**Abstract**

IscR, an Fe–S cluster-containing transcriptional factor, regulates genes involved in various cellular processes. In response to environmental stimuli such as oxidative stress and iron levels, IscR switches between its holo and apo forms to regulate various targets. IscR binding sequences are classified into two types: the type 1 IscR box that is specific for holo-IscR binding, and the type 2 IscR box that binds holo- and apo-IscR. Studying *Klebsiella pneumoniae* CG43S3, we have previously shown that iron availability regulates capsular polysaccharide (CPS) biosynthesis and iron-acquisition systems. The present study investigated whether IscR is involved in this regulation. Compared with that in CG43S3, the amount of CPS was decreased in AP001 (∆iscR) or AP002 (∆iscRCA), a CG43S3-derived strain expressing mutated IscR mimicked apo-IscR, suggesting that only holo-IscR activates CPS biosynthesis. Furthermore, a promoter-reporter assay verified that the transcription of *cps* genes was reduced in AP001 and AP002. Purified IscR:His<sub>6</sub> but not IscR<sub>CA</sub>:His<sub>6</sub>, was also found to bind to the predicted type 1 IscR box specifically in the *cps* promoter. Furthermore, reduced siderophore production was observed in AP004 (Δfur-iscR<sub>CA</sub>) but not in AP005 (Δfur-iscR<sub>CA</sub>), implying that apo-IscR activates iron acquisition. Compared with those in AP004, mRNA levels of three putative iron acquisition systems (*fhu*, *iuc*, and *sit*) were increased in AP005, and both purified IscR:His<sub>6</sub> and IscR<sub>CA</sub>:His<sub>6</sub> bound the predicted type 2 IscR box in the *fhuA*, *iucA*, and *sitA* promoters, whereas IscR<sub>CA</sub>:His<sub>6</sub> displayed a lower affinity. Finally, we analyzed the effect of external iron levels on *iscR* expression. The transcription of *iscR* was increased under iron-depleted conditions as well as in AP001 and AP002, suggesting an auto-repression exerted by apo-IscR. Our results show that in *K. pneumoniae*, IscR plays a dual role in the regulation of CPS biosynthesis and iron-acquisition systems in response to environmental iron availability.

**Introduction**

*Klebsiella pneumoniae* is a rod-shaped, gram-negative bacterium that causes community-acquired diseases including pneumonia, bacteremia, septicemia, and urinary and respiratory tract infections that occur particularly in immune-compromised patients [1]. In Asian countries, especially in Taiwan and Korea, *K. pneumoniae* is the predominant pathogen responsible for pyogenic liver abscesses in diabetic patients [2,3,4]. Among the virulence factors identified in *K. pneumoniae*, capsular polysaccharide (CPS) is considered the major determinant for *K. pneumoniae* infections. Pyogenic liver abscess isolates often carry heavy CPS loads that protect bacteria from phagocytosis and from being killed by serum factors [5,6]. Apart from its anti-phagocytic function, *Klebsiella* CPS also promotes bacterial colonization and biofilm formation at infection sites [7,8,9].

Our previous studies have demonstrated that CPS biosynthesis in *K. pneumoniae* is repressed in iron-replete conditions, and this regulation is controlled by an iron uptake regulator (Fur) [10]. Under iron-replete conditions, dimeric Fur in complex with Fe(II) indirectly activates *K. pneumoniae* CPS biosynthesis through transcriptional factors RmpA and RcsA and a small non-coding RNA, RyhB [10,11]. The transcription of *cps* genes is directly regulated by RmpA and RcsA but appears to be indirectly regulated by RyhB. These findings indicate that environmental iron availability influences *K. pneumoniae* CPS biosynthesis through multiple regulators.

To maintain iron homeostasis, Fur acts as a master regulator to control iron transport, storage, and metabolism in many gram-negative bacteria including *K. pneumoniae* [11–13]. We have previously reported that Fur directly represses at least six of the...
eight iron acquisition systems in *K. pneumoniae* CG43S3 [10]. In addition to Fur, the transcriptional regulator IscR plays a crucial role in iron metabolism. IscR regulates the biosynthesis of Fe-S clusters, which are key cofactors of proteins intervening in various cellular processes in bacteria [12,13]. Fe-S clusters can be generally classified into two types, rhombic [2Fe-2S] and cubic [4Fe-4S], which have either ferrous (Fe2+) or ferric (Fe3+) iron and sulphide (S2−) [14,15]. IscR is itself a [2Fe-2S] cluster-containing protein encoded by the first gene of the *isrSUA* operon. The switch between the [2Fe-2S] holo and apo forms of IscR is believed to be influenced by environmental conditions such as oxidative and nitric oxide stress and cellular iron levels [13,16,17,18]. Moreover, holo- and apo-IscR have been shown to be regulated by different target genes, suggesting that the presence of the [2Fe-2S] cluster affects the regulatory specificity of IscR [18,19,20,21].

Transcriptomic analysis has identified 40 genes in 20 predicted operons, which are regulated by IscR under aerobic and anaerobic conditions in *Escherichia coli* [19]. This analysis has also revealed two classes of IscR binding sites (IscR boxes). Type 1 IscR box consists of a 25-bp sequence interacted with holo-IscR, whereas type 2 IscR box consists of a 26-bp sequence interacted with apo-IscR [19]. Furthermore, a detailed analysis of the type 2 IscR box has verified an IscR binding motif for both holo and apo-IscR binding [21].

In this study, we investigated whether IscR participates in the regulation of CPS biosynthesis and the expression of iron acquisition systems in *K. pneumoniae*. We also analysed the expression of *iscR* in response to various iron levels.

**Results**

IscR activates *K. pneumoniae* CPS biosynthesis in an Fe-S cluster-dependent manner

To study whether IscR regulates *K. pneumoniae* CPS biosynthesis, we determined the amounts of K2 CPS in CG43S3 (wild type [WT]) and AP001 (*ΔiscR*) strains. Compared with the WT, AP001 produced significantly lower amounts of CPS (Fig. 1A), suggesting that IscR activates the biosynthesis of CPS. In *K. pneumoniae*, IscR contains three highly conserved cysteine residues (C92, C98, and C104 in *E. coli* IscR) which are thought to coordinate the [2Fe-2S] cluster [20].

To investigate the role of the [2Fe-2S] cluster in IscR regulation of CPS biosynthesis, we created an *iscR* mutant AP002 (*iscRSCA*) by replacing the three cysteines with alanines and tested whether this mutant, which is predicted to encode an IscR lacking an Fe-S cluster, affected CPS biosynthesis. As shown in Fig. 1A, we found that the amount of CPS decreased in the AP002 strain compared with that in WT, indicating that the regulation of IscR required the [2Fe-2S] cluster. Moreover, *iscR* and *iscRSCA* were respectively cloned into pACYC184, to yield pIscR and pIscRSCA, for complementation analysis. Compared with AP001 (pACYC184), AP001 (pIscR) produced a significantly higher amount of CPS, whereas the introduction of pIscRSCA into the AP001 strain did not change the CPS amount (Fig. 1B). These results confirmed that IscR has a positive role in the regulation of CPS biosynthesis and that the presence of the [2Fe-2S] cluster of IscR is essential for this regulation. On the other hand, the CPS amount appeared to obviously increased in AP001 (pIscR), compared with that in WT (pACYC184) (Fig. 1B), which may result from multicycop plasmids used for complementation. Therefore, we also used single copy constructs to complement the *iscR*-deletion (Methods S1), and the result showed that the expression of *iscR*, but not *iscRSCA*, could restore the CPS biosynthesis (Fig. S1).

The K2 *cps* gene cluster of *K. pneumoniae* contains 19 open reading frames (ORFs) organised into 3 transcription units—namely, *orf1–2*, *orf3–15*, and *orf16–17* [22]. To determine the role of IscR in regulating *cps* transcription, we used the reporter plasmids pOrf12 (pOrf1–2::lacZ), pOrf315 (pOrf3–15::lacZ), and pOrf1617 (pOrf16–17::lacZ), each carrying a promoterless lacZ gene transcriptionally fused to the putative promoter region of the K2 *cps* gene cluster [23], to transform the *K. pneumoniae* strains AP006 (ΔIsrZ), AP007 (ΔIsrZ-IscR), and AP008 (ΔIsrZ-iscRSCA). The measurements shown in Fig. 1C reveal that the promoter activity of *orf1–2* and *orf16–17* in AP007 and AP008 was lower than that in AP006 (P <0.01). Additionally, the promoter activity of *orf15* was apparently decreased in AP008 but slightly decreased in AP007 compared with that in AP006. These results indicated that IscR activates the transcription of *cps* genes in an Fe-S cluster-dependent manner.

IscR directly binds the promoter of *galF*

For further investigation of the mechanism of IscR regulation on *cps* transcription, the sequence of the putative IscR binding site was manually analysed in the three promoter regions of the K2 *cps* gene cluster. As shown in Fig. 2A, we found a putative type 1 IscR box with 52% (13/25 bp) homology to the consensus sequence located between −173 bp and −197 bp relative to the translational start codon of *galF* (*orf1* in the K2 *cps* gene cluster). In addition, the putative IscR binding sequence in *PgalF* was highly homologous to the IscR-binding motif (5′-AxxxCCxxAxxxxxx-TAxGxxGxxT-3′) reported by Nesbit et al. [21]. However, no typical IscR binding site was found in the upstream sequence of *wzi* (*orf3* in the K2 *cps* gene cluster) or *manC* (*orf16* in the K2 *cps* gene cluster), suggesting that IscR indirectly regulates the promoter activities of *orf3–15* and *orf16–17*, which remains to be studied. On the other hand, we hypothesised that IscR binds directly to the promoter region of *galF* to activate gene transcription, and we confirmed this by performing an electrophoretic mobility shift assay (EMSA). As shown in the upper panel of Fig. 2B, purified recombinant IscR::His6 protein was able to bind *PgalF* but not *PgalF*-2, in which the region containing the putative IscR binding site was deleted. In addition, compared with that of *IsrZ*:His6, the recombinant [2Fe-2S] clusterless IscR3CA::His6 had reduced *PgalF* binding activity. Furthermore, no obvious interaction between the recombinant IscR proteins and *PgalF*-1*+,* the *galF* promoter lacking only the 25-bp predicted IscR box, was found (the lower panel of Fig. 2B). Besides, *PgalF-1* and *PgalF-1* DNA showed a slightly different mobility in the gel. These results suggested a direct interaction between IscR and the *galF* promoter and that the [2Fe-2S] cluster of IscR plays a crucial role in this interaction. On the contrary, we also analysed whether recombinant IscR::His6 could bind the promoter regions of *wzi* and *manC*. As expected, EMSA showed no obvious DNA-protein complex (data not shown).

Effect of IscR on normal human serum resistance

Because CPS acts as a protectant for *K. pneumoniae* against serum factors, we hypothesized that through modulation of CPS levels, IscR affects the ability of *K. pneumoniae* to resist the bactericidal effects of serum. To test this hypothesis, we treated *K. pneumoniae* strains with 75% normal human serum and determined their survival rates. Compared with the WT, the AP001 and AP002 strains showed a slightly reduced survival rate (Fig. 3A), implying a positive role for [2Fe-2S]-IscR in the serum resistance of *K. pneumoniae*. To confirm this result further, we performed a complementation study. As shown in Fig. 3B, the introduction of pIscR, but not pACYC184 or pIscRSCA, into the
AP001 strain increased the bacterial survival rate, to a similar level compared with that of WT [pACYC184], after serum treatment. These results supported the hypothesis that [2Fe-2S]-IscR activates the expression of CPS to increase \textit{K. pneumoniae} resistance to normal human serum.

IscR has a regulatory role in iron acquisition systems

In \textit{E. coli}, both Fur and IscR play important roles in the maintenance of cellular iron homeostasis [12,13]. To analyse whether IscR regulates iron acquisition in \textit{K. pneumoniae}, we performed a chrome azurol S (CAS) assay to assess siderophore

**Figure 1. Holo-IscR positively regulates the biosynthesis of CPS.** (A) CPS levels of the WT, AP001, and AP002 strains grown in LB broth. (B) CPS levels in WT carrying pACYC184 and AP001 carrying pACYC184, plscR, or plscR\textsubscript{Ca} were determined in LB. Bacterial glucuronic acid content was determined after 16 h of growth. (C) $\beta$-Galactosidase activities of \textit{K. pneumoniae} AP006 and isogenic strains (AP007 and AP008) carrying the reporter plasmid pOrf12 (P\textsubscript{orf1-2}:lacZ), pOrf315 (P\textsubscript{orf3-15}:lacZ), or pOrf1617 (P\textsubscript{orf16-17}:lacZ) were determined using log-phase cultures grown in LB medium. Error bars indicate standard deviations. *P<0.01 compared with the indicated groups.

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Figure 2. IscR binds directly to \( P_{galF} \). (A) Diagrammatic representation of the \( galF \) loci. The primer sets used in PCR amplification of the DNA probes are indicated, and the numbers denote the DNA amplified length. DNA probes are listed on the left. The box in grey indicates the predicted type 1 IscR box. The dashed box indicates the DNA sequence alignment among the predicted type 1 IscR box, the IscR binding motif, and the putative IscR binding sequence in \( P_{galF} \), and the numbers denote the positions relative to the translational start site. Deletion of the predicted IscR
secretion in *K. pneumoniae* strains. As shown in Fig. 4A, no apparent siderophore secretion was detected in the WT, AP001, or AP002 strains. Moreover, as in our previous report [10], deletion of *fur* clearly increased halo formation on the CAS plate. However, the halo was reduced in the AP003 (*Δfur*-*iscR*) strain background by the further deletion of *iscR*, indicating the positive role of IscR in iron acquisition. Furthermore, no obvious difference in siderophore secretion was found between the AP003 and AP005 (*Δfur-iscR*<sub>3CA</sub>) strains, suggesting that the IscR regulation of iron acquisition does not require the [2Fe-2S] cluster.

To verify whether apo-IscR activates iron acquisition, we introduced pACYC184, pIscR, or pIscR<sub>3CA</sub> into the AP004 (*Δfur-*IscR) strain and performed a CAS assay. As shown in Fig. 4B, the introduction of both pIscR and pIscR<sub>3CA</sub> increased the halo phenotype on the CAS plate compared with that of the vector-only control. These results confirmed that IscR activates siderophore secretion in a [2Fe-2S] cluster-independent manner.

To further investigate the regulatory effect of IscR on iron acquisition, we used quantitative reverse transcription polymerase chain reaction (qRT-PCR) to measure the expression of genes corresponding to the eight putative iron acquisition systems in the indicated *K. pneumoniae* strains. As shown in Table 1, messenger RNA (mRNA) levels of genes (*fhuA*, *iucA*, and *sitA*) corresponding to three iron acquisition systems were increased more than 2-fold in the AP005 strain as compared with that in the AP004 strain. To further confirm this result, pACYC184, pIscR<sub>3CA</sub>, or pIscR were respectively introduced into the AP004 strain, to avoid the effects of Fur, and the transcription of *fhuA*, *iucA*, and *sitA* were measured. The introduction of pIscR<sub>3CA</sub> into AP004 apparently increased the transcription of *fhuA*, *iucA*, and *sitA* compared with that in the AP004 strain carrying pACYC184 only (Table 1). These results implied that apo-IscR activates the transcription of *fhu*, *iuc*, and *sit* to increase iron acquisition in *K. pneumoniae*. Besides, the introduction of pIscR into AP004 also slightly increased transcription of *fhu*, *iuc*, and *sit* (Table 1).

**IscR<sub>3CA</sub> directly binds the promoter region of *fhuA*, *iucA*, and *sitA***

Apo-IscR has been demonstrated to bind the type 2 IscR box in IscR-regulated promoter sequences directly in *E. coli* [19]. Analysis of the promoter regions of *fhuA*, *iucA*, and *sitA* revealed consensus sequences of the *E. coli* type 2 IscR box. As shown in Fig. 5A, the predicted type 2 IscR boxes are located at −154 to −130 relative to the translation start site of *fhuA* and −67 to −43 relative to the translation start site of *iucA*. The predicted type 2 IscR boxes in *P<sub>fhuA</sub>* and *P<sub>iucA</sub>* have 50% (13/26 bp) and 46% (12/26 bp) homology, respectively, with the consensus sequence. In addition, two putative type 2 IscR boxes (R1 and R2) located at −112 to −87 and at −53 to −28 relative to the translation start site of *sitA* were found in *P<sub>sitA</sub>*. The R1 and R2 sites contain 50% (13/26 bp) and 61.5% (16/26 bp) homology, respectively, with the consensus sequence.

To verify whether apo-IscR binds to these predicted type 2 IscR boxes, we performed an EMSA. As shown in Fig. 5B-D, both the purified IscR::His<sub>6</sub> and IscR<sub>3CA</sub>·His<sub>6</sub> were able to bind with the promoter regions of *fhuA*, *iucA*, and *sitA*, and IscR::His<sub>6</sub> appeared to contain higher binding activities. Furthermore, IscR<sub>3CA</sub>·His<sub>6</sub> did not bind *P<sub>fhuA</sub>*−2 and *P<sub>iucA</sub>*−2, which lacked a region containing a putative IscR box (Fig. 5B–C). We also found that IscR<sub>3CA</sub>·His<sub>6</sub> did not bind *P<sub>sitA</sub>*−2, which contained the R1 site but not the R2 site (Fig. 6D). To further confirm the importance of these predicted IscR boxes, recombinant IscR proteins were respectively interacted with these promoters lacking only the 26-bp predicted type 2 IscR box (*P<sub>fhuA</sub>*-1*, *P<sub>iucA</sub>*-1*, and *P<sub>sitA</sub>*-1*), and no obvious interaction was found. These results suggested that apo-IscR interacts directly with the promoters of *fhuA*, *iucA*, and *sitA* via the predicted type 2 IscR boxes.

**Regulatory control of *iscR* transcription in *K. pneumoniae***

To analyse whether environmental iron availability affects *K. pneumoniae* iscR expression, we grew AP006 in Luria-Bertani (LB) broth supplemented with increasing amounts of the iron chelator 2, 2-dipyridyl (Dip) and monitored the promoter activity of *iscR*.

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**Figure 3. Deletion effect of *iscR* on *K. pneumoniae* susceptibility to normal human serum.** The susceptibility to normal human serum of each bacterial mutant (A) and the complement strains (B) indicated in the margin was determined. Bacterial serum resistance was determined using log-phase cultures grown in LB medium. *P*<0.01 compared with the indicated groups.

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using a LacZ reporter system [23]. As shown in Fig. 6A, the addition of 250 or 500 μM Dip to the growth medium increased iscR promoter (PiscR) activity by approximately 2-fold and 3.9-fold, respectively, indicating that the transcription of the iscRSUA operon was activated by iron limitation.

In K. pneumoniae, Fur and RyhB reportedly play crucial roles in gene regulation in response to cellular iron levels [10,11]. Thus, we investigated whether Fur and RyhB regulate the activity of PiscR. As shown in Fig. 6B, the deletion of fur and the further deletion of ryhB in AP006 strain had no obvious effects on PiscR activity, whereas the deletion of iscR in AP006 strain activated PiscR activity by approximately 4.5-fold. In E. coli, IscR has been demonstrated to exert negative auto-regulation which requires the [2Fe-2S] cluster [20]. Thus, we measured PiscR activity in an AP006-derived strain, AP008 (ΔlacZ-iscR3CA), expressing a mutated IscR predicted to be defective in cluster binding [20]. As shown in Fig. 6B, the PiscR activity in AP008 was increased approximately 4.5-fold. This increase was comparable to that in AP007 (ΔlacZ-iscR). Our results suggested that IscR inhibits the

![Figure 4. Deletion of iscR decreases K. pneumoniae Δfur siderophore production assessed using CAS assay.](image)

All assayed bacterial mutants (A) and the complement strains (B) are indicated. The halos around the colonies correspond to the iron-chelating activity of siderophores in bacteria were measured after 24 h of incubation at 37 °C. The assay was independently repeated at least five times, and the differences among strains are consistent.

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| Systems | Gene          | RNA expression ratio b |
|---------|---------------|------------------------|
|         | AP005/AP004 a | AP004 [pIscR3CA]/AP004 [pACYC184] | AP004 [pIscR]/AP004 [pACYC184] |
| Fe³⁺    | ferrichrome   |                         |                         |
|         | fhuA          | 2.17 ± 0.31             | 2.83 ± 0.14             | 1.78 ± 0.42             |
|         | aerobactin    | 2.24 ± 0.12             | 3.23 ± 0.15             | 1.84 ± 0.22             |
|         | enterobactin  | 1.07 ± 0.38             | ND c                    | ND                      |
|         | fepA          | 1.11 ± 0.09             | ND                      | ND                      |
|         | fepB          | 0.80 ± 0.09             | ND                      | ND                      |
|         | entC          | 1.23 ± 0.52             | ND                      | ND                      |
|         | fecA          | 1.74 ± 0.25             | ND                      | ND                      |
|         | iroB          | 1.92 ± 0.13             | ND                      | ND                      |
|         | hmuR          | 1.58 ± 0.22             | ND                      | ND                      |
| Fe²⁺    | feoB          | 3.16 ± 0.01             | ND                      | ND                      |
|         | sitA          | 3.68 ± 0.11             | 2.27 ± 0.23             |

a AP004, CG4353Δfur-iscR; AP005, CG4353Δfur-iscR3CA.
b Mean expression ratio (±SD) of AP005 relative to AP004, AP004 [pIscR3CA] relative to AP004 [pACYC184], or AP004 [pIscR] relative to AP004 [pACYC184].
c ND, not determined.

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Table 1. qRT-PCR analyses of the expression of iron-acquisition genes in K. pneumoniae strains.
Characterization of IscR in Klebsiella pneumoniae

A

Type 2 Isc box

\[ \text{AWARCCYTSNGTTTGNGKTKW} \]

|     |     |     |     |
|-----|-----|-----|-----|
| P_{fhuA} | -154 | AACTCCATAATACCTACGGTTATT | -129 | 13/26 |
| P_{iucA} | -67  | AATCCCATATATGATTGCTTATT | -43  | 12/26 |
| P_{sitA-R1} | -12  | ATACCCCACGCATGCAAATAAGAATT | -87  | 13/26 |
| P_{sitA-R2} | -53  | ATAGCAATGGCTATGACGCTGGAA | -28  | 16/26 |

B

\[ \text{flhA} \rightarrow \text{flhA01} \rightarrow \text{flhA02} \rightarrow \text{flhA03} \rightarrow \text{flhB} \]

\[ \text{flhA01} \leftrightarrow \text{flhA02} \rightarrow 314 \text{ bp} \ P_{flhA-1} \]

\[ \text{flhA01} \leftrightarrow \text{flhA02} \rightarrow 288 \text{ bp} \ P_{flhA-1^*} \]

\[ \text{flhA03} \leftrightarrow \text{flhA02} \rightarrow 169 \text{ bp} \ P_{flhA-2} \]

C

\[ \text{iucA} \rightarrow \text{iucB} \rightarrow \text{iucC} \rightarrow \text{iucD} \rightarrow \text{iutA} \]

\[ \text{iucA02} \leftrightarrow \text{iucA01} \rightarrow 227 \text{ bp} \ P_{iucA-1} \]

\[ \text{iucA02} \leftrightarrow \text{iucA01} \rightarrow 201 \text{ bp} \ P_{iucA-1^*} \]

\[ \text{iucA02} \leftrightarrow \text{iucA03} \rightarrow 114 \text{ bp} \ P_{iucA-2} \]

D

\[ \text{sitA} \rightarrow \text{sitB} \rightarrow \text{sitC} \rightarrow \text{sitD} \]

\[ \text{sitA03} \leftrightarrow \text{sitA02} \rightarrow 283 \text{ bp} \ P_{sitA-1} \]

\[ \text{sitA03} \leftrightarrow \text{sitA02} \rightarrow 257 \text{ bp} \ P_{sitA-1^*} \]

\[ \text{sitA03} \leftrightarrow \text{sitA05} \rightarrow 160 \text{ bp} \ P_{sitA-2} \]

\[ \text{P}_{\text{iucA}-1} \]

\[ \text{P}_{\text{iucA}-1^*} \]

\[ \text{P}_{\text{iucA}-2} \]

\[ \text{P}_{\text{iucA}-1} \]

\[ \text{P}_{\text{iucA}-1^*} \]

\[ \text{P}_{\text{iucA}-2} \]

\[ \text{P}_{\text{iucA}-1} \]

\[ \text{P}_{\text{iucA}-1^*} \]

\[ \text{P}_{\text{iucA}-2} \]
transcription of the iscRSUA operon in a [2Fe-2S] cluster-dependent manner in *K. pneumoniae*.

To investigate whether IscR is the sole regulator of *iscRSUA* transcription in response to iron availability, we monitored P_{iscR} activity in the AP007 and AP008 strains in LB broth containing various levels of iron. As shown in Fig. 6C, the P_{iscR} activity in AP006 was activated in LB broth supplemented with Dip, and the further addition of 100 or 250 μM FeSO₄ reversed the activation. In the AP007 and AP008 strains, P_{iscR} activity was increased compared with that of AP006. Nevertheless, the addition of Dip still activated P_{iscR} activity, suggesting the presence of unknown factors, and the further addition of FeSO₄ restored the effect.

To analyse whether Fur is responsible for this regulation, we measured P_{iscR} activity in AP011 (∆iscR-Δfur-∆iscR) compared with that of AP007 at various iron levels and noted no obvious effect. These results indicated that in addition to IscR and Fur, other factors modulate the transcription of the *iscRSUA* operon in response to environmental iron availability.

**Discussion**

Clinically isolated *K. pneumoniae* strains usually carry large amounts of CPS to resist engulfment by phagocytes and serum bactericidal factors [6,24]. Therefore, tightly controlling CPS biosynthesis is critical for successful infection by *K. pneumoniae* [10,11,25]. We have previously shown that Fur represses the expression of mucoid factors RmpA and RcsA as well as the small RNA RyhB in response to environmental iron to decrease CPS biosynthesis indirectly in *K. pneumoniae* [10,11]. In this study, we focused on IscR, a central regulator of iron metabolism, to analyse more thoroughly how external iron affects CPS biosynthesis. Our data indicated that IscR activates CPS biosynthesis in a Fe-S cluster-dependent manner (Fig. 1A and B). Moreover, IscR positively regulates the transcription of three transcription units in the *cfs* gene cluster (Fig. 1C). Purified IscR::His₆ also directly interacts with the promoter of *orf1–2*, possibly through the predicted type 1 IscR box (Fig. 2). These findings indicated that Fur and IscR exert negative and positive regulation, respectively, on CPS biosynthesis in response to external iron.

However, we hypothesize herein that Fur plays a major regulatory role because (i) *K. pneumoniae* grown under iron-replete conditions displayed decreased CPS levels, (ii) the Promoter of the *fhuA* (β), *iscA* (γ), and *sitA* (δ) loci. The large arrows represent the *iscRSUA* operon. The primer sets used in PCR amplification of the DNA probes are indicated, and the numbers denote the DNA amplified length. The predicted IscR boxes is deleted and indicated by a caret. The grey boxes indicate the predicted type 2 IscR box. Different concentrations of purified IscR::His₆ were incubated with 5 ng of various DNA fragments of the upstream regions of indicated genes. Following incubation at room temperature for 30 min, the mixtures were analysed on a 5% non-denaturing polyacrylamide gel. The gel was stained with SYBR Green I dye and photographed.

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*Figure 5. IscR::His₆ binds directly to P_{fhuA} P_{iscA} and P_{sitA} (A) DNA sequence alignment between the *E. coli* type 2 IscR box and the putative IscR binding sequence in the upstream regions of *fhuA*, *iscA*, and *sitA*. Positions identical to the consensus sequences are bolded. Diagrammatic representation of the *fhuA* (β), *iscA* (γ), and *sitA* (δ) loci. The large arrows represent the *iscRSUA* operon. The primer sets used in PCR amplification of the DNA probes are indicated, and the numbers denote the DNA amplified length. The predicted IscR boxes is deleted and indicated by a caret. The grey boxes indicate the predicted type 2 IscR box. Different concentrations of purified IscR::His₆ were incubated with 5 ng of various DNA fragments of the upstream regions of indicated genes. Following incubation at room temperature for 30 min, the mixtures were analysed on a 5% non-denaturing polyacrylamide gel. The gel was stained with SYBR Green I dye and photographed.*

An analysis of the upstream region of *iscS* in *K. pneumoniae*, also identified a conserved sequence paired with RyhB (data not shown). However, the regulatory effect on P_{iscS} activity mediated by RyhB was not obvious under our assay conditions (Fig. 6B). On the contrary, as shown in Fig. 6C, P_{iscS} activity in AP007 and AP008 was still activated by iron depletion, which prompted us to verify whether IscR is the sole iron-responsive regulator that controls P_{iscS} activity. However, in AP007, the addition of FeSO₄ to iron-depleted medium still led to a reduction in P_{iscS} activity. These results suggested that an unknown regulator, beside of IscR and Fur, represses the transcription of *iscS* in response to external iron. In addition to IscR, the [2Fe-2S] cluster is critical for regulation mediated by FNR and SoxR [27]. However, sequence analysis of the promoter region of *iscS* revealed no typical FNR and SoxR binding sites. On the contrary, we found a putative binding site of SoxS [28], an oxidative transcriptional regulator activated by SoxR, in P_{iscS}. This putative SoxS binding site displays 79% (15/19 bp) homology with the consensus sequence
Figure 6. Regulation of *K. pneumoniae* *iscR* expression. (A) β-Galactosidase activity of *K. pneumoniae* AP006 (ΔlacZ) carrying the reporter plasmid piscRZ15 (P<sub>iscR</sub>::lacZ) were determined using log-phase cultures grown in the indicated concentrations of Dip. (B) The β-galactosidase activity of piscRZ15 was determined in the AP006 and isogenic strains, AP009 (ΔlacZ-Δfur), AP010 (ΔlacZ-Δfur-ΔryhB), AP007 (ΔlacZ-ΔiscR), and AP008 (ΔlacZ-isc<sub>R</sub>3CA), using log-phase cultures grown in LB medium. (C) The β-galactosidase activity of piscRZ15 was determined in the AP006 and isogenic strains, AP007, AP008, and AP011 (ΔlacZ-Δfur-ΔiscR), using log-phase cultures grown in the indicated media. Error bars indicate standard deviations. *P*<0.01 compared with the indicated group.

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and is located at position −144 to −126 relative to the translation start site of *iscR*. Therefore, we hypothesized that SosX may be involved in the regulation of *iscR* in response to oxidative stress and will investigate this possibility in future studies.

IscR differs from other known Fe-S cluster-containing transcription factors such as FNR and SoxR because both apo- and holo-IscR regulate transcription and exhibit different DNA binding specificities [29]. Structural and biochemical studies have suggested that the ligation of the [2Fe-2S] cluster broadens the DNA binding specificity of IscR, thereby allowing holo-IscR to bind both type 1 and type 2 boxes, whereas apo-IscR binds only the type 2 box [29]. In the *K. pneumoniae* K2 eps gene cluster, we found a type 1 box in the promoter region of *galF*, and purified IscR::His6 could bind this motif *in vitro* (Fig. 2). In addition and as expected, the clusterless IscR_{3CA}:His6, mimicking apo-IscR, showed no obvious binding affinity to P_{galF}. On the contrary, all three promoters of the iron acquisition genes regulated by IscR contain predicted type 2 boxes, and both IscR::His6 and IscR_{3CA}:His6 appeared to bind these boxes (Fig. 5). Although apo- and holo-IscR have been demonstrated to bind the type 2 box with similarly high affinity [21], EMSA revealed that IscR_{3CA}:His6 displayed weaker binding (Fig. 5). In *E. coli*, the Fe-S cluster status of IscR is a key variable that regulates gene expression in response to iron availability [12]. Our results suggested that the transcription of eps and iron acquisition genes is regulated by not only the level of IscR but also the cellular ratio of apo- and holo-IscR.

Although IscR was first discovered as an auto-repressor of the *isc* operon, it is now known to be a global regulator that influences apo- and holo-IscR. Fe-S cluster status of IscR is a key variable that regulates gene transcription of genes regulated by IscR. Thus, in response to oxidative stress and iron acquisition systems through IscR in *K. pneumoniae*, we demonstrated that external iron levels regulate CPS biosynthesis and iron acquisition systems, and other virulence factors in *K. pneumoniae* to facilitate bacterial persistence in the host. In this study, we demonstrated that external iron levels regulate CPS biosynthesis and iron acquisition systems through IscR in *K. pneumoniae* and proposed a working model (Fig. 7). In response to iron availability, IscR and Fur control the expression of downstream targets in a parallel and cooperative manner, which is predicted to play a crucial regulatory role during infection.

### Materials and Methods

#### Bacterial strains, plasmids, and media

Bacterial strains and plasmids used in this study are listed in Table 2. Primers used in this study are listed in Table 3. Bacteria were routinely cultured at 37°C in LB medium supplemented with appropriate antibiotics. The antibiotics used include ampicillin (100 μg/ml), kanamycin (25 μg/ml), streptomycin (500 μg/ml), and tetracycline (12.5 μg/ml).

#### Construction of the deletion of *iscR* mutants

Specific gene deletion of *iscR* was introduced into *K. pneumoniae* CG43S3 using an allelic exchange strategy as previously described [33]. In brief, two approximately 1000 bp DNA fragments flanking both sides of *iscR* were cloned into the suicide vector pKAS46 [36], a suicide vector containing rpsL, which allows positive selection with streptomycin for vector loss.

The resulting plasmid was then mobilized from *E. coli* S17-1λpir [37] to *K. pneumoniae* CG43S3 or CG43S3-derived strains by conjugation. The transconjugants, with the plasmid integrated into the chromosome via homologous recombination, were selected with ampicillin and kanamycin on M9 agar plates. Several of the colonies were grown in LB broth supplemented with 500 μg/mL of streptomycin to log phase at 37°C and then spread onto an LB agar plate containing 500 μg/mL of streptomycin. The streptomycin-resistant and kanamycin-sensitive colonies were selected, and the deletion was verified by PCR and Southern hybridization (data not shown). The resulting *K. pneumoniae* mutants are listed in Table 2.

#### Construction of the pIscR complementation plasmid and the pIscR_{3CA} mutant plasmid

To obtain the complementation plasmid (pIscR), a DNA fragment containing the promoter and coding sequence of *iscR* was amplified by PCR using the primer pair GT138/GT139 (Table 3) and cloned into the pACYC184 shuttle vector. The pIscR_{3CA} plasmid, which carried the mutant allele encoding IscR with the C92A, C98A, and C104A mutations, was constructed using the inverse-PCR method. Briefly, the pIscR plasmid was used as the PCR template to generate the mutant allele with the primer pair GT206/GT207 (Table 3). The recovered PCR product was treated with DpnI for 2 h, subjected to T4 polynucleotide kinase treatment, and self-ligated with T4 DNA ligase. The ligation product was transformed into *E. coli* DH5α.

The pIscR_{3CA} plasmid was subsequently confirmed by sequence analysis.

#### Construction of a *K. pneumoniae* *iscR_{3CA}* mutant

A DNA fragment carrying *iscR* and approximately 1000 bp adjacent regions on either side was amplified by PCR using primer pairs GT241/GT242 (Table 3) and cloned into yT&A. The resulting plasmid was used as the template for the inverse-PCR with the primer pair GT206/GT207 (Table 3) to generate a mutant *iscR* allele encoding the C92A, C98A and C104A mutations. Subsequently, the mutant allele of *iscR* was subcloned into pKAS46 and confirmed by DNA sequencing. Then, the plasmid was mobilized from *E. coli* S17-1λpir to the *K. pneumoniae* AP001 strain by conjugation, and the subsequent selection was performed as described above.

#### Extraction and quantification of CPS

CPS was extracted and quantified as previously described [38]. The glucuronic acid content, represents the amount of *K. pneumoniae* K2 CPS, was determined from a standard curve of glucuronic acid (Sigma-Aldrich) and expressed as micrograms per 10^9 c.f.u. [39].

#### Measurement of promoter activity

The promoter-reporter plasmids, pOrf12, pOrf315, pOrf1617, and pIscRZ15 were individually mobilized into *K. pneumoniae* strains by conjugation from *E. coli* S17-1λpir. The bacteria were grown to logarithmic phase in LB broth or indicated medium, and
the β-galactosidase activity was measured as previously described [23].

Bacterial survival in serum

Normal human serum, pooled from healthy volunteers, was divided into equal volumes and stored at 270°C before use. Bacterial survival in serum was determined as previously described [35]. First, the bacteria were grown to log phase in LB broth and the viable bacterial concentration was adjusted to 1×10^6 c.f.u./ml. Next, 1 ml of the cultures was washed twice using phosphate-buffered saline (PBS) and resuspended in 1 ml PBS. A mixture containing 250 μl of the cell suspension and 750 μl of pooled human serum was incubated at 37°C for 15 min. The number of viable bacteria was then determined by plate counting. The survival rate was expressed as the number of viable bacteria treated with human serum compared with the number of viable bacteria pretreatment. The 0% survival of K. pneumoniae AP012 (ΔgalU) served as a negative control.

Purification of IscR::His₆ and IscR₃CA::His₆

The coding regions of iscR and iscR₃CA were amplified using the primer pair GT215/GT216 (Table 3) and cloned into the NheI/XhoI site in pET30b (Novagen, 205 Madison, Wis). The resulting plasmids (pET30b-IscR and pET30b-IscR₃CA, respectively) were then transformed into E. coli BL21(DE3)[pLysS] (Invitrogen, USA), and overproduction of the recombinant proteins IscR::His₆ and IscR₃CA::His₆, respectively, were induced by the addition of 1 mM IPTG for 3 h at 37°C. The cell pellets were washed and resuspended in cold binding buffer (20 mM sodium phosphate, 0.5 M NaCl, 5 mM imidazole, pH 7.4). The cells were then broken by sonication and the cell pellets were removed by centrifugation at 14000 rpm for 10 min at 4°C. The recombinant proteins were then purified from the soluble fraction of the total cell lysate by affinity chromatography using His-Bind resin (Novagen, Madison, Wis) according to the manufacturer’s instructions. The nonbinding proteins were washed away using binding buffer and the recombinant proteins were eluted by elution buffer (20 mM sodium phosphate, 0.5 M NaCl, 500 mM imidazole, pH 7.4). Finally, the purified protein was checked for purity by SDS-PAGE and stored for up to two weeks at 4°C. The purified protein was transparent and no obvious precipitation was observed after storage.

EMSA

DNA fragments of the putative promoter regions galF, fhuA, iucA, and sitA were amplified with Phusion polymerase using specific primer sets (Table 3) to generate DNA probes for EMSA (PgalF-1 and PgalF-2 for galF; PfhuA-1 and PfhuA-2 for fhuA; PiucA-1 and PiucA-2 for iucA; and PsitA-1 and PsitA-2 for sitA). To obtain probes that lacked a putative IscR box, the DNA fragments were respectively amplified with Taq polymerase using the above-described primer sets (Table 3) and then cloned into yT&A to generate the plasmids pgalF, pfhuA, piucA, and psitA for subsequent mutagenesis via the inverse-PCR method using the primer pairs GT201/galF01, fhuA04/fhuA05, iucA03/iucA04, and sitA01/sitA02.
and sitA05/sitA06, respectively. The resulting mutant plasmids, \( pgalF^* \), \( pfhuA^* \), \( piucA^* \), and \( psitA^* \), respectively, were amplified with specific primer sets to generate the DNA fragments \( PgalF-1^* \), \( PfhuA-1^* \), \( PiucA-1^* \), and \( PsitA-1^* \) for DNA probes in EMSA.

For the EMSA, the purified \( \text{IscR}::\text{His}6 \) and \( \text{IscR}3CA::\text{His}6 \) proteins were incubated with 5-ng DNA in a 10 \( \mu \)l solution containing 4 mM Tris-HCl (pH 7.4), 10 mM KCl, 100 mM dithiothreitol, and 10 \( \mu \)g/ml BSA at 37°C for 30 min. The samples were then loaded onto a native gel of 5% nondenaturing polyacrylamide in 0.5 \( \times \) TB buffer (45 mM Tris-HCl, pH 8.0, 45 mM boric acid). Gels were electrophoresed with a 20-mA current at 4°C and then stained with SYBR Green I dye (Invitrogen). The assay was repeated in at least 3 independent experiments.

### qRT-PCR

Total RNAs were isolated from early-exponential-phase grown bacteria cells by use of the RNaseasy midi-column (QIAGEN) according to the manufacturer’s instructions. RNA was DNase-treated with RNase-free DNase I (MoBioPlus) to eliminate DNA contamination. RNA of 100-ng was reverse-transcribed with the Transcriptor First Strand cDNA Synthesis Kit (Roche) using random primers. qRT-PCR was performed in a Roche.

### Table 2. Bacterial strains and plasmids used in this study.

| Strains or plasmids | Descriptions | Reference or source |
|---------------------|--------------|---------------------|
| **K. pneumoniae**   |              |                     |
| CG43S3              | CG43 Sm'     | [41]                |
| AP001               | CG43S3\text{iscR} | This study           |
| AP002               | CG43S3\text{iscR}_{3CA} | This study           |
| AP003               | CG43S3\text{Mur}   | [42]                |
| AP004               | CG43S3\text{Mur-iscR} | This study           |
| AP005               | CG43S3\text{Mur-iscR}_{3CA} | This study           |
| AP006               | CG43S3\text{lacZ} | [23]                |
| AP007               | CG43S3\text{lacZ-iscR} | This study           |
| AP008               | CG43S3\text{lacZ-iscR}_{3CA} | This study           |
| AP009               | CG43S3\text{lacZ-Mur} | This study           |
| AP010               | CG43S3\text{lacZ-Mur-\text{AryhB}} | This study           |
| AP011               | CG43S3\text{lacZ-Mur-\text{AryhB}} | This study           |
| AP012               | CG43S3\text{DgalU} | [35]                |
| **E. coli**         |              |                     |
| DH5'                | supE44\text{lacU169 (80 lacZAM1515)(169recA1 endA1 gyrA96 thi-1 relA1)} | [43]                |
| BL21(DE3)           | F\text{ ompT hsdS}\_{\text{de}}\_{\text{s}}\_m\_y\_gal dcm [DE3] | New England Biolabs |
| S17-1 \( \lambda \_\text{pir} \) | hsdR recA pro BP4-2 [Tc-Muc Km:Trn71 (\text{Lpir})] | [37]                |
| **Plasmids**        |              |                     |
| pKAS46              | Ap\_ Km', positive selection suicide vector, \text{rpsL} | [36]                |
| yT&A                | Ap\_, TA cloning vector | Yeastern            |
| pACYC184            | Y\_C\_\text{Cm'}, low copy number cloning vector | New England Biolabs |
| pIscR               | Cm', 980-bp fragment containing an \text{iscR} allele cloned into pACYC184 | This study           |
| pIscR_{3CA}         | Cm', 980-bp fragment containing C92A, C98A and C104A mutant allele of \text{iscR} cloned into pACYC184 | This study           |
| pIscZ15             | Cm', promoter selection vector, lacZ\_+ | [23]                |
| pIscZ15_{3CA}       | Cm', 312-bp fragment containing the region upstream of \text{iscR} cloned into pIscZ15 | This study           |
| pOrf12              | Cm', 500-bp fragment containing the region upstream of \text{Klebsiella K2} cps \text{orf1-orf2} cloned into pIscZ15 | [23]                |
| pOrf315             | Cm', 900-bp fragment containing the region upstream of \text{Klebsiella K2} cps \text{orf3-orf15} cloned into pIscZ15 | [23]                |
| pOrf1617            | Cm', 300-bp fragment containing the region upstream of \text{Klebsiella K2} cps \text{orf16-orf17} cloned into pIscZ15 | [23]                |
| pET30b-IscR         | Km', 654-bp fragment encoding full-length IscR cloned into pET30b | This study           |
| pET30b-IscR_{3CA}   | Km', 654-bp fragment encoding full-length C92A, C98A and C104A mutant allele of \text{iscR} cloned into pET30b | This study           |
| piscR-pKAS46        | Ap\_ Km', 2.0 kb fragment containing \text{iscR} and its flanking regions cloned into pKAS46 | This study           |
| piscR_{3CA}-pKAS46  | Ap\_ Km', 2.0 kb fragment containing full-length C92A, C98A and C104A mutant allele of \text{iscR} and its flanking regions cloned into pKAS46 | This study           |
**Table 3. Primers used in this study.**

| Primer | Sequence (5’→3’) | Enzyme cleaved |
|--------|------------------|----------------|
| GT138  | GGATCCTGCGATCGATTGTCAACC | BamHI |
| GT139  | AAGCTTCAATTCAAGAAATACGCGCA | HindIII |
| GT142  | GGATCCGCGGATACGCTCAGATAA | BamHI |
| GT143  | AGATCTAGTTGAATCCTGCGCGG | BglII |
| GT206  | CTGAGGCGGCCGCCCTCCGCGG | BglII |
| GT207  | GCGCGGATACGCTGAGCTCAATAGTGCAATCAATG | XbaI |
| GT215  | GCTCGAGGGCGGCGATGTTAAGCGTCAAT | NheI |
| GT216  | CATATGAGACTGACATCTAAAGGGCG | XhoI |
| GT217  | GCATATGAGACTGACATCTAAAGGGCG | XhoI |
| GT241  | GTGCTGCTGTGATCCTGCGACCGGAATAACC | BamHI |
| GT242  | GAAGCTTTCGCGGCCATGCGG | HindIII |
| GT243  | CGAGCTCTGCAGCGGCAACATGGG | BamHI |
| GT250  | GGATCCGCGGATACGCTCAGATAA | BamHI |
| GT246  | CTGAGGCGGCCGCCCTCCGCGG | BglII |
| GT251  | GCGCGGATACGCTGAGCTCAATAGTGCAATCAATG | XbaI |

**For RT-qPCR**

| Sequence (5’→3’) | TaqMan probes | Target |
|------------------|---------------|--------|
| RT111 GGTAGGGGACGGGTCTCTGTA | 67 | 23S rRNA |
| RT121 TCACGATCCGACTCGTCTGAA | 3 | entC |
| GT56 | ACCCCCAGCTCTTAACCC | 20 | fecA |
| GT57 | TGCTCCCTTTACGGCACG | 20 | feoB |
| GT58 | CAACCTGAAACGGATTTTCC | 20 | fepA |
| GT59 | TCAGGCGCTCTCTTACGATCTT | 20 | hmuR |
| GT60 | CATAGCTAGGGCGGATACGCTCAG | 3 | fhuA |
| GT61 | GGAGGGTGAATCCTGAGCACTCAG | 20 | iroB |
| GT62 | CGAGGCGAGCTCTGAGG | 84 | iroB |
| GT63 | TTCCGGAATCTAGCGCGG | 84 | iroB |

**Characterization of IscR in Klebsiella pneumoniae**
LightCycler 1.5 Instrument using LightCycler TaqMan Master (Roche). Primers and probes were designed for selected target sequences using Universal ProbeLibrary Assay Design Center (Roche-applied science) and listed in Table 3. Data were analyzed using the real time PCR software of Roche LightCycler 1.5 Instrument. Relative gene expressions were quantified using the comparative threshold cycle 2^{−ΔΔCT} method with 23S rRNA as the endogenous reference.

**CAS assay**

The CAS assay was performed according to the method described by Schwyn and Neilands [40]. Each of the bacterial strain was grown overnight in LB medium, and then 5 μl of culture was added onto a CAS agar plate. After 24 h incubation at 37°C, the effects of the bacterial siderophore production could be observed. Siderophore production was apparent as a halo around the colonies; the absence of a halo indicated the inability to produce siderophores.

**Statistical methods**

An unpaired t-test was used to determine the statistical significance and values of P<0.01 were considered significant. The results of CPS quantification, β-galactosidase activity, serum survival rate, and qRT-PCR analysis were performed in triplicate and independently repeated at least three times, and the mean activity and standard deviation are presented.

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