

# YdfD

Subjects: Microbiology

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ydfD is a lytic gene from the Qin cryptic prophage that encodes a 63-amino-acid protein, the ectopic expression of which in *Escherichia coli* can cause nearly complete cell lysis rapidly. The bacterial 2-C-methyl-D-erythritol 4-phosphate (MEP) pathway is responsible for synthesizing the isoprenoids uniquely required for sustaining bacterial growth.

Keywords: IspG ; YdfD

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## 1. Introduction

Cryptic prophages, also known as defective prophages, are segments of phage DNA integrated and maintained in bacterial chromosomes. It has been shown that bacterial genomes can harbor multiple cryptic prophages <sup>[1][2][3][4]</sup>. In fact, as much as 10–20% of bacterial chromosome segments are estimated to be prophage genes, and a growing body of evidence suggests that the proportion may be even greater <sup>[1][3]</sup>. For example, it was in the 1950s that the *E. coli* K-12 genomes were first found to carry phage genes <sup>[5]</sup> and now there are 10 known cryptic prophages residing in the chromosome of *E. coli* K-12 <sup>[4]</sup>. The expression of most of the prophage genes is usually suppressed under normal conditions, as certain proteins they encode are typically lytic and, therefore, toxic to the host <sup>[6]</sup>. However, there is evidence that some prophage genes can not only endow their hosts with traits and behaviors beneficial to surviving harsh environmental conditions but also expand the hosts' genetic diversity, which plays a key role in microbial evolution <sup>[7][8][9][10]</sup>. Despite recent progress made in understanding cryptic prophages, the relationship between cryptic prophages and hosts is still largely elusive.

Investigation of cryptic prophages may provide hints to understand the process of phage infection, especially phage-induced cell lysis. As far as presently known, phages establish canonical lytic pathways to escape from hosts by two proteins, holin and endolysin <sup>[11][12][13]</sup>. Holins form holes in the cell membrane, and endolysins are a class of hydrolases that degrade bacterial cell wall. Therefore, holins are considered to control the access of endolysins to the cell periplasm. Apart from degradation of the cell wall, some phage genes encode proteins to inhibit the key enzyme to prevent cell wall biosynthesis. These proteins were dubbed "single-gene lysis" (Sgl) <sup>[14]</sup>. Many phages encode lytic protein. However, there are only three phage targets that have been proposed: (1) the Leviviridae Q $\beta$  uses the protein A2 to bind and inhibit MurA, which is the first step in bacterial cell wall biosynthesis <sup>[14]</sup>; (2) the protein E from phage  $\Phi$ X174 inhibits peptidoglycan synthesis by interaction with MraY, which involves the peptidoglycan (PG) biosynthesis pathway <sup>[15]</sup>; and (3) LysM inhibits the translocation of the final lipid-linked PG precursor across the cytoplasmic membrane by interfering with MurJ <sup>[16]</sup>. Therefore, phages being natural antibiotics, exploring the lysis targets of phages is timely and of considerable importance.

Q-independent (Qin) prophage was the third prophage identified, found in *E. coli* K-12 <sup>[17]</sup>. As a part of Qin prophage, *ydfD* encodes a 63-amino-acid protein and *ydfD* is located downstream of *dicB*. Both *ydfD* and *dicB* are regulated by *dicA* <sup>[18]</sup>. Overexpression of *ydfD* induces cell lysis in *E. coli*, whereas lysis is prevented by coexpression with the cell division inhibitor *dicB* or *sulA*. It is now known that DicB as well as Sula prevent cell division by inhibiting FtsZ polymerization <sup>[19][20]</sup>. This indicates that the cell lysis caused by YdfD depends on functional cell division. Based on what is currently known, YdfD is speculated to promote cell lysis by targeting an unknown unique host protein, although the identity of the target protein of YdfD has not been established <sup>[21]</sup>.

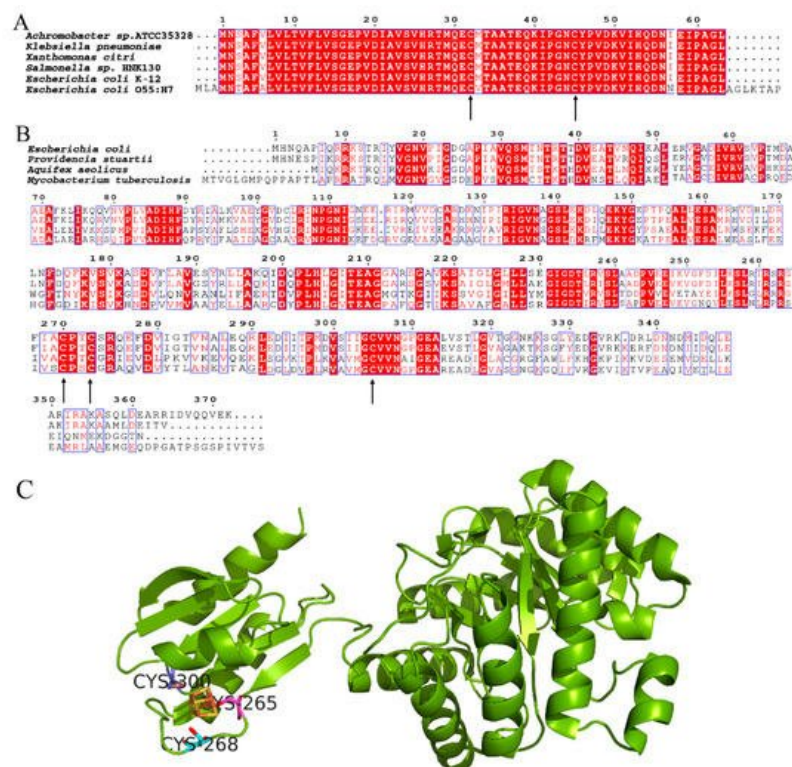
## 2. YdfD Is a Lysis Protein of the Qin Prophage

The number of prophages identified in bacterial genomes or plasmids has increased substantially in recent years <sup>[4][22][23]</sup>, although the underlying functional mechanisms of their host interactions have remained unclear <sup>[24]</sup>. Nevertheless, prophages appear to play key roles in host metabolism. Cryptic prophages have been shown to help bacteria resist antibiotic stress <sup>[24]</sup> and *dicB*, an upstream gene of *ydfD*, suppresses cell division in *E. coli* once induced <sup>[25]</sup>. Prophages can also prevent different categories of phages from infecting the host. DicB specifically inhibited infection by  $\lambda$  and other

phages that use ManYZ membrane proteins to prevent phage DNA entrance [26], provide population-level benefits by promoting biofilm formation, and influence the evolution of their hosts [27][28].

A previous study showed that overexpression of *ydfD* caused cell lysis of *E. coli* [21], although the molecular mechanism was not illustrated. In this study, the researchers attempted to characterize YdfD and its underlying mechanism of cell lysis activity. Herein, the researchers report that YdfD interacts with the essential MEP enzyme IspG. The MEP pathway synthesizes precursors of the cell wall and membrane, and a variety of inhibitors have already been designed and screened against it [27][28]. It is thus tempting to suggest that YdfD-mediated cell lysis may be caused by YdfD inhibition of MEP production. This phenomenon may be regarded as a strategy for the prophage precursor to escape from the host under adverse conditions. This is analogous to the lytic function of protein E from phage  $\Phi$ X174, which inhibits peptidoglycan synthesis by the *MraY* enzyme [45].

YdfD contains two well-conserved cysteine residues among bacterial species [21], and IspG has three conserved cysteine residues involved in the binding of the iron–sulfur cluster, responsible for the electron transfer of the catalytic process, as shown in **Figure 1**. The catalytic activity was significantly reduced when any cysteine that involved the cluster was replaced by serine [29]. However, the YdfD mutants lacking cysteine residue(s) had no effect on its binding to IspG, suggesting that the YdfD–IspG interaction is not through a disulfide bond. As the  $[4\text{Fe-4S}]^{2+}$  cluster is indispensable for IspG catalytic activity, all experimental procedures need to be performed under strict anerobic conditions [30]. In addition, ME-cPP, the substrate of IspG, is not commercially available. It is thus difficult to assay the effects of YdfD on IspG enzyme kinetics.



**Figure 1.** Conserved cysteine residues of YdfD and IspG. (A) Alignment of YdfD sequences from *Achromobacter* sp. ATCC35328, *Klebsiella pneumoniae*, *Xanthomonas citri*, *Salmonella* sp. , *E. coli* K-12, and *E. coli* O55:H7; the arrows indicate conserved cysteine residues. (B) Alignment of IspG sequences from *E. coli*, *Providencia stuartii*, *Aquifex aeolicus*, and *Mycobacterium tuberculosis*; the arrows indicate conserved cysteine residues. (C) *Aquifex aeolicus* IspG; three conserved cysteine residues binding the iron–sulfur cluster are responsible for the electron transfer of the catalytic process (PDB code: 3noy).

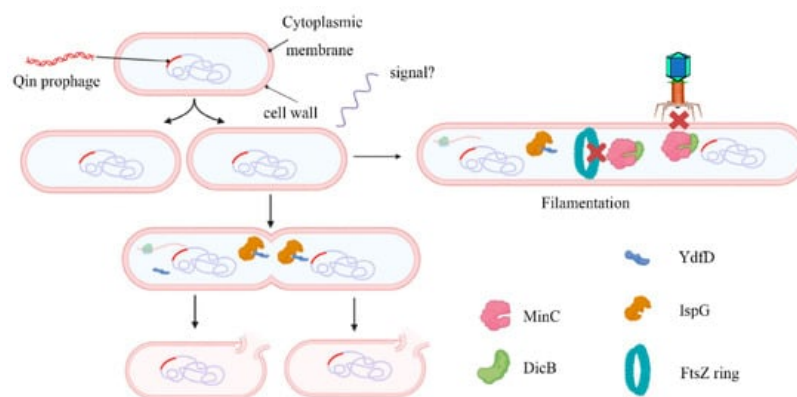
While it appears that the N-terminal domain of YdfD was responsible for binding to the N-terminal domain of IspG, intact YdfD was necessary for its full activity in the cell. It is possible that the N-terminal domain of YdfD recruits the C-terminal domain to IspG, which inhibits IspG's activity, which is why both N-terminal and C-terminal domains of YdfD are essential for its cellular activity. This agrees with previous studies showing that the C-terminal domain of YdfD is essential for lysis but that the killing activity of the C-terminal domain alone is significantly reduced compared to that of the intact protein [21]. The *Aquifex aeolicus* IspG is known to fold into two domains: an N-terminal and a C-terminal domain [30]. In the *Aquifex aeolicus* IspG catalytic process, the substrate binds to the positively charged surface region at the N-terminal domain [31]. The ProtParam model (<https://web.expasy.org/protparam/>, accessed on 10 November 2021) [32] shows that the C-terminal

domain of YdfD is negatively charged. Hence, it is possible that the C-terminal domain of YdfD competitively inhibits IspG by obstructing the binding of its substrate, ME-cPP, to the positively charged surface region of the IspG N-terminal domain. As a result, cooperation of both YdfD domains would be essential to inhibit IspG activity.

Under normal conditions, lysogenic phage levels remain steady. However, once the environment worsens, the transcription of lysis genes increases as the lysogenic phages look for ways to escape from the host [33][34][35]. Prophages also respond in this manner: although they cannot successfully assemble into viable phage particles, the lysis genes (*ydfD*) of Qin would increase transcription under adverse conditions. As expected, the RT-PCR of *E. coli* grown under stress conditions showed that mRNA levels of *ydfD* dramatically increases in non-acidic stress conditions and only slightly decreases under acidic conditions. The slight transcription decrease in the acidic medium is likely due to the bacteria's general preference for more alkaline conditions and the corresponding reduction in the growth rate in a medium with a pH of 5.0 [36][37]. Consequently, the transcription of all unessential genes would be expected to decrease when bacteria are grown in an acidic environment.

The products of the MEP pathway are IPP and DMAPP, both of which are sole precursors for vital cellular components, including hopanoids [38]; menaquinone [39]; and polyprenyl phosphates, particularly undecaprenyl diphosphate (Und-p), a lipid carrier to synthesize peptidoglycan cell wall precursors that are essential to maintain cell integrity [40][41][42]. No matter how complex the terpenoid compounds are, all of them share a common five-carbon isoprene building block. A likely mechanism for YdfD-induced lysis is that YdfD blocks the MEP pathway in the cell membrane and cell wall biosynthesis. This would result in a shortage of key precursors for the cell wall and the cell membrane, leading to the abortion of cell wall formation, especially during cell division. A previous study by Masudo et al. found that YdfD cannot induce cell lysis when cell division is restrained [21]. The researchers suppose that YdfD cuts off the MEP pathway by interaction with IspG, resulting in deficient cell components and cell wall membrane. When *E. coli* cells divide normally, as cytomembrane and cell wall element synthesis is most intense at sites of division, YdfD blocks terpenoid synthesis, thus leading to cell lysis. However, when cell division is slowed or stalled, the need for terpenoids (e.g., Und-P) is reduced. Therefore, YdfD cannot lyse the nondividing cells, i.e., YdfD-mediated effects would depend on cell division [21]. Further studies are needed to explore the direct effects of YdfD on IspG. HMBPP synthesis is a committed step in the MEP pathway, and IspG is strongly conserved across bacterial species [43], so it is unsurprising that phages would adapt to target IspG specifically to disrupt the host physiology and exert their function. IspG will be the fourth identified target of the phage lysis proteins; the details of how YdfD binds IspG and interferes with HMBPP will likely require structure studies of the YdfD/IspG complex.

It is clear that DicB binds to MinC and prevents FtsZ ring formation. Moreover, DicB specifically inhibits infection by  $\lambda$  and other phages that use ManYZ membrane proteins to prevent phage DNA entrance [26]. *dicB* and *ydfD* are located in the same operon, with partial overlap. As mentioned before, DicB was able to offset YdfD-induced lysis [21]. The researchers propose the following model: in normal conditions, DicB inhibits cell division and prevents phage infection; when the environment gets worse, the transcription and expression of *ydfD* are activated, eventually leading to cell lysis (**Figure 2**).



**Figure 2.** Working model for the regulation of host morphology by YdfD and DicB. The Qin cryptic prophage encodes the protein DicB and YdfD. When cell division is inhibited by DicB, demand for isoprenoid compounds is decreased and lysis is prevented (**right**). When YdfD is ectopically expressed, YdfD interacts with IspG, reducing isoprenoid production and causing the cells to lyse (**left**).

The growing severity of antibiotic resistance means there is a growing need to target novel bacterial pathways. The MEP pathway is absent in humans, making it an attractive target for antibiotic drug development [24]. The mechanisms used by prophages such as the Qin YdfD protein have proven to have a robust and widespread capacity to co-opt bacterial physiology. In addition to offering new insight into the particularities of host-prophage interactions, YdfD represents a promising new avenue for generating lead compounds for designing antibacterial agents.

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