



Fur-Dam Regulatory Interplay at an Internal Promoter of the Enteroaggregative *Escherichia coli* Type VI Secretion *sci1* Gene Cluster

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ABSTRACT The type VI secretion system (T6SS) is a weapon for delivering effectors into target cells that is widespread in Gram-negative bacteria. The T6SS is a highly versatile machine, as it can target both eukaryotic and prokaryotic cells, and it has been proposed that T6SSs are adapted to the specific needs of each bacterium. The expression of T6SS gene clusters and the activation of the secretion apparatus are therefore tightly controlled. In enteroaggregative Escherichia coli (EAEC), the sci1 T6SS gene cluster is subject to a complex regulation involving both the ferric uptake regulator (Fur) and DNA adenine methylase (Dam)-dependent DNA methylation. In this study, an additional, internal, promoter was identified within the sci1 gene cluster using +1 transcriptional mapping. Further analyses demonstrated that this internal promoter is controlled by a mechanism strictly identical to that of the main promoter. The Fur binding box overlaps the -10 transcriptional element and a Dam methylation site, GATC-32. Hence, the expression of the distal sci1 genes is repressed and the GATC-32 site is protected from methylation in iron-rich conditions. The Fur-dependent protection of GATC-32 was confirmed by an in vitro methylation assay. In addition, the methylation of GATC-32 negatively impacted Fur binding. The expression of the sci1 internal promoter is therefore controlled by iron availability through Fur regulation, whereas Dam-dependent methylation maintains a stable ON expression in iron-limited conditions.

IMPORTANCE Bacteria use weapons to deliver effectors into target cells. One of these weapons, the type VI secretion system (T6SS), assembles a contractile tail acting as a spring to propel a toxin-loaded needle. Its expression and activation therefore need to be tightly regulated. Here, we identified an internal promoter within the *sci1* T6SS gene cluster in enteroaggregative *E. coli*. We show that this internal promoter is controlled by Fur and Dam-dependent methylation. We further demonstrate that Fur and Dam compete at the -10 transcriptional element to finely tune the expression of T6SS genes. We propose that this elegant regulatory mechanism allows the optimum production of the T6SS in conditions where enteroaggregative *E. coli* encounters competing species.

KEYWORDS type VI secretion, epigenetism, methylation, microbial communities, regulation, repression

The fate of microbial communities is governed by communication, cooperation, and competition mechanisms between microorganisms (1–9). Bacteria have therefore developed an arsenal of signaling, sensing, and antagonistic activities. To eliminate competitors, bacteria evolved distinct mechanisms for release of antibiotics or bacteriocins in the extracellular medium, as well as delivery of toxins directly into the target cell (10–12). One of the delivery apparatuses, the type VI secretion system (T6SS),

Citation Brunet YR, Bernard CS, Cascales E. 2020. Fur-Dam regulatory interplay at an internal promoter of the enteroaggregative *Escherichia coli* type VI secretion *sci1* gene cluster. J Bacteriol 202:e00075-20. https://doi.org/10.1128/JB.00075-20.

Editor Thomas J. Silhavy, Princeton University

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Received 6 February 2020 Accepted 28 February 2020

Accepted manuscript posted online 9 March 2020 Published 27 April 2020 transports effectors into competing bacteria using a mechanism similar to that used by contractile injection systems such as bacteriophages and R-pyocins (13–19). This secretion apparatus comprises an ~800-nm long cytoplasmic needle-like structure composed of an inner tube tipped by a spike complex that is used to penetrate the membrane of the target cell (12, 14, 19). The inner tube is wrapped by an outer sheath that is assembled under an extended metastable conformation (20, 21). The tail tube/sheath complex is built on a baseplate that is anchored to the cell envelope by a membrane complex (22–29). Tail tube/sheath assembly, which can be visualized *in vivo* by fluorescence microscopy, is completed in a few tens of seconds (30–32). Contraction of the sheath powers the propulsion of the inner tube to deliver effectors into the target cell (15, 17, 31, 33–35). Effectors are usually charged within the inner tube lumen or loaded onto the spike complex via direct interactions with the VgrG/PAAR spike or via adaptor proteins (36–45).

The T6SS is a very efficient mechanism and hence is an important player in the regulation of microbiota (7, 46). Bacteria equipped with this apparatus colonize an environmental niche more efficiently and hence have better access to resources (47–51). Most of the T6SS gene clusters are not constitutively expressed and T6SS-dependent antagonistic activities are usually deployed once cells experience stress or nutrient starvation conditions (52–57). T6SS gene clusters are therefore subjected to a tight regulation that involves sensing of the environmental conditions (52, 53, 55). Most known regulatory mechanisms are hijacked by T6SSs for their own regulation, including transcriptional activators and repressors, alternate sigma factors, histone-like proteins, two-component transduction cascades, or quorum-sensing systems (52, 53). In addition, a number of T6SSs are posttranslationally activated by a threonine phosphorylation pathway in response to cell damage or envelope stress (58).

Enteroaggregative *Escherichia coli* (EAEC) is equipped with two functional T6SSs, named Sci1 (T6SS-1 subfamily) and Sci2 (T6SS-3 subfamily) (59, 60). These two T6SSs confer antagonistic activities but are not expressed under the same conditions, suggesting that T6SS-mediated antibacterial activities are required in two conditions that EAEC may encounter during its life cycle (31, 44). The *sci2* gene cluster is expressed during infection conditions and is activated in laboratory conditions when cells are grown in a synthetic medium mimicking the macrophage environment (59). This *sci2* gene cluster is under the control of the AraC-like AggR transcriptional regulator (59), which also modulates the expression of most biofilm determinants (59, 61), suggesting that the Sci2 T6SS is required for eliminating competing bacteria during aggregation, a phenomena that occurs during host colonization. In contrast, the *sci1* gene cluster is expressed in minimal synthetic media and has been shown to be under the dual control of the ferric uptake repressor (Fur) and Dam-dependent methylation (62).

To better understand the organization of the sci1 gene cluster, we defined its operon structure. Reverse transcriptase polymerase chain reaction (RT-PCR) experiments showed that all genes are contiguous, suggesting that all the genes are present on a single mRNA or on several overlapping mRNAs. Using +1 transcriptional mapping, we confirmed the existence of a promoter region upstream of the first gene of the cluster and revealed an additional promoter located upstream to the EC042_4532 gene, within the EC042 4531 coding sequence. We further identified a Fur-binding sequence overlapping with the -10 transcriptional box and demonstrated that Fur binds this sequence with high affinity, thereby preventing RNA polymerase from gaining access to the promoter. Sequence analyses showed that this Fur box overlaps with a GATC Dam methylation site, GATC-32. In vivo, we showed that Fur prevents methylation of the GATC-32 site when cells were grown in iron-replete conditions. In vitro competition experiments confirmed that Fur prevents GATC-32 methylation. In addition, we observed that Dam-dependent methylation of GATC-32 decreases the affinity of Fur for its Fur box. Taken together, our results demonstrate that a second functional, internal promoter controls the expression of T6SS sci1 genes and that this promoter is under a regulatory mechanism similar to the main promoter.

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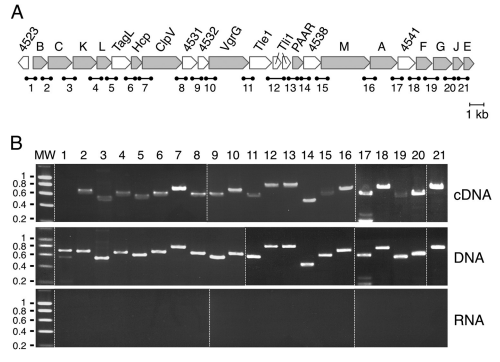


FIG 1 Operon structure of the EAEC *sci1* T6SS gene cluster. (A) Schematic organization of the EAEC *sci1* T6SS gene cluster (*EC042_4524* to *EC042_4545*). Genes encoding T6SS core components are indicated in gray. Accessory genes or genes of unknown function are represented in white. The fragments corresponding to gene junctions and amplified in the RT-PCR experiments are indicated below (1, 692 bp; 2, 672 bp; 3, 550 bp; 4, 618 bp; 5, 586 bp; 6, 643 bp; 7, 748 bp; 8, 629 bp; 9, 575 bp; 10, 654 bp; 11, 581 bp; 12, 768 bp; 13, 762 bp; 14, 459 bp; 15, 600 bp; 16, 673 bp; 17, 576 bp; 18, 720 bp; 19, 552 bp; 20, 591 bp; 21, 678 bp). (B) Operon structure of the EAEC *sci1* T6SS gene cluster. Agarose gel analyses of the indicated gene junctions (numbered 1 to 21 in panel A) amplified by PCR from cDNA, genomic DNA (middle panel; positive control), and total RNA (negative control). The presence of PCR fragments in the cDNA gels demonstrates cotranscription of the genes located 5' and 3' of the amplified region. Molecular weight markers (MW, in kilobases) are indicated on the left. Dashed lines separate different gels combined into a single image.

RESULTS AND DISCUSSION

Operon structure of the sci1 T6SS gene cluster. We previously reported that the promoter located upstream of the tssB gene, i.e., the first gene of the EAEC sci1 T6SS gene cluster, contains operator sequences for the ferric uptake regulator (Fur), as well as an overrepresentation of GATC motifs that are targets of the DNA adenine methylase (Dam). Using in vivo and in vitro Fur binding and methylation assays, we delineated the contribution of these two regulators on the expression of the tssB gene (62). However, whether additional or internal promoters exist and whether the entire gene cluster is subjected to this regulatory control remained undetermined. The EAEC sci1 gene cluster is an \sim 26-kb DNA fragment on the *pheU* pathogenicity island (Fig. 1A) (59). Prediction of the open reading frames (ORFs) within this fragment shows that it likely encodes 21 gene products, including the 14 T6SS core components, a toxin-immunity pair, and accessory genes or of unknown function (genes tssB to tssE [Fig. 1A]). With the exception of a large intergenic sequence (162 bp between the hcp and the clpV genes), most of the start and stop codons of contiguous genes overlap or are separated by few (<8) nucleotides (see Fig. S1 in the supplemental materials). This genomic organization suggests that translational coupling must occur and that the expression of these genes must be coordinated. To test whether the sci1 gene cluster is organized as a single genetic unit or constituted of several operons, we performed reverse transcriptase polymerase chain reactions (RT-PCR) using oligonucleotides designed for the amplification of each gene junction (numbered 1 to 21 [Fig. 1A]). RT-PCR experiments were performed on purified total RNAs extracted from cells grown in Sci1-inducing medium (SIM) (Fig. 1B, top panel). As controls, RT-PCRs were performed on purified genome

A P_{sci1}

<u>CCTGATTATTTGCATTATA</u>TC**GATC**GATGTATCTG TTATATTGAGATTTTTCA**GATC**TTCGTCC<u>TATAAT</u> <u>GATCAAAATTAAA</u>TCAGTGCACAAGGGGAGGCATC TGCGGTGATGGAACCCCTGAGATGCAGGTTTCACA GGAGAGAGCC<u>ATG</u>

B P₄₅₃₂

GCGAATATCCCATGGAGCAGCAGGCACAA<u>ATTATT</u> <u>GCTGATCATTTTA</u>CTTTGCAGGCTGAAGGATACGG GACATGGTGTGATATGAGAAGGGACGGTGATATCA CACTGGACGGAAATATGTCTGAGTATGTTATTCGC AGCCTGTATACCAGCACGTTGCCGGGGGTTCCC<u>ATG</u>

С

Fur1	TATAAT GAT CAAAATTAAA
Fur box	GATAATGATAATCATTATC
Fur-32	ATTATTGCT <mark>GATC</mark> ATTTTA

FIG 2 Regulatory elements of the *sci1* and *4532* promoters. (A and B) Nucleotide sequences of the *sci1* (A) and *EC042_4532* (B) promoters highlighting overlaps between the transcriptional elements, Fur binding boxes, and Dam methylation motifs. The +1 transcriptional site identified by 5'RACE is indicated in red. GATC Dam methylation sites are indicated in blue. The -10 elements are indicated in green. The underlined sequences indicate Fur binding boxes (italics) and translational start codons. (C) Sequence alignment of the *fur1 (sci1* promoter) and *fur-32 (EC042_4532* promoter) boxes with the *E. coli* Fur box consensus sequence. Identical bases are framed in gray. The -10 elements (green) and GATC motifs (blue) are indicated.

DNA (Fig. 1B, middle panel), as well as on the total RNA preparation in the absence of reverse transcriptase, to test for DNA contamination (Fig. 1B, bottom panel). As shown in Fig. 1B, RT-PCR products with the expected sizes were obtained for each gene junction of the *sci1* gene cluster from DNA or cDNA but not from RNA (Fig. 1B, lanes 2 to 21), suggesting that the 21 genes are cotranscribed. As expected, the *Ec042_4523* ORF, upstream of the first gene of the *sci1* cluster and in the reverse orientation compared to the *tss* genes, is not cotranscribed with *tssB* (Fig. 1B, lane 1). These results suggest that all the *sci1* genes are present on a unique polycistronic mRNA or that overlapping mRNAs are expressed from internal promoters.

An additional promoter is located upstream of EC042_4532. To identify a potential internal promoter(s), we used an in silico approach. Analysis of the T6SS sci1 gene cluster using the BProm algorithm (Softberry; http://linux1.softberry.com/berry .phtml) suggested the existence of an additional promoter with a σ^{70} -10 element upstream of the EC042_4532 gene. To test whether an internal promoter was present upstream of *Ec042_4532*, we used a 5' rapid amplification of cDNA ends (5' RACE) assay. mRNAs were extracted from EAEC cells grown in Sci1-inducing medium (SIM) and subjected to primer extension. The putative tssB promoter was also included in this assay. The results showed that transcription of the tssB mRNA starts at the A base located 73 bases upstream of the ATG start codon of tssB (colored red in Fig. 2A). The tssB transcription starts are therefore compatible with the putative -10 and -35transcription boxes identified through in silico analyses in our previous study (62) (Fig. 2A). A transcriptional start was also detected upstream of the EC042_4532 gene, suggesting the existence of an active internal promoter. The position of the identified transcriptional start (base G, located 117 bases upstream of the ATG of EC042_4532 [colored red in Fig. 2B]) is compatible with the location of the -10 element predicted by the BProm algorithm (Fig. 2B).

In silico sequence analyses of the *EC042_4532* promoter region identify Fur and **Dam sites overlapping with the -10 element.** Interestingly, the BProm computer program also identified a putative Fur-binding box in the *EC042_4532* promoter region

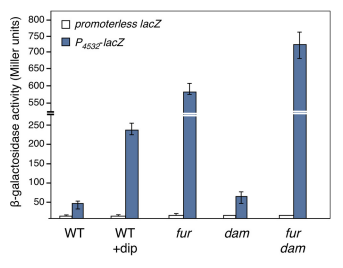


FIG 3 The 4532 promoter is under the control of iron levels, Fur, and Dam. β -Galactosidase activity (in Miller units) of a promoterless *lacZ* fusion and of the *P*₄₅₃₂-*lacZ* reporter fusion at OD₆₀₀ = 0.8 in the WT EAEC 17-2 strain after a 30-min treatment with 2,2'-dipyridyl (+dip; 100 μ M) or in the isogenic *fur*, *dam*, and *fur-dam* mutants.

(hereafter called Fur-32). This putative operator sequence overlaps with the -10 of transcription (Fig. 2B and C). This situation is reminiscent of the main promoter, which is repressed by the Fur protein in an iron-dependent manner (62). One of the Fur boxes contained in the *tssB* promoter contains a Dam-dependent methylation site (Fig. 2A), and we previously reported that Fur and Dam compete at this specific site to fine-tune the expression of the *sci1* gene cluster (62). Strikingly, a GATC motif is also found within the putative Fur-32 box of the *EC042_4532* promoter (Fig. 2C) (hereafter called GATC-32). Taken together, the *in silico* sequence analyses raised the question of whether the internal promoter was under a similar regulatory mechanism as the *tssB* main promoter.

The *P*₄₅₃₂-*lacZ* translational fusion is responsive to iron limitation and Fur. To test whether the expression of the internal promoter was regulated by Fur, we engineered a low-copy-number plasmid-borne translational fusion of a 570-bp fragment comprising the *EC042_4532* promoter (from -450 to +120 relative to the transcriptional +1, called here *P*₄₅₃₂) to *lacZ*. The β-galactosidase activity of this *P*₄₅₃₂-*lacZ* translational fusion was monitored in the EAEC *lacZ* strain or its *fur* isogenic mutant in the presence or absence of the iron chelator 2,2'-dipyridyl (dip). Figure 3 shows that the expression of the *P*₄₅₃₂ translational fusion increased ~6-fold in the wild-type (WT) strain upon treatment with the iron chelator. Compared to the WT strain in the absence of iron chelator, the activity of the *fur* mutant strain with 2,2'-dipyridyl had no additional effect on the activity of the *P*₄₅₃₂-*lacZ* translational regulator in an iron-dependent manner.

Fur binds to the *P*₄₅₃₂ **promoter and limits access to RNA polymerase.** To test whether Fur binds the *EC042_4532* promoter region *in vitro*, the purified *E. coli* Fur protein and the radiolabeled *P*₄₅₃₂ 570-bp fragment were used for electrophoretic mobility shift assays (EMSA). As controls and as previously published (62), Fur bound to the *sci1* promoter, yielding two bands due to the presence of two Fur boxes, but did not retard the Fur-independent *sci2* promoter (Fig. 4A, lanes 8 to 10). Fur also shifted the *P*₄₅₃₂ fragment in the presence of iron, its corepressor (Fig. 4A, lanes 1 to 5; Fig. 4B). This shift was strictly dependent on metal-bound Fur, as no band retardation could be observed when the fragment and the purified regulator were incubated in the presence of the metal chelator EDTA (Fig. 4A, lane 6). In contrast, control experiments showed that the σ^{54} enhancer binding protein NtrC did not bind the *P*₄₅₃₂ fragment (Fig. 4A,

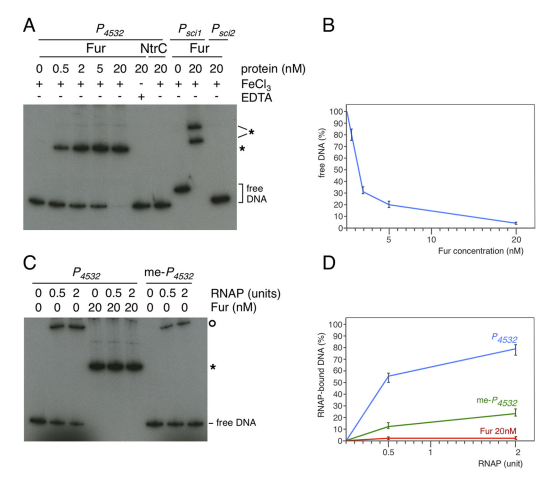


FIG 4 Fur binds to the 4532 promoter and prevents access to RNA polymerase *in vitro*. (A) Electrophoretic mobility shift assay of the *EC042_4532* promoter (P_{4532}) with the indicated concentration of Fur in the presence of FeCl₃ or in the presence of EDTA or using a purified NtrC transcriptional activator. Controls include Fur shift assays of the Fur-dependent *sci1* promoter or of the Fur-independent *sci2* promoter. DNA-Fur complexes are indicated by stars. (B) Densitometry analysis of Fur binding on the P_{4532} fragment, represented as free P_{4532} DNA as a function of Fur concentration. (C) Electrophoretic mobility shift assay of the unmethylated (P_{4532}) or methylated (me- P_{4532}) *EC042_4532* promoter with the indicated by the star and circle, respectively. (D) Densitometry analysis of RNAP binding on the unmethylated (red curve) P_{4532} fragment, represented as RNAP-bound DNA as a function of RNAP concentration.

lane 7). From these data, we conclude that Fur binds to the P_{4532} promoter *in vitro*, likely to the putative Fur-32 box.

Fur repression is usually caused by preventing access of the RNA polymerase (RNAP) to the promoter. We hypothesized that such a mechanism might be likely at promoter P_{4532} , as the putative Fur-32 box overlaps with the -10 RNAP-binding element (Fig. 2B). We therefore tested whether σ^{70} -RNAP holoenzyme binds to the P_{4532} promoter and whether Fur influences σ^{70} -RNAP binding. Figure 4C shows that the σ^{70} -RNAP complex binds to the P_{4532} promoter (Fig. 4C, lanes 1 to 3) and that preincubation of the P_{4532} fragment with Fur prevents binding of the σ^{70} -RNAP, demonstrating that Fur and RNAP complete for binding on P_{4532} (Fig. 4C, lanes 4 to 6; Fig. 4D).

Dam methylation at the GATC-32 site decreases RNAP binding to the P_{4532} promoter. To gain insight on the contribution of Dam to the regulation of *EC042_4532*, we measured the β -galactosidase activity of the P_{4532} -lacZ translational fusion in dam and fur-dam EAEC strains. Deletion of dam did not cause a significant variation in the activity of the promoter fusion compared to its parental wild-type strain (Fig. 3). In contrast, the activity of the promoter fusion in the fur-dam strain increased ~16-fold compared to the wild-type strain, and ~1.4-fold compared to the fur mutant. These

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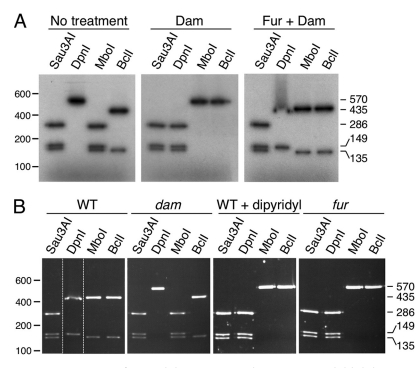


FIG 5 Fur protects GATC-32 from methylation *in vitro* and *in vivo*. (A) A radiolabeled PCR product corresponding to the 570-bp P_{4532} fragment was digested by the restriction enzymes indicated on top. Left panel, untreated PCR product; middle panel, PCR product treated with the Dam methylase; right panel, PCR product incubated with purified Fur (20 nM) prior to Dam methylation. Molecular weight markers (in base pairs) are indicated on the left. The sizes of the digestion products (in base pairs) are indicated on the left. The sizes of the digestion sof restriction sites and sizes of expected DNA fragments. (B) The P_{4532} promoters isolated from pGE573 vectors carrying the P_{4532} -lacZ fusion purified from the EAEC wild-type strain (WT) or its isogenic *dam* or *fur* mutant strain or from the WT strain treated with 2,2'-dipyridyl were digested by the restriction enzymes indicated on top. Molecular weight markers (in base pairs) are indicated on the right. The dashed lines indicate rorganization of the lanes from the same gel. See Fig. S2 for positions of restriction sites and sizes of expected DNA fragments.

results show that Dam and Fur have additive negative effects on regulation at the P_{4532} promoter and that the contribution of Dam is masked in the presence of Fur. Based on these results, we hypothesized that GATC-32 methylation affects RNAP binding. A Dam-methylated P_{4532} fragment was subjected to EMSA with the reconstituted σ^{70} -RNAP complex. As shown in Fig. 4C and D, σ^{70} -RNAP binding was diminished on the methylated P_{4532} fragment.

Fur-Dam competition at the P_{4532} **promoter.** The observation that the Dam effect was masked by Fur *in vivo* raised the idea that, similar to the P_{sci1} situation, Fur binding to the Fur-32 box prevents Dam methylation of the GATC-32 site. To test this hypothesis, *in vitro* and *in vivo* assays were conducted.

(i) Fur binding at the P_{4532} promoter prevents GATC-32 methylation *in vitro*. To test the impact of Fur binding on GATC-32 methylation *in vitro*, we added purified Dam methylase to radiolabeled P_{4532} fragments preincubated or not preincubated with purified Fur protein. The P_{4532} fragments were then used for enzymatic digestion using enzymes that cleave GATC motifs (Fig. S2). We used the fortuitous fact that the GATC-32 site is part of a larger palindromic sequence, TGATCA, which is the target for Bcll, a restriction enzyme that is sensitive to Dam methylation (Fig. S2). In addition to GATC-32, the P_{4532} fragment contains a GATC site at position 149 (GATC¹⁴⁹) that does not overlap a Fur box (Fig. S2). Figure 5A shows that, as expected, incubation with the Dam methylase caused methylation of the GATC sites as P_{4532} is cleaved into three fragments when incubated with Dpnl, an enzyme that specifically recognizes methylated GATC motifs. In agreement with this result, P_{4532} was resistant to Mbol and Bcll, two enzymes that are sensitive to GATC adenine methylation (Fig. 5A, middle panel).

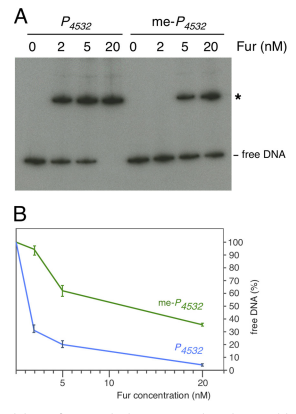


FIG 6 GATC-32 methylation influences Fur binding on P_{4532} . (A) Electrophoretic mobility shift assay of the unmethylated (P_{4532}) or methylated (me- P_{4532}) P_{4532} fragment with the indicated concentration of purified Fur. (B) Densitometry analysis of Fur binding on the unmethylated or methylated P_{4532} fragment, represented as free P_{4532} DNA as a function of Fur concentration.

When the P_{4532} fragment was preincubated with Fur, only the GATC¹⁴⁹ site was digested by Dpnl. In contrast, only the GATC-32 site was digested by Mbol or Bcll (Fig. 5A, right panel). These experiments demonstrate that in the presence of Fur, GATC¹⁴⁹ is methylated whereas GATC-32 is not, suggesting that Fur protects GATC-32 methylation by steric occlusion.

(ii) Fur binding at the P_{4532} promoter prevents GATC-32 methylation *in vivo*. The methylation status of the P_{4532} GATC sites was then tested *in vivo*. The pGE573 plasmid bearing the P_{4532} -lacZ fusion was extracted from various genetic backgrounds, the EcoRI-BamHI fragment comprising the P_{4532} promoter was purified and the methylation state of GATC-32 was assessed by restriction. In the WT strain grown in LB medium, the Mbol and BcII enzymes cleaved GATC-32 (Fig. 5B, left panel), revealing that this site is unmethylated. The absence of methylation is likely due to the presence of Fur bound to the Fur box overlapping with GATC-32, as GATC-32 was methylated in the *fur* isogenic background (Fig. 5B, right panel) or when WT cells were grown in the presence of the 2,2'-dipyridyl iron chelator (Fig. 5B, third panel from left).

Taken together, the results of the *in vitro* and *in vivo* Dam methylation assays demonstrate that Fur binding on the Fur-32 box prevents access of the Dam methylase to the GATC-32 site in iron-rich conditions. In contrast, Fur repression is relieved in iron limiting conditions and the GATC-32 site is then methylated.

(iii) GATC-32 Dam methylation decreases the affinity of Fur for the P_{4532} promoter. The observation that the GATC-32 site is methylated once Fur repression is relieved raised the question of whether methylation of the GATC-32 motif interferes with Fur binding. We therefore performed mobility shift assays with Fur using the P_{4532} fragment, methylated by Dam *in vitro*. Figure 6 shows that methylation of GATC-32 caused a significant decrease in the affinity of Fur for the P_{4532} promoter.

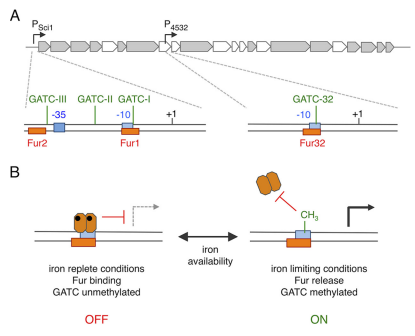


FIG 7 Schematic representation of *sci1* gene cluster regulation. (A) The *sci1* T6SS gene cluster is represented on top with the location of the main (P_{Sci1}) and internal (P_{4532}) promoters. Expanded genetic architectures of these promoters are shown below: +1, transcriptional start; -10 and -35 transcriptional elements (blue); Fur binding box (orange); Dam methylation GATC site (green). (B) Model of regulation of the *sci1* main and internal promoters by Fur and Dam. In iron-replete conditions (left), a Fur dimer (hexagons) complexed to iron (dots) is bound to the Fur box, preventing methylation of the GATC site and access by RNA polymerase. Expression from the promoter is repressed (off state). In iron-limiting conditions (right), Fur is released from the promoter, allowing GATC methylation by Dam and binding of RNA polymerase. Expression from the promoter is turned on (on state).

To summarize, we report in this study the presence of an internal promoter within the sci1 T6SS gene cluster of enteroaggregative E. coli. The presence of internal promoters that serve as transcriptional restarts or that are necessary for ensuring proper stoichiometric production is common in large gene clusters. It has been well documented for gene clusters encoding amino acid synthesis pathways such as histidine, tryptophan, threonine, or branched-chain amino acids (63–67). More recently, an internal promoter within the gene cluster encoding the ESX-3 type VII secretion system has been identified in Mycobacterium smegmatis (68). Here, we show that this internal promoter, P_{4532} , is under the control of a regulatory mechanism similar to that controlling the main promoter (Fig. 7). Expression from the P_{4532} promoter is repressed by the Fur protein, which binds to a Fur box overlapping with the -10 transcriptional element. In addition, a GATC site, GATC-32, which is a target of the Dam methylase, overlaps with the Fur-binding box. In iron-rich conditions, Fur binding to the promoter prevents methylation of this motif. However, during iron starvation, Fur removal allows methylation of the GATC-32 site and the methylation decreases the affinity of Fur for its binding box. Therefore, Fur controls the switch between on and off expression, whereas Dam methylation stabilizes the on phase (Fig. 7). This mechanism is therefore similar to that previously reported for the sci1 main promoter (62). However, differences can be noticed. First, the level of methylation and the activity of the Dam methylase might be slightly different on the main and the internal promoters, as the sequences flanking the GATC motifs have different AT content. Indeed, sequences flanking Dam sites have been previously shown to modulate the catalytic activity or the processivity of Dam (69). Second, an \sim 13-fold derepression of the internal promoter is observed in the absence of Fur, while a >25-fold derepression was observed for the main promoter (62). These results are consistent with the lower degree of consensus for the Fur-32 box compared to the Fur box overlapping with the -10 element of the main promoter

(Fig. 2C) and with the potential cooperativity of the two Fur-binding boxes at the main promoter (62).

The role of the Dam methylase in transcriptional gene regulation is well documented. In addition to its role in mismatch repair and replication initiation, Dam is involved in epigenetic control of the expression of many genes, including genes encoding type III secretion systems, adhesins, or fimbriae or those involved in lipopolysaccharide modifications (for reviews, see references 70 to 72). GATC sites can be found in intergenic regions, and in some cases these sites overlap transcriptional elements such as the -10 element (73). Hence Dam-dependent methylation may directly impact transcription. However, in most cases, GATC sites found in promoter regions do not overlap transcriptional elements, but rather overlap regulator-binding boxes. In these cases, the methylation status may control binding of the regulator, and reciprocally, regulator binding may prevent methylation of certain GATC sites. Several studies have reported competition between Dam-dependent methylation and regulator fixation, such as the OxyR repressor at the *agn43* promoter, or the Lrp repressor at the *pap* operon promoter (74–76). In general, competition between methylation and regulator binding results in the transition between off and on expression phases (72).

In conclusion, the *sci1* gene cluster is subjected to Fur-Dam regulation, and a transcriptional restart occurs after the eighth gene of the operon. Further experiments will be necessary to define whether this restart is necessary because transcription of the mRNA from the initial promoter stops before the last gene, or because the distal part of the operon requires additional copies of mRNA for proper stoichiometry.

MATERIALS AND METHODS

Bacterial strains, plasmids, medium, and growth conditions. *E. coli* K-12 strain DH5α was used for all cloning procedures. The EAEC strains used in this study are all derivatives of strain 17-2 and have been previously described (62). The plasmid-borne P_{4532} -*lacZ* fusion was engineered by ligating a blunt-end 570-bp fragment encompassing the 4532 promoter (corresponding to bases – 450 to +120, respective to the *EC042_4532* transcriptional start site [nucleotides 4892656 to 4893121], amplified from EAEC 17-2 chromosomal DNA using oligonucleotides 5'-CGCACCATGATCGTCTCTGTATCGC and 5'-CTGAAACGAAC TGCTCATGGCTCTCT) into the Smal-linearized pGE573, a vector that carries a promoterless *lacZ* gene (77). In this construct, the *lacZ* gene is under the control of the P_{4532} promoter. Proper insertion, orientation, and sequence of the fragment into the pGE- P_{4532} plasmid were verified by restriction, PCR, and DNA sequencing (MWG). *E. coli* cells were routinely grown in Luria broth (LB) or Sci1-inducing medium (SIM; M9 minimal medium supplemented with glycerol 0.25%, vitamin B₁ 200 µg · ml⁻¹, Casamino Acids 40 µg · ml⁻¹, MgCl₂ 2 mM, CaCl₂ 0.1 mM, and LB [10% vol/vol] [62]) supplemented with antibiotics when necessary (kanamycin 50 µg · ml⁻¹, ampicillin 100 µg · ml⁻¹ for K-12, or 200 µg · ml⁻¹ for EAEC).

RNA purification. EAEC total RNAs were extracted using the PureYield RNA midiprep system (Promega) from 8×10^9 cells grown in SIM and harvested in exponential growth phase (optical density at $\lambda = 600$ nm [OD₆₀₀] ~0.8). RNAs were eluted with 1 ml of water, cleared with DNase I (Ambion), and precipitated overnight at – 80°C by ammonium sulfate-ethanol procedures. The RNA pellet was washed and resuspended in 45 μ l of nuclease-free water. RNA quality and integrity were tested on agarose gels and by the absorbance ratio at $\lambda = 260/280$ nm. The absence of DNA contamination was further tested by PCR using 35 cycles of amplification. Quantifications gave an average RNA concentration of 70 μ g · ml⁻¹. Total RNAs were then subjected to reverse transcription-PCR (Access RT-PCR; Promega) or transcriptional +1 mapping (5' RACE; Invitrogen).

Reverse transcription-PCR. The reverse transcription and PCR assays were performed with the one-tube procedure, using the Access RT-PCR system (Promega), with 200 ng of total RNA and oligo-nucleotides allowing amplification of 550- to 750-bp regions overlapping the two contiguous genes (Fig. 1A) (primer sequences available upon request), following the supplier's guidelines. Briefly, both reverse transcriptase and Tfl *Taq* polymerase were added in each tube. The reverse transcription was carried out for 45 min at 45°C, and, after inactivation of the reverse transcriptase at 94°C for 5 min, a 30-cycle PCR was performed (denaturation at 94°C for 30 s, annealing at 55°C for 40 s, and amplification at 68°C for 50 s). As negative controls to test for DNA contamination, RT-PCRs were also performed in the absence of reverse transcriptase. As positive controls, the regions overlapping the two contiguous genes were amplified from 30 ng of genomic DNA.

5' RACE assay. Total RNAs (80 μ g · ml⁻¹) were subjected to transcriptional +1 mapping using the 5' RACE system (Invitrogen).

β-Galactosidase assays. β-Galactosidase activity was measured by the method of Miller (78) on whole cells harvested at an OD₆₀₀ of 0.8. Reported values represent the average from technical triplicates from three independent biological cultures, and standard deviations are shown on the graphs.

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Protein purification. The Fur and NtrC proteins were purified as described previously (62, 79). The σ^{70} -saturated RNAP holoenzyme was purchased from USB Corp. The Dam methylase and restriction enzymes were obtained from New England BioLabs and used as recommended by the manufacturer.

Electrophoretic mobility gel shift assay and Dam methylation assays. DNA radiolabeling, EMSA, Fur/RNAP competition EMSA, and *in vivo* and *in vitro* Dam methylation assays have been performed as previously described (62).

SUPPLEMENTAL MATERIAL

Supplemental material is available online only. **SUPPLEMENTAL FILE 1**, PDF file, 0.1 MB.

ACKNOWLEDGMENTS

We thank Emmanuelle Bouveret and Mireille Ansaldi for sharing strains, plasmids, and protocols; Laure Journet and the members of the Cascales, Lloubès, Bouveret, and Sturgis research groups for insightful discussions; and Isabelle Bringer, Annick Brun, and Olivier Uderso for technical assistance.

This work was supported by grants from the Agence Nationale de la Recherche to E.C. (ANR-10-JCJC-1303-03 and ANR-14-CE14-0006-02). Work in the E.C. laboratory is supported by the CNRS, the Aix-Marseille Université, the Fondation pour la Recherche Médicale (DEQ20180339165), and the Fondation Bettencourt-Schueller. Y.R.B. was a recipient of a doctoral fellowship from the French Ministry of Research.

Author contributions. Y.R.B. and E.C. conceived the study and designed the experiments. Y.R.B. performed all *in vivo* and *in vitro* experiments, with the help of C.S.B. for RNA analyses. Y.R.B. and E.C. analyzed the data. E.C. wrote the manuscript.

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