truncations of the * chuA*-sfGFP translational fusion (Figure 4A). Predicted secondary structure and the presence of Hfq distal face binding motifs [(ARN)x] were used as guides in making truncates of the *chuA* 5′UTR (Soper and Woodson, 2008; Zhang et al., 2013; Schu et al., 2015). A search through the 5′UTR of *chuA* identified 10 (ARN)x sites with one mismatch tolerated (ARN4m1). The first truncate (T1) begins at +57 nt and removes a section that forms two hairpins as well as 9 out of 10 ARN4m1 sites. The second (T2) and third (T3) begin at +116 and +177, respectively, and each one removes another hairpin from the overall predicted secondary structure while maintaining the RNA-thermometer. The second truncate T2 also removes the last ARN4m1 motif. The fourth truncate (T4) begins at +217 and leaves only the CyaR binding site, the RNA-thermometer, RBS and the start codon. In the absence of the CyaR overexpression plasmid, the *chuA* 5′UTR T1–T4 truncates caused 2.1-, 1.9-, 2.9-, and 9.2-fold increases in *chuA* translation, respectively. CyaR was able to activate
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**FIGURE 2 |** CyaR activates chuA translation through direct base-pairing. (A) IntaRNA prediction of the chuA–CyaR interaction. Compensatory point mutations predicted to disrupt the interaction are indicated by the arrows. (B) Fluorescence measurements of wild-type or mutant chuA-sfGFP translational fusions in the presence and absence of wild-type or mutant CyaR overexpression plasmid. Measurements are the mean median fluorescence intensity of three biological replicates (*p < 0.01, *p < 0.05).

**FIGURE 3 |** CyaR activates chuA translation independently of a FourU RNA-thermometer. (A) Diagram indicating the point mutation made to disrupt the formation of the chuA FourU RNA thermometer. (B) Fluorescence measurements of wild-type and RNA-thermometer disrupted chuA-sfGFP translational fusions in the presence and absence of CyaR overexpression plasmid. Measurements are the mean median fluorescence intensity of three biological replicates (**p < 0.001).

truncates T1, T2, and T4 between 20 and 40% consistent with CyaR activation of the wild-type construct (Figure 4B). These results indicate that CyaR does not act through alleviation of an inhibitory secondary structure or a regulatory sequence in the upstream +1 to +217 nt region of the 5’UTR. Notably, removing the region between +177 and +217 nt (T3–T4) of the chuA 5’UTR resulted in a dramatic increase in chuA-GFP expression (9.2-fold) indicating that the T3–T4 region of the 5’UTR contains sequences or RNA structures that strongly repress expression.

**chuA Is Subject to Premature Transcription Termination by Rho**

In *E. coli*, the average 5’UTR is approximately 25–35 nucleotides in length (Kim et al., 2012; Evfratov et al., 2017). The 5’UTR of chuA is 291 nucleotides, making it an unusually long UTR (Wyckoff et al., 1998; Kouse et al., 2013). In commensal *E. coli*, over half of all annotated genes with long 5’UTRs (defined as >80 nt) are prematurely terminated by the transcription termination factor Rho (Sediyarova et al., 2016). Further, horizontally transferred genes have been shown to be more susceptible to regulation by Rho-dependent termination (Cardinale et al., 2008; Peters et al., 2012). By analyzing our previously published Term-seq data, we noticed that there were termination sites at positions +129, +271, and +297 within the chuA mRNA (Figure 5A). We did not identify Rho-independent (intrinsic) terminators within the chuA 5’UTR using ARNold (Figure 5A; Naville et al., 2011; Sy et al., 2020). Collectively, this suggested that transcription of the pathogenicity island encoded chuA may be prematurely terminated by Rho.

To identify Rho-termination sites in EHEC, we sequenced total RNA extracted from cells treated with the Rho-inhibiting antibiotic bicyclomycin (BCM) (Zwiefka et al., 1993). Rho-termination sites were identified by using the approach adopted by Adams et al. (2021) that previously identified termination sites in BCM-treated *E. coli* cells. Consistent with earlier studies we found that horizontally acquired regions and antisense RNAs are enriched among genes that are up-regulated by BCM treatment (Figure 5B; Cardinale et al., 2008; Peters et al., 2009). Read-through scores were calculated to assess Rho-dependent termination using previously published scripts (Adams et al., 2021). Read-through scores were maximal for all
three replicates (±BCM) at positions +267–273 nt correlating well with the termination site at +271 nt identified by Term-seq. Read-through transcription at this site was up-regulated between 1.33- and 1.47-fold by BCM treatment indicating that Rho prematurely terminates transcription of chuA at position +271 (Figure 5A). To verify that chuA was subject to premature transcription termination by Rho, we measured the fluorescence of inducible full-length and truncated chuA-GFP translational fusions in the presence or absence of BCM (Figure 5A). Treatment with BCM resulted in a ~1.4-fold increase in fluorescence for the full-length and T1–T3 chuA-GFP constructs. No significant change was observed for chuA truncate T4 (Figure 5C). Rho-dependent termination requires recruitment of the Rho at upstream Rho utilization sites (rut) and we speculate that the rut site for Rho-dependent termination at +271 is removed in the T4 truncate. Taken together, these results indicate that chuA is negatively regulated by Rho-dependent termination at position +271 and this repression requires the 30 nt between +177 and +217 nt of the chuA 5′UTR.

Small RNAs can regulate Rho-dependent termination by altering the accessibility of Rho utilization sites, either as a by-product of translation inhibition, or by directly binding to the rut site itself (Bassi et al., 2012; Sedlyarova et al., 2016; Chen et al., 2019; Silva et al., 2019). We hypothesized that CyaR may activate chuA translation by preventing premature Rho termination within the 5′UTR. To test this, we measured the fluorescence of our chuA-GFP translational fusion in the presence or absence of BCM and CyaR. CyaR was able to activate translation of chuA in the presence of BCM, indicating that CyaR activates chuA independently of Rho termination (Figure 5D). Collectively our data demonstrate that chuA expression is independently regulated by CyaR sRNA, Rho terminator, and the FourU RNA thermometer.

Expression of CyaR Is Correlated With chuA Expression in vivo

In order to obtain a more biologically relevant representation of the relationship between CyaR and chuA expression, we analyzed previously published RNA sequencing data of EHEC str. EDL933 obtained during colonization of a bovine rectum, intestine and rumen (Segura et al., 2018, 2021). Previously, these datasets have been used to show that genes relating to iron acquisition are differentially regulated in each of the gastrointestinal niches when compared to minimal M9 media. We compared the expression of both chuA and CyaR between each of the gastrointestinal niches and we find that chuA is most highly expressed in the rectum, followed by the rumen, then the intestine (Figure 6). These findings are consistent with induction of chuA by environmental signals at this site, including iron-limitation, temperature and a gluconeogenic environment (Snider et al., 2009; Bertin et al., 2013). We also observe this trend with CyaR expression, with the sRNA being maximally expressed in the rectum. This suggests that CyaR contributes to the maximal expression of chuA in the rectum, allowing for better colonization of this environmental niche.

DISCUSSION

To successfully grow and colonize a host, pathogens utilize systems that allow them to retrieve trace minerals required for growth that are normally sequestered. Various pathotypes of E. coli, including EHEC and UPEC, utilize the ChuA outer membrane haem receptor to transport host derived haem. Transcription of chuA is repressed in the presence of iron by Fur (Torres and Payne, 1997), while translation is regulated by temperature through a FourU RNA-thermometer that occludes the RBS (Kouse et al., 2013). The combination of these two modes of regulation allows the pathogen to sense two signals associated with the host environment (low iron and high temperature) and activate expression of the haem receptor inside the host. Notably, each signal in isolation would generate many false positives when deciding on whether the cell had entered a mammalian host.

The natural order imposed by transcription and post-transcriptional regulation creates an AND-logic gate, where low iron levels AND high temperature are required for expression of chuA in the host. Here we have shown that chuA is controlled by two additional post-transcriptional signals: repression through Rho termination and activation through...
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Transcription of CyaR is activated by the global regulator Crp when cyclic-AMP levels are high, such as in the absence of glucose (De Lay and Gottesman, 2009). EHEC colonizes the colon where the primary source of carbon is mucus. Bacteroides thetaiotaomicron (Bt) can cleave mucus to release sugars that are utilized by EHEC (Xu et al., 2003; Fischbach and Sonnenburg, 2011). During colonization of the gastrointestinal mucus layer and epithelium, EHEC is likely to encounter varying niches that include both glycolytic and gluconeogenic environments (Miranda et al., 2004). Sensing variation in the availability of sugars and oxygen availability have been suggested to be key regulatory signals that allow EHEC to determine location in the gastrointestinal tract and proximity to the epithelium (Carlson-Banning and Sperandio, 2016). Close to the intestinal epithelium, the relative absence of microorganisms that metabolize and release sugars from mucus creates a gluconeogenic environment (Conway and Cohen, 2015). Lower levels of fucose, and higher levels of succinate reduces the level of the fucose-sensing two-component system FusKR that induces expression of Cra. In gluconeogenic environments, Cra can enhance binding of the cAMP receptor protein Crp to its targets (Ryu et al., 1995). Activation of Cra can induce expression of genes required for T3SS and adhesion to the epithelium (Njoroge et al., 2012; Curtis et al., 2014; Carlson-Banning and Sperandio, 2016). Expression of Crp can also be activated in the presence of Cra (Zhang et al., 2014). We propose that the gluconeogenic environment...
encountered at the gastrointestinal tract provides an activating signal for CyaR that has been incorporated into chuA regulation as an additional signal that indicates host colonization and haem availability. Consistent with this hypothesis, we find that both CyaR and chuA transcription is activated in bovine rectal samples (Figure 6).

The mechanism of CyaR activation of chuA expression remains unclear although we have ruled out three probable mechanisms: CyaR acts through direct base-pairing with the chuA 5′UTR rather than general titration of Hfq, CyaR does not disrupt the FourU RNA thermometer to override temperature-dependence, and CyaR does not inhibit premature Rho termination of chuA. While these results have uncovered two additional post-transcriptional regulators of chuA (CyaR and Rho) they also suggest that there exists an additional repressive element in the chuA 5′UTR that is overcome by CyaR binding. We note that our chuA-GFP translational fusion reports on multiple post-transcriptional processes including translation efficiency and mRNA stability, and that CyaR may be modulating either of these to affect chuA expression.

Our Term-seq data indicates that the chuA 5′UTR is terminated at positions +129, +271, and +297 with position +271 nt maximally sensitive to treatment with BCu. Using truncates of the chuA 5′UTR we have also found that the sequence between +177 and +217 nt (T3 and T4) represses chuA expression almost 10-fold. We suggest that Rho associates with the nascent chuA transcript between +177 and +217 nt to terminate transcription at +271 nt. Small RNAs can promote or occlude rut sites to control Rho termination in response to regulatory signals (Sedlyarova et al., 2016; Silva et al., 2019) and it remains plausible that Rho termination is modulated by yet another regulatory signal.

The chuA encoded haem receptor is transcribed as a bicistronic transcript with the haem oxygenase chuS. In previous work we showed that chuS expression is repressed by the sRNA FnrS that is induced under anaerobic conditions, consistent with ChuS requiring oxygen for activity. FnrS repression is over-ridden by the sRNA sponge AsxR transcribed from the Shiga toxin 2 encoding bacteriophage Sp5 (Tree et al., 2014). Collectively, expression of chuAS is subject to an impressive level of post-transcriptional regulation that provides complex integration of environmental signals beyond low iron (transcriptional regulation by Fur). These post-transcriptional signals appear to provide a much more accurate interpretation of the environment to indicate whether haem transport is required (i.e., whether EHEC is at the gastrointestinal epithelium). The logic of the regulatory circuit that controls chuAS expression is outlined in Figure 7.

Post-transcriptional regulation appears ideally suited to this layering of logic gates because the natural order (dependence) imposed by transcriptional and post-transcriptional signals creates AND gates. There are at least three ordered layers of AND gates in the chuA transcript that need to be satisfied before the next layer of signals are incorporated. Fur-dependent transcription, Rho-dependent termination, and post-transcriptional control through sRNAs or the FourU thermometer. Together these form an elegant set of AND, OR, and NOT gates that interpret the environment and genetic background of the host (the later through the EHEC-specific sRNAs AsxR and Esr41) (Figure 7). The extensive post-transcriptional logic of the chuAS operon suggests that layered post-transcriptional regulation can create complex regulatory logic gates to integrate and interpret environmental signals.

MATERIALS AND METHODS

Bacterial Strains and Growth Conditions

Bacterial strains, oligonucleotides and plasmids used for this study are listed in Supplementary Table 1. *E. coli* was routinely
grew at 37°C in liquid Luria-Bertani (LB) broth or on solid LB-agar plates. Bacterial media was supplemented with ampicillin (100 µg/mL) or chloramphenicol (34 µg/mL) where appropriate. For inhibition of Rho termination, bicyclomycin (BioAustralis, 50 µg/mL) was added to exponential phase (OD$_{600} = 0.6$) cultures for 30 min before being assayed.

**In silico Prediction of Interacting sRNAs**

A list of published sRNAs present in enterohemorrhagic *E. coli* was taken from the Bacterial Small Regulatory RNA Database (Li et al., 2013). sRNAs listed as not being Hfq-binding were filtered out, and the sequences of the remaining sRNAs were input into IntaRNA (Busch et al., 2008; Wright et al., 2014; Mann et al., 2017) to search for interactions with the 5'UTR of *chuA* sRNAs that were predicted to bind to regions of the *chuA* 5'UTR that were not Hfq-binding were removed from consideration.

**Construction of GFP-Translational Fusions and sRNA Expression Vectors for Testing sRNA–mRNA Interactions**

Plasmids pXG10SF containing the full-length and truncated *chuA*-GFP translational fusion and pZE12 carrying candidate sRNA or sRNA sequences were cloned according to method described in Urban and Vogel (2007). Briefly, the 5’UTR of *chuA* and candidate sRNAs were amplified from genomic DNA using primers identified in Supplementary Table 1. The *chuA*-GFP translational fusion was inserted using *NheI*-*NsiI* restriction cloning. Candidate sRNAs were inserted by first linearizing pZE12- luc using primers pLacO-B and pLacO-D. The linearized amplicon, as well as the candidate sRNA amplicon were digested using *XbaI* and ligated together using T4 DNA ligase.

Mutations were made in the *chuA* or sRNA sequences by using the Quickchange XL mutagenesis kit (Agilent) according to the manufacturer’s instructions. Primers for mutagenesis were designed using the Quickchange primer design program.

**Confirmation of sRNA–*chuA* Interactions Using the sfGFP 2-Plasmid System**

The expression of sfGFP was monitored and quantified with and without candidate sRNAs using a BD FACSCanto II or a BD LS Fortessa Special Order Research Product cell analyzer. Fluorescence was measured using a 530/30 nm bandpass filter. Forward scatter (FSC) and side scatter (SSC) were also measured.

**RNA Secondary Structure Prediction**

The secondary structure for the 5’UTR of *chuA* was predicted using the mfold2 and RNAfold webserver (Zuker, 2003; Lorenz et al., 2011). Figures were drawn on RNAstructure version 6.0.1 (Reuter and Mathews, 2010). ARN motifs were detected using custom scripts previously used in Tree et al. (2014).

**RNA Sequencing of Bicyclomycin-Treated Cells**

Three single colonies of *E. coli* O157:H7 str. Sakai stx- were grown for 16 h in LB broth. The following day, these subcultured 1/100 in MEM-HEPES supplemented with 0.1% glucose and 250 nM Fe(NO$_3$)$_3$. At an OD$_{600}$ of 0.75, cultures were split into two (treated and untreated), and 50 µg/mL of bicyclomycin was added to the treated sample. Cultures were incubated for a further 15 min, followed by addition of RNA stop solution (5% water-saturated phenol in ethanol). RNA was extracted using guanidinium thiocyanate-phenol as previously described in Tollervey and Mattaj (1987). Genomic DNA was digested using RQ1 DNase (Promega) and RNA was cleaned using another phenol-chloroform extraction. Total RNA was ribodepleted using the Zymo-Seq RiboFree Total RNA library kit and libraries were prepared using the Illumina NextSeq 500/550 Mid-Output Kit. Samples were 2 × 75 bp paired-end sequenced on an Illumina NextSeq 500 platform. Library preparation and sequencing were performed by the Ramaciotti Centre for Genomics in the University of New South Wales, Sydney, NSW, Australia.

**RNA Sequencing Analysis and Identification of Rhod-Dependent Transcripts**

RNA sequencing data for in vivo analysis was taken from SRA accession number SRP136076. Alignment and DSeq analysis of RNA sequencing output was done using the “align,” “coverage,” “gene-quant,” and “deseq” modules of READemption v1.0 (Förstner et al., 2014) using default settings.

To identify Rho-terminated transcripts, fastq files were re-aligned to the *E. coli* O157:H7 str. Sakai genome (accession number: NC_002695.2) using BWA-MEM (v0.7.17) (Li and Durbin, 2010). BAM files were generated using samtools (v1.10) (Li et al., 2009) and 3' end read counts were calculated using the genomcove tool from bedtools (v2.27.1) (Quinlan and Hall, 2010). Rho readthrough scores were calculate using the program RhoTerm-Peaks using a window size of 250 nt (Adams et al., 2021).

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1https://www.agilent.com/store/primerDesignProgram.jsp

2https://unafold.rna.albany.edu/?q=mfold
DATA AVAILABILITY STATEMENT

RNA sequencing datasets are deposited at the NCBI Gene Expression Omnibus (GEO) under accession numbers GSE143631, GSE46118, and GSE197379.

AUTHOR CONTRIBUTIONS

BS and JT conceived and designed the experimental work, analyzed the data, and wrote the manuscript. BS performed the experimental work. Both authors contributed to the article and approved the submitted version.

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