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**Citation:** Yuan, Yifeng et al. "Identification of the minimal bacterial 2#-deoxy-7-amido-7-deazaguanine synthesis machinery." *Molecular microbiology* 110 (2018): © 2018 The Author(s)

**As Published:** 10.1111/mmi.14113

**Publisher:** Wiley

**Persistent URL:** <https://hdl.handle.net/1721.1/124831>

**Version:** Author's final manuscript: final author's manuscript post peer review, without publisher's formatting or copy editing

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Published in final edited form as:

*Mol Microbiol.* 2018 November ; 110(3): 469–483. doi:10.1111/mmi.14113.

## Identification of the minimal bacterial 2'-deoxy-7-amido-7-deazaguanine synthesis machinery

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### Abstract

The 7-deazapurine derivatives, 2'-deoxy-7-cyano-7-deazaguanosine (dPreQ<sub>0</sub>) and 2'-deoxy-7-amido-7-deazaguanosine (dADG) are recently-discovered DNA modifications encoded by the *dpd* cluster found in a diverse set of bacteria. Here we identify the genes required for formation of dPreQ<sub>0</sub> and dADG in DNA and propose a biosynthetic pathway. The preQ<sub>0</sub> base is a precursor that in *Salmonella* Montevideo, is synthesized as an intermediate in the pathway of the tRNA modification queuosine. Of the eleven genes (*dpdA* - *dpdK*) found in the *S. Montevideo dpd* cluster, *dpdA* and *dpdB* are necessary and sufficient to synthesize dPreQ<sub>0</sub>, while *dpdC* is additionally required for dADG synthesis. Among the rest of the *dpd* genes, *dpdE*, *dpdG*, *dpdI*, *dpdK*, *dpdD* and possibly *dpdJ* appear to be involved in a restriction-like phenotype. Indirect competition for preQ<sub>0</sub> base led to a model for dADG synthesis in which DpdA inserts preQ<sub>0</sub> into DNA with the help of DpdB, and then DpdC hydrolyzes dPreQ<sub>0</sub> to dADG. The role of DpdB is not entirely clear as it is dispensable in other *dpd* clusters. Our discovery of a minimal gene set for introducing 7-deazapurine derivatives in DNA provides new tools for biotechnology applications and demonstrates the interplay between the DNA and RNA modification machineries.

### Graphical abstract

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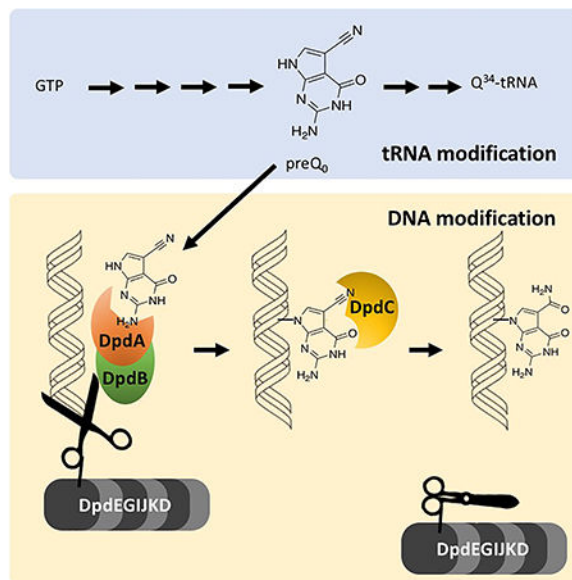
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Y.Y, G.H., J.G.V., J.H., R.H., P.C.D., D.I.-R., V.d.C.-L. designed research; Y.Y, G.H., J.G.V., J.H., J.H., R.H., and A.G. performed the experiments. J.G.V., J.H., R.H. and P.C.D. performed detection of modified nucleosides; Y.Y, G.H., J.G.V., J.H., R.H., J.H., P.C.D., D.I.-R., V.d.C.-L. analyzed the data and wrote the paper.

The authors declare no conflict of interest.



DNA modifications allow Bacteria to recognize self from non self. We describe a novel pathway responsible for the insertion of a complex modification that was thought to be found only in RNA.

## Abbreviated Summary

DNA modifications allow Bacteria to recognize self from non self. We describe a novel pathway responsible for the insertion of a complex modification that was thought to be found only in RNA.

## Keywords

DNA modification; 7-deazapurine; restriction

## Introduction

Although the first discovery of a non-canonical nucleoside in a nucleic acid was the identification of 5-methylcytosine (5mC) in DNA in the early 1950s (Hotchkiss, 1948; Wyatt, 1950), it is in RNA that both the quantity and complexity of nucleic acid modifications have been realized. More than one-hundred RNA modifications have been found to date (Machnicka et al., 2013), mainly in transfer RNA (tRNA) and ribosomal RNA (rRNA) (Grosjean, 2009). In contrast, only twenty modifications have been identified in DNA and most of the known DNA modifications are simple methylated derivatives of canonical nucleosides (Grosjean, 2009; Weigele & Raleigh, 2016). The few structurally complex DNA modifications are mainly found in phages where they serve to evade the host-encoded restriction systems (Warren, 1980; Weigele & Raleigh, 2016). Indeed, numerous bacteria have acquired restriction and modification (R-M) systems to defend against foreign DNA (Vasu & Nagaraja, 2013). Well-studied phage DNA modifications include 5-methylcytosine in phage XP12 (Feng, Tu, & Kuo, 1978; Huang, Farnet, Ehrlich, & Ehrlich, 1982); 5-hydroxymethylcytosine in phage T4gt,  $\beta$ -glucosyl-5-hydroxymethylcytosine in phage T4 (Cao, Huang, Farnet, & Ehrlich, 1983), N6-(1-acetamido)-adenine in

bacteriophage Mu (Hattman, 1979) and 5-hydroxymethyluracil in *Bacillus* SP8 (Vilpo & Vilpo, 1995). In addition, recently discovered 5-(2-aminoethoxy)methyluridine in *Salmonella* phage VII, 5-(2-aminoethyl)uridine in *Pseudomonas* phage M6 (Lee et al., 2018) and phosphorothioate (PT) modification of the DNA backbone (Chen et al., 2017; Wang et al., 2007) expanded the list of naturally-occurring DNA modifications.

Until recently, 7-deazapurine modifications were thought to be limited to tRNA. Queuosine (Q, 7-{5-[(1S,4S,5R)-4,5-Dihydroxy-2-cyclopenten-1-yl]amino}methyl}-7-deazaguanosine) is found at the wobble base (position 34) of a subset of bacterial and eukaryotic tRNAs (histidine, asparagine, aspartate and tyrosine) (Fergus, Barnes, Alqasem, & Kelly, 2015). Archaeosine (G<sup>+</sup>, 7-formamidino-7-deazaguanosine) is found at position 15 in the D-loop of many archaeal tRNAs (Gregson et al., 1993; Jühling et al., 2009; Sprinzl, Horn, Brown, Loudovitch, & Steinberg, 1998). However, this paradigm was changed by the recent discovery of 2'-deoxy-7-amido-7-deazaguanosine (dADG) and 2'-deoxy-7-cyano-7-deazaguanosine (dPreQ<sub>0</sub>) in bacterial genomic DNA and of 2'-deoxy-G<sup>+</sup> (dG<sup>+</sup>) in phage DNA (Thiaville et al., 2016). [Note that the standard nomenclature uses “Q” to represent the ribonucleoside, “preQ<sub>0</sub>” and “ADG” to represent the corresponding base, “dPreQ<sub>0</sub>” and “dADG” to represent the corresponding 2'-deoxyribonucleosides.] In phages such as the *Escherichia coli* phage 9g, the presence of dG<sup>+</sup> confers resistance to many restriction enzymes (Kulikov et al., 2014; Tsai, Corrêa, Xu, & Xu, 2017). The eleven-gene deazapurine-in-DNA (*dpd*) cluster of *Salmonella enterica* serovar Montevideo ATCC BAA-710 (*S. Montevideo*) was shown to be involved in modification of genomic DNA with dPreQ<sub>0</sub> and dADG, and in restriction-like activity that caused low transformation efficiency when using unmodified DNA compared with modified DNA. Deletion of the whole cluster in *S. Montevideo* eliminated both modification and proposed restriction-like activities, and the presence of the cluster in other phylogenetically-diverse bacteria correlated with the presence of dADG in DNA (Thiaville et al., 2016). A R-M system encompassing a 20kb region is unusual, as classical systems are usually much smaller (Roberts, Vincze, Posfai, & Macelis, 2015; Wilson, 1991).

All bacterial *dpd* islands include a set of 11 common genes with minor variations. The signature gene for the island, *dpdA*, encodes a homolog of the archaeal tRNA-guanine transglycosylase (aTGT; EC 2.4.2.48), the enzyme involved in inserting the 7-cyano-7-deazaguanine (preQ<sub>0</sub>) precursor into archaeal tRNAs (Bai, Fox, Lacy, Van Lanen, & Iwata-Reuyl, 2000; Watanabe et al., 1997) (Fig. 1A). Analysis of DpdA sequences suggests that, like aTGT, these proteins bind preQ<sub>0</sub>. *dpdA* is usually surrounded by the *dpdB* and *dpdC* genes in a tight physical cluster. *dpdB* encodes a protein similar to DndB, a DNA binding protein involved in regulating expression of genes involved in phosphorothioate DNA modifications (He et al., 2015), although the exact molecular function of DndB is not clear. The Aravind group has proposed that DpdC might be similar to ArcS, the archaeal amidotransferase that modifies the preQ<sub>0</sub> ribonucleoside into G<sup>+</sup> (Phillips et al., 2010) (Fig. 1A) based on very weak homologies around the active site (Iyer, Zhang, Maxwell Burroughs, & Aravind, 2013). Further sequence analysis (Alva, Nam, Söding, & Lupas, 2016; Hildebrand, Remmert, Biegert, & Söding, 2009) showed that DpdC harbored a domain of unknown function DUF328 (Thiaville et al., 2016). The *dpd* clusters encode several putative DNA-binding enzymes, including a member of the DEAD/DEAH box

helicase family (*dpdJ*), a SNF2-type helicase (*dpdE*), and a RecQ-like Superfamily II DNA helicase (*dpdF*) (Fig. 1B). *dpdK* encodes a protein harboring a phospholipase D-like domain. Little information is available about the proteins encoded by *dpdD*, *dpdG* and *dpdI*, but sequence analyses revealed low-confidence similarities of DUF3479 with a small portion of DpdD, of PF09821 with DpdG, and of PF15967 with DpdI (Thiaville et al., 2016).

The *S. Montevideo* *dpd* cluster has been clearly shown to encode a R-M system (Thiaville et al., 2016), but it is still poorly understood, and the actual function of each gene has yet to be determined. Which genes are required for the formation of the modification in DNA, which genes are required for cleaving unmodified DNA, and why so many genes are involved, are all open questions.

PreQ<sub>0</sub> is the common precursor of Q in Bacteria and G<sup>+</sup> in Archaea (Fig. 1). It is synthesized from GTP in both kingdoms. The first step of the pathway, catalyzed by GTP cyclohydrolase I (GCHI; EC 3.5.4.16), is shared in Bacteria with the tetrahydrofolate synthesis pathway (Phillips et al., 2008), and produces dihydroneopterin triphosphate (H<sub>2</sub>NTP). H<sub>2</sub>NTP is then converted by 6-carboxytetrahydropterin synthase (QueD; EC 4.1.2.50) and 7-carboxy-7-deazaguanine synthase (QueE; EC 4.3.99.3) to give 7-carboxy-7-deazaguanine (CDG) (McCarty, Lin, Jacobsen, & Bandarian, 2009; Reader, Metzgar, Schimmel, & de Crécy-Lagard, 2004). The last step in preQ<sub>0</sub> synthesis is catalyzed by 7-cyano-7-deazaguanine synthase (QueC; EC 6.3.4.20) (Nelp & Bandarian, 2015; Reader, Metzgar, Schimmel, & de Crécy-Lagard, 2004) in two ATP dependent half-reactions. Adenylation of CDG, followed by the addition of ammonia, generates 7-amido-7-deazaguanine (ADG) as an amide intermediate. Subsequently, dehydration of ADG consumes a second equivalent of ATP and produces the nitrile product, preQ<sub>0</sub> (Nelp & Bandarian, 2015). The discovery of dADG and dpreQ<sub>0</sub> in the DNA of species like *S. Montevideo* that use preQ<sub>0</sub> in Q synthesis in tRNA raises many questions. Do the synthesis pathways for 7-deazapurine production in RNA and DNA share a common precursor? Is there any competition for pathway intermediates? Here we define the necessary genes and propose a synthetic pathway for incorporation of the PreQ<sub>0</sub> and ADG bases into DNA.

## Results

### The *S. Montevideo* *dpd* island is composed of two transcription units

The 11 genes of the *dpdC-D* island are encoded by the same DNA strand, some overlapping, and could hence be in one transcriptional operon. RT-PCR analysis was performed on cDNA synthesized from RNA extracted from the WT *S. Montevideo* strain. (Fig. S1 and Methods). Each intergenic region was successfully amplified with a unique pair of primers, except the ~580-bp intergenic region between *dpdB* and *dpdE* (Fig. S1B). This suggests the presence of two operons: the *dpdCAB* operon and the *dpdEFGHIJKD* operon (Fig. S1A).

Promoter and terminator prediction tools reinforce the RT-PCR results. The BPROM promoter prediction tool (Softberry Inc., Solovyev & Salamov, 2011) predicts one promoter ~20 bp upstream the *dpdC* gene in the *S. Montevideo* 20 kb island and another ~200 bp upstream *dpdE* (Fig. S1A). The ARNold terminator prediction tool (Université Paris-Sud web server, Gautheret and Lambert, 2001; Macke et al., 2001) predicted three rho-

independent terminators in the *dpd* gene cluster with both the Erpin and RNAmotif programs. The top-scoring one is located ~8 bp downstream of the *dpdD* gene. The one with the second-best score is located ~15 bp downstream of *dpdB* (Fig. S1A).

### The *dpdEFGHIJKD* operon may encode a restriction system inhibited by the dADG modification

We have previously shown that the transformation efficiency of the WT *S. Montevideo* strain with pUC19 DNA lacking dADG and dPreQ<sub>0</sub> was reduced compared to pUC19 DNA containing dADG modifications. The efficiency of transformation was fully recovered when YYF3022, a *S. Montevideo* derivative with a 21-kb deletion eliminating the entire *dpd* cluster, was used a recipient (Fig. 5 in Thiaville et al., 2016), suggesting the *dpd* island encodes a R-M system. Growth rate and yield of the YYF3022 strain did not show significant changes under standard laboratory conditions compared to the WT (Fig. S2).

The whole *dpdEFGHIJKD* operon was deleted as described in the Methods section to give strain YYF3345 (Fig. S3). When unmodified pUC19 DNA was transformed into this strain, a ~100-fold increase in transformation efficiency was observed compared to a WT recipient (Fig. 2A), suggesting that genes required for the restriction-like phenotype are part of the *dpdEFGHIJKD* operon. Individual deletions of every gene of this operon were then tested. Transformation efficiencies with unmodified pBAD33 DNA were low in the *dpdF* or *dpdH* mutants, as in the WT strain (Fig. 2C). On the other hand, deletion of *dpdE*, *dpdG*, *dpdI*, *dpdJ*, *dpdK* or *dpdD* resulted in a significant increase in transformation efficiency that was abolished with the expression of the corresponding gene *in trans* (with the exception of *dpdJ* encoding a Lhr-like helicase that we were unable to clone, hence in this case a polar effect cannot be eliminated). We also showed that the none of the *dpdEFGHIJKD* genes are involved in the synthesis of the modifications, as DNA extracted from the *dpdE-dpdD* strain still harbors the dADG modification (Fig. 2B and Table 1).

### *dpdCAB* are necessary for ADG insertion in DNA

We were able to delete the *dpdA* gene only in a *dpdD* strain (YYF3014). When the pUC19 DNA extracted from the *dpdA::kan dpdD* strain was used to transform WT cells, transformation efficiencies were low and comparable to the results with unmodified pUC19 DNA extracted from YYF3022 strain lacking the whole *dpd* cluster. This suggested that DpdA is required for the formation of dADG and/or dPreQ<sub>0</sub> modifications (Fig. 2B). This hypothesis was confirmed by mass spectrometric analysis of genomic DNA extracted from the *dpdA* mutant as both dADG and dPreQ<sub>0</sub> levels were below the detection limits (Table 1). Using a similar strategy, deletion of *dpdB* and *dpdC* was performed in the *dpdD* background, to give the YYF3401 (*dpdC::kan dpdD*) and YYF3402 (*dpdB::kan dpdD*) strains, respectively. Mass spectrometric analysis of genomic DNA from these strains revealed that they lacked dADG, demonstrating that the *dpdB* and *dpdC* genes were also required for ADG formation in DNA. However, dPreQ<sub>0</sub> was detected in DNA from the *dpdC* strain, but not from the *dpdA* or *dpdB* strains (Table 1), suggesting that *dpdC* is dispensable for dPreQ<sub>0</sub> formation.

To confirm that all three genes were indeed required for dADG synthesis and our results were not due to polar effects, we first showed that deletion of *dpdC* or *dpdA* did not affect transcription of *dpdAB* or *dpdB* respectively by RT-PCR (Fig. S4). In parallel, we transformed each individual mutant with a plasmid expressing the deleted gene *in trans* and analyzed the genomic DNA of the resulting strains LC-MS/MS. As shown in Table 1, the dADG deficiency phenotype was complemented in all three cases. As dPreQ<sub>0</sub> levels were low and not consistently detected in WT DNA, its absence in the complemented strains was not really conclusive.

### ***dpdAB* are sufficient for dPreQ<sub>0</sub> synthesis and *dpdC* is required for dADG formation**

The previous genetic results suggested that only DpdA and DpdB were necessary for the synthesis of dPreQ<sub>0</sub> (Table 1, strain YYF3401). To test this hypothesis, the YYF3022 strain that lacks the entire *dpd* island was transformed with plasmids expressing *dpdAB* or *dpdCAB*. Mass spectrometric analysis of the genomic DNA of these strains (Table 1) showed an increased amount of dPreQ<sub>0</sub> compared to WT when *dpdAB* was overexpressed (the YYF3324 strain), while dADG could be detected only in the YYF3425 strain expressing the three genes (*dpdCAB*).

To address the question of what modified base (preQ<sub>0</sub> and/or ADG) conferred protection from the potential restriction activity, we isolated plasmid DNA from a variety of strains and investigated the transformation efficiencies of each of the plasmid samples. Transformation efficiencies were high when pUC19 was extracted from *S. Montevideo* mutant strain YYF3022 (Fig. 3A), *Salmonella enterica* serovar Typhimurium LT2 and *E. coli* K12 (Fig. 3B) expressing *dpdCAB in trans*. These results suggest that *dpdC*, *dpdA* and *dpdB* represent the minimal set of genes required for ADG insertion. In contrast, the transformation efficiencies were low and comparable to the negative control when pUC19 was extracted from these strains expressing only *dpdAB* (Fig. 3). Taken together the data support the conclusion that it is dADG, and not dPreQ<sub>0</sub>, that confers protection from the potential *dpd* restriction machinery. The data are also consistent with *dpdC*, *dpdA* and *dpdB* comprising the minimal set of genes required for dADG formation in DNA, at least in organisms that synthesize the preQ<sub>0</sub> precursor as discussed below.

### **PreQ<sub>0</sub> is the precursor of the dADG modification**

PreQ<sub>0</sub> and ADG are intermediates in the tRNA Q modification pathway (Fig. 1), with QueC converting CDG to preQ<sub>0</sub> through the ADG intermediate (Nelp & Bandarian, 2015). *S. Montevideo* possesses all of the genes involved in the Q pathway (Fig. 1) (Reader, Metzgar, Schimmel, & de Crecy-Lagard, 2004; Thiaville et al., 2016) and we have previously shown that Q is present in tRNA extracted from this organism (Thiaville et al., 2016). The parsimonious hypothesis was therefore that *S. Montevideo* QueC synthesizes the precursors needed for the formation of dPreQ<sub>0</sub> and dADG in DNA as well as Q in tRNA. We were therefore surprised that deletion of the *S. Montevideo queC* in a WT background produced the viable strain YYF3346. Indeed, if this gene is essential to the modification, then it should be impossible to delete *queC* without also deleting the potential restriction system, since the *S. Montevideo* putative restriction system would cleave its own DNA.

Using a gel-based Q detection method, we showed that this strain lacked Q in tRNA<sup>Asp</sup><sub>GUC</sub> (Fig. 4A, left), consistent with disruption of the Q pathway, and that this phenotype was complemented by expressing the *queC* gene in *trans* (Fig. 4A, middle, see methods). To determine the modification status of DNA in the *queC* mutant pUC19 was extracted from this strain and used to transform WT host cells (Fig. 4B). The low efficiency of transformation suggested that the DNA lacked the dADG modification, and subsequent mass spectrometric analysis confirmed that neither dADG or dPreQ<sub>0</sub> were detected in DNA from the *queC* strain, but that both modifications were present in the complemented strain YYF3461 (Table 1). This demonstrates that *queC*, and the other genes required for preQ<sub>0</sub> biosynthesis, are required for the synthesis of the dADG/dPreQ<sub>0</sub> modifications.

The fact that YYF3346 was viable despite the absence of dADG suggested that the *queC* mutant might also be deficient in restriction. Testing the restriction-like phenotype confirmed this suspicion (Fig. 4C), explaining the viability of the strain. The restriction deficiency was not complemented by expressing the *queC* gene *in trans* (Fig. 4C), suggesting that this phenotype was due to second-site mutations that inactivated the restriction machinery. However, none of the *dpd* genes we tested complemented the restriction deficiency phenotype (Fig. S5), leading us to sequence the *dpdJ* gene (4518 bp), the only gene in the cluster that we could not clone. Sequencing showed a G-to-C mutation at position 272 (Proline to Arginine) in YYF3346 compared with the WT strain, indicating that a second-site mutation had occurred in the *dpdJ* gene.

### Role of DpdC in dADG synthesis

While the previous results showed that DpdC is required for dADG synthesis, several models for its role in the pathway can be proposed. DpdC could be channeling ADG away from QueC or inhibiting part of the enzyme (Fig. 5A and 5B), or it could be converting preQ<sub>0</sub> to ADG before or after its insertion in DNA (Fig. 5C and 5D). Providing the *queC* strain (YYF3346) with exogenous preQ<sub>0</sub> suppressed the Q<sup>-</sup> phenotype (Fig. 4A, right), which demonstrates that, like *E. coli*, *S. Montevideo* can transport exogenous preQ<sub>0</sub> (Zallot, Yuan, & de Crécy-Lagard, 2017). pUC19 extracted from the *queC* strain exhibited higher transformation frequencies in WT cells when the *queC* mutant was provided with exogenous preQ<sub>0</sub> (Fig. 5E). This was consistent with the presence of dADG, which was confirmed by mass spectrometric analysis (Table 1). These results suggests that DpdC can synthesizes dADG directly from preQ<sub>0</sub> and does not require prior ADG synthesis by QueC, hence disproving the two models shown in Figure 5A and 5B.

To confirm DpdC produces ADG from preQ<sub>0</sub>, we used an indirect assay in a heterologous system. We postulated that if DpdC converted preQ<sub>0</sub> to ADG before the incorporation of preQ<sub>0</sub> into DNA, then DpdC should compete with QueF for the available preQ<sub>0</sub> and the formation of Q in tRNA would be reduced when *dpdC* is overexpressed. To detect this potential competition, conditions of preQ<sub>0</sub> limitation had to be engineered. This is the case in an *E. coli queC* strain when 10 nM of exogenous preQ<sub>0</sub> is added and cells are harvested shortly after supplementation. When *dpdC* was overexpressed, we saw a reduction of Q formation in tRNA (Fig. 6A), suggesting that DpdC competes with QueF for the available preQ<sub>0</sub>. This reduction was not observed when *dpdA* or *dpdB* were overproduced



individually (Fig. 6B). However, when both *dpdA* and *dpdB* were overexpressed, Q formation in tRNA was lower than when *dpdC* was overexpressed (Fig. 6A), consistent with the DpdAB complex competing more effectively for free preQ<sub>0</sub> than DpdC. The fact that mass spectrometric analyses have shown that dADG, not dPreQ<sub>0</sub>, is the predominant form found in *S. Montevideo* DNA (Table 1) suggests that the conversion of ADG from preQ<sub>0</sub> might occur after incorporation of preQ<sub>0</sub> in DNA, as depicted in the model in Figure 5D.

### Interactions of Dpd proteins with DNA

To further address the specific functions of the Dpd protein in dADG synthesis, we investigated the DNA binding properties of the recombinant DpdA, DpdB, and DpdC proteins in a set of preliminary experiments as we have yet to identify the target sequence. The *dpdABC* genes were each subcloned into *E. coli* expression vectors to allow overproduction of recombinant C-terminal His<sub>6</sub>-affinity-tagged proteins. The cleaved recombinant proteins (Fig. S6), which retained a C-terminal Factor Xa sequence (IGER), were incubated at a range of concentrations with linearized pUC19 plasmid DNA followed by electrophoresis on 0.5% agarose gels. As can be seen (Fig. 7), all three of the proteins clearly exhibit binding to the DNA relative to the BSA negative control (Fig. 7A), with DpdA exhibiting the strongest interaction (Fig. 7B). DpdB, which on the basis of the genetic experiments we hypothesized might provide binding specificity for DpdA, exhibited the weakest interaction (Fig. 7C), and did not appear to enhance binding of DpdA or DpdC when present together with either of them (Fig. 7E and 7F, respectively). While initially surprising, this result is not inconsistent with the proposal that DpdB determines DpdA location if DpdA possesses a non-specific affinity for DNA and requires DpdB for targeting to the specific sites where preQ<sub>0</sub> is to be inserted. Indeed, while it is tempting to interpret the high affinity of DpdA binding to the potential formation of the covalent intermediate in the base exchange reaction, the requirement for both DpdA and DpdB for the incorporation of preQ<sub>0</sub> into DNA argues against this interpretation and suggests that the binding is non-specific. Notably, the distinct DNA binding exhibited by DpdC is consistent with the results of the competition experiment (*vide supra*), which suggested that DpdC catalyzes hydrolysis of the preQ<sub>0</sub> nitrile group in the context of preQ<sub>0</sub>-modified DNA (Fig. 5D), not the free base.

### Discussion

Here we have demonstrated that preQ<sub>0</sub> base is a shared intermediate in the tRNA and DNA 7-deazapurine derivative modification pathways, and that only three genes of the *S. Montevideo* *dpd* islands, part of the *dpdABC* operon, are required for the formation of preQ<sub>0</sub> and ADG in DNA. Based on these results, the model we favor is that the DpdAB complex inserts preQ<sub>0</sub> into DNA as the first step in modifying DNA, followed by DpdC-catalyzed hydration to dADG, although the alternate model in which DpdC acts at the level of the free base before ADG insertion in DNA by DpdAB cannot be rigorously ruled out at this time. The fact that in our indirect competition assay the DpdAB complex, and not DpdA or DpdB alone, effectively competes for preQ<sub>0</sub> was surprising, since DpdA, the TGT paralog predicted to contain a preQ<sub>0</sub> binding pocket, is the logical candidate for the enzyme responsible for the base exchange (Thiaville et al., 2016). However, the results are consistent with our genetic data, which demonstrated that the base exchange reaction in DNA only

occurs in the presence of both DpdA and DpdB, at least in the *S. Montevideo* system. Indeed, DpdB might not always be required, as we identified fused DpdCA proteins in several organisms in which *dpdB* is missing, such as *Meiothermus chliarophilus* DSM 9957, *Spirosoma spitsbergense* DSM 19989, *Haloarcula sinaiensis* ATCC 33800, *Herbaspirillum* sp. JC206 and *Paenibacillus polymyxa* SC2 (Thiaville et al., 2016, Table. S1). Only when the dADG insertion system can be reconstituted *in vitro* will we be able to fully confirm the predicted biochemical roles of the three DpdABC proteins. Trials are ongoing, the results of which will be reported in due course.

Results from the transformation assays with both unmodified DNA and DNA containing preQ<sub>0</sub> or ADG show that only ADG confers resistance to the predicted restriction activity encoded by the *dpd* cluster. Unlike dG<sup>+</sup> in 9g phage (Tsai et al., 2017), the presence ADG in pUC19 does not confer resistance to the Type II restriction enzymes we tested (Fig. S7). However we do know that the 9g phage is heavily modified while the plasmid carries only a few modified bases (Thiaville et al., 2016). Further work is needed to explore if high densities of dADG can like dG<sup>+</sup> protect from Type II restriction enzyme cleavage.

Genetic analyses suggest that at least six of the eight genes of the *dpdEFGHIJKD* operon are part of a nuclease system that restricts DNA lacking ADG. Here again, biochemical studies will be required to provide a definitive confirmation and to elucidate the specific role(s) of each protein. Though the classification scheme for R-M systems was initially designed specifically for methylations, the *dpd* system would be most similar to a type I RM system (Loenen & Raleigh, 2014). Indeed, we identified a “modification complex” (DpdABC), composed of a potential DNA sequence specificity protein and two modification proteins, and a “restriction complex” (potentially DpdEFGHIJKD) that contains SNF2 helicase homologs (see Thiaville et al., 2016 for description). However, the complexity of this system, with at least six different proteins involved in restriction, and the novelty of the modification, could warrant a new RM system category.

## Experimental Procedures

### Bioinformatics

Sequences were acquired from the PubSEED subsystem “dpd cluster”. Promoter prediction was performed by BPROM online server (Softberry Inc., Solovyev & Salamov, 2011). Terminator prediction was performed by ARNold web server (Université Paris-Sud web server, Gautheret and Lambert, 2001; Macke et al., 2001). For other sequence analyses, the BLAST tools (Altschul, Gish, Miller, Myers, & Lipman, 1990) and resources at NCBI (<http://www.ncbi.nlm.nih.gov/>) were routinely used. The protein sequences were retrieved from the NCBI using the following accession numbers: DpdA, AHW12286.1; DpdB, AHW12287.1; DpdC, AHW12295.1 and AHW12296.1; DpdD, AHW12295.1; DpdE, AHW12288.1; DpdF, AHW12289.1; DpdG, AHW12290.1; DpdH, AHW12291.1; DpdI, AHW12292.1; DpdJ, AHW12293.1; DpdK, AHW12294.1; QueC, AHW08582.1.

## Strains, Media, and Growth Conditions

Strains used in this study are listed in (Table S1). *S. Montevideo* strains were routinely grown in LB (Tryptone 10 g L<sup>-1</sup>, yeast extract 5 g L<sup>-1</sup>, NaCl 5 g L<sup>-1</sup>) at 37 °C. *S. Montevideo* deletion constructs were made using the linear recombination method described by Datsenko and Wanner (Datsenko & Wanner, 2000). Antibiotic resistance gene was eliminated also as described by Datsenko and Wanner (Datsenko & Wanner, 2000). Oligonucleotides used for RT-PCR (Fig. 2), mutants construction (Fig. 3 and Fig. S2) and plasmids construction are listed in Table S2. Primers SM\_deletion\_dpdA\_F and SM\_deletion\_dpdA\_R were used for the introduction of the *dpdA* deletion; SM\_deletion\_dpdB\_F and SM\_deletion\_dpdB\_R for *dpdB* deletion; SM\_deletion\_dpdC\_F and SM\_deletion\_dpdC\_R for *dpdC* deletion; SM\_deletion\_dpdD\_F and SM\_deletion\_dpdD\_R for *dpdD* deletion; SM\_deletion\_dpdE\_F and SM\_deletion\_dpdE\_R for *dpdE* deletion; SM\_deletion\_dpdF\_F and SM\_deletion\_dpdF\_R for *dpdF* deletion; SM\_deletion\_dpdG\_F and SM\_deletion\_dpdG\_R for *dpdG* deletion; SM\_deletion\_dpdH\_F and SM\_deletion\_dpdH\_R for *dpdH* deletion; SM\_deletion\_dpdI\_F and SM\_deletion\_dpdI\_R for *dpdI* deletion; SM\_deletion\_dpdJ\_F and SM\_deletion\_dpdJ\_R for *dpdJ* deletion; SM\_deletion\_dpdK\_F and SM\_deletion\_dpdK\_R for *dpdK* deletion. SM\_deletion\_dpdE\_F and SM\_deletion\_dpdD\_R were used for the introduction of the *dpdE-D* deletion. *S. Montevideo dpdA* was cloned into the sites *SbfI* and *HindIII* of pBAD24 using primers GO36 and GO37 following standard procedures. *dpdC* and *dpdB* was cloned into the same sites with GO34 and GO35 and GO38 and GO39 respectively. *dpdAB* was cloned with GO36 and GO39. *S. Montevideo dpdCAB* was cloned into the sites *NheI* and *HindIII* of pBAD33 using primers Sm-NheI-RBS-dpdC-F and Sm-dpdB-HindIII-R. *S. Montevideo queC* was cloned into the sites *EcoRI* and *HindIII* of pBAD24 using primers Sm-EcoRI-RBS-queC-F and Sm-HindIII-queC-R. *dpdE* was cloned into the sites *SbfI* of pBAD24 using primers GO136 and GO82. *dpdD*, *dpdF*, *dpdG*, *dpdH*, *dpdI*, and *dpdK* were cloned into the *SaI* and *SbfI* sites of pBAD24 using primers GO72 and GO73, GO83 and GO84, GO85 and GO86, GO87 and GO88, GO89 and GO90, and GO137 and GO106, respectively.

For recombinant protein production the *dpdABC* genes were subcloned from the relevant pBAD constructs into the *NdeI/NotI* sites of pET30-Xa to give C-terminal His<sub>6</sub>-affinity tagged proteins with a Factor Xa site preceding the affinity tag. The cloning utilized the primers Sm-TGTa5-F and SM-TGTa5-R to generate pAGI-42.1 (*dpdA*), Sm-DpdB-F and Sm-DpdB-R to generate pAGI-42.2 (*dpdB*), and Sm-DpdC-F and Sm-DpdC-R to generate pAGI-42.3 (*dpdC*). Expression and purification of the recombinant proteins followed standard protocols.

## Plasmid Restriction Test

Restriction of the pUC19 plasmid was tested using a modified transformation efficiency (TE) method (Xu, Yao, Zhou, Deng, & You, 2010). TE reflects the number of transformants obtained per microgram of DNA and is calculated by dividing the number of transformants by the amount of DNA used in the electroporation. The pUC19 plasmid or the mix of pUC19 plus pBAD33 plasmid was propagated through *S. Montevideo* WT, YYF3022 (*dpdC-dpdD::kan*) and other strains, and isolated from each strain using QIAprep Spin

Miniprep Kit (Qiagen). 0.4% arabinose was added 30 min before plasmid extraction to induce the  $P_{BAD}$  promoter. The presence or absence of the modification on pUC19 was confirmed as described (Thiaville et al., 2016). Plasmid (modified or unmodified) was transformed by electroporation into freshly prepared electrocompetent recipient strains. Transformants were recovered in 1 mL LB for 1 h at 37 °C and dilutions were plated onto LB agar plates with or without antibiotic selection for the transforming DNA: ampicillin ( $100 \mu\text{g mL}^{-1}$ ). Transformation efficiency was determined by calculating the number of transformants per  $10^6$  viable CFU and dividing by the ng of DNA transformed.

### DNA Preparation

Total DNA was extracted from bacteria with phenol-chloroform followed by alcohol precipitation as described (Thiaville et al., 2016). A 400 mL culture of each strain was grown in the proper conditions described above. 0.4% arabinose and  $500 \text{ ng mL}^{-1}$  aTET was added 30 minutes before harvest to induce the  $P_{BAD}$  and  $P_{TET}$  promoter respectively. Cells were harvested by centrifugation at 4,500 rpm for 20 min. Pellets were washed once in 40 mL TEN buffer (100 mM Tris, pH 8.0; 100 nM NaCl; 10mM EDTA, pH 8.0) and centrifuged again. Pellets were suspended in 20 mL TEN buffer with  $600 \mu\text{g mL}^{-1}$  lysozyme, 1 % SDS,  $6 \text{ mAU mL}^{-1}$  Proteinase K (Qiagen), and incubated at 56 °C for 1 h. Following incubation, an equal volume of Tris-buffered phenol, pH 7 was added to each sample and incubated with shaking at room temperature overnight. The samples were centrifuged at  $4,000 \times g$  at room temperature for 10 min and the aqueous phase was transferred to the new tubes. The extractions were repeated with the mixture of 25:24:1 phenol:chloroform:isoamyl alcohol followed by a chloroform extraction. After the final extraction, the aqueous phase was treated with  $200 \mu\text{g mL}^{-1}$  RNase A for 30 min at 37 °C. RNase was removed by another round of phenol:chloroform extraction. DNA was precipitated with 0.1 volume sodium acetate and 1 volume isopropanol and spooled with a glass rod. After washing briefly in 70% ethanol, the DNA was suspended in sterile deionized water.

### DNA Modification Analysis

Quantification of dADG and dPreQ<sub>0</sub> was achieved by liquid chromatography-coupled triple quadrupole mass spectrometry (LC-MS/MS) (Thiaville et al., 2016). DNA ( $100 \mu\text{g}$ ) was hydrolyzed in 10 mM Tris-HCl (pH 7.9) with 1 mM  $\text{MgCl}_2$  with Benzonase (20 U), DNase I (4 U), calf intestine phosphatase (17 U) and phosphodiesterase (0.2 U) for 16 h at ambient temperature. Following passage through a 10 kDa filter to remove proteins, the filtrate was lyophilized and resuspended to a final concentration of  $2 \mu\text{g } \mu\text{L}^{-1}$  (based on initial DNA quantity); a final concentration of  $0.2 \mu\text{g } \mu\text{L}^{-1}$  was used for dADG quantification. Aliquots of DNA hydrolysates ( $2 \mu\text{g}$ ) were injected on an Agilent 1290 series HPLC equipped with a Kinetex EVO C18 column (150 mm  $\times$  2.1 mm, 2.6  $\mu\text{m}$  particle size, 100Å pore size; Phenomenex) and a diode array detector (DAD). The column was heated to 40 °C and eluted at  $0.5 \text{ mL min}^{-1}$  with 100% solvent A (5 mM ammonium acetate, pH 5.3) for 2 min followed by a linear gradient to 10% solvent B (acetonitrile) over 8 min, 60% solvent B at 9 min and held for 1 min. The column eluent was monitored by UV absorbance at 260 nm to determine the retention times of canonical 2'-deoxynucleosides (dC, 1.2 min; dG, 3.8 min; dT, 4.2 min; dA, 5.5 min). The HPLC column was coupled to an Agilent 6430 triple

quadruple mass spectrometer with an electrospray ion source in positive ion mode and parameters as follows: gas temperature 300 °C, gas flow 12 L min<sup>-1</sup>, nebulizer pressure 40 psi, and capillary voltage 3800 V. 2'-Deoxynucleosides were quantified using selected reaction monitoring (SRM) for the following HPLC retention times and transitions involving loss of 2-deoxyribose: 6.7 min, *m/z* 292→176 for dPreQ<sub>0</sub>; and 6.4 min, *m/z* 310→194 for dADG. Quantification of dPreQ<sub>0</sub> and dADG was achieved using external calibration curves, with standards synthesized as described elsewhere (Thiaville et al., 2016). Nine calibration solutions were prepared with the lowest concentrations of 2 μM dPreQ<sub>0</sub> and 200 pM dADG. Calibration samples (10 μL) were interspersed with DNA hydrolysate samples. The limit of detection was determined to 1 fmol for dADG and dPreQ<sub>0</sub>, which corresponds to ~5 modifications per 10<sup>6</sup> nucleotides in 2 μg of injected DNA digest.

### Exogenous PreQ<sub>0</sub> feeding

Cells were cultured in M9 defined media with 1% glycerol. Arabinose was added to 0.4% after cells reached an optical density (*A*<sub>600nm</sub>) of 0.1 to induce the P<sub>BAD</sub> promoter. PreQ<sub>0</sub> (from Ark Pharm, Libertyville, IL, USA) and DMSO was added respectively after cells reached an optical density (*A*<sub>600nm</sub>) of 0.2 to 0.3. The transport reaction was stopped at time points of 0, 20, 40 and 60 min after supplementing with preQ<sub>0</sub> and DMSO by placing samples on melting ice, and then centrifuging, followed by tRNA extraction.

### tRNA Extraction

This extraction method was proven efficient for the purification of tRNA enriched fraction (Zallot, Yuan, et al., 2017). Cells were harvested by centrifugation at 16,000 × *g* for 2 min at 4 °C. Cell pellets were immediately resuspended in 1 mL of Trizol (Life). Small RNAs were extracted using Purelink miRNA Isolation kit (Life) according to manufacturer protocol. The purified RNAs were eluted in 50 μL of RNase free water.

### Detection of queuosine in bulk tRNA

This method of detection of Q in tRNA was originally developed by Igloi and Kossel (Igloi & Kössel, 1985) and adapted by Zaborske et al. (Zaborske et al., 2014). For each sample, bulk tRNAs were deacylated by incubation in 50 mM Tris-HCl (pH 9) for 30 min at 37°C. Deacylated tRNAs were precipitated at 16,000 *xg* at 4°C after incubation in ammonium acetate, isopropanol, and linear polyacrylamide overnight at -20 °C (Gaillard & Strauss, 1990). The pellet was washed with 70% ethanol and dissolved in RNase-free water (Life). Prepared tRNAs were quantified using a Nanodrop 1000 spectrophotometer. For each well, 150 ng of tRNAs were resuspended in 2× RNA Loading Dye (NEB) and loaded onto a denaturing 8 M urea, 8% polyacrylamide gel containing 0.5% 3-(acrylamido)phenylboronic acid (Sigma-Aldrich). The migration was performed at 4°C in 1× TAE. Migrated tRNAs were transferred onto a Biodyne B precut Nylon membrane (Thermo Scientific) using a wet transfer apparatus in 1× TAE at 150 mA at 4°C for 90 min. After the transfer, the membrane was baked in an oven for 30 min at 60°C and then UV irradiated in a UV Crosslinker (Fisher FB-UVXL-1000) at a preset UV energy dosage of 120 mJ cm<sup>-2</sup>. tRNA<sup>ASP</sup> was detected with the North2South Chemiluminescent Hybridization and Detection Kit (Thermo). The initial membrane blocking was realized with DIG Easy Hyb (Roche), because it drastically limits the background noise, compared to the membrane blocking buffer supplied with the

North2South kit. Hybridization was done at 60°C, with using the specific biotinylated primer for tRNA<sup>ASP</sup><sub>GUC</sub> (Phillips et al., 2012) (5'-biotin-CCCTCGGTGACAGGCAGG 3') at 50 ng mL<sup>-1</sup> final. The blot was exposed to X-Ray film (Thermo Scientific, CL-X Posure Film). The film was developed using a Film Processor (Konica QX-60A).

### DNA binding experiments

Purified plasmid pUC19 was linearized by cutting at the single *Sal*I site using FastDigest Sal I (Thermo) following manufacturer recommended protocols, purified on a 1% agarose gel, and isolated using a GeneJET gel extraction kit (Fermentas). A portion (0.5 µg) of linearized pUC19 was incubated in a 20 µL reaction at 37 °C for 10 min in the presence of the relevant Dpd protein(s) at a concentration of either 100 nM, 200 nM, 500 nM, 1 µM, 2 µM, or 4 µM, along with 20 mM HEPES (pH 7.5), 50 mM KCl, 10 mM MgCl<sub>2</sub>, and 2 mM 2-mercaptoethanol (BME). In experiments containing two proteins the total protein concentration was the same as in single protein experiments, each present at half. The incubations were then subjected to electrophoresis on a 0.5% agarose gel containing 0.33 µg/mL ethidium bromide made with a TBE buffer containing 89 mM Tris (pH 8.3), 89 mM boric acid, and 2 mM EDTA. The gels were run in the same TBE buffer for 40 min at 107 V.

### Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

### Acknowledgments

This work was funded by the National Institutes of Health (R01 GM70641 to V dC-L) and the National Research Foundation of Singapore through its Singapore-MIT Alliance for Research and Technology Antimicrobial Resistance IRG (to PCD). A portion of the mass spectrometric analyses were performed in the Bioanalytical Facilities Core of the MIT Center for Environmental Health Sciences, which is supported by a grant from the National Institute for Environmental Health Sciences (ES002109).

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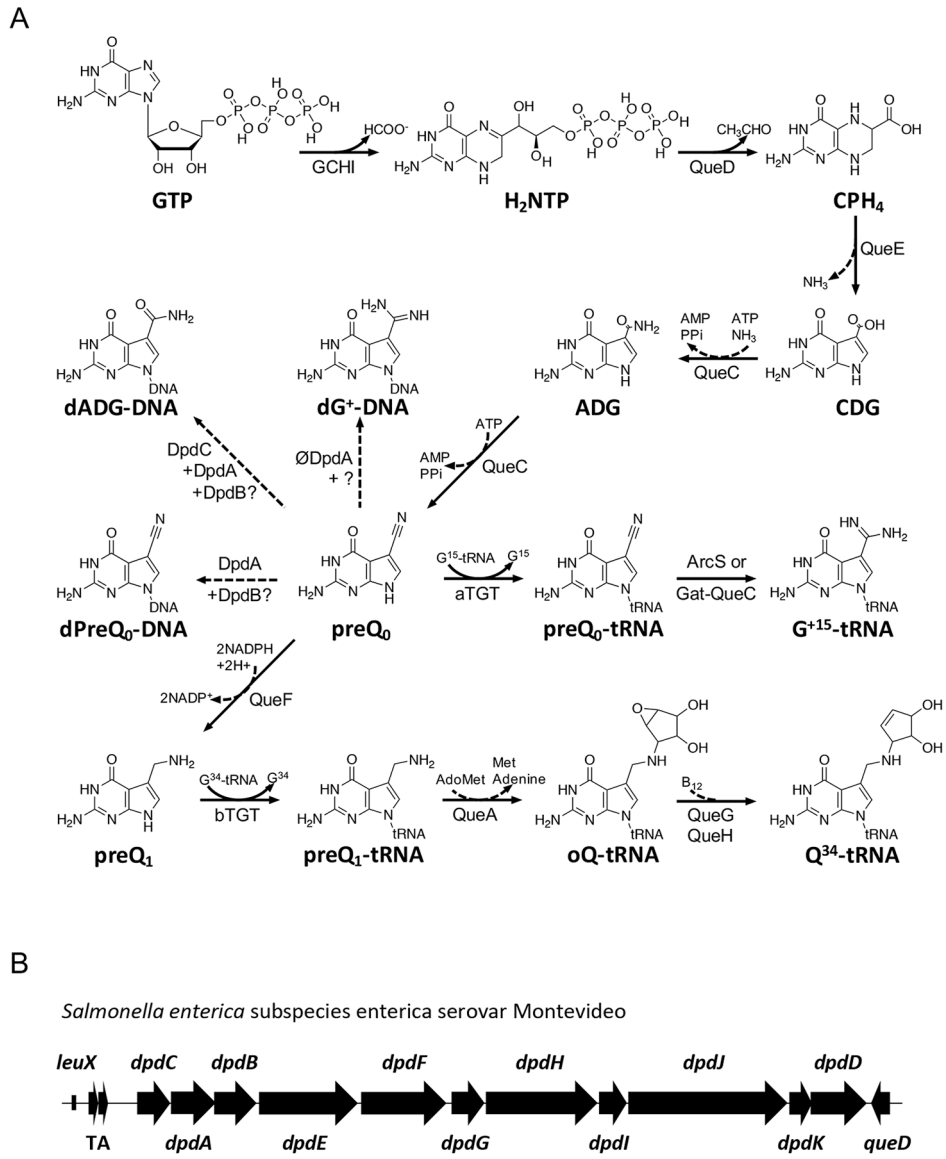
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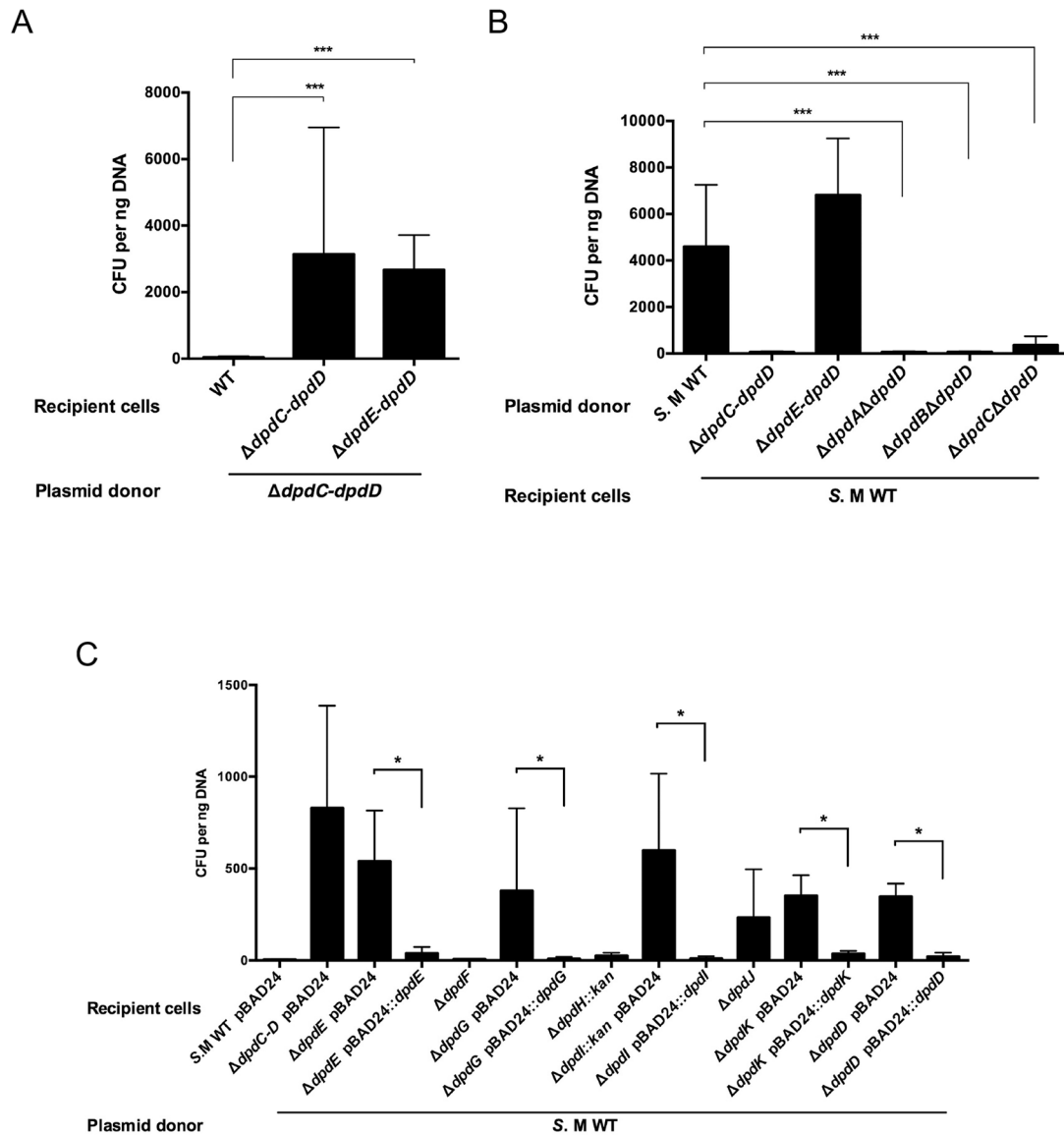


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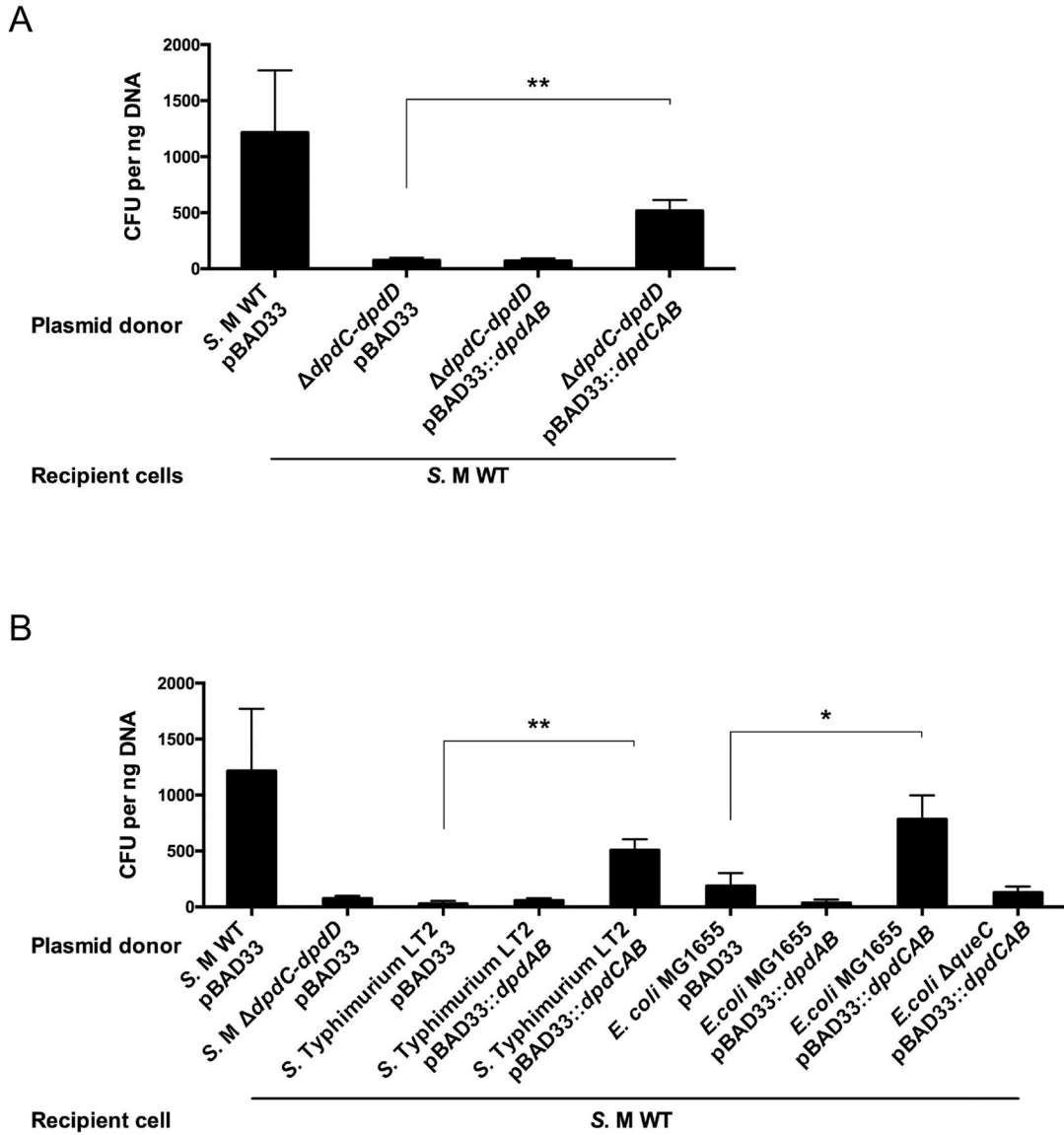
**Figure 1. Biosynthesis of 7-deazaguanine modifications in tRNA and DNA.**

(A) The biosynthetic pathways for queuosine and archaeosine in tRNA. Dashed arrows represent proposed reactions. CPH4, 6-carboxytetrahydropterin; bTGT, tRNA-guanine transglycosylase. Enzymes abbreviations are described in text except for QueA, QueG and QueH that are described in (Zallot, Harrison, Kolaczkowski, & de Crécy-Lagard, 2016) in Zallot, Ross, et al., 2017. (B) The gene cluster governing 7-deazapurine biosynthesis in *Salmonella enterica* subspecies *enterica* serovar *Montevideo* (GCA\_000238535.2; TgtA5 UniProt ID E7V8J4) (Thiaville et al., 2016). TA, toxin-antitoxin gene pair (*ccdA*, *ccdB*); *leuX*, tRNA-leu gene.

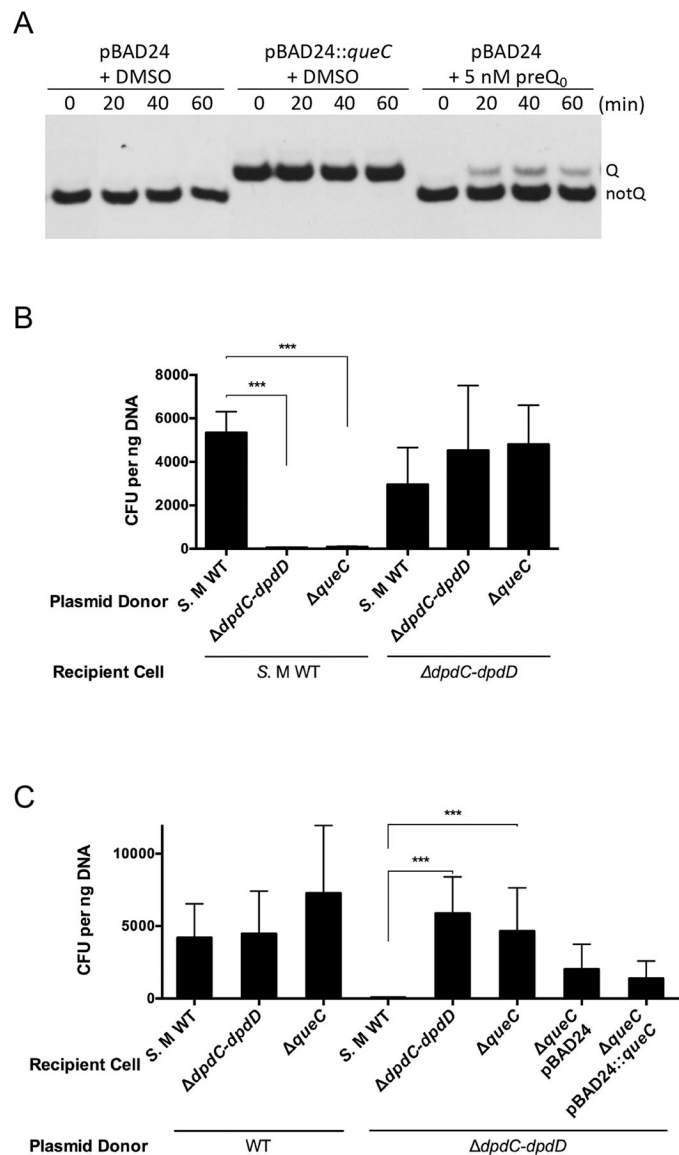


**Figure 2. The *dpdEFGHIJKD* mutant showed WT level transformation efficiency as donor and increased transformation efficiency as host.**

(A) Unmodified pUC19 DNA (10 ng) was transformed in the *S. Montevideo* WT and *dpdE-dpdD::kan* strains. (B) pUC19 DNA (10 ng) extracted from different mutant donor cells was transformed in *S. Montevideo* WT. (C) Unmodified pBAD33 DNA (10 ng) was transformed in the WT host and in each individual mutant strain containing pBAD24 or its derivative expressing the corresponding *dpd* gene *in trans* (with the exception of *dpdJ* that could not be cloned). The average of three experiments is shown, with error bars representing SD (\* $P < 0.05$ , \*\*\* $P < 0.001$ , two-tailed Student's *t*-test).

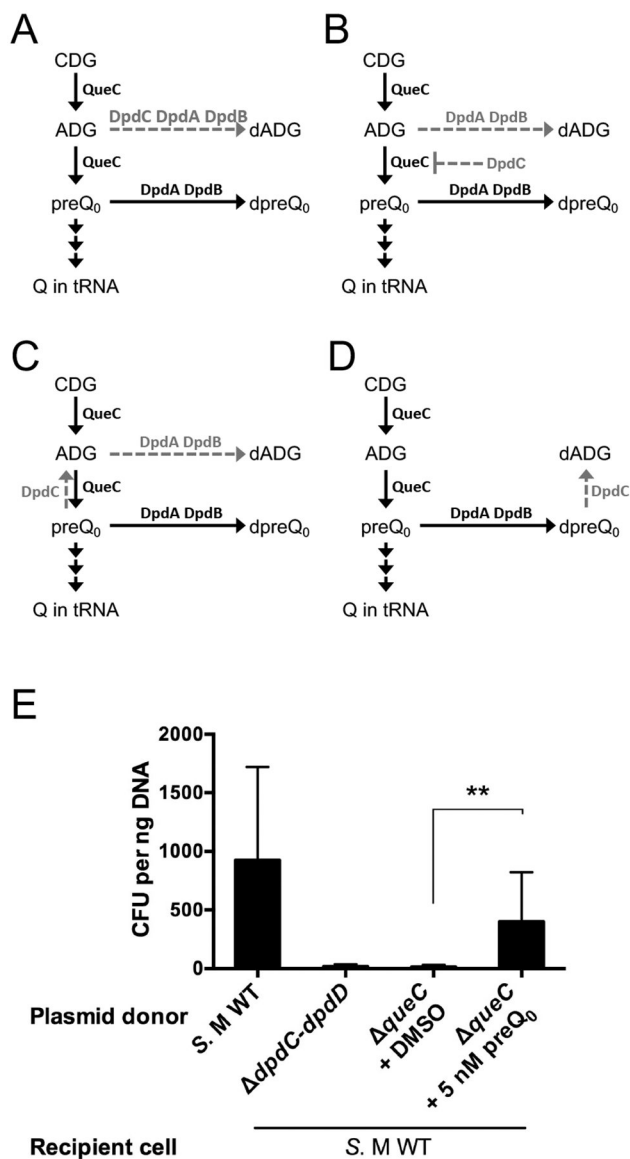


**Figure 3. Expression of *dpdCAB* genes restores high transformation efficiencies in the WT strain.** (A) pBAD33 and pUC19 DNA (20 ng of total) extracted from *S. Monteideo* strains carrying empty pBAD33 vector, pBAD33::*dpdAB* or pBAD33::*dpdCAB* was transformed in the WT strain. (B) pBAD33 and pUC19 DNA (20 ng of total) extracted from the indicated strains was transformed in the *S. Monteideo* WT strain. The average of three experiments is shown, with error bars representing SD (\*P < 0.05, \*\*P < 0.01, two-tailed Student’s t-test).



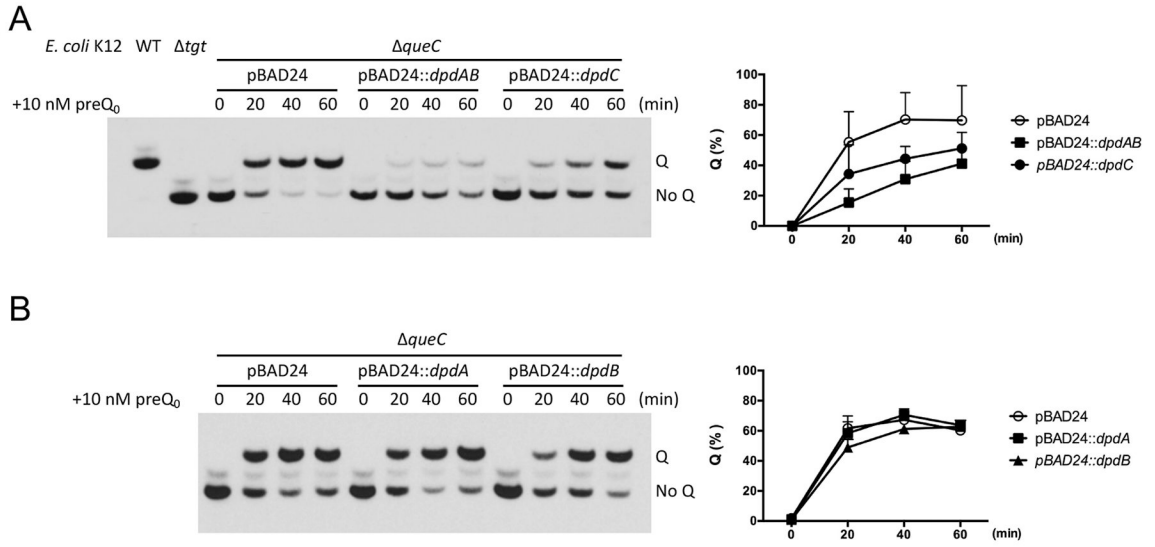
**Figure 4. The *queC* deletion strain show deficiency in Q modification and dADG-dependent modification and increased transformation efficiency with unmodified DNA.**

(A) Q<sup>-</sup> phenotype in the *queC* deletion strain can be complemented by preQ<sub>0</sub> feeding. Q incorporation into tRNA<sup>ASP</sup><sub>GUC</sub> tRNA was detected in total tRNA extracted at 0, 20, 40 and 60 min after supplementing *S. Montevideo* culture medium with 5 nM preQ<sub>0</sub> or DMSO in M9 broth containing 0.4% arabinose. tRNA<sup>ASP</sup><sub>GUC</sub> was visualized by hybridization with a specific biotinylated probe. tRNAs modified with Q (upper bands) migrate slower than the unmodified tRNA (lower bands) (Igloi & Kossel, 1985). (B) pUC19 DNA (10 ng) extracted from different *S. Montevideo* mutant donor strains was transformed in the WT host and YYF3022. (C) Modified and unmodified pUC19 DNA (10 ng) was transformed in different *S. Montevideo* mutant host cells. The average of three experiments is shown, with error bars representing SD (\*\*\*)P < 0.001, two-tailed Student's t-test).

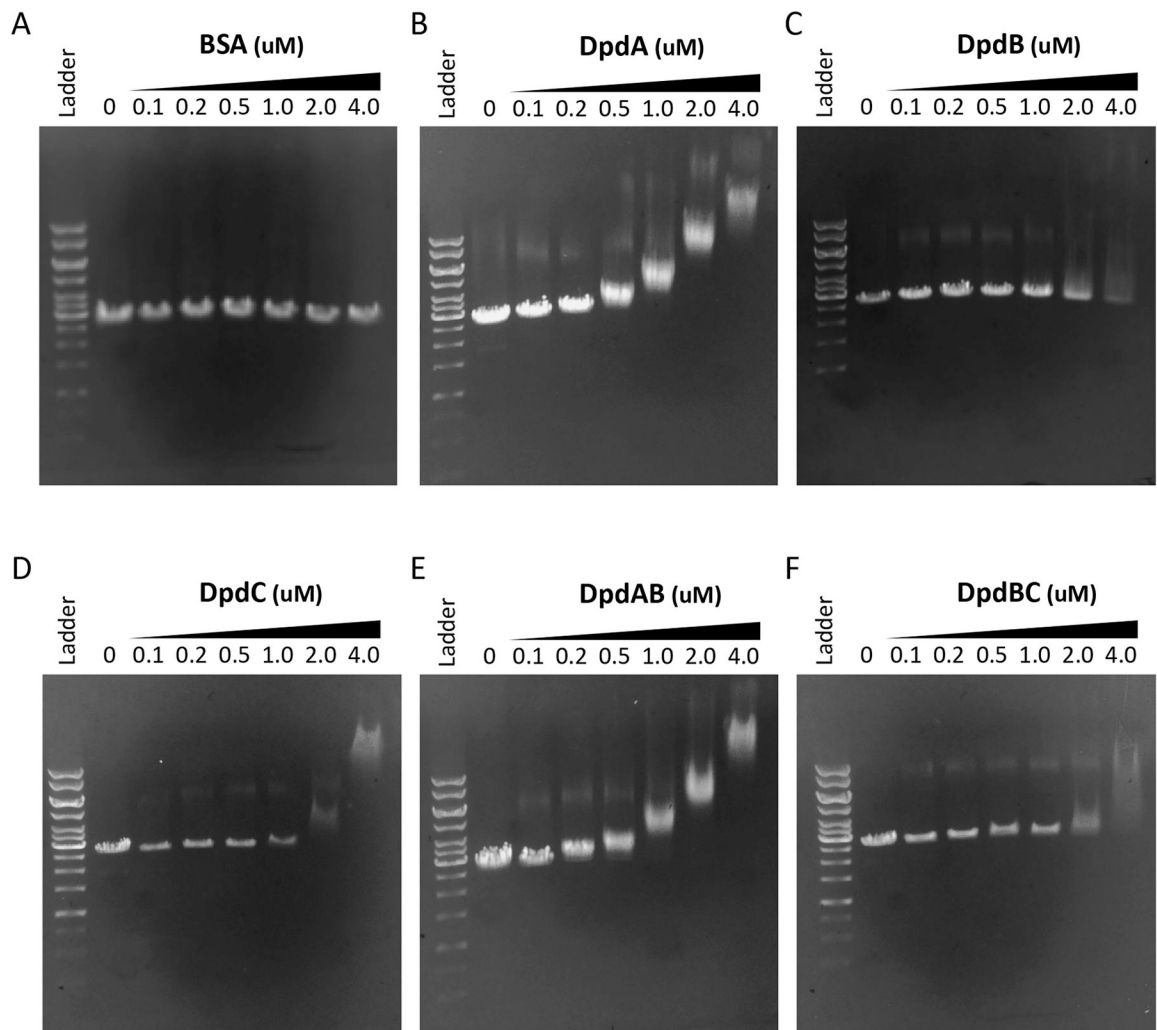


**Figure 5. dADG can be formed from exogenous preQ<sub>0</sub> favoring a role of DpdC after preQ<sub>0</sub> insertion.**

(A) Schematic representation of possible models for the roles of DpdA, DpdB and DpdC in dADG formation. Dashed arrows represent proposed reactions. All molecule abbreviations and protein names are described in the text. (B) pUC19 (10 ng) extracted from the *queC* deletion strain was transformed in the *S. Montevideo* WT host. The donor strain was fed either with DMSO (negative control) or preQ<sub>0</sub>. The WT and *dpdC-dpdD::kan* donors are respectively positive and negative control of the experiment. The average of three experiments is shown, with error bars representing SD (\*\*P < 0.01, two-tailed Student's t-test).



**Figure 6. Overexpression of DpdC in *E. coli* K12 reduces the rate of Q formation in tRNA<sup>ASP</sup>.** (A) (Left) The formation rate of Q was estimated by visualizing tRNA<sup>ASP</sup><sub>GUC</sub> extracted at different time points (0, 20, 40 and 60 min) after supplementing with preQ<sub>0</sub>. tRNA extracted from WT and *tgt* strains were used as control. (Right) The density of the bands was measured by ImageJ (Rueden et al., 2017). The percentage of Q modified tRNA<sup>ASP</sup><sub>GUC</sub> was plotted. The average of three experiments is shown, with error bars representing SD. (B) The effect of expression of DpdA and DpdB on the Q formation rate was estimated with the same approach.



**Figure 7. Dpd protein/DNA binding assays.**

Gel mobility shift analysis of Dpd proteins with DNA. In each panel the lanes correspond to the following: lane 1 (far left), kb ladder; lane 2, linear pUC19 alone; lane 3, protein at 0.1  $\mu\text{M}$ ; lane 4, protein at 0.2  $\mu\text{M}$ ; lane 5, protein at 0.5  $\mu\text{M}$ ; lane 6, protein at 1.0  $\mu\text{M}$ ; lane 7, protein at 2.0  $\mu\text{M}$ ; lane 8, protein at 4.0  $\mu\text{M}$ . The proteins in each panel are (A) BSA negative control, (B) DpdA, (C) DpdB, (D) DpdC, (E) DpdA/B, and (F) DpdB/C. pUC19 is present at 0.5  $\mu\text{g}$  in each lane.



**Table 1.**

Levels of dADG and dPreQ<sub>0</sub> in DNA from strains lacking or expressing various components of the *dpd* cluster.

Strains	Genotype	Condition	dADG per 10 <sup>6</sup> nt	dPreQ <sub>0</sub> per 10 <sup>6</sup> nt
YYF3048	<i>S. Montevideo</i> WT		330	<5 <sup>†</sup>
YYF3022	<i>dpdC-D::kan</i>		<5 <sup>1</sup>	<5
YYF3286	<i>dpdE-dpdD::kan</i>		308	<5
YYF3014	<i>dpdD::kan</i>		69	10
YYF3434	<i>dpdA::kan dpdD</i>		<5	<5
YYF3402	<i>dpdB::kan dpdD</i>		<5	<5
YYF3401	<i>dpdC::kan dpdD</i>		<5	28
YYF3422	<i>dpdA::kan dpdD</i> pBY279.1:: <i>dpdA</i> pUC19	500 ng mL <sup>-1</sup> aTET	86	<5
YYF3416	<i>dpdB::kan dpdD</i> pBY279.1:: <i>dpdB</i> pUC19	500 ng mL <sup>-1</sup> aTET	58	<5
YYF3418	<i>dpdC::kan dpdD</i> pBY279.1:: <i>dpdC</i> pUC19	500 ng mL <sup>-1</sup> aTET	475	<5
YYF3324	<i>S. M dpdC-dpdD::kan</i> pBAD33:: <i>dpdCAB</i>	0.4% arabinose	529	<5
YYF3425	<i>S. M dpdC-dpdD::kan</i> pBY279.1:: <i>dpdAB</i>	500 ng mL <sup>-1</sup> aTET	<5	334
YYF3346	<i>queC::Kan</i>		<5	<5
YYF3460	<i>queC::kan</i> pBAD24	0.4% arabinose	<5	<5
YYF3461	<i>queC::kan</i> pBAD24:: <i>queC</i>	0.4% arabinose	394	130
YYF3346	<i>S. M queC::kan</i>	5 nM preQ <sub>0</sub>	24	<6
YYF3297	<i>E. coli queC</i> pBAD33	5 nM preQ <sub>0</sub>	<6	<6
YYF3297	<i>E. coli queC</i> pBAD33:: <i>dpdAB</i>	5 nM preQ <sub>0</sub>	<6	75
YYF3303	<i>E. coli queC</i> pBAD33:: <i>dpdCAB</i>	5 nM preQ <sub>0</sub>	76	<6

<sup>†</sup>Limit of detection for both dADG and dPreQ<sub>0</sub>.