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Functional Promiscuity of the COG0720 Family

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Abstract

The biosynthesis of GTP derived metabolites such as tetrahydrofolate (THF), biopterin (BH₄), and the modified tRNA nucleosides queuosine (Q) and archaeosine (G⁺) relies on several enzymes of the Tunnel-fold superfamily. A subset of these proteins include the 6-pyruvoyl-tetrahydropterin (PTPS-II), PTPS-III, and PTPS-I homologs, all members of the COG0720 family, that have been previously shown to transform 7,8-dihydroneopterin triphosphate (H₂NTP) into different products. PTPS-II catalyzes the formation of 6-pyruvoyltetrahydropterin in the BH₄ pathway. PTPS-III catalyzes the formation of 6-hydroxymethyl-7,8-dihydropterin in the THF pathway. PTPS-I catalyzes the formation of 6-carboxy-5,6,7,8-tetrahydropterin in the Q pathway. Genes of these three enzyme families are often misannotated as they are difficult to differentiate by sequence similarity alone. Using a combination of physical clustering, signature motif, and phylogenetic co-distribution analyses, *in vivo* complementation studies, and *in vitro* enzymatic assays, a complete reannotation of the COG0720 family was performed in prokaryotes. Notably, this work identified and experimentally validated dual function PTPS-I/III enzymes involved in both THF and Q biosynthesis. Both *in vivo* and *in vitro* analyses showed that the PTPS-I family could tolerate a translation of the active site cysteine and was inherently promiscuous, catalyzing different reactions on the same substrate, or the same reaction on different substrates. Finally, the analysis and experimental validation of several archaeal COG0720 members confirmed the role of PTPS-I in archaeosine biosynthesis, and resulted in the identification PTPS-III enzymes with variant signature sequences in *Sulfolobus* species. This study reveals an expanded versatility of the COG0720 family members and illustrates that for certain protein families, extensive comparative genomic analysis beyond homology is required to correctly predict function.

Keywords

Queuosine; archaeosine; tetrahydrofolate; biopterin; tRNA modification; riboflavin; 6-pyruvoyl-tetrahydropterin synthase

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GTP is a precursor of RNA, DNA, and a number of other fundamental metabolites. Among these are riboflavin and the deazaflavin derivatives related to F₄₂₀, the pterin related coenzymes tetrahydrobiopterin (BH₄), tetrahydrofolate (THF), methanopterin, and molybdopterin, and a variety of 7-deazaguanine derivatives such as queuosine (Q) and archaeosine (G⁺), found in tRNA, and toyacamycin and tubercidin which are secondary metabolites produced in *Streptomyces*.

Many of the enzymes involved in the synthesis of these GTP derived metabolites are members of the same structural superfamily, the Tunnel-fold or T-fold superfamily (1). This superfamily is comprised of a functionally diverse set of enzymes that assemble through oligomerization of a core domain comprised of a pair of two-stranded anti-parallel β -sheets and two helices to form a $\beta_{2n}\alpha_n$ barrel (1). Two barrels associate in a head-to-head fashion, and bind planar substrates such as purines or pterins at the interface using a conserved Glu/Gln residue to anchor the substrate. Illustrating the diversity of the reactions catalyzed by T-fold enzymes, two enzymes of the BH₄ synthesis pathway (Figure 1A) belong to the T-fold superfamily GTP cyclohydrolase IA (GCYH-IA or FolE) and 6-pyruvoyl-tetrahydropterin synthase (PTPS-II or PtpS) (2); GCYH-IA catalyzes the first step of the pathway producing 7,8-dihydroneopterin triphosphate (H₂NTP) from GTP (3, 4). H₂NTP is then converted to 6-pyruvoyl-tetrahydropterin (PTP) by PTPS-II (5, 6) (Figure 1A). PTP is then reduced to BH₄ by sepiapterin reductase (SR encoded by the *spr* gene and part of the dehydrogenase-reductase (SDR) superfamily) (7–9).

GCYH-IA is also the first enzyme of the THF biosynthetic pathway (3). It is replaced in some organisms by GTP cyclohydrolase IB (GCYH-IB or FolE2) (10), another T-fold enzyme (11). In most plants and Bacteria the THF pathway contains a second T-fold enzyme, dihydroneopterin aldolase (DHNA) encoded in *Escherichia coli* by *folB* (12) (Figure 1A). furthermore, in *Plasmodium falciparum* and various bacteria, the DHNA step is bypassed by yet another T-fold enzyme, PTPS-III, a homolog of PTPS-II, that cleaves the side chain of H₂NTP to form 6-hydroxymethyl-7,8-dihydropterin (6HMDP) (13–15) (Figure 1A).

Queuosine (Q) is a 7-deazaguanosine derivative found at position 34 of several bacterial and eukaryal tRNAs (16–18), while archaeosine (G⁺), a related derivative, is found specifically at position 15 of archaeal tRNA (19). Like the flavin and folate pathways, the Q/G⁺ pathways are populated by T-fold enzymes and GCYH-IA (or GCYH-IB) catalyzes the first biosynthetic steps (20). The second enzyme of the pathway, PTPS-I or QueD, is homologous to PTPS-II and catalyzes the formation of 6-carboxy-5,6,7,8-tetrahydropterin from DHNTP *in vitro* (21) (Figure 1A). Finally, the enzyme QueF, an oxidoreductase that reduces the nitrile side chain of 7-cyano-7-deazaguanine (preQ₀) the last common intermediate in the Q and G⁺ pathways (22, 23), to the aminomethyl side chain of 7-aminomethyl-7-deazaguanine (preQ₁), is also a T-fold enzyme (24).

Functional diversity is found not only between the different T-fold sub-families but also within a given subfamily. As alluded to above, three members of the COG0720 subfamily, PTPS-I, II, and III have been shown to catalyze different reactions in different pathways (Figure 1A), and a fourth COG0720 member, PTPS-IV, whose structure was recently determined, has an as yet unknown function (25).

By combining comparative genomic with biochemical and genetic characterization, we provide evidence that the COG0720 family is an example of a family of enzyme containing functionally promiscuous members. This functional promiscuity is exploited *in vivo* with single enzymes contributing different reactions to different pathways.

RESULTS

Separation of Four COG0720 Subfamilies by Comparative Genomics

Because of its functional diversity, the COG0720 family of enzymes is particularly difficult to annotate. Out of 810 bacterial COG0720 sequences in the NCBI database as of July 2010, 516 are annotated in RefSeq (26) as 6-pyruvoyltetrahydropterin synthase, or PTPS-II. However, with the exception of specific cyanobacteria that synthesize glycosylated BH₄ derivatives (27), the BH₄ pathway is absent in most of these organisms, and thus these enzymes likely have activities other than that of a PTPS-II. To illustrate the difficulty of annotating COG0720 members using sequence similarity (BLAST score) alone, the rat PTPS-II protein (NP_058916.1) was used as input to search the *Synechococcus* sp. PCC7942 genome using default BLASTP parameters (28), and two COG0720 proteins were retrieved. The one with the lowest similarity (YP_400201.1; E-value: 5e-20) has robust canonical PTPS-II activity *in vitro*, whereas the one with the highest similarity (YP_400970.1; E-value 6e-31) exhibits only low activity, and deletion of the corresponding gene does not affect BH₄ levels *in vivo* (27). Transferring the function of the experimentally characterized member of the family, the rat PTPS-II, to the best scoring homolog in *Synechococcus* sp. PCC7942 therefore results in an erroneous annotation. In order to better annotate the COG0720 family other types of association evidence were required. Using the SEED database (29), we performed a comparative genomic analysis of the four PTPS families (I to IV). A subsystem named “Experimental-PTPS” was constructed that included all the COG0720 homologs in the database. More than one copy of the gene was found in 114 of the 918 genomes analyzed, confirming that the misannotation risk is indeed very high.

Physical clustering analysis revealed that specific members of the COG0720 subfamilies could be efficiently separated by analyzing their genomic context. Twenty four COG0720 genes cluster with other genes of the BH₄ pathway such as *folE* and *spr* (Figure 1B and see “Experimental PTPS” subsystem in SEED) and hence were annotated as *ptpS* genes encoding PTPS-II enzymes. Another 283 COG0720 genes cluster with queuosine genes (*queCEF*) (30) and *folE* or *folE2* (20) (Figure 1B and See “Experimental PTPS” subsystem in SEED) and were therefore annotated as *queD* encoding PTPS-I enzymes. Finally, 16 COG0720 genes cluster with folate biosynthesis genes such as *folE*, *folK*, and *folP* (Figure 1B and see “Experimental PTPS” subsystem in SEED) and were annotated as encoding PTPS-III enzymes.

We derived signature motifs for these three subfamilies of enzymes using the PRATT tool from PROSITE suite (31) as well as Web Logo 3.0 (32). The sequences of the experimentally characterized enzymes, as well as sequences from additional members predicted through physical clustering, were used as input for each COG0720 subgroup. Previous sequence and structural analysis of the PTPS-III family had shown that it could be distinguished from the PTPS-II family by the presence of specific motifs surrounding the catalytic residues (13); {CX(5)-H-G-H} for PTPS-II enzymes, and {E-X(2)-H-G-H} for PTPS-III enzymes. We confirmed and slightly expanded these two signature motifs and identified the signature motif for the PTPS-I family as {C-X(3)-H-G-H} (Supplemental Figure 1).

PTPS-IV encoding genes are found only in a few halophilic Archaea and Actinomycetes (a total of 14 organisms), and members of the PTPS-IV family are still of unknown function, although physical clustering suggests a link with riboflavin. Indeed, in Archaea PTPS-IV genes cluster with GTP-cyclohydrolase III (GCYH-III) genes (*arfA*) (33) and in Bacteria with GTP-cyclohydrolase II genes (*ribA2*) that contain mutations conferring GTP-cyclohydrolase III type activity (34), and in both they further cluster with a formamide

hydrolase gene (*arfB*) that encodes the subsequent enzyme in this GCYH-III dependent riboflavin pathway (35). We therefore predict that PTPS-IV proteins are involved in the synthesis of a product derived from 2,5-diamino-6'-riboseamino-4(3H)-pyrimidinone 5'-phosphate (APy), the riboflavin/F₄₂₀ biosynthetic intermediate generated by the ArfA and ArfB proteins. A PTPS-IV signature motif, {F-X(0,1)-G-X-[ANTV]-[NPQST]} was identified using the 14 PTPS-IV sequences as input (Figure 1 and Figure S1). Finally, two groups of COG0720 proteins (PTPS-V and VI) that do not contain any of the motifs identified in the PTPS-I/II/III/IV families were found in Crenarchaea (listed in Supplemental data 1). The PTPS-V members are found in all sequenced *Pyrobaculum* sp., in *Vulcanisaeta* sp. and in *Thermoproteus neutrophilus* and contain a {S-X(2)-W-X(3)-H-G-H} motif. The PTPS-VI members are found in *Metallosphaera sedula* DSM 5348 and in all sequenced *Sulfolobus* species and contain a {S-S-X(4)-Q-X-H-G-H} motif. The functions of these last two families of COG0720 proteins are unknown.

The combination of physical clustering and motif analysis allowed us to differentiate six COG0720 subfamilies and derive signature motifs that were used to propagate the annotations in the SEED database (see “Experimental-PTPS” subsystem).

PTPS-I/III Enzymes Can Function in Both Folate and Q Pathways

Genomic analysis revealed that a group of bacteria (Supplemental Table 1) contained only one COG0720 encoding gene but were predicted to require both PTPS-III and PTPS-I activities as they possessed Q biosynthetic genes (*queCEF*) as well as the signature folate genes *folK* and *folP* (examples are shown in Figure 2A), but lacked *folB* (encoding DHNA). Closer analysis of the COG0720 proteins encoded in these organisms revealed that they contained a hybrid PTPS-I/PTPS-III motif {C-E-X-[ILPV]-H-G-H} (Supplemental Table 1). We had previously shown that the predicted PTPS-I/III_{Sa} enzyme from *Syntrophus acidotrophicus* (YP_462286.1) exhibited PTPS-III activity *in vivo* as the corresponding gene complemented the thymidine (dT) auxotrophy of an *E. coli* Δ *folB* strain (15). However, physical clustering linked the corresponding gene to the Q biosynthesis pathway (Figure 1B). More generally, out of 38 genes encoding proteins containing the dual motif, 7 of them cluster with folate biosynthesis genes and 14 of them cluster with Q biosynthesis genes (See “Experimental PTPS” subsystem in SEED). To test whether the PTPS-III proteins containing hybrid motifs also exhibited PTPS-I activity, we examined the nucleoside constituents of bulk tRNA purified from cultures of WT *E. coli* transformed with pBAD24 (VDC3339) and of Δ *queD* strains transformed with pBAD24 (VDC3321) or with derivatives expressing the *PTPS-I/III*_{Sa} gene (VDC3335) or the *PTPS-I/III*_{Sa}Cys26Ala gene (VDC3365). Bulk tRNAs were enzymatically hydrolyzed, dephosphorylated, and the ribonucleosides analyzed by LC/MS-MS as described previously (20). The 410 *m/z* ion that corresponds to the molecular ion (MH⁺) of Q was detected by MS at 20.31 min in the WT background, while no 410 *m/z* ion was detected in the Δ *queD* strain (Figure 2B). Expression of the *PTPS-I/III*_{Sa} gene complemented the Q⁻ phenotype of the *E. coli* Δ *queD* mutant (Figure 2B). Mutating Cys26 of the {C-E-X-[ILPV]-H-G-H} motif to Ala in the *S. acidotrophicus* protein abolished complementation of the Q⁻ phenotype but not the dT auxotrophy phenotype (Table 1). Expressing a canonical *PTPS-III*_{Li} gene from *Leptospira interrogans* (NP_712930.1) did not lead to complementation of the Q⁻ phenotype (Table 1), whereas the same clone was effective in complementing the dT auxotrophy phenotype of the *folB* strain (15). In addition, the *PTPS-I*_{Cb} (YP_0011383898.1) and *PTPS-I/III*_{Cb} (YP_0011383205.1) genes from *Clostridium botulinum* strain 19397 were analyzed in both complementation tests. In this organism, the *PTPS-I*_{Cb} gene clusters with Q biosynthesis genes and the *PTPS-I/III*_{Cb} gene clusters with folate biosynthesis genes (Figure 1B). Both complemented the Q⁻ phenotype and thus were active as PTPS-I enzymes (Table 1). Only *PTPS-I/III*_{Cb} complemented the dT auxotrophy phenotype, and thus exhibited PTPS-III

activity (Figure 2C). These results show that COG0720 members that contain hybrid PTPS-III/I motifs are active in both folate and Q biosynthesis pathways and that the conserved cysteine in that motif is critical for PTPS-I activity but not PTPS-III activity.

Role of COG0720 Family Proteins in Archaea

All Euryarchaeota predicted to synthesize preQ₀ because of the presence of the *queCE* genes encode a PTPS-I like protein (See “Experimental PTPS” subsystem in SEED). PTPS-I genes are surprisingly absent in many Crenarchaea, some of which are known to make G⁺ (36), although many have the PTPS-V or PTPS-VI variant. Finally, several archaeal genomes encode more than one COG0720 protein (Experimental data 1), such as *Haloferax volcanii*, which contains both putative *PTPS-I* (*HVO_1718*) and *PTPS-IV* genes (*HVO_1282*).

Because *H. volcanii* is known to contain G⁺-modified tRNA (37) and many of the genes encoding the biosynthetic enzymes have been identified (*folE2*, *HVO_2348*; *queC*, *HVO_1717* and *queE*, *HVO_1716*) and shown to be essential for G⁺ formation (20, 38), it was logical to propose that *HVO_1718* encodes the QueD/PTPS-I protein involved in G⁺ synthesis. A Δ *HVO_1718* derivative was constructed as described in the methods section. Subsequent analysis of ribonucleosides from bulk tRNA extracted from the *H. volcanii* WT and Δ *HVO_1718* strains grown in minimal medium showed that the peak observed for the molecular ion of G⁺ (325 *m/z*), corresponding to the 26.2 min peak detected on the UV trace present in the WT *H. volcanii* H26 strain, was absent in the mutant strain (Figure 3A). Expression of the PTPS-I gene from *Halobacterium* sp. NRC1 (*Vng6306*) on the shuttle vector pJAM202 complemented the G⁺ deficiency phenotype of the Δ *HVO_1718* strain (Supplemental Figure 2) clearly establishing *HVO_1718* as encoding a functional PTPS-I.

The function of the PTPS-IV protein in *H. volcanii* is less clear. *H. volcanii* is rare among Archaea in having a complete folate pathway (39, 40), but genes encoding FolB or PTPS-III could not be identified in this organism (41). One possibility is that while PTPS-IV lacks the PTPS-III signature motif, it functionally replaces FolB. To test this hypothesis a Δ *HVO_1282* strain was constructed, but unlike the *H. volcanii* Δ *folE2* mutant previously constructed (42) it did not require dT for growth, suggesting that PTPS-IV is not involved in folate biosynthesis (Figure 3B). Physical clustering suggested a possible link between PTPS-IV and riboflavin biosynthesis (Figure 1B). Some of the canonical riboflavin biosynthetic genes are known to be missing in Archaea (35), thus we tested the involvement of *HVO_1282* in riboflavin biosynthesis. As a control, we constructed a *H. volcanii* Δ *ribA* deletion strain (Δ *HVO_1284*); as shown on Figure 3C, no growth defect in the absence of riboflavin was observed in the Δ *HVO_1282* strain, whereas the Δ *HVO_1284* strain required riboflavin to grow as expected. At this point however, even if comparative genomic analysis links the PTPS-IV family to the synthesis of an APy derivative, the family does not appear to be directly involved in the synthesis of riboflavin.

Finally, we investigated whether the variant COG0720 members found in *Sulfolobus solfataricus* or in *Pyrobaculum calidifontis* had QueD or FolB activity in *E. coli* complementation tests. We found that the *SSO2412* gene (PTPS-VI family) could complement the dT auxotrophy of the *folB* mutant (Figure 3D) but not the Q deficiency of the *queD* strain (Table 1). We failed to see any complementation of either phenotype by the *Pcal_1063* gene (PTPS-V family) in the conditions tested.

In Vitro and *In Vivo* Flexibility of the Bacterial and Archaeal PTPS-I Catalytic Site

An exhaustive comparative analysis of the Zur regulon (43) revealed that certain bacteria contained two copies of the *queD/PTPS-I* gene, *queD* and *queD2*, with *queD2* predicted to

be under the control of negative regulator Zur (Supplemental Table 2). Zur is a repressor that responds to zinc levels and upregulates genes under its control when zinc is low (44). These include genes encoding high-affinity zinc transporters such as ZnuABC, and paralogs of zinc dependent enzymes that can use metals other than zinc (45). Analysis of the QueD2 sequences showed that these harbored a {C-X(4)-H-G-H} motif instead of the {C-X(3)-H-G-H} found in canonical QueD/PTPS-I enzymes (Supplemental Table 2). Furthermore, certain organisms, such as *Acinetobacter baylyi* sp. ADP1, contain only a *queD2* gene and no *queD* gene (Supplemental Table 2). To test whether QueD2 proteins were functional PTPS-I enzymes we expressed the *A. baylyi* sp. ADP1 *queD2_{Ab}* gene (YP_046954.1) in the *E. coli* Δ *queD* strain (VDC4660). As shown in Table 1, complementation of the Q⁻ phenotype was observed, confirming QueD2 does possess PTPS-I activity. Interestingly, introducing the Lys23Cys and Cys24Ser mutations in the *E. coli* QueD protein, thereby changing the {KC-X(3)-H-G-H} motif to a {C-X(4)-H-G-H} motif, also allowed functional complementation (Table 1), suggesting that the PTPS-I catalytic pocket is quite plastic. To probe this idea further, we tested if PTPS-II enzymes, with {C-X(5)-H-G-H} signature motifs, also exhibited PTPS-I activity. Previous studies had shown that PTPS-I from *Synechococcus* sp. PCC7942 did possess PTPS-II activity *in vitro* (albeit only 10% of the activity of the canonical PTPS-II from the same organism) (27), but the reverse scenario had never been tested. As shown in Figure 2B and Table 1, expressing the rat *ptpS_{Rn}* gene in the *E. coli* Δ *queD* mutant (strain VDC3331) did restore the production of Q, demonstrating that the rat PTPS-II exhibited enough PTPS-I activity to functionally replace the chromosomal encoded *queD*, at least when expressed on a multicopy plasmid. Finally, we tested whether a motif containing a shortened spacer {C-X(2)-H-G-H} still led to a functional PTPS-I enzyme by transforming the *E. coli* Δ *queD* and Δ *folB* strains with a plasmid expressing a mutant PTPS-I/III_{5a} enzyme possessing the Cys26Ala and Glu27Cys double mutation. Like the Cys26Ala single mutant, this clone failed to complement the Q⁻ phenotype, but unlike the single mutant, it also failed to complement the dT auxotrophy (Table 1).

The plasticity of the PTPS active-site had already been suggested by the report that the *E. coli* PTPS-I/QueD enzyme displayed relaxed substrate selectivity *in vitro*, utilizing H₂NTP or sepiapterin to form 6-pyruvoyl-tetrahydropterin (PTPS-II activity) or 7,8-dihydropterin, respectively, the latter a fundamentally new activity of side-chain cleavage (21, 46). Using a direct *in vitro* assay for activity that eliminates potential artifacts introduced through post-reaction work-up, the activity of the *E. coli* enzyme was subsequently shown to produce only 6-carboxy-5,6,7,8-tetrahydropterin (CPH₄) (21), the biologically relevant product. We show here that this enzyme is able to produce CPH₄ from several substrates, including H₂NTP, sepiapterin, and dihydroneopterin monophosphate (H₂NMP) (Figure 4A–E). Collectively these results demonstrate that the *E. coli* QueD enzyme is able to catalyze the formation of a common product from multiple substrates.

To test if other members of the PTPS-I family shared this substrate plasticity, we investigated the catalytic activity of the predicted archaeal PTPS-I enzyme from *Methanocaldococcus jannaschii*, MJ1272. We first confirmed that MJ1272 was a member of this family as it catalyzed the formation of CPH₄ from the expected biological substrate, dihydroneopterin-2',3'-cyclic phosphate (H₂NMPc) (47) (Figure 4F). The generation of CPH₄ in these reactions was established by detection of two of its oxidative decomposition products, 6-carboxypterin and pterin. The pterin was labeled with deuterium when the incubation was conducted in deuterated water, consistent with the incorporation of deuterium at C-6 during retroaldol reaction leading to the formation of CPH₄. This was retained and measured in the pterin produced by its subsequent oxidative decomposition. Furthermore, MJ1272 also exhibited good activity with a variety of other pterin substrates, including dihydroneopterin, sepiapterin and H₂NMP (as shown on Figure 4F). These results

demonstrate that, like the *E.coli* QueD, MJ1272 can catalyze the formation of a common product from multiple substrate pterins.

Structural Analysis of the COG0720 Family

The PTPS-II family has been the most thoroughly investigated structurally, and a detailed chemical mechanism has been proposed (48). The 3D structure of the rat liver PTPS-II exhibits a homohexameric structure formed by a dimer of trimers with 3-fold symmetry (5). Based on molecular modeling, site-directed mutagenesis, and refined crystal structures of the enzyme alone and in complex with natural substrate, it was shown that substrate binding occurs at the interface of 3 subunits, comprising two subunits from one trimer and one subunit from the opposing trimer (48). Because of this structure, catalytically important residues are contributed from multiple subunits. For example, the active site residue Cys42 (*Rattus norvegicus* PTPS-II numbering) is contributed from a subunit of one trimer, while Asp88 and His89, which complete the catalytic triad with Cys42, and the Zn²⁺ binding residues His23, His48, and His50 are contributed by a subunit in the adjacent trimer (48, 49). The proposed reaction involves base-catalyzed isomerization followed by triphosphate elimination (48, 50), with Zn²⁺ serving to activate the substrate and stabilize the intermediates so as to disfavor breaking of the C1–C2 bond in the pyruvoyl side-chain (48).

To gain insight into the structural basis for plasticity of the PTPS active sites, the structure of rat PTPS-II that produces 6-pyruvoyltetrahydropterin (PDB: 1B66) was compared with the structure of the *Pseudomonas aeruginosa* PTPS-I that produces CPH₄ (PDB: 2OBA) using the Accelrys Discovery Studio 2.5 tool. Structural superimposition was made based on monomers of each protein structure. The best global fit was obtained when using the PTPS-II structure from *R. norvegicus* as a reference with the following three homology regions as anchoring sites: the region from Thr66 to Gly67, the region from Thr105 to Glu107, and the third from Glu133 to Tyr134 (*R. norvegicus* numbering) (Figure 5A). Overall, the two structures superimpose with a rmsd of 0.4. The spatial locations of the canonical T-fold Glu residues, which interact with the exocyclic 2-amino group of the pterin, are conserved in both structures (Glu 133/Glu107) (48). The catalytic residues Asp88 and His89 (*R. norvegicus* numbering) of the active site are superimposable with the catalytic residues Asp67 and His68 of the *P. aeruginosa* PTPS-I (PDB: 2OBA). The Zn²⁺ binding His residues of PTPS-II (His23, His48 and His50) (48) occupy the same coordinates as the Zn²⁺ binding residues (His13, His28 and His30) of PTPS-I (Figure 5B). Although the catalytic Cys residues involved in catalysis do not completely overlap, the distances from Cys to Zn²⁺ are similar in both enzymes (Figure 5 Table inset). Despite these shared active site characteristics, the *R. norvegicus* PTPS-II has two extra helix domains around the active-site region, one found on the N-terminal side adjacent to Cys42 and the other around Asp88/His89 (Figure 5A).

The superimposition of the PTPS-I from *P. aeruginosa* (PDB: 2OBA chainA) and PTPS-III from *Plasmodium falciparum* (PDB: 1Y13 chain A) performed with the rigid FATCAT algorithm showed that the two structures superimpose with a rmsd of 2.21 (Figure 5C). As was observed in the comparison of the PTPS-I and PTPS-II structures, most of the conserved catalytic residues overlap well; thus, Asp67 and His68 (in PTPS-I overlap with the Asp79 and His80 in PTPS-III, the spatial coordinates of the three Zn²⁺ binding His residues are conserved in both structures, and the spatial locations of the canonical T-fold Glu residues are conserved in both structures (Glu107/Glu161) (Figure 5C and D). Furthermore, while the key catalytic residue Cys24 in PTPS-I is replaced by Glu38 in the active site of PTPS-III (Figure 5D), the distances of the side chain groups of Cys24 and Glu38 to the respective Zn²⁺ are similar, thus the spatial positions of the two putative general acid/base residues with respect to Zn²⁺ are comparable in the two enzymes (Figure 5 Table inset). However, there are noteworthy differences between the two structures; the

loops containing the active site do not overlap completely, and PTPS-III has an extra loop close to the active site region (Figure 5C). Despite the active sites similarities among the three PTPS enzymes, the notable differences around the active sites could influence substrate binding allowing the production of different products.

DISCUSSION

The combination of comparative genomic analyses and experimental validation allowed separation of the COG0720 family into at least six functional subfamilies (PTPS-I through PTPS-VI), illustrating the diversity of its catalytic potential. The subfamilies have similar but distinct catalytic activities and are involved in different pathways, three of which have been identified: folate biosynthesis for PTPS-III, Q/G⁺ biosynthesis for PTPS-I, and biopterin biosynthesis for PTPS-II. Of particular interest is the observation that some members of the COG0720 family have both PTPS-I and PTPS-III activity *in vivo*, and that these proteins harbor a hybrid catalytic motif {C-E-X(2)-H-G-H}. This is an example where catalytic promiscuity of a single enzyme satisfies different metabolic needs in different pathways. An example of this phenomenon is observed with *o*-succinylbenzoate synthase (OSBS), a member of the enolase superfamily, which functions both as succinylbenzoate synthase in menaquinone biosynthetic pathway and as an N-succinylamino acid racemase (NSAR) in racemization of N-acetylmethionine (51, 52). Only an amino acid change (changing {C-X(3)-H-G-H} to {C-E-X(2)-H-G-H}) is necessary to transform a PTPS-I into a PTPS-I/III (15) and only one additional mutation (the Cys to another amino acid) is required to transform a PTPS-I/III to a PTPS-III. This latter phenomenon, in which the identity of a single residue can change the predominant product of the enzyme, is similar to observations of the GCYH II/III enzymes, where a single Tyr or Met residue at a key position results in an enzyme with type-II or type-III activity, respectively (34).

While PTPS-I is able to accept a variety of substrates *in vitro*, the actual biological substrate for PTPS-I enzymes remains uncertain. In Archaea the product of GCYH-I is H₂NMPc (47), while in Bacteria it is H₂NTP, so it is possible that both are further processed by a phosphatase to form H₂NMP as the relevant substrate *in vivo*. Further work is needed to differentiate between these two possibilities.

The functions the COG0720 family members might also be dependent on the identity of the bound metal. PTPS-II enzymes are thought to be zinc dependent (49), and while the metal dependence of PTPS-III enzymes have not been rigorously determined, the structures of the *P. falciparum* (PDB: 1Y13) and *Plasmodium vivax* (PDB: 2A0S) proteins show a bound Zn²⁺. Interestingly, in the PTPS-I family the archaeal member MJ1272 appears to be Fe²⁺ dependent (see Supplemental material). This might be a methanogen specific trait as *M. jannaschii* FOLE2 (or MptA), another T-fold superfamily member, is also Fe²⁺ dependent (47) whereas in bacteria such as *B. subtilis* and *Neisseria gonorrhoeae* the enzyme is most active with Mn²⁺ (11). Finally, genes encoding QueD2 variants with a different active-site signature [{C-X(4)-H-G-H} instead of {C-X(3)-H-G-H}] were predicted to be under Zur regulation, consistent with the canonical QueD requiring zinc whereas the QueD2 enzymes use another metal. Several cases are now known in which paralogs of zinc dependent enzymes that use metals other than zinc are upregulated when zinc is low (see (53) for examples). Nevertheless, while the existing data is compelling, detailed investigation of the metal dependence of these enzymes will be required to determine if varied metal dependence in fact contributes to the catalytic potential of COG0720 members.

Consideration of the transformations catalyzed by PTPS-I, PTPS-II, and PTPS-III reveal that each can be explained by the involvement of general acid/base catalysis in mediating isomerization, elimination, retroaldol, cleavage/elimination, and hydration reactions (Figure

6). Thus PTPS-III requires only a retroaldol cleavage and isomerization to produce dihydro-6-hydroxymethylpterin, PTPS-II requires an isomerization followed by an elimination of triphosphate to produce 6-pyruvoyltetrahydropterin. The fact that PTPS-I can utilize sepiapterin as well as H₂NTP (and the di- and monophosphates) led to the proposal (21) of a mechanistically complex series of reactions in which the first three reactions are the same as that used by PTPS-II to produce PTP, followed by two isomerization reactions, a hydration and finally a retroaldol reaction to produce CPH₄. However, given the observed promiscuity of these enzymes it is possible that the activity with sepiapterin does not reflect an actual role in the reaction catalyzed with H₂NTP, as one can envision a much simpler route to CPH₄ in two steps *via* similar acid/base catalyzed transformations (Figure 6); differentiating these 2 pathways may be possible by evaluating the kinetic competence of sepiapterin once PTPS- I has been kinetically characterized with the dihydroneopterin substrates. Overall, the products of the various PTPS enzymes are all accessible *via* subtle changes in the expression of general acid/base catalysis operating in each of these systems.

Even after this extensive characterization of the COG0720 family many questions remain. The function of the PTPS-IV subfamily is still unknown: it was already known not to function as a PTPS-II (25). We have shown here that it possesses neither PTPS-III nor PTPS-I activity, and that it has no direct role in riboflavin synthesis. PTPS-I is missing in many Crenarchaea that make G⁺, such as *Sulfolobus* or *Pyrobaculum* species (19, 36, 54) (see “Experimental-PTPS” subsystem in SEED), and possible candidates for this activity are the PTPS-V and PTPS-VI families that are found in many Crenarchaea (Supplemental data 1). However, while expressing the PTPS-VI from *S. solfataricus* SSO2412 complemented an *E. coli folB* mutant, it did not complement a *queD* mutant, so it is unlikely that the PTPS-VI family harbors the missing PTPS-I activity in Crenarchaea. Notably, while the PTPS-VI enzymes exhibit PTPS-III activity (based on the *folB* complementation result), they lack the PTPS-III signature motif. Further biochemical work will be required to fully characterize this complex COG0720 family.

METHODS

Chemicals

All chemicals were obtained from Aldrich/Sigma unless otherwise indicated.

Bioinformatics

Analysis of the phylogenetic distribution and physical clustering was performed in the SEED database (29). Results are available in the “Experimental-PTPS” subsystem on the public SEED server (<http://theseed.uchicago.edu/FIG/subsys.cgi>). We also used the BLAST tools and resources at NCBI (28). The *H. volcanii* genome sequence was accessed through the UCSC archaeal genome browser (55). Multiple alignments were built using the ClustalW tool (56). The PRATT tool (31) from Prosite website (<http://expasy.org/prosite/>) was used to derive the specific protein motifs. These motifs were used to examine the COG0720 family using ScanProsite (57). The same Prosite motifs were also used to search for pattern occurrence in protein queries using Phi-BLAST at NCBI (58). Web logo (<http://weblogo.berkeley.edu/logo.cgi>) (32) was used to create sequence logos for the four PTPS families. Structure based alignments were performed using the Esript platform (<http://esript.ibcp.fr/ESPrpt/ESPrpt/>) (59).

Three Dimensional (3D) Structure Superimposition

First, the sequences of PTPS-I from *P. aeruginosa* (PDB: 2OBA) and PTPS-II from *R. norvegicus* (PDB: 1B6Z) were aligned using ClustalW2 (56) to find conserved regions along the aligned sequences that were used as anchoring points for the superimposition. The 3D

superimpositions were performed using the superimposition tool of the software “Discovery Studio 2.5” (<http://accelrys.com/>) (60, 61) and the superimposition tools of Protein Data Bank (PDB) (<http://www.pdb.org>).

Strains, Media, Growth, and Transformation

All strains and plasmids used are listed in Supplemental Table 3. *E. coli* derivatives were routinely grown at 37°C in LB (BD Diagnostic System) or minimal M9 medium (62) supplemented with 0.4% glycerol as a carbon source. Growth media were solidified with 15 g l⁻¹ agar (BD Diagnostic System) for the preparation of plates. Transformations of *E. coli* were performed following standard procedures (62) Ampicillin (Amp, 100 µg ml⁻¹), Thymidine (dT, 300 µM), Kanamycin (Kan, 50 µg/ml), isopropyl-beta-D-thiogalactopyranoside (IPTG, 1 mM) and L-arabinose (0.2%) were added when needed. *H. volcanii* derivatives were grown at 45 °C in Hv-YPC rich medium (63) and Hv-minimal medium (64). Riboflavin (20 µg ml⁻¹), uracil (50 or 10 µg ml⁻¹), novobiocin (Nov, 0.2 µg ml⁻¹) and 5-fluoroorotic acid (FOA, 50 µg ml⁻¹) were added when needed. *H. volcanii* growth media were solidified with 20 g l⁻¹ agar (BD Diagnostic System) for preparation of the plates. Transformations of *H. volcanii* were performed essentially as described (65).

Plasmid and Strain Constructions for Genetic Studies

All primers used for plasmid and strain construction, site directed mutagenesis and strain verification are listed in Supplemental Table 4. Polymerase chain reactions (PCRs) were performed using Phusion™ Hot Start (Finnzymes, Espoo, Finland) as previously described (66). The genomic DNA from *H. volcanii* and *Halobacterium* sp. NRC.1 were prepared as previously described (64). *E. coli* and *A. baylyi* sp. ADP1 genomic DNA were prepared as previously described (62).

The *ygcM* gene (NP_417245.1) was amplified from *E. coli* genomic DNA using primers *ygcM.ol1* and *ygcM.ol2* bearing *EcoRI* sites and cloned into pBAD24. The *PTPS-I2_{Ab}* gene (YP_046954.1) was amplified from *A. baylyi* genomic DNA using primers *QueDADP1_EcoRI_Fw* and *QueDADP1_XbaI_Re* and cloned into pBAD24 after digestion with the appropriate enzymes. The *PTPS-II_{Rn}* gene (NP_058916.1) was sub-cloned from pSTV28MPS (67) into pUC19 (68) using the *BamHI* and *EcoRI* restriction sites. The *VNG6306* gene (NP_395805.1) was amplified from the *Halobacterium* sp. NRC1 genomic DNA using primers *HsQueD_NdeI_Fw* and *HsQueD_BlpI_Rev* bearing *NdeI* and *BlpI* and cloned into pJAM202 (69) after digestion with appropriate enzymes. The *HVO_1282* (YP_003535334.1) gene was amplified from the *H. volcanii* DS70 genomic DNA using primers *HvPTPSIV_NdeI_Fw* and *HvPTPSIV_BlpI_Rev* bearing *NdeI* and *BlpI* and cloned into pJAM202 after digestion with the appropriate enzymes. The *SSO2412* (NP_353770.1) and *Pcal_1063* (YP_001055954.1) genes were amplified from genomic DNA from *S. solfataricus* and *P. calidifontis* respectively using primers *SsQueD2QHGH_Fw* and *SsQueD2QHGH_Rev* bearing *NcoI* and *BamHI* restriction sites for *SSO2412* and *PcQueD2WHGH_Fw* and *PcQueD2WHGH_Rev* bearing *NcoI* and *BamHI* restriction sites for *Pcal_1063*. The obtained PCR fragments were cloned into pBAD24 after digestion with appropriate enzymes.

Site directed mutagenesis was performed as described previously (66) using the specific oligonucleotides pairs listed in Supplemental Table 4. The resulting plasmids were verified by Sanger sequencing at the U of F core facility.

The $\Delta ygcM::Kan^r$ deletion was transferred by P1 transduction (70) from the *E. coli* JW2735-2 strain from the Keio collection (71) into *E. coli* K12 MG1655. The Kan^r marker was then excised using the F1p recombinase (72) to create the $\Delta ygcM$ strain (VDC2043).

The $\Delta folB::Kan^r$ deletion was transferred by P1 transduction (70) from the *E. coli* JW3030-2 strain from the Keio collection (71) into *E. coli* K12 MG1655 yielding strain VDC3276. Deletion of both *folB* and *ycgM* and excision of the Kan^r gene were confirmed by PCR. The *H. volcanii* ΔHVO_1718 deletion strain was constructed as described previously (42). The *H. volcanii* ΔHVO_1282 and ΔHVO_1284 strains were constructed as previously described (38). Deletions of the genes were checked by PCR as shown in Supplemental Figure 3.

Cloning and Expression of the *M. jannaschii* MJ1272 Gene

The *M. jannaschii* gene MJ1272 (NP_248268.1) was amplified by PCR from genomic DNA by using oligonucleotide primers. The primers used were MJ1272-Fwd and MJ1272-Rev. PCR amplification was performed using a 55°C as annealing temperature. The PCR product was purified by using a QIAquick spin column and digested with *Nde*I and *Bam*HI restriction enzymes and then ligated into compatible sites in plasmid pT7-7 to make the recombinant plasmid pMJ1272. DNA sequence was verified by dye-terminator sequencing at the DNA Facility of Iowa University. The resulting plasmid was transformed into *E. coli* strain BL21-Codon Plus (DE3)-RIL. The transformed cells were grown in Luria-Bertani medium supplemented with 100 $\mu\text{g mL}^{-1}$ Amp at 37°C with shaking until they reached an OD₆₀₀ of 1.0. Recombinant protein production was induced by addition of lactose to a final concentration of 28 mM. After an additional 2 hours of culture, the cells were harvested by centrifugation (4,000 $\times g$, 5 min) and frozen at -20°C. *E. coli* cells expressing recombinant protein were resuspended in 4 mL extraction buffer (50 mM *N*-tris(hydroxymethyl)methyl-2-aminoethanesulfonic acid (TES) pH 7.0, 10 mM MgCl₂, 20 mM DTT) and lysed by sonication. After precipitating the majority of *E. coli* proteins by heating the cell lysate to 80°C for 10 min, the MJ1272-derived protein was purified by anion exchange chromatography on a MonoQ HR column (1 \times 8 cm, Amersham Bioscience) with a linear gradient from 0–1 M NaCl in 25 mM TES pH 7.5 over 55 mL at 1 ml min⁻¹. The purified protein ran as a single band at approximately 18 kDa, consistent with the predicted molecular weight of 18.8 kDa, and was >98% pure as judged by SDS-polyacrylamide gel electrophoresis with coomassie blue staining. Protein concentration was determined by Bradford analysis (73).

Enzyme Assay of *M. jannaschii* MJ1272

The standard assay for MJ1272 consisted of 5–15 μg MJ1272, 25 mM TES/K⁺ buffer pH 7.0, 5 mM MgCl₂, 10 mM DTT, and ~1.0 mM substrate (sepiapterin, H₂neopterin, H₂neopterin-P) in a total volume of 50 μL . When required, 2 mM Fe⁺² was also included in the incubation mixture. For assays utilizing H₂neopterin 2',3'-cyclic phosphate, 1.9 μg MptA, 2 mM MnCl₂ and 2 mM GTP were also included in the incubation mixture (47). The product of the MptA reaction was confirmed by the absorbance using a diode array detector. 6-Carboxypterin was also confirmed by the formation of the methyl ester with HCl/Methanol and reanalysis by HPLC. To test the involvement of cysteine in the reaction MJ1272 (~10 μg) was preincubated in reaction buffer with 2 mM iodoacetamide for 1 hr at room temperature and then assayed for activity with sepiapterin as the substrate according to the standard assay procedures.

Assays were typically incubated for 30 min at 70°C and quenched by the addition of 60 μL methanol. Reaction products were oxidized to the fluorescent pterins by addition of 5 μL iodine in MeOH (50 mg/mL) and the samples were incubated at room temperature for 30 min. Excess iodine was reduced by addition of 5 μL 1 M NaHSO₃. Following centrifugation (14,000 $\times g$, 10 min), 4 μL portion was separated for TLC analysis and the remainder combined with 600 μL H₂O for HPLC analysis. For samples incubated with D₂O, 50% of the water in the incubation mixture was replaced with D₂O.

Chromatographic separation and analysis of pterins was performed on a Shimadzu HPLC (High performance liquid chromatography) System with a C18 reverse phase column (Varian PursuitXRs 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, the pterins were eluted in the following order (min): D-neopterin 2',3'-cyclic phosphate (7.6), 6-carboxypterin (8.4), neopterin (11.0), pterin (16.8), 6-hydroxymethylpterin (16.8) and 6,7-dimethylpterin (27.0). The pterins were also identified by TLC (Thin layer chromatographies) with fluorescence detection. The following R_f were observed using acetonitrile-water-formic acid (88%), 40:10:5 v/v/v as the developing solvent: pterin-6-carboxylate methyl ester, 0.63; pterin, 0.465; 6-hydroxymethylpterin, 0.45; biopterin, 0.44; neopterin, 0.31; and pterin-6-carboxylate, 0.24.

In order to separate 6-hydroxymethylpterin from pterin a polyfluorophenyl column (Varian Pursuit-PFP column 250 × 4.6 mm 5 μ) was used with the HPLC system. The elution profile was isocratic 25 mM sodium acetate buffer pH 6.0 over 45 min at 0.5 ml min⁻¹. With these conditions pterin eluted at 27 min and 6-hydroxymethylpterin at 29 min.

For GC-MS analysis, pterins were purified from incubation mixtures by retention on a Dowex 50W-8X-H⁺ column followed by elution with 7 M aqueous ammonia. Pterins was analyzed for deuterium incorporation either by direct insertion (DI) mass spectral of the pterin, M⁺ = 163 *m/z* or by DI or GC-MS analysis of the pterin (TMS)² derivative with M⁺ = 307 *m/z* and M⁺ - 15 = 292 *m/z*. For the GC-MS analysis of the pterins, the individual pterins were purified by preparative TLC from the ammonia eluted material and analyzed as their TMS derivatives as previously described (74).

Enzyme Assays of *E. coli* QueD

The *ycmM* gene from *E. coli*, which encodes QueD (PTPS-I), was amplified from *E. coli* genomic DNA using primers ECygmS and ECygmAS, and cloned using ligation-independent cloning into the plasmid pET30-Xa (Novagen) following the manufacturer's instructions. The integrity of the sequence was confirmed by sequencing (PSU-Keck Genomics Facility, Portland State University). Recombinant QueD was over-produced in *E. coli* BL21(DE3) using standard protocols, and purified as an N-terminal His₆-affinity tagged fusion protein *via* Ni-NTA affinity chromatography, then cleaved *via* Factor Xa to give the native wild-type enzyme.

Assays of recombinant *E. coli* QueD activity were carried out in reactions containing 100 mM Tris-HCl (pH 8.0), 100 mM KCl, 2 mM MgCl₂, 2 mM DTT, 0.5 mM 7,8-dihydroneopterin monophosphate (H₂NMP) and 50 μM QueD, and were monitored by LC-ESI-MS at 360 nm. Coupled reactions involving GCYH-IB (50 μM) from *N. gonorrhoeae*, 50 μM QueD, and 0.5 mM GTP were also carried out. Reactions were incubated at 37°C for 60 min in the dark under anaerobic conditions. The protein was removed from the solution using a YM-10 micron centrifugal device, and the eluent subjected to LC-ESI-MS analysis on a Bruker MicroTOF-Q equipped with a Supelco Discovery C18 column (25 cm × 2.1 mm, 5 μm particles) eluted at 0.3 mLmin⁻¹ with a solvent gradient of 5% solvent A (0.1% formic acid in water) for 5 min, then 5–50% of solvent B (0.1% formic acid in acetonitrile) in 25 min.

tRNA Extraction and Analysis

Bulk tRNA was prepared, hydrolyzed, and analyzed by liquid chromatography tandem mass spectrometry (LC-MS-MS) as described in (75) from cells grown in chemically defined medium. To evaluate tRNA concentrations, we measured the levels of the m¹G-modified base (298 *m/z* at 21 min.) by integrating the peak area from the extraction ion chromatograms and compared them between samples. The MS-MS fragmentation data were also used to confirm the presence or absence of the nucleosides, Q and G⁺. All tRNA extractions and analysis were performed at least twice independently.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Abbreviations

6HMDP	6-hydroxymethyl-7,8-dihydropterin
APy	2,5-diamino-6-ribosylamino-4(3 <i>H</i>)-pyrimidinone 5'-phosphate
<i>arfA</i>	GTP-cyclohydrolase III gene
<i>arfB</i>	formamide hydrolase gene
BH₄	biopterin
CPH₄	6-carboxy-5,6,7,8-tetrahydropterin
DHNA	dihydroneopterin aldolase
dT	thymidine
FolB	dihydroneopterin aldolase
FolE	GTP cyclohydrolase I
FolE2	GTP cyclohydrolase IB
G⁺	archaeosine
GCYH-IA	GTP cyclohydrolase IA
GCYH-IB	GTP cyclohydrolase IB
GCYH-III	GTP-cyclohydrolase III
GTP	guanosine triphosphate
H₂NMP	7,8-dihydroneopterin monophosphate
H₂NMPc	dihydroneopterin-2',3'-cyclic phosphate
H₂NTP	7,8-dihydroneopterin triphosphate
LC/MS-MS	liquid chromatography – tandem mass spectrometry
preQ₀	7-cyano-7-deazaguanine
preQ₁	7-aminomethyl-7-deazaguanine
PTP	6-pyruvoyl-tetrahydropterin
PTPS-I	COG0720 subfamily involved in queuosine biosynthesis
PTPS-I/III	COG0720 subfamily involved in both queuosine and folate synthesis
PTPS-III	COG0720 subfamily involved in folate biosynthesis

PTPS-IV	COG0720 subfamily that clusters with riboflavin genes in <i>Streptomyces</i> sp.
PTPS-V	COG0720 subfamily of unknown function
PTPS-VI	COG0720 subfamily of unknown function
Q	queuosine
QueD	COG0720 subfamily involved in queuosine biosynthesisPTPS-II/ PtpS 6-pyruvoyl-tetrahydropterin synthase, COG0720 subfamily involved in bioppterin synthesis
ribA2	GTP-cyclohydrolase II gene
SDR	dehydrogenase-reductase
SR	sepiapterin reductase
T-fold	Tunnel-fold
THF	tetrahydrofolate
Ec	<i>Escherichia coli</i>
Sa	<i>Syntrophus acidotrophicus</i>
Li	<i>Leptospira interrogans</i>
Cb	<i>Clostridium botulinum</i>
Ab	<i>Acinetobacter baylyi</i> sp. ADP1
Rn	<i>Rattus norvegicus</i>
Ss	<i>Sulfolobus solfataricus</i>
Pc	<i>Pyrobaculum calidifontis</i>

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