Structural Insights into the Function of the Nicotinate Mononucleotide:phenol/p-cresol Phosphoribosyltransferase (ArsAB) Enzyme from Sporomusa ovata

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Supporting Information

ABSTRACT: Cobamides (Cbas) are cobalt (Co) containing tetrapyrrole-derivatives involved in enzyme-catalyzed carbon skeleton rearrangements, methyl-group transfers, and reductive dehalogenation. The biosynthesis of cobamides is complex and is only performed by some bacteria and archaea. Cobamides have an upper (Coβ) ligand (5′-deoxyadenosyl or methyl) and a lower (Coα) ligand base that contribute to the axial Co coordinations. The identity of the lower Coα ligand varies depending on the organism synthesizing the Cbas. The homoacetogenic bacterium Sporomusa ovata synthesizes two unique phenolic cobamides (i.e., Coα-(phenolyl/p-cresolyl)-cobamide), which are used in the catabolism of methanol and 3,4-dimethoxybenzoate by this bacterium. The S. ovata ArsAB enzyme activates a phenolic lower ligand prior to its incorporation into the cobamide. ArsAB consists of two subunits, both of which are homologous (∼35% identity) to the well-characterized Salmonella enterica CobT enzyme, which transfers nitrogenous bases such as 5,6-dimethylbenzimidazole (DMB) and adenine, but cannot utilize phenolics. Here we report the three-dimensional structure of ArsAB, which shows that the enzyme forms a pseudosymmetric heterodimer, provide evidence that only the ArsA subunit has base:phosphoribosyl-transferase activity, and propose a mechanism by which phenolic transfer is facilitated by an activated water molecule.

Cobamides are cobalt-containing modified cyclic tetrapyrole-derivatives, which are members of a broad family that include all forms of heme (iron), chlorophylls (magnesium), and coenzyme F430 (nickel). Several features distinguish cobamides from other members of this family of molecules. As depicted in Figure 1, a cobamide has an upper (Coβ) ligand covalently bound to the cobalt ion of the ring, and a lower (Coα) ligand interacting with the Co ion via a coordination bond.

Vitamin B12 is a cobamide (Cba) that contains a cyano (CN) group as the Coβ ligand (Figure 1). When DMB is the Coα ligand, it is known as cyanocobalamine. Cobamides are in their coenzymic form when the Coβ ligand is 5′-deoxyadenosine (AdoCba). AdoCba participates in radical mediated molecular rearrangement such as in diol dehydratase and methylmalonyl-CoA mutase. Cobamides also serve as transient methyl-group carriers in Co(I)Cba-dependent methyltransferases. In nature, the nucleoside base of cobamides varies, depending on the microorganism synthesizing it. Purines and purine analogues linked to the ribosyl group via an N-glycosic bond can form coordination bonds with the cobalt ion, while bases linked to the ribosyl group via an O-glycosic bond cannot (Figure 1B,C).

Only some bacteria and archaea synthesize cobamides, and unique among cobamides are those with phenol or p-cresol as the lower ligand (Figure 1C). Phenolic cobamides were first described in the homoacetogenic bacterium Sporomusa ovata. Two features of the S. ovata cobamides are of note. First, the phenolic compound is covalently attached to the ribosyl group via an O-glycosic bond, rather than the N-glycosic bond found in all other known cobamides; second, unlike any other cobamides, phenolic cobamides cannot exist in the base-on conformation because neither phenol nor p-cresol contains an atom that can establish a coordination bond with the Co ion of the corrin ring. Enzymes that require cobamides in the base-on conformation for catalysis (e.g., glycerol dehydratase, diol dehydratase, ethanolamine ammonia-lyase) cannot use or are inhibited by phenolic cobamides.

Recently, the S. ovata arsAB genes encoding the two subunits of the enzyme responsible for the conversion of phenol/p-cresol to the corresponding α-O-glycosidic riboside monophosphate were identified. The arsAB-encoded enzyme was isolated to homogeneity, and initial analyses of its activity were performed in vivo and in vitro. The enzyme functions as a heterodimer; i.e., neither subunit is active by itself. The two
subunits are evolutionarily related and share 37% and 57% amino acid identity and similarity, respectively. Furthermore, in addition to phenol and p-cresol, the enzyme converts DMB to α-DMB riboside monophosphate (α-RP). The latter feature of ArsAB is noteworthy because, to date, none of the enzymes known to transfer DMB or any other nitrogenous base can utilize phenol or p-cresol. This feature of ArsAB is even more intriguing in light of the fact that both ArsA and ArsB proteins are homologues of the well-characterized nicotinate mononucleotide (NaMN):DMB phosphoribosyltransferase (CobT) enzyme of Salmonella enterica (ScCobT). ArsA and ArsB share 37% and 36% amino acid sequence identity with ScCobT, but prior to this study it was unknown whether the phosphoribosyltransferase activity of ArsAB was provided by one or both subunits of the inferred heterodimer. The general scheme for the activation of the base is shown below in eq 1.

\[
\text{base + nicotinate mononucleotide (NaMN) + enzyme} \\
\rightarrow \alpha\text{-riboside-5'-P + nicotinate}
\]  

(eq 1)

Crystallographic analyses of ScCobT (a homodimer) in complex with diverse substrates provided structural explanations for how the enzyme phosphoribosylates most of the nitrogenous bases found in cobamides, with the exception of phenol and p-cresol.

Here we report the three-dimensional crystal structure of S. ovata ArsAB in its substrate-free form (2.1 Å), and in complex with DMB (1.5 Å), p-cresol (2.1 Å), phloroglucinol (1.6 Å), and phenol (2.4 Å). Comparisons of the ArsAB complexes with the corresponding ScCobT complexes shed light onto the subtle but profound evolutionary changes required for ArsAB to phosphoribosylate phenolic bases. Remarkably, only the active site in ArsA was occupied by substrate. Analysis of the ArsB active site identified an arginyl side chain that may block access to the site. The apparent inactivity of the ArsB subunit suggests a strictly structural role for ArsB in the phosphoribosyl transferase activity of this enzyme, though this does not preclude some other unidentified activity for that active site.

### EXPERIMENTAL PROCEDURES

**Construction of Plasmids Encoding ArsAB and Variants.** The *arsA* and *arsB* coding sequences were amplified from plasmid pARSAB using primers listed in Table 1. The fragment was cut with *Smal* and *NheI* enzymes and was ligated into the *StuI* and *NheI* sites of plasmid pH6T, a pTEV5 derivative, yielding plasmid pARSAB22. Plasmid pARSAB22 directed the synthesis of H6-ArsA and tag-less ArsB. Variants of ArsAB were constructed using the QuikChange method (Stratagene) using pARSAB22 as template. The plasmids constructed and primers used are listed in Table 1.

**ArsAB Enzyme Isolation.** ArsAB was overproduced using plasmid pARSAB22 in strain JE13607, a derivative of *E. coli* BL21(λ DE3) that lacks the *cobT* gene encoding the NaMN:5,6-dimethylbenzimidazole (DMB) phosphoribosyltransferase (CobT) enzyme of this bacterium. Strain JE13607 was used to ensure that native *E. coli* CobT protein did not contaminate the ArsAB heterodimer during subsequent kinetic analyses or interfere with protein folding. Strain JE13607 harboring pARSAB22 was grown in six 2-L super broth (SB) medium cultures (tryptone 35 g, yeast extract 20 g, NaCl 5 g, NaOH 2N, 2.5 mL per liter of deionized H2O). Expression of

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*Figure 1. Chemical structures of cobamides, vitamin B12, and phenolic cobamides. The upper Coβ ligand (R) varies. When R = cyano (CN) group, the cobamide is in its vitamin form; when R = S-deoxyadenosine, the cobamide is in its coenzymic form. The chemical nature of the lower (Coα) ligand base (B*) is diverse.*
Table 1. List of Plasmids and Primers Used in This Study

<table>
<thead>
<tr>
<th>plasmid</th>
<th>genotype and description</th>
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<td>pHF6T</td>
<td>overexpression vector that fuses the N-terminus of the protein of interest to a His tag, which can be removed by rTEV protease, bla +</td>
</tr>
<tr>
<td>pARSAB22</td>
<td>So arsAB′ in pHF6T, bla·</td>
</tr>
<tr>
<td>pARSAB31</td>
<td>So arsAB′ (ArsA M87Q) in pHF6T, bla·</td>
</tr>
<tr>
<td>pARSAB32</td>
<td>So arsAB′ (ArsA I321S) in pHF6T, bla·</td>
</tr>
<tr>
<td>pARSAB33</td>
<td>So arsAB′ (ArsA M87Q; I321S) in pHF6T, bla·</td>
</tr>
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</table>

arsA was induced by the addition of isopropyl-β-D-1-thiogalactopyranoside (IPTG, 0.5 mM) when cultures reached log-phase (~0.5 OD600). After induction, cultures were shifted to 20 °C for 16 h. Cells were harvested by centrifugation (Beckman/Coulter Avanti J2-XP1 refrigerated centrifuge, equipped with a JLA-8.1000 rotor; 15 min at 4 °C; 6000g), and frozen at −80 °C until used. Frozen pellets were resuspended in buffer A [50 mM 4-(2-hydroxyethyl)-1-piperazine-N′-(2-ethanesulfonic acid) (HEPES), pH 8], containing 300 mM NaCl, and 10 mM imidazole; 1 mg/mL lysozyme and DNase were added to the cell suspension prior to disruption by sonication (60 s, 28% duty, 2 s pulses, setting 9) using a 500 Sonic Dismembrator (Fisher Scientific). Cell debris was removed by centrifugation (Beckman/Coulter Avanti J25-1 refrigerated centrifuge, equipped with a JA-25.50 rotor; 45 min at 4 °C; 43000g). Clarified extract was applied onto an AKTA FPLC Purifier system (Amersham Biosciences) equipped with a 5-mL Ni-charged HisTrap fast flow (FF) column (GE Healthcare) equilibrated at a flow rate of 2 mL/min. After loading, the column was washed with 50 mL of buffer A before application of a 50-mL linear gradient of imidazole from 10 to 300 mM. Fractions containing ArsAB were pooled and His-tagged recombinant tobacco etch virus (His7-rTEV) protease was added at 1:100:His7-rTEV:ArsAB ratio to cleave the His6 tag fused to ArsA.16 Tag-less ArsAB was reapplied onto the Ni-charged column to separate His7-rTEV and other contaminants; tag-less ArsAB was found in the flow through. The ArsAB enzyme was dialyzed into 50 mM HEPES, pH 8.0, 100 mM NaCl, and 15% v/v glycerol then flash frozen in liquid nitrogen and stored at −80 °C until used.

Preparation of Variant Proteins. ArsAB variants encoded in pARSAB31, pARSAB32 and pARSAB33 were overexpressed in strain JE13607 in lysogenic broth (LB) as described above in 1-L scale. Clarified extracts in buffer A were applied onto a 1-mL bed volume of Ni-NTA Superflow resin (Qiagen) equilibrated with buffer A and washed with 10 column volumes of buffer A containing 20 mM imidazole. The variant enzymes were eluted with 10 column volumes of buffer A containing 300 mM imidazole. The His6 tag fused to ArsA was cleaved as described above. Variant proteins were stored at −80 °C in the same buffer as wild type ArsAB until used.

Preparation of ArsAB Proteins for Crystallography. For crystallographic studies, the concentration of NaCl and HEPES in ArsAB solutions was gradually reduced by dialysis (5 steps), from 50 mM HEPES pH 8.0 and 300 mM NaCl to 10 mM HEPES pH 7.5; the resulting solution was concentrated to 10 mg/mL before drop-freezing in liquid nitrogen. The concentration of ArsAB was determined using the combined calculated extinction coefficients of ArsA and ArsB at 280 nm (21150 cm⁻¹ M⁻¹) utilizing a NanoDrop 1000 spectrophotometer (Thermo).

Crystallization of ArsAB in the Substrate-Free State. ArsAB heterodimers were screened for initial crystallization conditions by vapor diffusion at 25 and 4 °C with a 144-condition sparse matrix screen developed in the Rayment laboratory. Single, diffusion-quality crystals were grown by hanging drop vapor diffusion by mixing 2 μL of 10 mg/mL ArsAB heterodimers in 10 mM HEPES pH 7.6 with 2 μL of reservoir solution containing 100 mM 3-(N-morpholino)-propanesulfonic acid (MOPS pH 7.1), 12.5% methyl ether polyethylene glycol 5000 (MEPEG5K), 20 mM NaCl, 12% ethylene glycol, and 10 mM phloroglucinol. Hanging droplets were immediately nucleated from an earlier spontaneous crystallization event with a cat’s whisker. Crystals grew to approximate dimensions of 200 × 200 × 400 μm within 7 days. Phloroglucinol was identified from an additive screen as a compound that dramatically improved the size, stability, and diffraction properties of the crystals. Subsequent structural studies showed that phloroglucinol binds in the active site of ArsA, which in hindsight is not surprising since it shares structural similarity to phenolic substrates. Additionally, phosphogluconol was found at low-occupancy in a different orientation in the corresponding ArsB site (Figure S1, Supporting Information). In order to prepare crystals of the substrate-free protein, the crystals were soaked in a solution containing 90 mM MOPS pH 7.1, 11.25% MEPEG5K, 25 mM NaCl, 12% ethylene glycol, and 25 mM imidazole for 1 week. The crystals of ArsAB were unstable in a synthetic mother liquor that lacked an aromatic base, however imidazole was not observed in the crystal lattice. The soaked crystals were transferred in two steps into a cryoprotectant solution which contained 100 mM MOPS pH 7.1, 12.5% MEPEG5K, 25 mM NaCl, 17% ethylene glycol, and 25 mM imidazole and rapidly plunged into liquid nitrogen. Substrate-free ArsAB crystallized in the space group P2₁2₁2₁ with unit cell dimensions of a = 51.3 Å, b = 78.0 Å, and c = 151.8 Å where there was a single ArsAB heterodimer in the asymmetric unit.

Crystallization of ArsAB Complexed With DMB. Crystals were grown by mixing 2 μL of ArsAB at 10 mg/mL containing 5 mM DMB, and 10 mM MOPS pH 7.6 with 2 μL of reservoir solution containing 100 mM MOPS pH 7.1, 12.5% MEPEG5K, 20 mM NaCl, 12% ethylene glycol and 5 mM diethylenetriamine. Hanging droplets were nucleated immediately by streak-seeding where after the crystals grew to approximate dimensions of 200 × 200 × 400 μm within 7 days. The crystals were transferred stepwise into a cryoprotectant solution containing 100 mM MOPS pH 7.1, 12.5% MEPEG5K, 20 mM NaCl, 15% ethylene glycol, and 5 mM diethylenetriamine and rapidly plunged into liquid nitrogen. ArsAB complexed with DMB crystallized in the space group P2₁ with unit cell dimensions of a = 52.7 Å, b = 77.4 Å, c = 152.3 Å, α = 90.0°, β = 90.2°, and γ = 90.0° where there were two heterodimers of ArsAB in the asymmetric unit.
Crystallization of ArsAB Complexed With p-Cresol.
Crystals of ArsAB in complex with p-cresol were prepared in an similar fashion to those required to obtain ArsAB in the substrate-free state with the exception that 20 mM p-cresol replaced the 25 mM imidazole in the soaking and cryoprotecting solutions. Crystals of ArsAB complexed with p-cresol belong to the space group $P2_12_12_1$ with unit cell dimensions of $a = 52.5 \, \text{Å}, b = 77.9 \, \text{Å},$ and $c = 152.2 \, \text{Å}$ and contained a single ArsAB heterodimer in the asymmetric unit.

Crystallization of ArsAB Complexed With Phenol.
Crystals of ArsAB in complex with phenol were grown by combining 2 $\mu$L of 10 mg/mL ArsAB heterodimers in 10 mM HEPES pH 7.6 with 2 $\mu$L of reservoir solution containing 100 mM MOPS pH 7.1, 13.0% MEPEG, 20 mM NaCl, 12% ethylene glycol, and 25 mM phenol. Crystals were transferred stepwise into a cryoprotecting solution composed of 100 mM MOPS pH 7.1, 13.2% MEPEG, 17% ethylene glycol, and 25 mM phenol and frozen by rapidly plunging into liquid nitrogen. The crystals of ArsAB complexed with phenol belong to the space group $P2_12_12_1$ with unit cell dimensions of $a = 52.7 \, \text{Å}, b = 77.4 \, \text{Å}, c = 152.3 \, \text{Å}, \alpha = 90.0^\circ, \beta = 90.0^\circ,$ and $\gamma = 90.0^\circ$ and two ArsAB heterodimers in the asymmetric unit.

Data Collection and Structure Determination for ArsAB Substrate-Free, DMB, p-cresol, and Phenol Complexes. X-ray data for substrate-free ArsAB and ArsAB complexed with DMB, p-cresol, and phenol-glucinol were collected at 100 K on the Structural Biology Center beamline 19BM at the Advanced Photon Source in Argonne, IL. Diffraction data were integrated and scaled with HKL3000. Data collection statistics are given in Table 2. The ArsAB heterodimer structure was determined using the structure of CobT from Salmonella enterica (PDB entry 1L4B) as a molecular replacement search model in the program Molrep.

Final models were generated with alternate cycles of manual model building and least-squares refinement using the programs Coot and Refmac. Refinement statistics are presented in Table 2.

Data Collection and Structure Determination for ArsAB Complexed With Phenol. X-ray data for the ArsAB complexed with phenol were collected at 100 K with a Bruker AXS Platinum 135 CCD detector equipped with Montel optics and controlled by the Proteum software suite (Bruker AXS Inc.). All data sets were integrated with SMART version 7.06A and internally scaled with SADABS version 2005/1. The structure was determined by molecular replacement using PHASER in which ArsAB from the DMB complex structure was used as a search model. Model refinement was performed by alternate cycles of manual building with Coot and restrained refinement with Refmac. Refinement statistics are presented in Table 2.

ArsAB Kinetic Assays. Pseudo-first-order kinetics of the ArsAB-catalyzed reaction was performed using saturating concentrations of NaMN (5 mM), varying amounts of DMB or phenol (0.010–2 mM), and ArsAB protein at 0.25 $\mu$M. Stock solutions of DMB and phenol substrates were made in 30% (v/v) ethanol. All reaction mixtures contained 100 mM glycine pH 9 and 3% (v/v) ethanol at 37 °C. Triplicate assays were performed and analyzed as described elsewhere.

ArsAB Activity Assays. Activities of ArsAB and variants on DMB and p-cresol were measured with 3 mM of substrates (DMB/p-cresol and NaMN) and 0.2 $\mu$g/$\mu$L enzyme in 100 mM glycine, pH 9.0. The amount of enzyme used in this assay was normalized to the least amount of variant protein needed to give detectable activity. Twenty-five microliters of each reaction was taken at 1, 6, and 13 min and boiled to stop the reaction. An equal volume of 20 mM ammonium acetate buffer

Table 2. X-ray Data Collection and Refinement Statistics

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</table>

$^a$Values in parentheses are for the highest-resolution shell. $^bR_{merge} = \frac{\sum|I(hkl)| - \langle \sum I(hkl) \rangle}{\langle \sum |I(hkl)| \rangle}$ where the average intensity $I$ is taken over all symmetry equivalent measurements and $I(hkl)$ is the measured intensity for a given observation. $^cR_{merge} = \frac{\sum|I(hkl)| - F_{calc}(hkl) \times 100.}{\sum |I(hkl)|},$ where $R_{merge}$ refers to the $R_{merge}$ for the data utilized in the refinement and $R_{free}$ refers to the $R_{free}$ for 5% of the data that were excluded from the refinement.
that included a hexahistidine (His6) tag and a rTEV protease (pARSAB7), was used to isolate an MEVI residues of ArsA but retained the His6 a arsAB
Expression of whether the true start codon was the alternative methionine. more than 3 mg/mL (data not shown). This led to the question conditions tested, and the enzyme could not be concentrated to production of ArsAB from pARSAB7 was poor under the fi MEVI residues. ArsB copuri
fold higher than that of the enzyme containing the additional (nmol of product/min/mg of protein) of this material was 10-
under multiple conditions. Furthermore, the speci
icity (nmol of product/min/mg of protein) of this material was 10-
fold higher than that of the enzyme containing the additional MEVI residues (274 vs 24, respectively). Finally, the resulting protein could be concentrated readily to more than 10 mg/mL. Open reading frame (ORF) prediction software GeneMark.hmm also predicted the downstream translational start codon as the likely ORF for the gene.23 The plasmid pARSAB22 was used to prepare ArsAB enzyme for all subsequent crystallographic and kinetic studies.

Structure of the Substrate-Free ArsAB Heterodimer. Crystals of substrate-free ArsAB were obtained by soaking out the crystallization additive phloroglucinol, which bound in the ArsA active site. The final substrate-free-ArsAB model contains 334 of 348 expected amino acids for the ArsA subunit which extend from Ser2-Asn339. For the ArsB subunit the model contains 328 of 350 expected amino acid residues, which extend from Leu2-Ala333. The electron density is continuous for ArsA except for a break between Ser208 and Leu212 in ArsA. This disordered region consists of a loop that folds over the putative binding site for the substrate NaMN. In the ArsA subunit this loop is displaced by the N-terminus of a symmetry-related ArsB molecule in what is presumably a crystal packing interaction. The length of the break in this region varies among the substrate complexes determined. There is also one break in the electron density for ArsB between Ala70 and Met78. This break occurs in a surface loop that spans the active site cavity. The structure of substrate-free-ArsAB is shown in Figure 2.

Despite sharing only 37% sequence identity, ArsA and ArsB exhibit highly similar tertiary structures with a root-mean-square deviation (RMSD) of 1.3 Å for 311 structurally equivalent α-carbon atoms. As expected, ArsA and ArsB have a very similar fold to that described for SeCobT24 with an overall RMSD of 1.25 Å with respect to either ArsA or ArsB relative to PDB coordinates 1D0S. The overall fold contains two domains where the large domain consists of a six-stranded parallel β-sheet surrounded by α-helices in the β6, β5, β4, β1, β2, β3 arrangement characteristic of the classic Rossmann dinucleotide binding motif. The small domain is defined by a three-helix bundle, which is built from two N-terminal helices and a longer C-terminal helix that spans both domains.

The ArsAB heterodimer exhibits the same quaternary arrangement as the SeCobT homodimer, with numerous

**RESULTS AND DISCUSSION**

Identification of the Initiating Methionine of ArsA. The initial characterization of the S. ova ta arsAB genes identified two potential translational initiation sites in the arsA-coding region.12 The difference between the two putative ArsA proteins was four amino acids (i.e., MEVI). In the previous report on ArsAB,12 an overexpression vector (pARSAB7), was used to isolate an N-terminally tagged protein that included a hexahistidine (His6) tag and a rTEV protease site followed by the MEVI residues mentioned above. The same vector also directed the synthesis of ArsB without a tag. After isolation of His6-ArsA, the His6-tag was removed with rTEV protease,15 leaving two additional glycines prior to the MEVI residues. ArsB copurified with ArsA, but the overproduction of ArsAB from pARSAB7 was poor under the conditions tested, and the enzyme could not be concentrated to more than 3 mg/mL (data not shown). This led to the question of whether the true start codon was the alternative methionine. Expression of arsAB from plasmid pARSAB22 eliminated the MEVI residues of ArsA but retained the His6 affinity-tag and the rTEV enzyme recognition site. Protein expression driven by this construct yielded ArsAB enzyme that was readily purified under multiple conditions. Furthermore, the specific activity (nmol of product/min/mg of protein) of this material was 10-fold higher than that of the enzyme containing the additional

**Figure 2.** Stereo representation of ArsAB heterodimer. ArsA is colored in blue, ArsB is colored in green. DMB is shown in yellow.
contacts along the dimer interface. The total buried surface area of the dimer is about 2300 Å²/subunit as computed with the program AREAIMOL in the CCP4 package.25 In SeCobT and ArsAB the active site is located at the dimer interface and is built from both subunits of the dimer. The C-terminus of one subunit and a small N-terminal loop of the second subunit24 form the site and result in two structurally equivalent active sites per dimer. Due to the heterodimeric nature of ArsAB, the two active sites in the dimer are structurally distinct.

There are notable structural differences in the surface loops that surround the active site cavities of ArsA and ArsB. The loop between helices 2 and 3 in ArsB (Leu27-Leu34) that forms part of the active site for ArsA curls away at Pro30 in ArsB in a fashion similar to that of SeCobT and provides sufficient room for substrate binding in the ArsA active site. Conversely, in ArsB the corresponding loop from ArsA penetrates further into the analogous site in ArsB (Figure 3). ArsB also contains an additional C-terminal helix spanning Phe327-Glu331 that projects from the three helix bundle of the small domain and is directed away from the ArsA active site. There is no observable electron density for the C-terminal 17 amino acids of ArsB. These differences, along with others discussed later, indicate that DMB phosphoribosyltransferase activity has been lost from the ArsB active site.

The observation that ArsAB heterodimers crystallized supports the earlier suggestion that neither ArsA nor ArsB can assemble independently to form an active enzyme.25 ArsAB is likely the result of a gene duplication event onto which selective forces resulted in the coevolution of ArsA and ArsB into functional a heterodimer. Why functional homodimers were not favored remains unclear. At present, information available from genome databases shows that the tandem organization of arsAB is only found in some members of Veillonellaceae family, suggesting that the evolution of heterodimeric SeCobT homologues may be rooted in these bacteria.

**Structure of ArsAB Heterodimer in Complex With DMB.** The ArsAB heterodimer in complex with DMB crystalized in the space group P2₁ with two heterodimers in the asymmetric unit. No significant differences in tertiary structure were observed between the ArsAB heterodimer in either crystal system (RMSD: 0.45 Å), and inspection of the unit cell revealed no substantial change in crystal packing interactions. It is unclear whether the shift in space group was due to the presence of DMB, or by a change in crystallization conditions which included the crystallization additive diethylenetriamine.

There is unambiguous electron density for DMB in the ArsA active sites for both heterodimers in the asymmetric unit (Figure 4A). The overall RMSD between the two heterodimers in the asymmetric unit is 0.24 Å. Given the similarity between the two heterodimers, the discussion and figures are based on chains A and B in the asymmetric unit. The substrate binding site lies in a cavity at the interface of the ArsA and ArsB subunits, near the periphery of the dimer in a manner that is similar to that seen for SeCobT.24 The cavity is defined by helices 5 and 6 and their connecting loop in ArsA, the loop leading into helix 16 in ArsA, and β4 and its adjoining loop in ArsA. Additionally, the loop between helices 2 and 3 of the ArsB subunit forms part of the cavity (Figure 4B). The cavity in which DMB binds is hydrophobic. DMB sits atop Met87 and Met177, Ile179, and Leu317 in ArsA and Pro30 in ArsB. An imidazole nitrogen on DMB forms a hydrogen bond with the proposed active site residue Glu319 (2.9 Å) (Glu317 in SeCobT).26

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Figure 3. Structural alignment of ArsA with ArsB in stereo representation. ArsA (blue) with p-cresol bound (yellow) was superposed onto ArsB (green). The original positions of ArsA and p-cresol are depicted in light gray. Arrows indicate areas of greatest divergence: 1 the loop that folds over the second substrate [ArsA: Gly206-Thr218, ArsB: Gly189-Gly197], 2 the loop that projects into the active site on the adjacent subunit [ArsA: Val33-Gly41, ArsB: Leu27-His35], and 3 additional C-terminal helix in ArsB.
DMB Binding Is Conserved. Alignment of DMB-bound structures for ArsAB and ScCobT reveals that DMB binds in a nearly identical orientation in both enzymes (Figure 5). This is consistent with the sequence and structural conservation of the two enzymes. The character of most of the side chains in the DMB binding site are conserved between ArsA and ScCobT, many of which are hydrophobic. Notable exceptions are the substitution of both Gln88 and Leu175 in ScCobT to methionine in ArsA. These methionine residues form a large part of the hydrophobic binding surface for DMB in ArsA. The most apparent difference in first shell interactions between ScCobT and ArsAB is the replacement of Ser80 in ScCobT with Tyr79 in ArsA. This substitution likely compensates for the absence of hydrophobic surface contributed by the C-terminus of ScCobT. There is no evidence that the largely disordered C-terminus of ArsB makes contacts with the active site of ArsA. Tyr79 of ArsA may play a role in the activation of phenolics as discussed later. A key conserved interaction in the active site of

Figure 4. (A) Stereoview of the electron density for DMB in the ArsA active site. The electron density (3.5σ) was calculated from coefficients of the form $F_o - F_c$ where DMB was omitted from the phase calculation and refinement. ArsB has been omitted for clarity. (B) The DMB binding site is formed by contributions from both subunits: ArsA is shown in blue and ArsB is shown in green.

Figure 5. Alignment of ScCobT and ArsAB in complex with DMB. Coordinates for ScCobT-DMB were derived from PDB ID: 1D0S.24 DMB from ScCobT is colored violet. DMB from ArsAB is colored yellow.
ArsA is the hydrogen bond between Glu319 (Glu317 in SeCobT) and one of the imidazole nitrogen atoms of DMB. In addition to its role in coordinating and orienting the substrate, this residue has been proposed to act as an active site base24 in SeCobT. This constellation of interactions is also observed in ArsAB providing strong evidence that ArsAB enzyme may utilize the same catalytic mechanism as SeCobT to catalyze phosphoribosyl transfer.

The second substrate, NaMN, was not captured in complex with ArsAB. Indeed, the portion of the NaMN binding site in SeCobT that lies atop the phosphate moiety is disordered in the structurally equivalent region in ArsAB. This disordered loop
spans helices 9 and 10 in ArsA, and may undergo a disorder/order transition upon binding the second substrate. Alternatively, crystal-packing interactions may have displaced these residues as there are numerous contacts between the N-terminus of a symmetry related molecule of ArsB (Glu3 and Glu4) and the NaMN binding pocket in ArsA (Lys215). Despite the lack of structural information in this region, many key interactions, such as those between side chain of SeGlu174 and the amide nitrogen of SeGly176 with the ribose moiety are observed to be in structurally equivalent positions in ArsA (Glu179 and Gly 178). The cavity in which the nicotinate moiety binds is also maintained, however a slight rearrangement is observed around Ser293 in ArsA as compared to SeCobT, where SeSer291 interacts with an oxygen of the carboxylate group of nicotinate. In ArsA, the peptide backbone of the Ser293 has rotated nearly 180° leaving the side chain directed toward the solvent rather than the nicotinate binding cavity.

DMB Phosphoribosyl Transferase Activity Is not Maintained in ArsB. While there is a strong conservation of active site architecture and substrate binding between SeCobT and ArsA, the equivalent site in ArsB shows considerable divergence (Figure 6). The most striking difference is the position of the side chain of Arg83, which lies directly within the DMB binding site. Additionally, the hydrophobic loop that is contributed by the adjacent ArsA subunit lies much deeper in the cavity and is not curled away from the site by Pro30 (ArsB) as is observed in the ArsA DMB binding site. Considered together, these structural features of ArsAB likely preclude the binding of DMB in a conformation that is catalytically competent. Interestingly, the proposed catalytic residue Glu303 is maintained in a structurally equivalent position in ArsB. The NaMN-binding site has also undergone several structural rearrangements as depicted in Figure 6. The loop that folds over the phosphate moiety in NaMN in SeCobT is shortened in ArsB and tracks along the outer surface of the cavity rather than toward the interior. A sequence alignment of SeCobT and ArsB shows an 11 residue gap in this region in ArsB. Lastly, the loop that includes Tyr79 in ArsA and spans β1 and helix 4 is disordered in ArsB but is directed inward toward the active site cavity before becoming disordered. The structural divergence of the active site in ArsB could be the result of selective pressures leading to heterodimer formation, or to the ability of ArsA to transfer phenolic bases. At this point, it is unknown whether the ArsB site has any catalytic activity whatsoever. It is conceivable that the ArsB active site has evolved to perform another function. In this respect, although the ArsB active site does not have base:phosphoribosyl transferase activity, the presence of phloroglucinol (Figure S1, Supporting Information) in the ArsB active site suggests that the capacity to bind phenolic bases has been maintained.

Structure of ArsAB Heterodimer in Complex with Phenol and p-Cresol. To gain insight into the mechanism by which ArsAB generates phenolic α-ribose monophosphates, crystal structures were determined for ArsAB in complex with both p-cresol and phenol. There is clear electron density in the ArsA active site for both substrates (Figure 7). The phenolic substrates bind in the same location in the active site cavity at
the interface between the ArsA and ArsB subunits. The constellation of side chains surrounding the bound substrate remains unchanged relative to the complex with DMB with the notable exception of the hydrogen bonding pattern about Glu319 and the loop between helices 4 and 5, most notably at Tyr79. While Glu319 participates in a direct hydrogen bond with the imidazole nitrogen in complex DMB (2.9 Å), it has been replaced with a well-ordered water molecule with phenolic substrates bound. In the case of p-cresol, this water molecule (W1) is 2.8 Å from Oε2 of Glu319 and 2.4 Å from the hydroxyl moiety on p-cresol (Figure 8). The presence of W1 results in a rotation of the planar substrate away from Glu319 and toward the NaMN binding site. This position is additionally maintained by the rotation of Tyr79 3.6 Å inward from that of the DMB complex, resulting in a close interaction with p-cresol (Figure 8). Taken together, these interactions likely explain how p-cresol is positioned for a nucleophilic attack at the C1 carbon of ribose following deprotonation, which is facilitated by the close hydrogen bond with W1, as it is ideally positioned to transfer a proton to Glu319.

Comparison of SeCobT with ArsAB Bound to Phenolic Substrates. SeCobT is unable to catalyze phosphoribosyl transfer to phenolic substrates, whereas this is readily accomplished by ArsAB. The presence of a water molecule in the active site of the ArsAB-phenolic complexes compared to the absence of a water molecule in the phenolic complexes with SeCobT suggests that a water molecule plays a key role in the positioning and deprotonation of p-cresol for attack at the C1 carbon of ribose. In SeCobT, p-cresol hydrogen bonds directly (2.6 Å) with Oε2 of Glu317. This close hydrogen bond between p-cresol and Glu317, places the phenolic hydroxyl group hydroxyl group in a position that is most likely too far from the C1 carbon of ribose (>4.0 Å) for the reaction to occur. The reason that SeCobT does not allow inclusion of a water molecule in the equivalent position of W1 in ArsAB is not immediately obvious, but careful examination of the comparison of the active sites in ArsAB and SeCobT suggests several possibilities.

Superposition of the active sites (Figure 9) shows that in order for ArsA to accommodate a water molecule the catalytic glutamate Glu319 (Glu317 in SeCobT) must move away from its position in SeCobT. This would appear to be facilitated by the replacement of Gln88 in SeCobT by Met87 in ArsA. This change removes a hydrogen bonding interaction which is predicted to stabilize the position of the catalytic glutamate in the active site. There is also a difference in the position and conformation of the backbone atoms for the polypeptide chain that surrounds the catalytic base. This change appears to be coupled to a number of changes in this region. In particular, replacement of Ser319 in SeCobT by Ile321 in ArsA requires a rearrangement in the backbone to accommodate the loss of a hydrogen bond.

The hypothesis that Met87 and Ile321 play a role in enabling ArsAB to catalyze phosphoribosyl transfer to phenolic substrates was tested by preparing the reverse mutations M87Q and I321S (Table 3). The effect of the M87Q mutation is most pronounced with respect to phenolic specificity, as the specific activity with p-cresol drops by 2.6 fold while a slight increase (1.2 fold) in activity with DMB was observed. Introduction of serine at Ile321 was deleterious to both phenolic and DMB activation, suggesting that the role of this amino acid residue in ArsA is not well understood. From these experiments it is clear that specificity for phenolic substrates is influenced by more than one residue of the base-binding pocket. A full understanding of the structural basis for the utilization of phenolic substrates by ArsAB must await further structural and kinetic studies of variants.

Efficiency of the ArsAB Enzyme as a Function of its Base Substrate. Earlier work by Stupperich et al. showed that p-cresolyl-Cba and phenolyl-Cba comprised >90% of the cobamides synthesized by S. ovata, yet data shown in Table 4 indicate that the ArsAB enzyme efficiently uses DMB as its base substrate. However, the $K_m$ of ArsAB for DMB reported herein (415 μM) is much higher than other CobT homologues, namely S. enterica ($K_m$ < 10 μM) and Pseudomonas denitrificans (16 nM). The dramatic increase in $K_m$ for DMB in ArsAB may be a consequence of the evolutionary changes required for this enzyme to phosphoribosylate phenolic bases. From this perspective the higher $K_m$ for DMB observed with ArsAB suggests that increased flexibility needed to accommodate a water molecule might be a compromise to maintain both activities. The crystallographic snapshots of ArsAB and SeCobT show DMB bound in the same position and orientation in both enzyme active sites so there is no easy structural explanation for the observed difference in $K_m$ between ArsAB and SeCobT. Interestingly, the interaction between the C-terminal residues contributed by the neighboring subunit in SeCobT and DMB are apparently lacking in ArsAB. This could affect on-rates and off-rates of substrate and thereby $K_m$.

The catalytic efficiency of the activation of DMB and phenol for ArsAB differ only by a factor of 2 (Table 4), suggesting that,
under some conditions, this bacterium may synthesize cobalamin (Co\(^{5,6}\)-dimethylbenzimidazolyl-Cba) or other cobamides containing purines or purine analogues.\(^5\)\(^,\)\(^12\) Notably, while the apparent \(K_M\) for phenol was 8-fold lower than the apparent \(K_M\) for DMB, the turnover number when phenol was the substrate was 4-fold slower than the one calculated for the reaction when DMB was the substrate (Table 4). It is noteworthy that phenolyl-Cba comprises only 20% of the reaction when DMB was the substrate (Table 4). It is known that phenolyl-Cba comprises only 20% of the reaction when DMB was the substrate (Table 4).

### Table 4. Kinetic Parameters of the ArsAB-Catalyzed Reaction\(^a\)

<table>
<thead>
<tr>
<th>substrate</th>
<th>(K_M) (μM)</th>
<th>(k_{cat}) (s(^{-1}))</th>
<th>(k_{cat}/K_M) (s(^{-1}) M(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>5,6-dimethylbenzimidazole</td>
<td>415 ± 56</td>
<td>1.6 ± 0.08</td>
<td>3.9 \times 10(^4)</td>
</tr>
<tr>
<td>phenol</td>
<td>50 ± 11</td>
<td>0.4 ± 0.05</td>
<td>8 \times 10(^4)</td>
</tr>
</tbody>
</table>

*Reaction mixtures contained either 1.9 mM phenol + [\(^{14}\)C-\(U\)]-phenol (0.1 mM, 60 mCi mmol\(^{-1}\)), or 1.98 mM 5,6-dimethylbenzimidazole (DMB) + [\(^{14}\)C-2]-DMB (0.02 mM, 43 mCi mmol\(^{-1}\)). Other components present in the reaction mixture and the protocol for the assessment of product formation are described under Experimental Procedures.

\(^a\)These authors contributed equally to this work.

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