Repression of \textit{btuB} gene transcription in \textit{Escherichia coli} by the GadX protein

Guang-Sheng Lei\textsuperscript{1}, Wan-Jr Syu\textsuperscript{1}, Po-Huang Liang\textsuperscript{2}, Kin-Fu Chak\textsuperscript{3}, Wensi S Hu\textsuperscript{4}, Shiau-Ting Hu\textsuperscript{1,5*}

\textbf{Abstract}

\textbf{Background:} BtuB (B twelve uptake) is an outer membrane protein of \textit{Escherichia coli}, it serves as a receptor for cobalamines uptake or bactericidal toxin entry. A decrease in the production of the BtuB protein would cause \textit{E. coli} to become resistant to colicins. The production of BtuB has been shown to be regulated at the post-transcriptional level. The secondary structure switch of 5\textsuperscript{'} untranslated region of \textit{butB} and the intracellular concentration of adenosylcobalamin (Ado-Cbl) would affect the translation efficiency and RNA stability of \textit{btuB}. The transcriptional regulation of \textit{btuB} expression is still unclear.

\textbf{Results:} To determine whether the \textit{btuB} gene is also transcriptionally controlled by trans-acting factors, a genomic library was screened for clones that enable \textit{E. coli} to grow in the presence of colicin E7, and a plasmid carrying \textit{gadX} and \textit{gadY} genes was isolated. The \textit{lacZ} reporter gene assay revealed that these two genes decreased the \textit{btuB} promoter activity by approximately 50\%, and the production of the BtuB protein was reduced by approximately 90\% in the presence of a plasmid carrying both \textit{gadX} and \textit{gadY} genes in \textit{E. coli} as determined by Western blotting. Results of electrophoretic mobility assay and DNase I footprinting indicated that the GadX protein binds to the 5\textsuperscript{'} untranslated region of the \textit{btuB} gene. Since \textit{gadX} and \textit{gadY} genes are more highly expressed under acidic conditions, the transcriptional level of \textit{btuB} in cells cultured in pH 7.4 or pH 5.5 medium was examined by quantitative real-time PCR to investigate the effect of GadX. The results showed the transcription of \textit{gadX} with 1.4-fold increase but the level of \textit{btuB} was reduced to 57\%.

\textbf{Conclusions:} Through biological and biochemical analysis, we have demonstrated the GadX can directly interact with \textit{btuB} promoter and affect the expression of \textit{btuB}. In conclusion, this study provides the first evidence that the expression of \textit{btuB} gene is transcriptionally repressed by the acid responsive genes \textit{gadX} and \textit{gadY}.

\textbf{Background}

BtuB (B twelve uptake) is a 614 amino acid outer membrane protein of \textit{Escherichia coli}. It is responsible for the uptake of cobalamins [1], such as vitamin B\textsubscript{12} including cyanocobalamin, hydroxocobalamin, methylcobalamin, and adenosylcobalamin[2]. It also serves as the receptor for bacteriophage BF23 [3]. The synthesis of the BtuB protein in \textit{E. coli} is regulated at the translational level by adenosylcobalamin (Ado-Cbl) which is produced by the BtuR protein (CobA in \textit{Salmonella typhimurium} and CobO in \textit{Pseudomonas denitrificans}) [4-6]. BtuR is an ATP:corrinoid adenosyltransferase and converts cobalamins to Ado-Cbl [4]. In the presence of Ado-Cbl, the stability of the \textit{btuB} mRNA is reduced with a half-life of only 2 - 4 minutes [7]. In addition, Ado-Cbl binds to the leader region (5\textsuperscript{'} untranslated region, 5\textsuperscript{'} UTR) of the \textit{btuB} mRNA and suppresses its translation [8,9]. A 25-nucleotide sequence designated as the B\textsubscript{12}-box located +138 - +162 nucleotides downstream from the transcription initiation site of \textit{btuB} in \textit{E. coli} has been suggested to be the binding site of Ado-Cbl [10]. A B\textsubscript{12}-box is also present in the 5\textsuperscript{'} UTR of both \textit{btuB} and \textit{cbiA} genes of \textit{S. typhimurium} [11]. The \textit{btuB} gene of \textit{S. typhimurium} is highly homologous to that of \textit{E. coli}. The CbiA protein is a cobyrinic acid a, c-diamide synthase using cobyrinic acid as substrate [10,12]. Binding of Ado-Cbl to the 5\textsuperscript{'} UTR of the mRNAs of these genes may interfere with ribosome binding and thus decrease their translation [7-9,13].
It is unknown whether BtuB synthesis is also controlled by regulatory proteins at the transcriptional level. Results of this study suggest that GadX (Glutamic acid decarboxylation) is a transcriptional regulator of btuB. GadX has been shown to suppress the expression of perA encoded by a plasmid of enteropathogenic E. coli [14], but activate gadX, gadA, gadB, and gadC in response to acid stress [15-19]. GadA and GadB are isozymes of glutamate decarboxylases that convert glutamate to γ-aminobutyric acid (GABA) which is then exported by the antiporter protein GadC [20,21]. An intracellular proton is consumed during GABA production [22], but the released acid (GABA) which is then exported by the antiporter GadC [20,21]. An intracellular proton is consumed during GABA production [22], but the released GABA, which is less acidic than glutamate, provides local buffering of the extracellular environment.

The expression of gadX is activated by the alternative sigma factor RpoS during the stationary phase of growth [15,19,21], but is repressed during the exponential phase by the nuclear protein H-NS due to its binding to the gadX promoter or its destabilizing effect on RpoS [23-25]. However, the acid stress increases the RpoS level and thus induces gadX expression even during the exponential phase of growth [26]. GadW, like GadX, belongs to the family of AraC-like regulators. It represses the expression of gadX and inhibits the activation of gadA and gadBC by GadX [15,18,27]. In addition to these trans-acting proteins, the production of GadX is also controlled by gadY that is located between gadX and gadW in an opposite orientation to gadX [28,29]. The gadY gene has no known protein products. It produces three RNA species of 105, 90, and 59 nucleotides with a common 3′ end [28]. The 3′ ends of gadX and gadY RNAs overlap by at least 30 nucleotides and are complementary to each other. Annealing of gadY RNA to the 3′ end of gadX mRNA stabilizes gadX mRNA, resulting in an increased production of the GadX protein [28].

BtuB is also involved in the import of colicins such as colicin E7 (ColE7) [30-34]. ColE7 is composed of three domains responsible for the translocation of ColE7 through the OmpF porin, binding of ColE7 to BtuB, and cleavage of DNA [35,36], respectively. The import of ColE7 is dependent on the Tol (tolerance to colicin) system that is composed of TolQ, TolR, TolA, and TolB proteins [35,36]. Deletion or mutation of BtuB, OmpF, or any of the Tol proteins renders E. coli resistant to ColE7 [33,37,38]. Based on this information, we used a ColE7 resistance assay in this study to search for transcriptional regulators of btuB from a genomic library of E. coli strain DH5α and found that gadX and gadY genes down regulate the expression of btuB.

**Results**

**Screening of genes conferring E. coli resistance to ColE7**

To search for genes that can confer E. coli resistance to ColE7, plasmids in the genomic library were transformed into the ColE7-sensitive E. coli strain DH5α, and the transformants were plated on LB agar plates containing 50 μg/ml of ampicillin and 5.0 ng/ml of His6-tagged ColE7/ImE7. Two colonies were seen after incubation at 37°C overnight. The plasmids of each colony were isolated after culturing in 3 ml LB medium containing 50 μg/ml of ampicillin then retransformed into DH5α. The new transformants were plated on agar plates containing 0, 1.3, 2.6, 3.9, or 5.2 ng/ml of His6-tagged ColE7/ImE7 to confirm their resistance to ColE7. The insert in the plasmid that conferred DH5α resistance to 5.2 ng/ml His6-tagged ColE7/ImE7 was sequenced. A 1,470-bp DNA region on the chromosome at position 3662617 to 3664086 was analyzed that contains both complete gadX and gadY genes. The plasmid was thus named pGadXY (Figure 1).

To determine whether gadX or gadY was responsible for ColE7 resistance, pGadX, pGadY, and pGadXY that contain gadX, gadY, and gadXY, respectively, were separately introduced into E. coli strain DH5α and then assayed for their ability to confer ColE7 resistance. 1 × 10⁵ cells containing pGadX, pGadY, or pGadXY were plated on LB agar containing 1.3, 2.6, 3.9, or 5.2 ng/ml of His6-tagged ColE7/ImE7. Cells containing the vector pGAD10 were also plated to serve as controls. The percent survival of cells containing pGAD10, pGadXY, pGadX, and pGadY in the presence of 1.3 ng/ml of His6-tagged ColE7/ImE7 were 41.7, 95.5, 71.4, and 73.5%, respectively, and 1.5, 63.9, 3.6, and 9.1%, respectively, in the presence of 2.6 ng/ml of His6-tagged ColE7/ImE7. Only pGadXY conferred ColE7 resistance to 3.9 and 5.2 ng/ml of His6-tagged ColE7/ImE7 with 29.1 and 17.1% survival rates, respectively (Table 1).

**Detection of protein whose expression is affected by gadXY**

To investigate the mechanism by which gadXY affects ColE7 resistance, the expression levels of BtuB, TolQ, TolR, TolA, TolB, Pal, and OmpF that are involved in ColE7 import were determined by Western blotting, and BtuB was the only protein found to be affected. Its expression level was reduced by 93% in the presence of gadXY (Figure 2) as determined by densitometry.

**Effect of gadXY on btuB promoter**

To determine whether gadXY affects the transcription of btuB, the β-galactosidase reporter assay was performed. The 461-, 673-, 913-, and 1285-bp DNA fragments (Figure 3) containing the promoter of btuB were fused with the lacZ coding sequence to generate pCB461lacZ, pCB673lacZ, pCB913lacZ, and pCB1285lacZ, respectively. Each of these single copy plasmid together with pGAD10 or pGadXY was transformed into E. coli strain
DH5α. The transformed cells were grown in LB medium with 50 μg/ml of chloramphenicol and ampicillin to OD600~0.8 then assayed for β-galactosidase activity as described by Miller [39]. The β-galactosidase activity of cells containing pGadXY and a pCB derivative with the btuB promoter-lacZ fusion was divided by that of cells containing the control plasmid pGAD10 and the same pCB derivative to determine the percent decrease in btuB promoter activity in the presence of gadXY. The btuB promoter in the 461-, 673-, 913-, and 1285-bp DNA fragment was found to be decreased by 45.7, 47.1, 54.5, and 56.7%, respectively in the presence of gadXY, and was about 6 fold more active in the 1285-bp fragment than in other fragments (Table 2).

To investigate the effect of gadX or gadY alone on the promoter activity of btuB, the same experiment was performed using DH5α cells containing pCB1285lacZ and pGAD10, pGadXY, pGadX, or pGadY. The β-galactosidase activity of cells containing pCB1285lacZ and pGadXY, pGadX, or pGadY was compared to those containing pGAD10 and pCB1285lacZ. The results indicated that btuB promoter activity was decreased 20.5% by

### Table 1 Effects of gadXY, gadX, and gadY on ColE7 resistance

| ColE7 conc./Bacteria | pGAD10/DH5α | pGadXY/DH5α | pGadX/DH5α | pGadY/DH5α |
|----------------------|-------------|-------------|-------------|-------------|
| 1.3 ng/ml            | 41.7%       | 95.5%       | 71.4%       | 73.5%       |
| 2.6 ng/ml            | 1.5%        | 63.9%       | 3.6%        | 9.1%        |
| 3.9 ng/ml            | 0           | 29.1%       | 0           | 0           |
| 5.2 ng/ml            | 0           | 17.1%       | 0           | 0           |

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gadX and 20.3% by gadY, but was decreased 54.4% by gadXY (Table 3).

Binding of GadX to btuB promoter
GadX has been shown to be a DNA binding protein and can bind to the gadA or the gadB promoter. To determine whether GadX also binds to the btuB promoter, the DNA mobility shift assay was performed. Only GadX was assayed because gadY does not encode any proteins. The 461-bp DNA fragment containing the btuB promoter was labeled with $^{32}$P and incubated with 2, 4, or 6 pmoles of purified GadX protein (MalE-GadX) that was fused to the maltose binding protein. The DNA fragment containing the promoter of gadA or gadB was used as the positive control for GadX binding, and the DNA fragment containing the pal promoter was used as the negative control. As shown in Figure 4, DNA band shift was observed on gadA and gadB promoter fragments but not on the negative control. Band shift was also observed on the btuB promoter fragment in a dose-dependent manner, indicating that GadX binds to the btuB promoter.

Identification of binding sequence of GadX on btuB promoter
DNase I footprinting was then performed to determine the binding sequence of GadX on the btuB promoter. The 461-bp DNA fragment containing the btuB promoter was labeled with $^{32}$P and incubated with 0, 2, 4, or 8 pmoles of purified MalE-GadX protein and then digested with DNase I. Results shown in Figure 5 revealed three MalE-GadX protein binding sites that included nucleotide positions +56 - +81 (I), +96 - +105 (II), and +183 - +188 (III).

Table 2 Effect of gadXY on btuB promoter

| Plasmid                | $\beta$-galactosidase activity | % inhibition |
|------------------------|-------------------------------|-------------|
| (A): pGAD10 + pC-lacZ  | 0                             | 0           |
| (B): pGadXY + pC-lacZ  | 0                             | 0           |
| (A): pGAD10 + pCB461lacZ| 8.4 ± 0.2                     | 45.7        |
| (B): pGadXY + pCB461lacZ| 3.5 ± 0.2                     | 47.1        |
| (A): pGAD10 + pCB673lacZ| 7.2 ± 0.1                     | 47.1        |
| (B): pGadXY + pCB673lacZ| 3.8 ± 0.1                     | 47.1        |
| (A): pGAD10 + pCB913lacZ| 4.8 ± 0.2                     | 54.5        |
| (B): pGadXY + pCB913lacZ| 2.2 ± 0.5                     | 56.7        |
| (A): pGAD10 + pCB1285lacZ| 3.75 ± 0.7                    | 56.7        |
| (B): pGadXY + pCB1285lacZ| 16.2 ± 0.5                    | 56.7        |

*Miller unit.

*Calculated according to the following equation: $\frac{1 - \beta$-galactosidase activity of (B) - $\beta$-galactosidase activity of (A)] \times 100%.
(II) and +123 - +137 (III) on the 5’ untranslated region of btuB.

Decreased btuB expression under acidic conditions
It is known that gadX and gadY are highly expressed under acidic environments in stationary phase [15-19,28]. To determine whether the expression of btuB is also repressed in an acidic condition, wild type BW25113 cells were cultured in LB medium pH 7.4 or buffered with 100 mM MES pH5.5. Stationary phase cells grown in different culture media were collected and then assayed for the transcriptional level of btuB by quantitative real-time PCR. The cDNA amplification comparison results showed the transcription of gadX with 1.4-fold increase but the level of btuB was reduced to 57% (Table 4).

Discussion
Although it has been suggested that the expression of btuB in E. coli is also regulated at the transcriptional level, the trans-acting regulators of btuB had not been identified [40,41]. In this study, we used the ColE7 resistance assay to search for such regulators and found that both gadX and gadY genes can repress the production of BtuB rendering E. coli DH5α cells resistant to ColE7. Introduction of pGadX, which contains a gadX gene, into DH5α cells caused 3.6% of the cells to become resistant to 2.6 ng/ml of ColE7. In a similar experiment, pGadY which contains the gadY gene enabled 9.1% of the cells to grow in the presence of the same concentration (2.6 ng/ml) of ColE7 (Table 1). Although gadY does not encode any proteins, it had a greater effect on making E. coli resistant to ColE7 than gadX. This is probably due to the binding of gadY RNA derived from pGadY to the gadX mRNA produced by the gadX gene in the chromosome. This binding stabilizes gadX mRNA so that more GadX protein is produced to suppress the production of BtuB, making the cells resistant to ColE7. The greatest effect (63.9% survival in 2.6 ng/ml ColE7) on ColE7 resistance was seen when pGadXY, which contains both gadX and gadY genes, was introduced into the cells. Since pGadXY is a high copy number plasmid, more gadX and gadY mRNAs would be produced and thus more GadX protein would be synthesized to suppress BtuB synthesis. However, excess GadX had adverse effects as over expression of GadX with a strong promoter, such as the T5-lacO promoter, was found to have toxic effect to E. coli [19]. Therefore, expression of gadX and gadY in this study was driven by their own promoters.

Since GadX is a known transcriptional regulator [14-16,18,19,42], the decrease in BtuB expression is due

| Plasmids   | β-galactosidase activity | % inhibition |
|------------|--------------------------|--------------|
| (A): pGAD10/pC-lacZ | 0                        | 0%           |
| (B): pGAD10/pCB265lacZ | 48.8 ± 3.9               | 48.8%        |
| (C): pGadXY/pCB265lacZ | 22.3 ± 0.7                | 54.4%        |
| (D): pGadX/pCB265lacZ  | 38.9 ± 2.6                | 20.5%        |
| (E): pGadY/pCB265lacZ  | 38.9 ± 2.0                | 20.3%        |

*Miller unit.
*Calculated according to the following equation: 1 - [β-galactosidase activity of (C), (D), or (E) - β-galactosidase activity of (A)] × 100%.
to its transcriptional repression by GadX. Our data showed that the *btuB* promoter activity was reduced by approximately 50% in the presence of *gadXY* (Table 2), most likely due to binding of GadX to the 5′ untranslated region of *btuB* as DNase I footprinting experiment revealed binding of GadX to nucleotides positions +56 - +81 (I), +96 - +105 (II) and +123 - +137 (III) downstream from the *btuB* transcription initiation site (Figure 5).

From the sequence alignment of GadX binding sites on *btuB*, *gadA*, and *gadBC* regulatory regions,[42] we found that sequence in the region I (the 31 nucleotides) has 62.5% identity (+52-AGCGGTAAGGAAAGGTGCGATGATTGCGTTAT-+82, underlined nucleotides indicate the protected region) with *gadBC*. Sequence in the region III (the 26 nucleotides) has 60.7% identity (+106-AAGTCATCATCTCTTGATATCTTAGATA-+133, underlined nucleotides indicate the protected region) with *gadA* regulatory region. From the footprinting result, the GadX binding sites on 5′ untranslated region of *btuB* share only partial homology with the 42 nucleotides consensus sequence which was reported by Tramonti *et al.*[42]. The sequence analysis also revealed the *btuB* expression was regulated by the binding of GadX from the *btuB* transcription initiation site (Figure 5).

Table 4: Fold changes of transcripts of *gadX* and *btuB* attribute to different pH medium (pH 5.5/pH 7.4) from early stationary phase

| Gene | Fold increasea |
|------|----------------|
| *gadX* | 1.43 ± 0.07 |
| *btuB* | 0.57 ± 0.13 |

*aExperiments were performed in triplicate and the data are presented as mean values ± SD.*

Figure 5: Binding sequence of GadX on *btuB* promoter. (A) The 461-bp DNA fragment containing *btuB* promoter was labeled at 5′ end with 32P, incubated with 0, 16, 24, 32, or 40 pmoles of MalE-GadX, and then subjected to DNase I footprinting. A Sanger’s DNA sequencing reaction was also done on the 461-bp fragment to reveal GadX binding sequences. All reactions were electrophoresed in a 6% urea-acrylamide gel, and the DNA bands were detected by autoradiography. The GadX bound regions are indicated with vertical lines, and the binding sequence of GadX are shown. (B) Sequence of the *btuB* promoter region. The boxed sequences are GadX binding sequences determined by the DNase I footprinting. The shaded sequences are -10 and -35 regions of the *btuB* promoter. The initiation codon of *btuB* is underlined.
on its 5’ untranslated region. Binding of transcriptional regulator to the 5’ untranslated region to regulate gene expression is also seen in the glp regulon of *E. coli*, in which four repressor binding sites are located at -41 to -60, -9 to -28, +12 to -8, and +52 to +33 of the glpACB genes [43]. In addition, two IHF binding sites are present downstream from the glpT transcriptional start site at positions +15 to +51 and +193 to +227 [44].

In the *btuB* promoter assay experiment, different lengths of DNA fragments containing *btuB* promoter were fused to *lacZ*. The minimum length of DNA fragment with *btuB* promoter activity was 461 bp spanning -219 to + 242 nucleotides relative to the translation initiation site of *btuB*. No significant difference in promoter activity was observed when the 5’ end of these fragments was extended to -671. However, a 6 fold (37.5 vs. 6.4 β-galactosidase units, Table 2) increase in promoter activity was detected when the DNA fragment was extended to -1043 with a total length of 1,285 bp as compared to that of the 461-bp fragment. It is very likely that a certain transcription regulator binds to the region between -1043 and -671 and enhances the expression of *btuB*. The β-galactosidase activity in these assays was not very high because the *lacZ* fusions were constructed using the single copy plasmid vector pCC1Bac™ (Epitcentre). The purpose of using the single copy number plasmid in this experiment was to mimic the natural state of *btuB* expression in *E. coli*. In fact, the promoter activity of *btuB* is lower than other membrane protein, we have determined the *ompC* promoter activity, under the same test condition the Miller’s Units of *lacZ* driven by *ompC* promoter is 8 folds higher than that of *btuB* (data not shown).

Although the results of footprinting and reporter assay revealed that the GadX binding sites on *btuB* 5’ untranslated region share only partial homology with the GadX binding consensus sequence[42] and showing 50% down regulation in the reporter assay, the expression of *btuB* was indeed controlled by GadX.

Both *gadX* and *gadY* genes belong to a group of genes that are induced by acid stress under stationary growth phase [44]. Our data showed that the expression of *btuB* was indeed reduced when *E. coli* cells were grown to stationary phase in an acidic medium as compared to the same cells grown in neutral medium (Table 4). The reduction in the production of *btuB* in response to acid stress probably represents a physiological regulatory mechanism of bacteria facing environmental challenges such as low pH. Under stress environment, bacteria need to alter their metabolism to adapt to the environmental change. The transportation of cobalamin by BtuB receptor is driven by proton motive force (PMF) [45]. Since the PMF of bacteria is increased at low pH [46], the cobalamin transportation may be enhanced by increased PMF. The higher concentration of cobalamin in cytoplasm will initiate riboswitch mechanism to repress the translation of BtuB receptor, which is in good accord with the repression of *btuB* transcription by the acid-induced GadX for bacteria to decrease the production of BtuB in response to this acidic stress.

**Conclusions**

Through biological and biochemical analysis, we have demonstrated the GadX can directly interact with *btuB* promoter and affect the expression of *btuB*. When bacteria were grown to stationary phase in an acidic medium, the increased *gadX* expression would repress the *btuB* transcription to help bacteria to adapt to acidic shock. In conclusion, this study provides the first evidence that the expression of *btuB* gene is transcriptionally repressed by the acid responsive genes *gadX* and *gadY*.

**Methods**

**Plasmid constructions**

To produce the His6-tagged ColE7/lm7 protein complex for the ColE7 resistance assay, pQE30ColE7-lm7 was constructed. The *cea7-cei7* genes encoding the colicin E7 and immunity proteins, that form an active ColE7 complex, were amplified from plasmid K317 [47] by PCR using primers F/cea7-BamHI and R/cei-PstI (Table 5). The 1,996-bp PCR product thus generated was inserted between BamHI and PstI sites of pQE30 (Qiagen), fusing the His6-tag to the N terminus of ColE7. For searching transcriptional regulators of *btuB*, a genomic library of *E. coli* K-12 strain constructed with the pGAD10 vector (Figure 1) was purchased from Clontech (catalog number XL4001AB) and transformed into *E. coli* strain DH5α. The plasmid pGadXY (Figure 1) was isolated from the library in this study. To investigate the effect of GadX on *btuB* expression, pGadX was constructed as follows. A 1,077-bp DNA fragment containing *gadX* was generated by PCR using pGadXY (Figure 1) as the template and the MATCHMAKER 5’ insert screening sequence 5’-TACCACCTACAATGGATG-3’ (Clontech) and R/gadX-PstI (Table 5) as primers. This 1.1-kb PCR fragment was inserted into pGEM-T Easy (Promega) by TA cloning, generating pGEMgadX. The 1.1-kb fragment was then isolated from pGEMgadX by EcoRI digestion and inserted into the EcoRI site of pGAD10, resulting in pGadX (Figure 1). To investigate the effect of *gadY* on *btuB* expression, pGadY (Figure 1) was constructed by deleting the *NcoI*-DraI fragment containing *gadX* from pGadXY.

To assay *btuB* promoter activity, DNA fragments (461, 673, 913, and 1,285 bp) containing different portions (Figure 3) of the *btuB* promoter was fused to *lacZ*. These fragments were generated by PCR using primers.
Table 5 Oligonucleotide primers used in this study

| Primer Name | Sequence 5’ – 3’ |
|-------------|------------------|
| F/cea7- BamHI | GGATCCATGACGCGTGGAGATGGACC |
| R/cea7- PstI | CTGACGTCGACGGCTTTTAATTACCC |
| F/btuB-219-Xbal | GGCTCTAGAAGACGCGTCTATGATTTGG |
| R/btuB+242-HindIII | GCGGACGGTATCTGTTTTGAGATGGACC |
| F/btuB-767 | GTTACGGTIGCCTGACGATAC |
| R/btuB-1087 | TCGATAGATGCGCCGATAC |
| F/btuB-431-Xbal | GCTCTAGAACGGGATACCCATATTTGG |
| R/btuB-671-Xbal | GCTCTAGATCATCTTCTCTCCC |
| F/btuB-1043-Xbal | GCTCTAGACCCGCTCGGCGGG |
| R/lacZ | TTATTTTTGACACCAGACC |
| F/gadA-176 | GATGCGCGGAAAGACAAC |
| R/gadA+77 | CGTGAAAGCAGATGACC |
| F/gadB-173 | AATAACAGCATAAAAAC |
| R/gadB+77 | CGTGAAAGCAAGTATGG |
| R/pal-Xbal | TGATAGATGCCGGTATTACG |
| R/pal-HindIII | AAGCTTATTATTATGATTTATTTG |
| F/gadX-BamHI | GGATCCATGACGCGTGGAGATGGACC |
| R/gadX-PstI | CTGACGTCGACGGCTTTTAATTACCC |
| F/gadX-393 | TATACCGCTGTCTGAAAG |
| R/gadX-726 | TCGCTCTGAATCTTCTGG |
| F/ma-A483 | CGTTACCCGCGAAGAAGC |
| R/mr-A808 | GTGGACTACCGGAGATCTAACTC |

The underlined letters indicate restriction sites.

F/btuB-219-Xbal, F/btuB-431-Xbal, F/btuB-671-Xbal, and F/btuB-1043-Xbal were amplified with the three’ primer R/btuB +242-HindIII (Table 5). The resulting PCR products were digested with XbaI and HindIII and then inserted into corresponding sites on pKM005 that carries a promoterless lacZ gene [48], generating pKMbtuB461-lacZ, pKMbtuB671-lacZ, pKMbtuB913-lacZ, and pKMbtuB1285-lacZ. To mimic native expression of btuB inserted into pCC1 vector was used generating pKMbtuB461-lacZ, pKMbtuB671-lacZ, pKMbtuB913-lacZ, and pKMbtuB1285-lacZ. The fragments containing btuB promoter and lacZ on pKM005 derivatives were amplified with primers F/btuB-219-Xbal, F/btuB-431-Xbal, F/btuB-671-Xbal, and F/btuB-1043-Xbal paired with the three’ primer R/lacZ (Table 5), and the resulting 3.3, 3.5, 3.74, and 4.1-kb DNA fragments were separately inserted into pGEM-T<sub>Easy</sub> (Promega) by TA cloning. The 3.3, 3.5, 3.74, and 4.1-kb fragments were then isolated from these pGEM-T<sub>Easy</sub> derivatives by NotI digestion and inserted into the NotI site of pCC1 vector, generating pCB<sub>461</sub>-lacZ, pCB<sub>671</sub>-lacZ, pCB<sub>913</sub>-lacZ, and pCB<sub>1285</sub>-lacZ. The plasmid pCB<sub>lacZ</sub> that contains a promoterless lacZ gene inserted into pCC1 vector was used as a negative control. To produce GadX for DNA binding assay, pMalE-GadX that contains maltose-binding protein fused to GadX (MalE-GadX) was constructed. The 825-bp DNA fragment containing gadX was generated by PCR using pGadXY as the template and F/gadX-BamHI and R/gadX-PstI (Table 5) as primers and then ligated between the BamHI and PstI sites of pMAL-C2X (New England Biolab), resulting in pMalE-GadX.

Production of ColE7
To produce the His<sub>6</sub>-tagged ColE7/ImE7 complex, E. coli strain XL1-Blue containing plasmid pQE30<sub>ColE7-ImE7</sub> was cultured in LB medium containing ampicillin (50 μg/ml) and tetracycline (20 μg/ml). When the bacterial growth reached OD<sub>600</sub> ~1.0, IPTG was added to a final concentration of 1 mM. After a 2-hr induction, bacteria were harvested by centrifugation at 6,500 × g for 20 min and then resuspended in HB buffer (20 mM Tris, 150 mM NaCl, 30 mM imidazole, pH 8.0). The resuspended bacteria were lysed with a French Pressure Cell (SLM Instruments, Inc. Urbane, IL), and the cell lysate was centrifuged at 48,000 × g for 1 hour. The clarified supernatant was passed through a ProBond™ nickel-nitrilotriacetic acid resin affinity column (Invitrogen, Carlsbad, CA) to purify the His<sub>6</sub>-tagged ColE7/ImE7 according to manufacturer’s protocol (Invitrogen, Carlsbad, CA).

Antibody preparation for detection of protein whose expression is affected by gadXY
To prepare antibodies against envelope proteins BtuB, TolQ, TolR, TolA, TolB, Pal, and OmpF, the coding region of each protein was fused inframe with His<sub>6</sub>-tag in the plasmid pQE30 (Qiagen), respectively. The His<sub>6</sub>-tagged proteins were then expressed and purified using the same method as described for His<sub>6</sub>-tagged ColE7/ImE7 and sent to the company to prepare polyclonal antibodies. The specificities of these antibodies were confirmed by Western blotting using these antibodies as reported by Pan et. al.[49].

DNA binding assay
The electrophoretic mobility shift assay was performed to investigate binding of GadX to the btuB promoter. To obtain purified MalE-GadX protein, E. coli strain XL-1 Blue containing pMalE-GadX was grown in 100 ml of LB containing ampicillin (50 μg/ml) and 0.2% glucose to OD<sub>600</sub> ~1.0. IPTG was then added to a final concentration of 1 mM. After 2 hr of incubation, the cells were pelleted, resuspended in maltose binding buffer (20 mM Tris-HCl pH 8.0, 200 mM NaCl), and lysed with a French Pressure Cell. The cell lysate was centrifuged at 48,000 × g for 1 hour. The clarified supernatant was subjected to an amylose resin affinity chromatography (New England BioLabs) to purify the MalE-GadX protein.

To make probes for the DNA binding assay, a 461-bp (Figure 3, -219 - +242) DNA fragment containing the


**btuB** promoter was amplified with primers F/btuB-219-XbaI and R/btuB+242-HindIII (Table 5) by PCR. The DNA fragment containing the promoter of *gadA* (-176 to -77, 253 bp) or *gadB* (-173 to -77, 250 bp) was used as the positive control, which were amplified with primer pairs F/gadA-176 and R/gadA+77 and F/gadB-173 and R/gadB+77 (Table 5), respectively, as described by Tramonti et al. [19]. The DNA fragment containing the upstream noncoding region of *pal* was used as the negative control, which was amplified with primers F/pal-XbaI and R/pal-HindIII (Table 5). These DNA fragments were end-labeled with [γ-32P] ATP by T4 polynucleotide kinase (New England BioLabs). The labeled DNA fragments (6 fmol) were incubated with the MalE-GadX protein at room temperature for 20 min in 10 μl of binding buffer [19]. Samples were then loaded on a 5% nondenaturing polyacrylamide gel in 0.5X TBE buffer and electrophoresed for 35 min at room temperature. The gels were then dried and autoradiographed.

**DNase I footprinting**

DNase I footprinting was performed to determine the binding sequence of MalE-GadX on *btuB* promoter as described by Tramonti et al [19]. Thirty μl of reaction mixture that contains 5 ng of 32P-labeled 461-bp *btuB* promoter fragment, various amounts of the MalE-GadX protein, and reaction buffer (40 mM HEPES pH 8.0, 100 mM potassium chloride, and 10 mM magnesium acetate) was incubated at room temperature for 20 min. At the end of the incubation, 0.5 U DNase I (Roche Biochemicals, Indianapolis, IN) was added to each reaction mixture and then incubated at 37°C for 1 min followed by addition of 3 μl of quench solution (0.1% xylene cyanol, 4% SDS, and 50% glycerol) to stop the DNase I digestion. The partially digested product was passed through a Sephadex G25 spin column (GE Healthcare), and the eluate was subjected to 30 cycles of asymmetric PCR (SequiTherm Excel™II, Epicentre) using 5'-end 32P-labeled primer R/btuB+242-HindIII (Table 5). The PCR-generated products were electrophoresed on a 6% sequencing gel. The gel was then dried and autoradiographed. To determine the binding sequence of GadX, the 461-bp *btuB* DNA probe was sequenced by the Sanger’s sequencing method using the 5'-end 32P-labeled primer R/btuB+242-HindIII (Table 5).

**Quantitative Real-Time Polymerase Chain Reaction**

Total RNA of wild type *Escherichia coli* strain BW25113 grown under LB (pH 7.4) or LB/MES (LB buffered with 100 mM MES, pH 5.5) to early stationary phase were isolated using a modified hot-pheno1 extraction method [21]. This was followed by further purification using RNAspin Mini RNA purification kit (GE) to remove contaminating genomic DNA and enhance the quality of RNA. Each cDNA sample was synthesized from 0.1 μg total RNA with specific primers of *rrsA*, *gadX* and *btuB* using RevertAid™ First strand cDNA synthesis kit (Fermentas). Following reverse transcription, specific gene transcription levels were determined by quantitative real-time PCR using the ABI PRISM 7700 Sequence Detection System (Applied Biosystem). Real-time PCR was performed with each specific primer pair using SYBR Green PCR Master mix (MBI). For *rrsA*, primer pair *rrsA* F and *rrsA* R was used; for *gadX*, primer pair *gadX* F and gadX R was used; and for *btuB*, primer pair *btuB* F and *btuB* R was used (Table 5). The levels of 16S rRNA was chosen as the normalizing gene. The expression levels of *gadX* and *btuB* of cells grown in medium with different pH and different growth were compared.

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**Authors’ contributions**

GSL designed and performed most of the experiments, analyzed data and wrote the manuscript. WJS and WSH gave suggestions and analyzed data for this research. WJS and WSH gave suggestions and analyzed data for this research. All the authors have read and approved the final manuscript.

**Competing interests**

The authors declare that they have no competing interests.

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