Comparative Genomics Guided Discovery of Two Missing Archaeal Enzyme Families Involved in the Biosynthesis of the Pterin Moiety of Tetrahydromethanopterin and Tetrahydrofolate

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Supporting Information

ABSTRACT: C-1 carriers are essential cofactors in all domains of life, and in Archaea, these can be derivatives of tetrahydromethanopterin (H₄-MPT) or tetrahydrofolate (H₄-folate). Their synthesis requires 6-hydroxymethyl-7,8-dihydropterin diphosphate (6-HMDP) as the precursor, but the nature of pathways that lead to its formation were unknown until the recent discovery of the GTP cyclohydrolase IB/MptA family that catalyzes the first step, the conversion of GTP to dihydroneopterin 2’,3’-cyclic phosphate or 7,8-dihydroneopterin triphosphate [El Yacoubi, B.; et al. (2006) J. Biol. Chem. 281, 37586–37593 and Grochowski, L. L.; et al. (2007) Biochemistry 46, 6658–6667]. Using a combination of comparative genomics analyses, heterologous complementation tests, and in vitro assays, we show that the archaeal protein families COG2098 and COG1634 specify two of the missing 6-HMDP synthesis enzymes. Members of the COG2098 family catalyze the formation of 6-hydroxymethyl-7,8-dihydropterin from 7,8-dihydronopterin, while members of the COG1634 family catalyze the formation of 6-HMDP from 6-hydroxymethyl-7,8-dihydropterin. The discovery of these missing genes solves a long-standing mystery and provides novel examples of convergent evolutions where proteins of dissimilar architectures perform the same biochemical function.

The availability of over 3000 published genome sequences has enabled the use of comparative genomic approaches to drive the biological function discovery process. Classical methods, such as phylogenetic distribution profiles, physical clustering, gene fusion, coexpression profiles, structural information and other genomics data and postgenomic resources are now used to make very strong functional hypotheses that can be quickly validated by simple genetic and/or biochemical tests. The whole procedure can occur in just weeks, taking advantage of the constantly growing available postgenomic resources such as gene deletion or expression libraries. Here, we illustrate this paradigm shift with the discovery of two archaeal protein families involved in the synthesis of 6-hydroxymethyl-7,8-dihydropterin diphosphate (6-HMDP), the precursor of the pterin containing moiety of the essential C₁-carriers tetrahydrofolate (H₄-folate) and tetrahydromethanopterin (H₄-MPT) (Figure 1). These enzymes had eluded classical genetic and biochemical approaches and had been missing for decades.

Most organisms use H₄-folate (Figure 1) as the essential carrier of C₁ fragments in both anabolic and catabolic reactions. The known exceptions are the methanogenic Archaea that use H₄-MPT (Figure 1) and methylo trophic bacteria that use dephospho-H₄-MPT. The situation in Archaea is quite diverse. Halophilic Archaea such as Halobacterium species harbor folates. Hyperthermophiles like Pyrococcus or Sulfolobus species use C₁-carriers lacking the C-7 methyl group on the pterin as seen in methanop therms. Methanogenic Archaea such as Methanobacterium thermoautotrophicum ΔH (now called Methanobacterium thermoautotrophicus) use H₄-MPT, whereas Thermococcus litoralis and Pyrococcus furiosus use only a more exotic derivative of methanopterin containing poly-β-(1→4)-N-acetylglucosamine as side chains on their C₁-carrier coenzyme. Certain Archaea such as Methanosarcina barkeri contain both H₄-MPT and H₄-folate derivatives.
**Figure 1.** Early steps of tetrahydrofolate and tetrahydromethanopterin pathways in Bacteria and Archaea. Most bacteria use the FolE (or FolE2)/FolB/FolK route (in blue) to 6-HMDP even if some use the bacterial PTPS-III shunt (in green). Several routes to the common 6-HMDP intermediate in tetrahydrofolate and tetrahydromethanopterin are found in Archaea. A common pathway is the FolE2/MptD/MptE route (in red) such as in *H. volcanii* paralleling the bacterial pathway. However, some methanogens such as *M. jannaschii* use the MptA/MptB/MptD/MptE route, whereas *P. furiosus* uses the archaenal PTPS-III shunt. Phosphatases still to be identified are noted by a question mark (?). FolE/FolE2, GTP cyclohydrolase IA/IB (GCYH-IA/B); FolB, 7,8-dihydroneopterin aldolase (DHNA); FolK, 7,8-dihydro-6-hydroxymethylpterin diphosphokinase (6-HMDPK); MptA, archaenal GTP cyclohydrolase I (Fe(II)-dependent enzyme); MptB, Fe(II) dependent-cyclic phosphodiesterase; MptD, archaenal specific DHNA; MptE, archaenal specific 6-HMDPK; PTPS-III/PTPS-V/PTPS-VI, pyruvyltetrahydropterin synthase paralogs involved in 6-HMDP synthesis.

*Sulfolobus solfataricus* contains a hybrid coenzyme C1-carrier coenzyme harboring a nonmethylated pterin and the same arylamine moiety found in methanopterin.16 Although numerous variations in the C1-carrier structures exist among the various archaeal lineages, the early steps in the syntheses of H$_4$-folate and of H$_4$-MPT and its derivatives, leading to the formation of the 6-HMDP intermediate have been predicted to be similar17 (Figure 1). The 6-HMDP pathway is well characterized in bacteria, plants, and fungi. GTP cyclohydrolase IA (GCYH-IA or FolE) or GTP cyclohydrolase IB (GCYH-IB or FolE2) catalyze the first step of the pathway producing 7,8-dihydroneopterin triphosphate (H$_2$NTP) from GTP.18–20 H$_2$NTP produces 7,8-dihydroneopterin (H$_2$Neo) after the lost of a diphosphate and a phosphate. Then, 7,8-dihydroneopterin aldolase (DHNA) encoded in *Escherichia coli* by *folB*21 catalyzes the formation of 6-hydroxymethyl-7,8-dihydropterin (6-HMD) from H$_2$Neo. A derivation from the classical bacterial 6-HMDP synthesis pathway occurs in *Plasmodium falciparum* and various bacteria. The DHNA step is bypassed by PTPS-III that cleaves the side chain of H$_2$NTP to form 6-HMD22–24 (Figure 1). In all cases, 6-HMD is then diposphorylated with ATP by a 7,8-dihydro-6-hydroxymethylpterin diphosphokinase (6-HMDPK); encoded in *E. coli* by *folK*25 to form 6-HMDP.

*Methanocaldococcus jannaschii* was the first Archaea with a sequenced genome. It was immediately apparent that this organism lacked homologues of FolE, FolB, and FolK and used nonorthologous enzymes to catalyze the same reactions.26 This prediction was confirmed as more archaeal genomes became available (Figure 2). As shown in Figure 2, a minority of Archaea (16 out of 58 analyzed) contained homologues of the canonical FolE and expression of the corresponding gene from *Sulfolobus solfataricus* (*sso0364*) complemented the deoxythymidine (dT) auxotrophy of an *E. coli* Δ*folE* mutant.27 Most Archaea (40/58 analyzed) contained homologues of the more recently discovered FolE2 (Figure 2) that were experimentally validated in a few species. The *folE2* mutant of *Haloflexx volcanii* (*ΔHVO_2348*) is a ΔT and hypoxanthine auxotroph,28 and the *M. jannaschii* FolE2 homologue MptA (*MJ0775*) is a unique Fe(II)-dependent GTP cyclohydrolase IB that forms...
7,8-dihydroneopterin 2′,3′-cyclic phosphate. In M. jannaschii, MptB (MJ0837), a cyclic phosphodiesterase, is required to cleave the cyclic phosphate to form a mix of 7,8-dihydroneopterin 2′-monophosphate and 7,8-dihydroneopterin 3′-monophosphate. This pathway, involving a 7,8-dihydroneopterin 2′,3′-cyclic phosphate intermediate, could be specific to a subset of methanogens because homologues of MptB are mostly found in Methanococcales (Figure 2). Even if the first archaeal 6-HMDP biosynthesis enzymes have been characterized, the remaining steps encoded in bacteria by FolB and FolK remain to be discovered in most Archaea. The identification and characterization of these missing gene families is the focus of this study.

Only two sequenced Archaea (Sulfolobus acidocaldarius DSM 639 and Caldivirga maquilingensis IC-167) contain homologues of bacterial FolB proteins fused with homologues of bacterial FolK proteins (Saci_1101 and Cmaq_0517, respectively) (Figure 2) that certainly derive from a lateral gene transfer event (the closest homologue to these two proteins is the fused FolKB from Pneumocystis carinii f. sp. macacae (AAN38834.1) with a Blastp E-value of 9e-40). A few Archaea such as P. furiosus or Methanosarcina barkeri str. fusaro harbor proteins of the PTPS-III family (Figure 2) that in bacteria function in a DHNA bypass where H2NTP is converted directly to 6-HMDP (Figure 1). Surprisingly, in vitro, the PTPS-III homologue from P. furiosus, PF1278, catalyzed the cleavage of 7,8-dihydroneopterin monophosphate (H2NMP) to 6-HMD, but H2NTP was not a substrate (Supporting Information and Figure 1). Finally, we recently showed that close homologues of PTPS-III with a slightly different active site motif named PTPS-VI were found in a few Sulfolobus species (Figure 2). Expressing the PTPS-VI gene from Sulfolobus acidocaldarius (sso2412) partially complemented the ΔfolB E. coli mutant suggesting a role of PTPS-VI proteins in 6-HMDP
synthesis even if the substrate specificity of this family is yet to be experimentally determined (Figure 1). In summary, 56 out of 58 of the archaeal genomes analyzed lacked a FolK homologue, and 47 out of 58 lacked a FolB, PTPS-III, or PTPS-VI homologue. Hence, we set out to identify these missing archaeal 6-HMDP synthesis enzymes using a combination of comparative genomic approaches.

We first searched for genes that physically clustered with pterin related genes using the clustering tool of the SEED platform\(^2\) and identified the COG1634/DUF115 gene family as a candidate (Figure 3A). Members of COG1634 are uncharacterized proteins found in most Archaea (Figure 2) and are part of the thiamin pyrophosphokinase (TPK, thiamin diphosphokinase) catalytic domain superfamily.\(^3\) TPK is a thiamin salvage enzyme that transfers the diphosphate group of ATP to thiamin to form thiamin diphosphate, the active form of the cofactor.\(^4\) TPK consists of two domains: the N-terminal catalytic domain that binds ATP and the C-terminal substrate-binding domain that binds thiamin.\(^5,6\) The COG1634 family members show sequence similarity to the TPK catalytic domain but not to the C-terminal domain (Supplemental Figure 1). Moreover, fold recognition servers, e.g., FFAS,\(^7\) predict the TPK catalytic domain being a good template for the COG1634 subunit fold; they also suggest even higher-scoring hits to the structures of bacterial GST-II-like sialyltransferases\(^8\) (Supplemental Figure 1). These bacterial enzymes use CMP-NeuAc as a sugar donor and have a distinct fold that is also found in mammalian sialyltransferases.\(^9\) Comparison of the sialyltransferase and TPK structures revealed a common structural core and similar binding modes of their respective products, CMP and AMP, suggesting a distant evolutionary relationship of these protein families (Supplemental Figure 2). The COG1634 members are predicted to share the NMP-binding site (Supplemental Figure 1). In addition, COG1634 and TPK family members share the metal (Mg(II)) ion-binding site, involved in binding and transfer of the diphosphate group (Supplemental Figures 1 and 2). On the basis of the physical clustering evidence and fold homology, we predicted that COG1634 was the missing archaeal 6-HMDPK family. The homology with the sialyltransferase family opens the possibility that members of the COG1634 family may utilize other nucleoside triphosphates, e.g., CTP. There are documented cases in archaeal biosynthetic pathways where CTP substitutes for ATP; for example, the Archaeon-specific riboflavin kinase uses CTP as its phosphoryl donor,\(^10\) and the archaeal FAD synthetase (RibL) catalyzes the cytidylation of FMN with CTP.\(^11\)

Figure 3. Comparative genomic evidence. (A) Clustering of COG1634 and COG2098 genes with pterin and cofactor biosynthetic related genes. Abbreviation not found in the text: FolP-like, dihydropteroate synthase-like enzyme homologous to the bacterial folate enzyme FolP but of unknown function;\(^5\) MptG, β-ribofuranosylaminobenzene 5′-phosphate synthase;\(^6\) FoLM, alternative dihydrofolate reductase;\(^7\) F420-lig, coenzyme F420: l-glutamate ligase.\(^8\) (B) The archaeal DHNA (MptD) tetramer with bound pterin ring mimic (PDB 2OGF). The individual subunits of MJ0408 are shown with differently colored cartoons, the bound ligand 8-oxoguanine with orange carbons. (C) Putative active site of the archaeal DHNA with manually docked neopterin. The MptD structure is from PDB 2IEC, and the neopterin ligand (orange carbons) is from PDB 2O90 (in alternative conformation B). The active site residues contributed by three different subunits are shown with green, cyan, and magenta carbons, respectively (as in panel B and Supplemental Figure 3).
We then observed that, in Desulfurococcus kamchatkensis, a gene in the COG2098 family was in a predicted operon with both the folE2 and COG1634 genes (Figure 3A). Physical clustering in only one organism is not very strong evidence; nonetheless, further structural analysis suggested that COG2098 was the missing archaeal DHNA family. The COG2098 family previously was targeted by Structural Genomics Initiatives resulting in the determination of three representative structures: one from Picrophilus torridus (PTO0218; PDB: 2I52), one from M. jannaschii (MJ0408; PDB: 2OGF), and one from Methanopyrus kandleri (MK0786; PDB: 2IEC). The subunit fold comprising two $\alpha$-helices and a four-stranded $\beta$-sheet somewhat resembles the fold of bacterial DHNA in architecture but differs from it in topology, as its secondary structure elements are connected in a very different order. Moreover, the COG2098 subunits assemble in a compact homotetramer, unlike the tunnel-like architectures of the canonical DHNA octomer. The tetramer is an apparent biological unit of the COG2098 family. The most conserved residues are scattered across the subunit surface but come together in the subunit interfaces. There are four equivalent putative active sites in the tetramer, each formed by the residues from three different subunits (Figure 3B). Fortuitously, in one of the determined structures, 2OHG, there is a ligand bound to each of the four sites that was tentatively identified as 8-oxoguanine (8-oxoG). Using the bound ligand as a guide, we manually docked the predicted substrate molecule in the COG2098 active site (Figure 3C). The dihydroneopterin molecule is in essentially the same conformation as in the structures of canonical DHNA complexes and fits almost...
perfectly in the active site pocket when its pterin ring is aligned with the 8-oxoG mimic. The environment of bound substrate is also similar to that of the canonical FolB, suggesting a similar enzymatic mechanism for the predicted archaeal DHNA.43

A genetic approach was first used to validate these predictions. The COG2098 and the COG1634 encoding genes from M. janaschii and represented, respectively, by mj0408 and mj1634, were cloned into pBAD24 under the P_BAD promoter44 and tested for complementation of the dT auxotrophy phenotype of the ΔfolB::kan^R E. coli strain (VDC326731) and of the ΔfolK::tet^R E. coli strain (C600 ΔfolK::tetB45), respectively. E. coli strains deleted in H4-folate biosynthesis genes can grow on rich medium if dT is added to the medium, albeit poorly because of the absence of the formylation of the inhibitor tRNA.46 As shown on Figure 4A, expression of mj0408 complemented the dT auxotrophy phenotype of the folB deletion as did expression of the E. coli folB positive control. Overexpression of mj0408 seemed to be toxic with cells showing better growth with no arabinose (Figure 4A). Similarly, overexpression of mj1634 complemented the Δt auxotrophy phenotype of the folK deletion as did expression of the E. coli folK positive control (Figure 4B).

A biochemical validation strategy was then used to confirm the genetic results. mj0408, mj1634, and its homologue from P. furiosus (pf0930) were all expressed in E. coli. The respective gene products, MJ0408, MJ1634, and PF0930 were purified by heating the extract to 80 °C followed by anion exchange chromatography. The resulting proteins were greater than 95% pure as judged by polyacrylamide gel electrophoresis with coomassie staining (Supplemental Figure 4). The identity of the purified proteins was confirmed by MALDI MS of the cryptic-digested protein band. The protein product from mj0408 was confirmed to be a DHNA as it was found to catalyze the formation of 6-HMD from H$_2$Neo (Figures 1 and 4C). The retention time of the 6-hydroxymethylpterin was identical to that of the 6-hydroxymethylpterin standard under two separate chromatographic systems utilizing either a Varian PursuitXRs C18 column or a Varian Pursuit polyfluorophenyl (PPF) column. The formation of 6-hydroxymethylpterin was linear with respect enzyme concentration, and no product was observed in control samples that were incubated in the absence of enzyme. In order to confirm that the observed activity was due to the mj0408 gene product and not a result of the E. coli DHNA activity, the activity of a cell extract and identical purified fractions from E. coli expressing the mj0408 gene to those of E. coli expressing a different gene (mj0929) were compared. The DHNA activity was greater than 4-fold higher in cell extracts of E. coli expressing the mj0408 gene. The identical MonoQ-purified fraction of mj0929 exhibited no activity relative to the purified mj0408 gene product.

The protein product from pf0930 was confirmed to be a 6-HMDPK as it was found to catalyze the formation of 6-HMDP from 6-HMD and ATP (Figures 1 and 4D). Of the four nucleotide phosphates tested, the maximum activity was observed with ATP. Relative to ATP, CTP, UTP, and GTP had, respectively, 41%, 40%, and 12% of the ATP activity. The fact that the oxidized product peak was 6-hydroxymethylpterin diphosphate (6-hydroxymethylpterin-PP) was confirmed by the following observations. The product peak had the same UV/visible absorbance and fluorescence spectra as a known sample of 6-hydroxymethylpterin-PP. Attempts to confirm the identity of the product peak by LC-ESI-MS analysis of crude incubation mixtures was not successful most likely due to ion suppression from the many salts in the sample. Thus, a reaction mixture was applied to a DEAE-Sephadex HCO$_3^-$ column (2 x 5 mm), and the column was washed with 0.5 mL of 0.1 M NH$_4$HCO$_3$ and the 6-hydroxymethylpterin-PP eluted with 0.5 mL of 0.4 M NH$_4$HCO$_3$. After evaporation of the NH$_4$HCO$_3$, LC-ESI-MS analysis of this sample showed the expected MH$^+$ ion at 354.1 m/z and (M – H)$^-$ ion at 352.1 m/z for 6-hydroxymethylpyrin-PP. The identity of the compound was further confirmed by the measurement of the MRM 354/125 and 354/176 fragments, the same as observed for the known sample of 6-hydroxymethylpyrin-PP.

The combination of genetic and biochemical data presented here strongly validates our comparative genomic derived predictions solving the long-standing mystery of 6-HMDP biosynthesis in most Archaea. We therefore renamed the two families MptD for the archaeal specific DHNA and MptE for the archaeal specific 6-HMDPK. The discovery of the missing archaeal 6-HMDPK synthesis genes completes the picture for the initial steps in the pterin pathways in the third kingdom of life (Figures 1 and 2). A great diversity in the metabolic and enzymatic solutions used to produce this molecule is observed, and the picture may be even more complex as some genes are still missing in specific lineages. MptE homologues are found in almost all archaeal genomes (Figure 2). The handful of organisms that lack this gene like Nanoarchaeum equitans, or Staphylothermus marinus have lost all other 6-HMDP biosynthesis genes47 and must certainly salvage pterin cofactors, even if one cannot rule out missing genes without further studies. For MptD, the situation is more complex. As discussed above, if the majority of archaeal genomes analyzed encode an MptD homologue, a minor subset does not. It seems most Thermococcales use a PTPS-III dependent bypass (Figures 1 and 2 and Supplemental Information), whereas Sulfolobus solfataricus uses a PTPS-VI dependent one.31 A few archaea species known to synthesize pterin containing C1-carrier coenzymes such as Archaeoglobus fulgidus48 or Pyrobaculum species49 lack homologues of MptD, FolB, PTPS-III, or PTPS-VI. A few of these (mainly Thermoproteales) encode members of the PTPS-V family (Figure 2), but initial validation tests with the Pyrobaculum caldifontis PTPS-V encoding gene, Pcal_1063, were negative.31 Some archaeal genomes such as M. barkeri encode both MptD and PTPS-III homologues and others such as S. solfataricus encode both FolB and PTPS-VI homologues (Figure 2). Of note, these organisms synthesize both H$_4$-folate and H$_4$-MPT derivatives or hybrid molecules.13,15,49 There are differences between the uses H$_4$-folate and H$_4$-MPT as C1 donors since H$_4$-MPT derivatives do not form N$^{15}$-formyl derivatives as a result of thermodynamic differences in the chemical properties of the arylamine nitrogen N$^{15}$ in H$_4$-folate and H$_4$-MPT. In organisms that use both cofactors, it might be necessary to use two different gene families in order to independently control the production of these molecules.

The discovery of MptD and MptE provides new examples of both divergent and convergent enzyme evolution.50 Further characterization of these enzymes therefore will be of interest for structural biologists and biochemists. Finally, folate biosynthesis genes are traditional antibacterial targets and recently methanopterin biosynthesis has been proposed as a target to eliminate the dominant archaeon in the human gut Methanobrevibacter smithii.51 Both MptD and MptE represent new targets found neither in human nor other members of the bacterial flora. As new roles of the gut Euryarchaeota emerge,52
inhibiting methanopterin pathway enzymes might be a viable solution to selectively eliminate Archaea from the flora.

**METHODS**

Bioinformatics. Analysis of the phylogenetic distribution and physical clustering was performed in the SEED database on the 58 genomes available at the time of the analysis (Dec/2011). Results are available in the "Pterin Biosynthesis Archaea" subsystem on the public SEED server (http://pubseed.theseed.org/SubsysEditor.cgi). A subset of the analysis is summarized Figure 2. We also used the BLAST tools and resources at NCBI.53 Multiple sequence alignments were built using the ClustalW tool.54 Structure based alignments were performed using the Espript platform (http://espript.ibcp.fr/ESPript/ESPript/).55 Visualization and comparison of protein structures and manual docking of ligand molecules were performed using PyMOL (The PyMOL Molecular Graphics System, Version 1.4.1, Schrödinger, LLC).

**Chemicals.** 7,8-Dihydroxynopterin, 6-hydroxymethylpterin-monophosphate (6-hydroxymethylpterin-P), 6-hydroxymethyl-P, 6-hydroxymethyl-7,8-dihydroxypterin, 6-hydroxymethyl-7,8-dihydroxypterin, P, 6-hydroxypterin-7,8-dihydroxypterin-PP, and D-neopterin were supplied by Sigma. CTP, UTP, and 6-hydroxymethylpterin was supplied by Schircks Laboratories, Jona, Switzerland. ATP, GTP, GPN, 6-hydroxymethyl-7,8-dihydropterin-PP, and D-neopterin were shown in Supplemental Figure 4 for MJ1634 as prototypical of the enzymes described herein. Protein concentrations were determined by Bradford analysis.

**Cloning of the P. furiosus pf0930 and pf1278 Genes and Expression of Their Gene Products.** The recombinant plasmids pPF0930 and pPF1278 were constructed as described in Sugar et al.54 Briefly, the primers used for pf0930 are (5′-GTTGACATTCTGTTGAAGAGTTGTTGTTATGT-3′) and (5′-CAGTGGTAAGGTCAAGGTTTGA-3′) for MJ1634 and (pf0930Fwd (5′-GCTGGATCCTTATTTTAAAAATTCAATCTC-3′) and PF1278Rev (5′-AAGCTCGAGGGCGGGCATTTGAAGCTTGAATAC-3′) and PF0930Rev (5′-AAGCTCGAGGGCGGGCATTTGAAGCTTGAATAC-3′) and PF1278Rev (5′-AAGCTCGAGGGCGGGCATTTGAAGCTTGAATAC-3′) and PF1278Rev (5′-AAGCTCGAGGGCGGGCATTTGAAGCTTGAATAC-3′) and PF1278Rev (5′-AAGCTCGAGGGCGGGCATTTGAAGCTTGAATAC-3′). These primers were used to amplify the genes by PCR and cloned into a modified pET24 vector using BamHI and NotI restriction enzymes. Proteins encoded by these plasmids were expressed and purified as described.58

**Enzymatic Assay of DHNA Activity.** The standard assay used for the measurement of DHNA enzymatic activity was conducted in 200 μL reaction volume and included 7 ng of M. jannaschii MJ0408, 40 mM TES/KCl buffer pH 7.4, 8 mM MgCl₂, 16 mM DTT, and 110 μM H₃Neo. Samples were sealed under argon and incubated for 10 min at 70 °C. Following incubation, the reactions were quenched by the addition of 20 μL of 1 M NaOH and excess iodine removed by reduction with 8 μL of 1 M NaHSO₃. Following centrifugation, the samples were combined with water for a final volume of 1 mL and analyzed by HPLC as described below.

Enzymatic Assay of 6-HMDPK Activity. The standard assay used for the measurement of 6-HMDPK enzymatic activity was conducted in 200 μL reaction volume and included 3.5 ng of the PF0930 (or MJ1634) enzyme, 40 mM TES/KCl buffer pH 7.4, 8 mM MgCl₂, 16 mM DTT, 100 μM 6-HMD, and 1 mM ATP. Samples were sealed under argon and incubated for 10 min at 70 °C. Following incubation, the reactions were quenched by the addition of 20 μL of 1 M HCl. 6-HMD and 6-HMDP in the incubation mixture were oxidized to fluorescent neopterin and 6-hydroxymethylpterin by the addition of 8 μL of a saturated solution of iodine in methanol and incubated at RT for 30 min. Following oxidation, the samples were neutralized by the addition of 20 μL of 1 M NaOH and excess iodine removed by reduction with 8 μL of 1 M NaHSO₃. Following centrifugation, the samples were combined with water for a final volume of 1 mL and analyzed by HPLC as described below.

HPLC Analysis of Pterins. Chromatographic separation of pterins was performed on a Shimadzu HPLC System with a C18 reverse phase column (Varian PursuitXR 250 × 4.6 mm, 5 μm particle size). The elution profile consisted of 5 min at 95% sodium acetate buffer (25 mM, pH 6.0, 0.02% NaN₃) and 5% MeOH followed by a linear gradient to 20% sodium acetate buffer/80% MeOH over 40 min at 0.5
mL/min. Pterins were detected by fluorescence using an excitation wavelength of 356 nm and an emission wavelength of 450 nm. Under these conditions, pterins were eluted in the following order (min): 6-hydroxymethylpterin-PPP (4.982), 6-hydroxymethylpterin-PP (5.243), 6-hydroxymethylpterin-P (6.605), D-neopterin (10.10), monapterin (12.012), and 6-hydroxymethylpterin (16.490). Alternately, pterins were separated on a Varian Pursuit polyfluorophenyl (PF) column (250 × 4.6 mm, 5 μm particle size). The elution profile was isocratic with 95% formic acid in water (0.1%) and 5% MeOH. Pterins were detected by fluorescence using an excitation wavelength of 356 nm and an emission wavelength of 450 nm.

**ASSOCIATED CONTENT**

Supporting Information

- Ezymatic and HPLC assays. This material is available free of charge via the Internet at http://pubs.acs.org.

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**Notes**
The authors declare no competing financial interest.

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