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Biochemical and spectroscopic studies of epoxyqueuosine reductase: A novel iron-sulfur cluster and cobalamin containing protein involved in the biosynthesis of queuosine

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Abstract

Queuosine is a hypermodified nucleoside present in the wobble position of tRNAs with a 5'-GUN-3' sequence in their anticodon (His, Asp, Asn, and Tyr). The 7-deazapurine core of the base is synthesized de novo in prokaryotes from guanosine-5'-triphosphate in a series of eight sequential enzymatic transformations, the final three occurring on tRNA. Epoxyqueuosine reductase (QueG), catalyzes the final step in the pathway, which entails the 2-electron reduction of epoxyqueuosine to form queuosine. Biochemical analyses reveal that this enzyme requires cobalamin and two [4Fe-4S] clusters for catalysis. Spectroscopic studies show that the cobalamin appears to bind in a base-off conformation, whereby the dimethylbenzimidazole moiety of the cofactor is removed from the coordination sphere of the cobalt but not replaced by an imidazole sidechain, which is a hallmark of many cobalamin-dependent enzymes. The bioinformaticallyidentified residues are shown to have a role in modulating the primary coordination sphere of cobalamin. These studies provide the first demonstration of the cofactor requirements for QueG.

Keywords

queuosine; RNA modification; EPR; cobalamin

RNA is one of the most chemically diverse biological building-blocks with over 110 modifications reported to date¹. Modifications can be as complex as the tricyclic ring structure observed in wybutosine^{2,3}, or as simple as acetylation or thiolation of various positions on the purine or pyrimidine core. These modifications contribute to a myriad of functions including, but not limited to, translational efficiency and accuracy, structural stabilization of the tRNA, enhanced recognition between the tRNA and its cognate

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The sequence of the codon optimized gene and the sequences of the primers used for constructing the site-directed variants are reported in the Supporting Information. This material is available free of charge via the Internet at http://pubs.acs.org.

aminoacyl-tRNA synthetase, and decoding of degenerate codons^{4,5}. However, even simple modifications, such as methylation, are garnering attention as regulators of RNA processes. Only recently has the importance of such modifications come to light through identification of enzymes that reverse this methylation process, in an analogous fashion to DNA, adding a layer of regulation described as RNA epigenetics^{6,7}.

Queuosine is one of the most highly decorated RNA modifications identified to date and has been observed in RNA across all domains of life⁸. The unique 7-deazapurine core of queuosine, which is also found in a variety of secondary metabolites⁹, is further adorned with a cyclopentenediol moiety derived from S-adenosyl-L-methionine (SAM)¹⁰. Queuosine is present in the wobble position of tRNAs containing a 5'-GUN-3' anticodon sequence that encodes for Asp, Asn, His, and Tyr amino acids¹¹. In bacteria, it is synthesized from the precursor GTP through an eight step biosynthetic pathway (Fig. 1)^{10,12–22}, which entail biosynthesis of 7-aminomethyl-7-deazaguanine, exchange into the wobble position, followed by two additional steps that occur on tRNA. Higher order organisms obtain queuine, the free base of queuosine, from dietary sources and exchange this base into mature tRNA in place of guanine $en\ masse^{23}$. While no discrete function is known for this modification, absence has been correlated with numerous biological phenomena such as cancer pathology^{24–28}, pathogenicity^{29,30}, symbiosis³¹, and neurological disorders³².

Our understanding of the complete *de novo* biosynthetic pathway was completed recently with the discovery of epoxyqueuosine reductase (QueG), which catalyzes the final epoxide reduction, converting epoxyqueuosine to queuosine²² (see Fig. 1). QueG is homologous to reductive dehalogenase (RDH) enzymes that are essential to bacteria that utilize halogenated compounds, such as tetrachloroethylene, as their terminal electron acceptors^{33,34}. RDHs are known to contain multiple iron-sulfur clusters and corrinoids as cofactors, and require strong reductants for activity^{35–39}. QueG retains the eight Cys residues that coordinate the two 4Fe-4S clusters in RDHs (Fig. 2). Recent X-ray crystal structures of two RDHs provide views of the active sites of these enzymes^{40,41}. The epoxide reduction reaction catalyzed by QueG is analogous to RDHs in that it is a two-electron reduction requiring a strong reductant. Prior to identification of oQ reductase, bacterial feeding experiments had demonstrated a cobalamin requirement for the conversion of oQ to Q⁴². *In vitro*, oQ reduction was shown to be stimulated by addition of exogenous cobalamin²². To date, however, there have been no systematic studies on the role(s) of the cofactors and conserved residues in catalysis in RDHs or QueG.

Herein we report a biochemical and spectroscopic analysis of the cofactor requirements of QueG. We have established a method to obtain cofactor replete, active recombinant protein in an anaerobic environment allowing the cofactor stoichiometry of the protein to be established unambiguously. The results demonstrate the presence of two iron-sulfur clusters and a cobalamin that are absolutely required for activity. In addition, an *in vivo* alanine scanning experiment has identified residues that are critical for catalysis. Analysis by electron paramagnetic resonance (EPR) spectroscopy of a subset of conserved residues that are essential for activity has revealed an interesting role for these residues in modulating the coordination environment of the cobalamin cofactor. Taken together, these analyses highlight the unique cofactor requirements necessary to facilitate the novel epoxide

reduction catalyzed by QueG on RNA and show that despite the substantially different reaction catalyzed by oQ reductase, it utilizes similar cofactors, suggesting that reductive dehalogenation and epoxide reduction likely follow similar mechanistic pathways.

METHODS

Materials

All materials were purchased commercially (unless otherwise noted) and were of the highest purity. All assays and purification steps were carried out in a Coy anaerobic chamber in an atmosphere of 95% N_2 , 5% H_2 . All buffers and materials were deoxygenated in the chamber several days prior to use and were made RNase-free when possible.

Cloning of B. subtilis queG

The codon-optimized gene encoding *B. subtilis* QueG was obtained from Genscript (Piscataway, NJ). The sequence of the synthetic gene is available in the Supporting Information. The gene was excised from a supplied pUC57 vector by digestion with *NdeI* and *HindIII* and ligated into a similarly cut pET28a vector to obtain pZM419. The gene was then amplified from this vector using the primers: 5′-

AAAGGATCCATGAATGTTTACCAACTGAAAGAAGAAC-3' (forward) and 5'-AAAGGATCCGGACAGACCTTGTTTCGTCATACC-3' (reverse) at an annealing temperature of 43 °C and included flanking *Bam*HI restriction sites. The amplified fragment was digested using *Bam*HI and cloned into a pASK-IBA43plus vector (IBA) that was similarly digested to yield pZM471 for expression of N-terminal His₆-tagged and C-terminal Strep-tag II QueG. Mutants of pZM419 and pZM471 were created using Stratagene Mutagenesis Kit according to the manufacturer's recommended protocol and using the primers listed in Supplementary Table 1.

Expression of B. subtilis QueG

E. coli BL21(DE3) cells containing pZM471 were grown in 12 L of LB containing 0.1 mg/mL ampicillin at 37 °C to an OD_{600nm} of approximately 0.3 at which time ferric chloride was added to a final concentration of 50 μM and the flasks were cooled to 18 °C. The cells were grown further to an OD_{600nm} of approximately 0.6 and protein expression was induced with 20 μL of 10 mg/mL anhydrotetracycline hydrochloride in dimethylformamide (Acros Organics) per 1 L of LB. Cells were grown overnight and harvested the next day by centrifugation (5,000 × g), frozen in liquid N₂, and stored at -80 °C.

Purification of QueG

All steps were carried out in an anaerobic chamber as described in the materials. Cells (~20 g) were suspended in buffer containing 100 mM Tris•HCl (pH 8.0), 150 mM NaCl, 10 mM dithiothreitol (DTT), and 1 mM PMSF, and sonified using a Branson digital sonifier (45% amplitude). The lysate was centrifuged for 30 min at $18,000 \times g$ (4 °C) to pellet insoluble material. The cleared lysate was loaded onto a 5 mL StrepTrap HP column (GE Healthcare) that had been equilibrated with buffer containing 100 mM Tris•HCl (pH 8.0) and 150 mM NaCl (buffer A). The column was washed with 25 mL of buffer A and QueG was eluted with 25 mL of buffer A containing 2.5 mM d-desthiobiotin (Sigma-Aldrich). Fractions

containing QueG were identified using SDS-PAGE and pooled. The concentration of the pooled protein was determined using the Bradford method with bovine serum albumin (ThermoScientific) as the standard.

Reconstitution of QueG

The reconstitution was carried out on ice and the solution was stirred constantly. First, solid DTT was added to a final concentration of 10 mM and the solution was allowed to mix for 15 min. Ten molar equivalents of ferric chloride dissolved in anaerobic water was added slowly over the course of 5 min and the solution was allowed to mix for an additional 10 min. Ten equivalents of sodium sulfide dissolved in anaerobic water was added slowly over 5 min and the solution was allowed to mix for another 3 h. The protein solution was buffer exchanged using a Bio-Rad DG-10 column into 20 mM Tris•HCl (pH 8.0), 100 mM KCl, and 2 mM DTT (buffer B). The protein was concentrated by centrifugation at $6,000 \times g$ (4 °C) using an Amicon Ultra-4 10 kDa cutoff concentrator to a final volume of less than 1 mL. To the resulting concentrated protein, 100 µL of 20 mM hydroxocobalamin acetate salt (Sigma-Aldrich) was added and the solution was allowed to incubate at room temperature for 5 min. The protein was then loaded onto a Sephacryl S300 HR gel filtration column (16 × 60 cm, GE Healthcare) equilibrated in buffer B, and eluted at a constant flow rate of 1 mL/ min. Fractions containing QueG were identified using SDS-PAGE, pooled, and concentrated by centrifugation at 4,000 × g (4 °C) using an Amicon Ultra-15 10 kDa cutoff concentrator. Aliquots were flash frozen in liquid N₂ and stored at -80 °C. Protein concentration was determined using the Bradford method with bovine serum albumin (ThermoScientific) as the standard.

Amino acid, iron, labile sulfide, and cobalamin content analyses

All subsequent analyses were completed on at least three independent protein preparations and values are reported as averages.

Amino acid analysis was performed at the Molecular Structure Facility University of California-Davis (Davis, CA). The protein was exchanged by gel filtration using a NICK Sephadex G-50 column (GE Healthcare) into 10 mM NaOH. A Bradford assay was performed on the eluate and 0.3 mL of the remaining protein solution was utilized for the analysis. The amino acid correction factor was used to calculate the resulting stoichiometries during the cofactor content analyses.

Iron analysis was performed at the University of Arizona Department of Hydrology and Water Resources using a Perkin-Elmer Optima 5300 DV ICP-OES. QueG was diluted into 1% (v/v) nitric acid to a concentration of 1 μ M for analysis. The concentration of labile sulfide was determined using the Beinert method⁴³.

The cobalamin content of QueG was determined by thermal denaturation of the protein and derivatization of cobalamin to dicyanocob(III)alamin by addition of potassium cyanide as follows. To a solution containing QueG at a known concentration, potassium cyanide was added to a final concentration of 10 mM in a total reaction volume of 0.1 mL. The reaction was allowed to incubate at room temperature in the dark for 30 min and was placed at 99 °C

for 15 min. The mixture was placed directly on ice for 5 min and then centrifuged at 21,000 \times g for 5 min to pellet the precipitated protein. The concentration of dicyanocob(III)alamin in the supernant was determined by measuring the absorbance at 580 nm using an extinction coefficient of 10.13 mM⁻¹ cm⁻¹ ^{44,45}.

Size-exclusion chromatography to determine oligomerization state

Purified QueG (4 mg/mL) was loaded onto a Sephacryl S200 HR column (16×60 cm, GE Healthcare) equilibrated in 20 mM Tris•HCl (pH 8.0), 100 mM KCl, and 2 mM DTT. The column was run at a constant flow rate of 1 mL/min and the fractions were analyzed by SDS-PAGE to observe the presence of the standards and QueG. The standards (Bio-Rad) used to determine the molecular weight of QueG were: thyroglobulin (670 kDa), γ -globulin (150 kDa), ovalbumin (44 kDa), myoglobin (17 kDa) and vitamin B_{12} (1.35 kDa).

In vitro activity assays to determine requirements of iron-sulfur clusters and cobalamin

The conversion of oQ to Q by QueG was assayed using a synthetic 17-mer oQ-containing stem loop substrate corresponding to the anticodon loop of Tyr-tRNA and prepared as previously described 22 . The assays contained 20 mM PIPES NaOH (pH 7.4), 10 mM sodium dithionite, 0.75 mM methyl viologen, 0.75 mM hydroxocobalamin (if present in assay), and 5 μ L of ~0.5 mM oQ stem loop. The reactions were initiated by addition of 5 μ M QueG enzyme in a total volume of 0.1 mL. The reaction was allowed to proceed in the dark at room temperature for 5 h and quenched using QIAzol reagent (Qiagen), which contains phenol. RNA from the assay was purified, digested to nucleosides, and analyzed by LC/MS as previously described 22 .

In vivo alanine scanning complementation experiment

The *yjeS E. coli* strain from the Keio Collection⁴⁶ was lysogenized using the λ DE3 Lysogenization Kit (Novagen) following the manufacturer's recommended protocol. The kanamycin resistance cassette, which was inserted in place of the yieS gene in the Keio Collection, was removed by transformation with the pCP20 plasmid⁴⁷ following a published method⁴⁸. Briefly, 30 µL of stationary phase overnight culture of the lysogenized knockout strain was used to inoculate 3 mL of LB containing 34 µg/mL kanamycin. The culture was then left to grow at 37 °C, 200 rpm, to an OD_{600nm} of ~0.6–0.7. The cells were then made electrocompetent by placing 1 mL of the culture in a pre-chilled, sterile 1.5 mL tube and spun at $13,000 \times g$ (4 °C) for 1 min. to pellet the cells. The pellet was resuspended in 1 mL pre-chilled sterile water and was again spun at $13,000 \times g$ (4 °C) for 1 min to pellet the cells. This process was repeated. The pellet from the final spin was resuspended in 50 µL of the residual supernatant and transformed with the pCP20 plasmid, resuspended in 1 mL room temperature SOC media, and left to shake at 30 °C, 200 rpm for 1 h. The culture was then added to 100 mL of LB containing 100 µg/mL ampicillin and 34 µg/mL chloramphenicol, and allowed to shake at 30 °C, 200 rpm overnight. An aliquot (30 µL) of the overnight growth was used to inoculate a 3 mL LB culture and it was allowed to shake at 30 °C, 200 rpm until the OD_{600nm} reached ~0.1, at which time the temperature was shifted to 42 °C and the culture was allowed to grow to an OD_{600nm} of $\sim 0.8-1.0$. The culture was then streaked onto an LB/agar plate and left overnight at 37 °C. Colonies were selected that could no

longer grow on either kanamycin or ampicillin and verified to be the correct *yjeS E. coli* strain by colony PCR. Strains verified to be Kan^{-r} and *yjeS* were made competent and transformed with a pET28a plasmid containing wild-type QueG or QueG variants. Site directed variants were prepared using the primers listed in Table S1. A single colony was used to inoculate 50 mL of LB containing 34 μ g/mL kanamycin and was grown at 37 °C, 200 rpm to an OD_{600nm} of ~0.5–0.7. Then, the culture was induced by addition of 5 μ L of 1 M IPTG and left to continue growing for 16 h total. The cells were harvested the next day and total RNA was extracted and analyzed on the LC/MS as previously described²². Each variant was analyzed in this manner at least in triplicate.

Electron paramagnetic resonance spectroscopy of wild-type QueG and variants

EPR samples were prepared in anaerobic conditions by mixing concentrated, purified protein with a final concentration of 15% glycerol (v/v) to a total volume of \sim 0.15 mL and frozen in liquid nitrogen in an EPR tube. Continuous wave (CW) X-band EPR experiments were preformed on a Bruker Biospin EleXsys E500 spectrometer equipped with a cylindrical TE_{011} -mode resonator (SHQE-W), an ESR-900 liquid helium cryostat, and an ITC-5 temperature controller (Oxford Instruments). Spectra were recorded at 40 K at a microwave frequency of 9.323 GHz (WT), 9.333 GHz (D104A), 9.321 GHz (Y105A), 9.373 GHz (H106A), and 9.376 GHz (D134A). The power and modulation amplitude were 0.2 mW and 2 G, respectively. Simulations were carried out in MatLab using the EasySpin suite of programs⁴⁹.

RESULTS and DISCUSSION

Purification of QueG

Previous studies on purified QueG confirmed that the enzyme was sufficient for conversion of oQ to Q and demonstrated that addition of exogenous cobalamin stimulated activity²². However, the protein used in those studies was recalcitrant to expression and purification to a more homogenous state, and the observation that cobalamin stimulated activity was presumably due to the lack of cofactor replete protein. Therefore, we obtained a codon-optimized gene encoding the *B. subtilis* homolog and developed a method of expression in *E. coli* and purification to obtain >95% pure protein (Fig. 3A). QueG is a monomer as judged by size exclusion chromatography (Fig. 3B). Based on the sequence homology to reductive dehalogenases, we hypothesized that the enzyme would contain iron-sulfur clusters and cobalamin as cofactors. Indeed, the UV-visible spectrum of QueG (Fig. 3C) includes features common to both [4Fe-4S] clusters at ~420 nm⁵⁰ and cob(II)alamin at ~475 nm⁵¹.

Cofactor stoichiometry

After establishing the protein purification and cofactor incorporation scheme, we turned our attention to determining whether the iron-sulfur clusters and cobalamin are required for activity. The basic strategy for obtaining replete protein entailed reconstitution of the purified protein in two stages with iron and sulfide first, followed by cobalamin. Excess cofactors were removed by gel filtration prior to activity and stoichiometry measurements.

To determine the cofactor requirements, an aliquot of the reconsitution mixture was assayed for activity following the initial affinity purification steps, after reconstitution with iron and sulfide, and after addition of cobalamin. As shown in Fig. 4, the as isolated protein has no activity whereas protein that has undergone iron, sulfur and cobalamin additions is active. However, reconstitution with iron and sulfide is not sufficient for activity; therefore, oQ reductase activity requires FeS clusters and cobalamin. To determine the stoichiometry of the clusters and cobalamin present, amino acid analysis was carried out on reconstituted QueG after the final size-exclusion step to obtain an accurate protein concentration. Based on the amino acid analysis, the protein concentration obtained using the Bradford method overestimates [QueG] by ~1.7-fold. QueG was subjected to ICP-OES to determine the amount of iron and the Beinert method⁴³ was used to determine the amount of labile sulfide. Finally, a cyanolysis was carried out to quantify cobalamin content^{44,45}. The iron, sulfide and cobalamin stoichiometry derived from the above analyses are summarized in Table 1. Purified and reconstituted QueG contains 6.5 moles of iron, 8 moles of labile sulfide, and one mole of cobalamin per monomer. Recently, two structures of reductive dehalogenase proteins have been reported that are relevant in this discussion^{40,41}. Both structures show that RDHs have two [4Fe-4S] clusters and a cobalamin. While there is limited sequence similarity between QueG and RDHs, the Cys residues that coordinate the two iron-sulfur clusters are conserved in QueG (shown in red in Fig. 2). Intriguingly, oQ reductase has the same complement of cofactors as RDHs, despite the differences in the overall reactions catalyzed by these enzymes.

In vivo screening for essential residues

The initial difficulties in expression and purification of QueG and the complicated reconstitution methods that are required to obtain cofactor-replete active protein made it necessary to develop an alternative method to rapidly assess the importance of residues in catalysis. In these experiments we utilized the queG strain of E. coli from the Keio collection 46 and examined the oO/O content of the cells upon expression of wild-type and site-directed variants of QueG on a plasmid. Potential catalytic residues were identified from a multiple sequence alignment that was constructed with ~1100 sequences obtained through a protein BLAST search^{40,41,52,53} of the non-redundant database with the E. coli QueG homolog (YjeS; b4166) as the initial search sequence. The candidates obtained from the BLAST search had 30-80% sequence identity. Analysis of the extensive alignment identified multiple absolutely conserved motifs (Fig. 2). Two of the conserved motifs, CX₂CX₂CX₃C and CX₂CX₃C with another distally conserved Cys, are likely involved in forming the iron-sulfur clusters as both of these motifs are also the sequence signatures associated with enzymes involved in reductive dehalogenation, where they had been proposed to coordinate two [4Fe-4S] clusters that are necessary for electron transfer and reduction of the cobalamin cofactor^{35,54–56}. Another absolutely conserved sequence was a DYH motif. This DYH motif was of great interest as it was reminiscent of the DxHxxG motif commonly found in cobalamin-dependent enzymes, specifically, enzymes that bind cobalamin in its base-off/His-on conformation, such as methionine synthase⁵⁷ and glutamate mutase^{58,59}. In this conformation, a histidine side chain displaces the dimethylbenzimidazole moiety of cobalamin, which in solution coordinates the cobalt atom through the N3 atom in the lower axial position. Based on the conserved residues observed

in the multiple sequence alignment, mutations were made in a pET28a plasmid harboring the B. subtilis QueG enzyme (pZM419). In all, 18 single mutation variants were generated where the residue of interest was mutated to an alanine. The yjeS E. coli Keio Collection strain was then lysogenized with $\lambda DE3$ and made electrocompetent to allow for expression of the wild-type or variant queG gene using T7 polymerase after transformation with the corresponding pET28a plasmid. Cells were induced in the log phase of growth and allowed to grow overnight. Total RNA was isolated, digested to nucleosides, and analyzed for the presence of oQ vs Q via LC/MS. The effects of the mutations on the production of both oQ and Q in vivo are shown in Fig. 5. Mutation of almost all of the cysteine residues thought to be involved in cluster formation completely negated the ability to convert oQ to Q. Curiously, the point mutation C240A did not seem to have any effect on activity in comparison to the wild-type protein. However, there have been examples of iron-sulfur cluster containing enzymes where a single mutation in a cysteine residue comprising an iron-sulfur cluster is not sufficient to abolish cluster formation and activity⁶⁰. Mutations in residues comprising the DYH motif showed differing results. The introduction of Y105A and H106A variants of QueG into the knockout strain reduced the extent of cellular Q relative to oQ. By contrast, the D104A mutation did not alter the oQ/Q ratio relative to wildtype.

Interestingly, the D134A and R141A variants significantly reduced the Q-content of the strain relative to wild-type. The importance of the Cys residues to catalysis in QueG is easy to justify due to the observation that analogous Cys residues in RDH proteins coordinate the two [4Fe-4S] clusters in the protein. The significance of D104, Y105, H106 and D134, however, was not immediately obvious.

EPR spectroscopy of QueG and variants

To further probe the role of residues that appeared to be essential by the *in vivo* experiments, we carried out electron paramagnetic resonance studies to investigate whether these residues are in the vicinity of the cobalamin cofactor. Cobalamin can exist in three oxidation states with distinct UV/visible spectral signatures. However, in the +2 oxidation state, the cofactor is paramagnetic and may be studied by EPR.

Reconstituted wild-type, D104A, Y105A, H106A, and D134A proteins were analyzed by EPR spectroscopy. The EPR spectrum of the wild-type protein (Fig. 6A) is best simulated by a rhombic g- and A-tensor (Table 2). Interaction of the unpaired spin with cobalt (I=7/2) introduces extensive fine structure, which is best fit with a rhombic A-tensor (A_x =186, A_y =133, A_z =340 MHz). We observe no additional strongly coupled nuclei. If a nitrogen (I=I) from the dimethylbenzimidazole or His imidazole sidechain had been present, we would have observed triplet splittings of the g_z features, as is observed in other cobalamindependent enzymes^{57,61,62}. Therefore, the data clearly show that upon binding of cobalamin to QueG, the dimethylbenzimidazole ligand that coordinates the cobalt in solution is dissociated, but *not* replaced with a His sidechain. This observation is similar to what has been observed in the structures of the RDH proteins^{40,41}.

To determine if the DYH motif is in the vicinity of the cofactor, the residues in this motif were mutated to Ala, the purified proteins were reconstituted with cobalamin, iron and

sulfide and EPR spectra of each variant were recorded (Fig. 6C, E, G). The cobalamin content of the variants was quantified by cyanolysis to be 0.85–1.2 per monomer. The spectra of the wild-type and H106A variant are nearly identical, and can be simulated with very similar sets of parameters (Table 2). The spectrum of the D104A variant, however, shows subtle differences with the wild-type that are best captured in simulations where nearly axial g-and A-tensors are included. By contrast, there is a very dramatic change in the spectrum of the Y105A variant relative to wild-type, whereby there is significantly less gtensor anisotropy and A-tensors. The differences between the wild-type, D104A, H106A, and Y105A variants are reminiscent of the observations with the H759G variant of cobalamin-dependent methionine synthase (MetH). In MetH, binding of the cobalamin leads to interchange of the dimethylbenzimidazole for the imidazole sidechain of the His759 residue⁵⁷. Mutation of the His residue reveals two sets of EPR cobalamin spectra, one of which corresponds to a cob(II)alamin interacting with a water molecule (5-coordinate), and another which is essentially 4-coordinate. The EPR spectra of wild-type QueG and all the variants reported here are consistent with 5-coordinate geometry around the cobalamin, but the differences in the g- and A-tensors suggest that in all but the Y105A variant, the interaction between the putative water ligand and the cobalamin is weakened⁶³.

In addition to the residues in the conserved DYH motif, we also examined the effect of mutating the absolutely conserved D134 to A on the EPR spectrum of the variant. Interestingly, the EPR spectrum of this variant (Fig. 6I) exhibits features of both the wild-type and Y105A variants. We did not attempt to simulate the complex pattern; however, the spectrum of the protein is reasonably reproduced (Fig. 6J) by mathematically adding experimental spectra of the wild-type and Y105A proteins, suggesting that the cobalamin in this variant exists as a 50:50 mixture of the forms that are present in the wild-type and Y105A proteins.

The molecular mechanisms by which these conserved residues modulate the environment of the cobalamin remain to be established by structural methods, and we cannot exclude the possibility that they are distant from the active site. However, taken in the context of the *in vivo* analysis of oQ to Q ratios in a *queG* knockout strain, it seems likely that these residues are in the active site of the protein.

CONCLUSIONS

The identity of QueG, which catalyzes the final step in the biosynthesis of queuosine, was surprising in that it revealed the protein to be homologous to reductive dehalogenases. In this paper we demonstrate that despite the drastic differences in the reactions catalyzed by these enzymes, QueG and RDHs utilize the same set of cofactors to accomplish their respective chemical transformations. Reconstitution experiments with QueG demonstrate that iron-sulfur clusters and cobalamin are required for function. This data establishes a direct link between cobalamin and RNA modification for the first time since the feeding experiments noted a role for cobalamin in oQ reduction *in vivo*⁴². Based on the similarity of the EPR spectroscopic parameters to MetH^{61–63}, the cobalamin cofactor in QueG is bound in a base-off conformation but is 5-coordinate, presumably by interactions with a ligand, such as water, that does not have a nuclear spin. Intriguingly, the conserved DYH motif does

not provide an imidazole sidechain to coordinate the cofactor. However, the significant perturbations of the electronic environment of the cobalamin observed in these variants suggest that these residues are in the active site of the protein. Therefore, our findings place QueG in the class of enzymes that displace the dimethylbenzimidazole of the cobalamin cofactor upon binding to the protein. Understanding the mechanism of QueG, the role of the conserved residues in activity, and the correlations between the reactions catalyzed by QueG and RDHs must await high-resolution structures of QueG.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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ABBREVIATIONS

EPR electron paramagnetic resonance

oQ epoxyqueuosine

Q queuosine

QueG epoxyqueuosine reductase

RDH reductive dehalogenase

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Figure 1. Biosynthesis of queuosine.

```
-MSEPLDLNQLAQKIKQNGLELGFQQVGITDTDLSESEPK-LQAWLDKQYHGEMDWMARHGMLRARPHELLPGTLRVISVRMN"LPANAAFASTLKNPKLGYVSRYALGRDYHKLLRNRLKKLGEMIQQHCVSLNFRPFVDSAPIL®RPL
                  ----MDYQQLANQIKQWAIELGFEKVGICDVDLSEHEPA-LQAWLDAGYHGEMDWMARHGMMRARPAELLPGTLRVISARIN LPPQAQFASNLRDPNQAYISRYALGRDYHKLVRNQLKKLGEKIEQEVGKLGYRPFVDSAPIL®RPL
Vibrio cholerae
Pseudomonas putida MSASTPDLAQLAQSIKIWGQELGFAHVGIAGVDLGEHEHH-LQRWLDAGYQEMEYLGAHGSKRSHPDQLIPGTVRVVSLRMD LPGDTQMAQRLAQPEKAYVSRYALGRD
                                                                                                                                HKLVRKRVQFLADRIQEAIGPFGYRAFVDSAPVLEKAL
Mesorhizobium loti mrtstsdaaklralidaeahragfdaiavttpdaiplaparlaefvadgfhgsmdwiaetiarrsepstlwpdvrsivvlamn gpdhdprvloarh-drgaisvyaonrd
                                                                                                                                HDVMKGRLKEIAGKIVARAG-GDVKVFVDTAPVMEKPL
                                                                                                                                                                        148
MATICAULIS MATIS ----DADATLS-----LALELGFSTARICRADEAWAAGDRLAETVADGHHGSMAWMEETLERRQHPTAMWPEAKSAVVVGLN: GPEQDPLPLLERR-SEGVVSVYAQNGDYHDLLKKRLKHLARAFAGKTG-KQVKVFVD-TAPLMERPL
                                                                                                                                                                        139
Bacillus subtilis
                  ----MNVYQLKEELIEYAKSIGVDKIGFTTADTFDSLKDRLILQESLGYLSGFE--EPDIEKRVTPKLLLPKAKSIVAIALA PSRMKDAPRSTRTERRGIFCRASMGKD
                                                                                                                                HDVLREKLDLLEDFLKSKHEDIRTKSMVDTGELSDRAV
                                                                                                                                                                        143
                                                                                 ::*:***:**
                                                                                               * !* * * !***!!*** * ** *!
                   *: **:*: .*: : : **:.:*. :: . : * . : ** *: *** *:
Escherichia coli
                  AEKAGLGWTG/HSLILNREAGSFFFLGELLVDIPLPVDQPVE-EGCGKCVACMTICPTGAIVEPYTVDARRCISYLTIELEGAIPEELRPLMGNRIYGC:DCQLICPWNRYSQLTTEEDFSPRKPLHAPELIELFAWSEEKFLKVTEGSA
                                                                                                                                                                        297
Vibrio cholerae
                  AQKAGLGWTGHSLILDKENGSWFFLGELLVDIPLPVDEPSE-NQCGKCTACITSCPTNAIVAEGVVDARRCISYLTIEYSGVIPLEFRRAMGNRIYGCDCQLVCPWNRFAPLTQQSDFHRRQSLNNADLVVLFEWDEATFLKNMEGSA
                                                                                                                                                                        293
Pseudomonas putida AEQAGLGWIG«NTLLLNRKAGSYFFLAELFVDLPLPVDEATTSEHCGRCQACLDICPTQAFVGFYVLDARRCISYLTIELRGAIPVELRAKMGNRVFGC©DCQIVCPWNRFAKHSQEQFQPRHGLENTDLAAMFLWDERTFLRKTEGGF
                                                                                                                                                                        299
Mesorhizobium loti aqaaglgwqghtnlvsrehgswlflgtiftaelapdraei-dhcgscracldacptdafpapyrldarrcisyltienkgpiphefrekignriygcddclaacpwnkfaraaseaklaarddlrappladllelddaafrvffsgsp
                                                                                                                                                                        297
Maricaulis maris
                  AEKAGIGWOG HTNLVSREFGSWLFLGVMLTEAELEPDEAER-DTCGSCRACLDICPTNAFPOPYRLDARRCISYLTIEHKGVIPVEYROPMGNRIYGCDCLAVCPWNKFAQTSTEMAFHPRPELKGPLLAELAALDDAAFROVFTASP
                                                                                                                                                                        288
Bacillus subtilis AERAGIGFSAWNCMITTPEYGSYVYLAEMITNIPFEPDVPIE-DMCGSCTKCLDACPTGALVNPGQLNAQRCISFLTQT-KGFLPDEFRTKIGNRLYGCUTCQTVCPLNKGKDFHLHPEMEPDPEIAKPLLKPLLAISNREFKEKFGHVS
                                                                                                                                                                        291
                            ** :. .:
                  IRRIGHLRWL®NIAVALGNAPWDETILTALESRKG-EHPLLDEHIAWAIAQQIERRNACIVEVQLPKKQRLVRVIEKGLPRDA---
Escherichia coli
                                                                                                                  379
                  IRRIGHQQWQRNLIIAMGNAPYSPRIIDTLQRHLG-QSELLDEHIHWALDEQNQK-----TSTARQHARLIRIIEKGLPRDA----
Vibrio cholerae
Pseudomonas putida LRRAGYERFLENLAVGLGNAPSTIPVIEALKARRDDESELVREHVEWALARHGAQ-----
                                                                                                                  354
Mesorhizobium loti IKRIGRDRFIRNVLIAAGNS-GDVVLAVPVRALVGDPSPLVRGAAIWALARLVPDAEYSERAATGLKAENDAAVREEWRLARPDRAHA-----
                                                                                                                  384
                  VKRIGRDRFV@NVMIAIGNS-GDHELVSVVEAALDDGSALVRGMAVWALGQLSP-LRVSEMHEMKVAAEPDLDVVTEWQALSQKIS-----
Maricaulis maris
                                                                                                                  372
Bacillus subtilis GSWRGKKPIO NAILALAHFKDASALPELTELMHKDPRPVIRGTAAWAIGKIGDPAYAERLEKALEKEKDERAKLEIEKGIELLKASGMTKOGLS
```

Figure 2. A representative multiple sequence alignment of QueG homologs and *in vivo* alanine scanning of conserved residues of QueG

Conserved cysteine residues required for iron-sulfur cluster formation are in red, the conserved DYH motif is shown in orange, and other conserved residues mutated in the *in vivo* alanine scanning experiment are highlighted in blue. The accession numbers for the sequences are as follows: *E. coli* (EGI43095), *Vibrio cholerae* (WP_000386321), *Pseudomonas putida* (WP_014754621), *Mesorhizobium loti* (WP_010912587), *Maricaulis maris* (WP_011642061), and *Bacillus subtilis* (NP_388772).

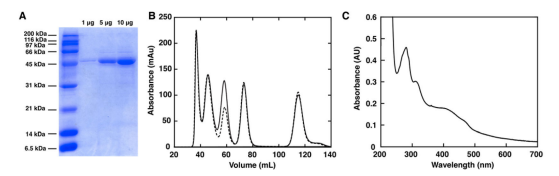


Figure 3. Purification, UV/visible spectrum, and quaternary state determination of QueG A. An SDS-PAGE gel of purified QueG. Based on the gel, QueG is >95% pure with a molecular weight consistent with the predicted weight of 48.7 kDa. B. A size exclusion chromatogram of protein standards (shown in dotted black line) superimposed with a chromatogram of protein standards including purified QueG enzyme (shown in solid black line). Based on the elution profiles, only the peak pertaining to the ~44 kDa standard changes with inclusion of QueG in the solution mixture. QueG is ~48.7 kDa leading to the increase in only the absorbance of the concurrent standard peak. The standard peaks (from left to right) are: thyroglobulin (670 kDa), γ -globulin (150 kDa), ovalbumin (44 kDa), myoglobin (17 kDa) and vitamin B₁₂ (1350 Da). C. A UV/visible spectrum of the purified enzyme contains features consistent with inclusion of both iron-sulfur clusters (~420 nm) and cob(II)alamin (~475 nm) as cofactors.

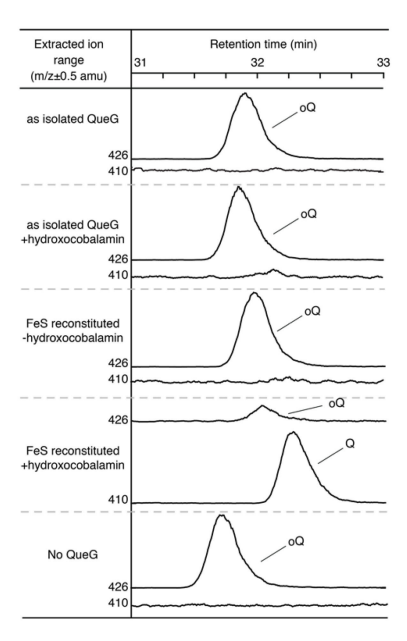


Figure 4. Iron-sulfur clusters and cobalamin are required for QueG activity in vitro Extracted ion chromatograms of digested nucleosides of oQ (m/z = 426) and Q (m/z = 410) from oQ stem loop used as the substrate in an *in vitro* activity assay of QueG. Protein was assayed for activity before and after reconstitution in the presence or absence of exogenous hydroxocobalamin. The appearance of turnover in only enzyme with reconstituted iron-sulfur clusters and added hydroxocobalamin demonstrates the necessity of both for activity.

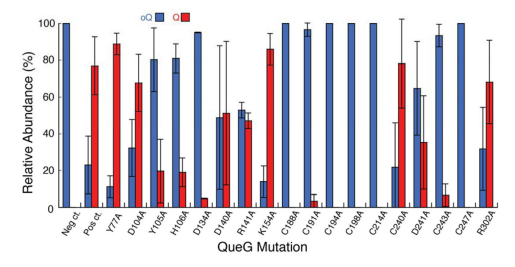


Figure 5. In vivo QueG mutagenesis

The relative abundances of Q and oQ were calculated from the extracted ion chromatograms of oQ (blue) and Q (red) present in total RNA nucleosides from overnight growths of *queG E. coli* overexpressing the indicated variant form of QueG (*B. subtilis* numbering). All values are normalized to the sum of both oQ and Q and given as a percentage of the total.

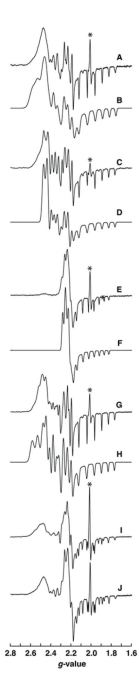


Figure 6. EPR of wild-type QueG and variants

The EPR spectra and simulations of **A.** wild-type and variants **C.** D104A, **E.** Y105A, **G.** H106A, and **I.** D134A of QueG in the cob(II)alamin form. Simulations are shown in **B**, **D**, **F**, **H**, and **J** for wild-type and variants in the same order as above. The * denotes signal that results from degradation of the iron-sulfur clusters during purification and reconstitution.

Table 1

Analysis of the cofactor content of QueG.

Cofactor	Stoichiometry (mol/monomer QueG) ^a
Iron	6.54 ± 1.61
Labile sulfide	8.01 ± 1.08
Cobalamin	1.01 ± 0.11

^aThe values are consistent with the presence of two [4Fe-4S] clusters and one cobalamin cofactor per monomer of QueG. The concentration of QueG protein used to obtain the stoichiometric values was calibrated with amino acid analyses as described in Methods.

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Table 2

EPR simulation parameters for cob(II)alamin in wild-type QueG and site-directed variants.

Drotoin	6	6	6	$\mathbf{A}_{\mathbf{x}}$	$\mathbf{A}_{\mathbf{y}}$	$\mathbf{A}_{\mathbf{z}}$
LIONEIII	x S	Š	22	J	(MHz)	
wild-type a	2.437	2.355	2.061	186a	133	340
H106A	2.435	2.354	2.064	186	130	330
D104A	2.396	2.349	2.065	125	125	333
Y105A	2.282	2.250	2.063	6	11	235

^aSimulation of the wild-type spectrum better matched the experimental when an A-tensor strain of [48.6 17 0] (MHz) was included.

 $^b\mathrm{Simulation}$ of the H106A variant was best fit with an A-tensor strain of [30 0 0] (MHz).

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