

Characterization of Two Recombinant 3-Hexulose-6-Phosphate Synthases from the Halotolerant Obligate Methanotroph *Methylomicrobium alcaliphilum* 20Z

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Abstract—Two key enzymes of the ribulose monophosphate (RuMP) cycle for formaldehyde fixation, 3-hexulose-6-phosphate synthase (HPS) and 6-phospho-3-hexulose isomerase (PHI), in the aerobic halotolerant methanotroph *Methylomicrobium alcaliphilum* 20Z are encoded by the genes *hps* and *phi* and the fused gene *hps-phi*. The recombinant enzymes HPS-His₆, PHI-His₆, and the two-domain protein HPS–PHI were obtained by heterologous expression in *Escherichia coli* and purified by affinity chromatography. PHI-His₆, HPS-His₆ (2 × 20 kDa), and the fused protein HPS–PHI (2 × 40 kDa) catalyzed formation of fructose 6-phosphate from formaldehyde and ribulose 5-phosphate with activities of 172 and 22 U/mg, respectively. As judged from the k_{cat}/K_m ratio, HPS-His₆ had higher catalytic efficiency but lower affinity to formaldehyde compared to HPS–PHI. AMP and ADP were powerful inhibitors of both HPS and HPS–PHI activities. The two-domain HPS–PHI did not show isomerase activity, but the sequences corresponding to its HPS and PHI regions, when expressed separately, were found to produce active enzymes. Inactivation of the *hps-phi* fused gene did not affect the growth rate of the mutant strain. Analysis of annotated genomes revealed the separately located genes *hps* and *phi* in all the RuMP pathway methylotrophs, whereas the *hps-phi* fused gene occurred only in several methanotrophs and was absent in methylotrophs not growing under methane. The significance of these tandems in adaptation and biotechnological potential of methylotrophs is discussed.

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Aerobic bacteria utilizing reduced C₁-compounds such as methane and its oxidized and substituted derivatives as growth substrates (methylotrophs) are ubiquitous in various ecosystems where they regulate C₁-compound contents. Studies of methylotrophy as a microbial nutrition mode are essential for understanding of evolutionary scenario of the Earth's life, as well as for use in biotechnologies as catalysts for production of various useful metabolites from renewable (and commonly toxic) C₁-

substrates. Methylotrophic bacteria assimilate C₁-compounds via three specific biochemical pathways, where the key enzymes perform first formation of a C–C bond. In the serine pathway, the primary product serine is formed via condensation of formaldehyde and glycine catalyzed by serine transhydroxymethylase (EC 2.1.2.1), and further phosphosugar biosynthesis proceeds by gluconeogenesis. The key enzymes of the ribulose monophosphate (RuMP) cycle, 3-hexulose-6-phosphate synthase (HPS, EC 4.1.2.43) and 6-phospho-3-hexulose isomerase (PHI, EC 5.3.1.27), catalyze condensation of formaldehyde with ribulose 5-phosphate (Ru5P) and isomerization of D-arabino-3-hexulose 6-phosphate into D-fructose 6-phosphate [1–4]. Some aerobic methylotrophs oxidize C₁-substrates into CO₂, which further is fixed via the Calvin cycle.

Abbreviations: FA, formaldehyde; F6P, fructose 6-phosphate; GPD, glucose-6-phosphate dehydrogenase; HPS, 3-hexulose-6-phosphate synthase; LB, Luria–Bertani medium; ORF, open reading frame; PGI, phosphoglucoisomerase; PHI, 6-phospho-3-hexulose isomerase; RuMP, ribulose monophosphate; Ru5P, ribulose 5-phosphate.

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The RuMP cycle is the most efficient C₁-assimilation pathway [5]. Despite its key role in carbon utilization, HPS has been characterized from only a few methylophilic bacteria. The enzyme from the thermotolerant methanotroph *Methylococcus capsulatus* Bath was found to be a unique hexameric membrane protein [1]. Later, the genome of this bacterium was found to contain three copies of the *hps* genes: in the structure of the duplicated *hps/phi* operon and another gene *hps-phi* encoding HPS–PHI fused protein. As follows from analysis of the molecular mass of the enzyme, only the HPS–PHI fused protein has been characterized. The functionality, properties, and role of HPS remained beyond researchers' vision.

Orthologs of HPS and PHI have been found in a variety of bacteria not growing on C₁-substrates. In *Bacillus subtilis* and *Burkholderia cepacia* TM1, these enzymes are involved in formaldehyde detoxification, whereas in some archaea they perform pentose phosphate formation from fructose 6-phosphate, thus compensating for the absence of the pentose phosphate pathway [6–8]. In some archaea, HPS and PHI are expressed as a two-domain protein [9, 10]. It was suggested that bifunctional HPS–PHI enhances efficiency of sequential reactions and stability against thermal denaturation in comparison to the individual enzymes.

The halotolerant methanotroph *Methylobacterium alcaliphilum* 20Z is a promising organism for biotechnological application for synthesis of various useful metabolites from methane or methanol because this strain demonstrates high growth rate and ability to grow in a wide range of salinity, pH, and methanol concentrations; also, it can ferment formaldehyde under oxygen-limited conditions [11, 12]. Recently, in the genome of this methanotroph putative *hps* and *phi* genes and the additional *hps-phi* fused gene were found. The goal of this work was to elucidate functionality and properties of the two HPSs, of the two forms of the hexulose phosphate synthase from *M. alcaliphilum* 20Z obtained by heterologous expression in *Escherichia coli*.

MATERIALS AND METHODS

Bacteria and growth conditions. *Methylobacterium alcaliphilum* 20Z (VKM B-2133T = NCIMB 14124T) was grown at 30°C in a methane–air atmosphere (1 : 1) or in the presence of 0.3% methanol (v/v) on mineral 2P medium containing 0.1 M NaHCO₃ and 0.3 M NaCl [11]. *Escherichia coli* Rosetta (DE3) obtained from Stratagene (USA) was routinely grown at 37°C in LB medium [13]. Ampicillin (100 µg/ml), kanamycin (100 µg/ml), and chloramphenicol (25 µg/ml) were added to the growth medium if required.

Cloning and expression of the *hps* and *hps-phi* genes. DNA was isolated from *M. alcaliphilum* 20Z as described

previously [14]. The *hps* (CCE25609), *phi* (CCE25608), and *hps-phi* fused (CCE23598) genes from the genomic DNA were amplified using the appropriate primers (Table 1). The PCR products were digested by endonucleases NdeI and HindIII and ligated into the expression vector pET30a(+) treated with the same restrictases. The *hps-phi* gene was cloned into the pHUE vector [15] between the SacII and HindIII endonuclease sites. Analogously, DNA fragments containing ORFs coding for either the synthase (*hps-2*) or isomerase (*phi-2*) domain of the HPS–PHI fused protein were amplified using HPS–HPI (NdeI)F and Delta-HPS–R(HindIII) or Delta-HPI–F and Delta-HPI–R primers, respectively (Table 1). All resultant plasmids were transformed into *E. coli* (DE3) Rosetta cells, and protein expression was induced by adding 0.5 mM isopropyl-1-thio-β-D-galactopyranoside (Sigma-Aldrich, USA) in the logarithmic phase of the culture (OD₆₀₀ of 0.6–0.8). After 18-h growth at 18°C, the cells were harvested by centrifugation for 30 min at 5000g and 4°C.

Purification of the recombinant enzymes. The recombinant HPS, HPS-2, PHI, and PHI-2 were purified by affinity chromatography on a Ni²⁺-nitrilotriacetic acid (Ni-NTA) column as described [16]. A modified protocol was used to purify the recombinant HPS–PHI. The *E. coli* cells were suspended in 100 mM Tris-HCl buffer, pH 8.0, containing 0.8 M NaCl, 2 mM imidazole, and 5 mM MgCl₂. The cells were disrupted by sonication for 1 min with 10-s bursts, with 30-s cooling on ice between bursts, using an MSE sonicator (MSE, UK), and centrifuged for 30 min at 11,000g at 4°C. The supernatant was loaded into a Ni-NTA column equilibrated with the same buffer. The column was washed with 100 mM Tris-HCl buffer (pH 8.0) containing 0.8 M NaCl, 5 mM MgCl₂, and 4 mM imidazole. The ubiquitin–His₆-tagged protein was eluted with the same buffer containing 150 mM imidazole. The recombinant His₆-Ub–HPS–PHI was additionally treated with deubiquitinase (overnight at 4°C), dialyzed against 100 mM Tris-HCl buffer, pH 8.0, with 0.8 M NaCl, and loaded again into the Ni-NTA column equilibrated with this buffer. The protein not bound with the column was collected and stored at 4°C.

Determination of quaternary structure. The quaternary form of the enzyme was analyzed by nondenaturing gel electrophoresis using pore-limiting gradient polyacrylamide (4–30%) [17]. Ferritin (440 kDa), amylase (200 kDa), alcohol dehydrogenase (150 kDa), and bovine serum albumin (66 kDa) from Sigma-Aldrich Group were used as reference proteins.

Enzyme assays. Two methods were used to determine HPS activity. Method I based on the measurement of formaldehyde concentration was used to determine the pH and temperature optima. The reaction mixture (0.5 ml) contained 100 mM Tris-HCl buffer, pH 8.5 (for HPS–PHI), or 100 mM NaOH-glycine buffer, pH 9.5 (for HPS), 5 mM MgCl₂, 2.5 mM Ru5P, and the enzyme.

Table 1. Primers used in this work. Restriction sites are underlined

Primer	5'-3' sequence	Cloning region
HPS1-F(NdeI)	TACATATGGCAAGACCATTAATACAAATGGC	<i>hps</i>
HPS1-R(HindIII)	TTAAGCTTATGCACTCGCTGCATCAACC	<i>hps</i>
HPS-HPI (Nde)F	TCATATGGCAAACCATTTGATTCAATT	<i>hps-phi</i>
HPS-HPI (HindIII)R	TAAGCTTGAACGCCGTAAACCCAT	<i>hps-phi</i>
HPI1-F(NdeI)	TACATATGCATCAGCAACTGATCATTGAC	<i>phi</i>
HPI1-R(HindIII)	TTAAGCTTCATTCCAGGTTTCGCGTGCC	<i>phi</i>
Delta-HPS-R(Hind)	TAAGCTTAGCCCTTGCTATTGACCAAGCGT	synthase domain of <i>hps-phi</i> gene
Delta-HPI-F	TCATATGACGCATCGTGAATTTGTCTGTC	isomerase domain of <i>hps-phi</i> gene
Delta-HPI-R	TAAGCTTATTCCAGGTTTCGCATGTCTC	isomerase domain of <i>hps-phi</i> gene
dHPSupF(EcoRI)	TGAATTCTGCCATAAAACGCCATC	upstream region of <i>hps</i> gene
dHPSupR(Acc65I)	AGGTACCGGTCTTGCCATGTTTCT	upstream region of <i>hps</i> gene
dHPSdownF(ApaI)	AGGGCCCGTCAACTGGTTGATGCA	downstream region of <i>hps</i> gene
dHPSdownR(SacI)	TGAGCTCCAGGAAGAAGACGCGTCG	downstream region of <i>hps</i> gene
dHPIupF(EcoRI)	TGAATTCTGTTGTTGGTTGACCTGA	upstream region of <i>phi</i> gene
dHPIupR(Acc65I)	TGGTACCTTGTCATGATCAGTTGCT	upstream region of <i>phi</i> gene
dHPIdownF(ApaI)	TGGGCCCCGACGCTGTTCTTCCTGGA	downstream region of <i>phi</i> gene
dHPIdownR(SacI)	TGAGCTCGGGTTAGTCGGGTTAT	downstream region of <i>phi</i> gene
dHPS/HPIupF(BglII)	AAGATCTATCGCATCCAAAGTGTG	upstream region of <i>hps-phi</i> gene
dHPS/HPIupR(Acc65I)	AGGTACCTTTATGCGCTCCTGATTA	upstream region of <i>hps-phi</i> gene
dHPS/HPIdownF(ApaI)	TGGGCCCCGAGACATGCGAACCTGG	downstream region of <i>hps-phi</i> gene
dHPS/HPIdownR(SacI)	TGAGCTCACCGCCGTTTTTAGA	downstream region of <i>hps-phi</i> gene

The mixture was incubated for 3 min at 30°C, and the reaction was started by adding 0.5 mM formaldehyde. After 10 min, 0.5 ml of the Nash reagent [18] was added, the mixture was incubated at 50°C for 10 min, and the optical density at 420 nm was measured.

In method II, PHI-His₆ purified from *M. alcaliphilum* 20Z, phosphoglucosomerase (PGI), and glucose-6-phosphate dehydrogenase (GPD) were used as the coupling enzymes. The reaction mixture (see method I) additionally contained 10 U of PHI, 10 U of PGI and GPD, and 0.5 mM NADP⁺. The rate of NADP⁺ reduction was assayed at 340 nm with a Shimadzu UV-1700 spectrophotometer. Method II was used to determine the kinetic parameters of the enzymes.

The apparent values K_m and A_{max} were calculated using the Enzyme Kinetics Module for SigmaPlot 12. The following buffers were used to test the effect of pH on the activity: potassium phosphate (pH 6.0-8.0), Tris-HCl (pH 7.0-9.0), and NaOH-glycine (pH 9.0-10.5). The following compounds were tested as potential effectors: pyruvate, citrate, serine, phosphoenolpyruvate, glycerate, 2-ketoglutarate, malate, AMP, ADP, ATP, pyrophosphate, and orthophosphate at a final concentration of 1 mM; glucose, glucose 1-phosphate, glucose 6-phosphate, fructose, fructose 1-phosphate, fructose 6-phosphate (F6P), and fructose 1,6-bisphosphate at final concentration 5 mM.

The reverse reaction catalyzed by HPS was assayed by detecting formaldehyde formation from F6P with the Nash reagent. The reaction was performed in 100 mM Tris-HCl buffer, pH 8.5 (in the case of HPS-PHI) or 100 mM NaOH-glycine buffer, pH 9.5 (in the case of HPS). The mixture (0.5 ml) contained 10 mM F6P, 10 mM MgCl₂, HPS (or HPS-PHI), and PHI-His₆.

Mutant generation. Insertions in the *hps* and *phi* genes and the *hps-phi* fused gene were generated using the suicide vector pCM184 as described [19]. Briefly, 3'- and 5'-fragments of each gene to be mutated were PCR amplified using the appropriate primers (Table 1) and inserted into pCM184 upstream and downstream of the kanamycin (Km) resistance gene. The resulting plasmids were transformed into *E. coli* S17-1, and the resulting donor strain was mated with wild-type *M. alcaliphilum* 20Z in biparental mating. Mutants were selected on 0.5% methanol plates with kanamycin (100 µg/ml). Double-crossover mutants were identified by diagnostic PCR tests.

Phylogenetic analyses. The full-length amino acid sequences from the protein databases of NCBI were used for phylogenetic analyses. The sequences were aligned using the Alignment tool of the MEGA 6 software [20]. Minor corrections of the alignments were made manually. There were 208 informative positions in the final dataset. The phylogenetic tree was generated with MEGA 6 using the Neighbor-Joining method.

RESULTS

Cloning of the *hps*, *phi*, and *hps-phi* genes and purification of the recombinant enzymes. BLAST analysis of the *M. alcaliphilum* genome revealed two ORFs (CCE25609 and CCE25608) sharing noticeable sequence identities with HPS and PHI characterized from *Methylomonas aminofaciens* 77a (64.7 and 53.6%) and from *Mycobacterium gastri* MB19 (43.3 and 33.2%) [3, 8]. An additional ORF (CCE23598) coding for the putative bifunctional enzyme having both HPS and PHI domains was found. The *hps* and *phi* genes were cloned into the pET30a(+) vector designed for the expression of the C-terminal His₆-tagged fusion proteins under the control of the T7 promoter. The *hps-phi* fused gene was cloned into the pHUE vector allowing overproduction of the recombinant enzyme with histidine-tagged ubiquitin (His₆-Ub) fusion at the N-terminus [15]. The genes were expressed in *E. coli* Rosetta (DE3), and the recombinant enzymes were purified by affinity chromatography on Ni-NTA agarose. His₆-Ub-HPS-PHI was additionally treated with a deubiquitinase to obtain the protein in its native form. The subunit molecular masses of the proteins determined by SDS-PAGE were in a good agreement with the theoretically calculated values (22.2, 19.2, and 41.9 kDa for HPS, PHI, and HPS-PHI, respectively) (Fig. 1). Nondenaturing gel electrophoresis in pore-limited gradient of acrylamide revealed single bands of about 40 kDa for HPS and PHI and one band of 70 kDa for HPS-PHI, thereby indicating that all the enzymes are homodimers (data not shown).

Kinetic parameters of recombinant HPS and HPS-PHI. Due to high instability of D-arabino-3-hexulose 6-phosphate, we were unable to characterize PHI from *M. alcaliphilum* 20Z. Nevertheless, the recombinant PHI demonstrated high activity and was used for measuring the HPS activity as a coupling enzyme at low concentration 0.01 mg/ml.

In the presence of PHI-His₆, HPS-His₆ catalyzed F6P formation from Ru5P and formaldehyde. The recombinant *M. alcaliphilum* HPS had a more alkaline pH optimum compared to the enzyme purified from *M. capsulatus* Bath (Fig. 2 and Table 2). This property correlated with the pH optimum for growth of *M. alcaliphilum* 20Z [11]. The enzyme activity was 70% higher at 50°C than at 30°C (Fig. 3). Like other studied HPSs, the *M. alcaliphilum* HPS was a Mg²⁺-dependent enzyme and strongly specific to Ru5P, since it did not use glucose, glucose 6-phosphate, glucose 1-phosphate, fructose, F6P, fructose 1,6-bisphosphate, ribose 5-phosphate, ribose 1-phosphate, or erythrose 4-phosphate as formaldehyde acceptors.

The activity of HPS was completely inhibited by 1 mM AMP. In the presence of 1 mM ADP in the reaction mixture, HPS retained 37% activity, while 1 mM ATP had only a minor inhibitory effect (Table 3). The enzyme obeyed Michaelis-Menten kinetics. Although

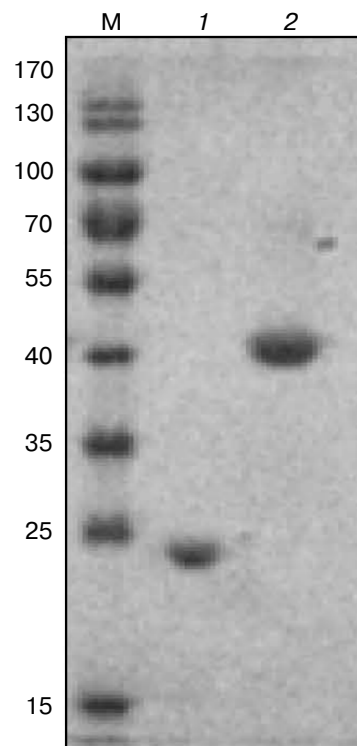


Fig. 1. A 12% SDS-PAGE of HPS-His₆ (1) and HPS-PHI (2) from *M. alcaliphilum* 20Z. M, markers of molecular masses (kDa).

the enzyme had a higher temperature optimum, all kinetic assays were carried out at 30°C corresponding to the optimal growth temperature for the strain. At this temperature, the maximal activity was 172 U/mg protein and the apparent K_m values for formaldehyde and Ru5P were 0.98 ± 0.40 and 0.13 ± 0.01 mM, respectively. In the presence of PHI, HPS catalyzed formaldehyde formation from F6P with activity of 0.95 U/mg protein.

The two-domain enzyme HPS-PHI displayed HPS activity only in the presence of PHI-His₆, but it showed no isomerase activity *per se*. The maximal activity (28 U/mg protein) was observed at 50°C and pH 8.5. The apparent K_m values for formaldehyde and Ru5P measured at 30°C and pH 8.5 were 0.64 ± 0.06 and 0.24 ± 0.03 mM, respectively. Like the single-domain HPS, HPS-PHI was a Mg²⁺-dependent enzyme and was strongly specific to Ru5P. HPS-PHI was inhibited by ADP and completely inactivated by AMP, and it was not inhibited by ATP. In addition, PP_i remarkably inhibited the enzyme activity (Table 3). The k_{cat}/K_m calculation revealed that HPS was 13-fold more efficient with Ru5P and 5-fold more efficient with formaldehyde than the HPS-PHI fused enzyme (Table 2). The reverse activity of the HPS-PHI fused enzyme (0.19 U/mg protein) was detected only in the presence of PHI, thereby confirming that the isomerase domain was unable to catalyze the reaction in either direction.

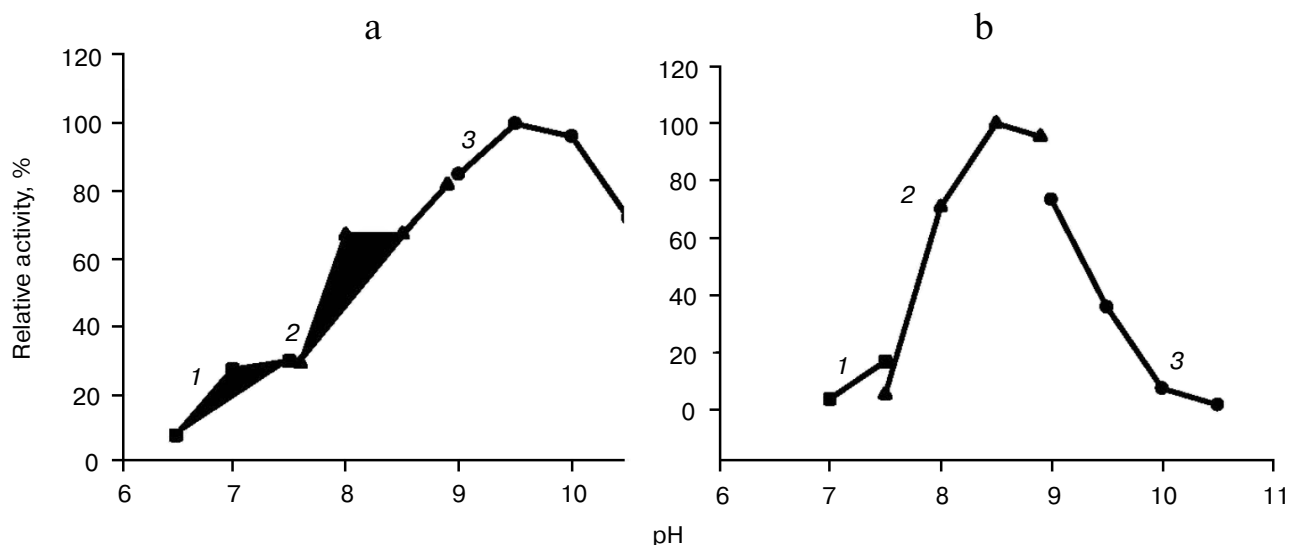


Fig. 2. Dependence of specific activity of recombinant HPS (a) and HPS-PHI (b) on pH. Curves: 1) potassium phosphate buffer; 2) Tris-HCl buffer; 3) NaOH-glycine buffer.

Testing of activities of HPS and PHI domains of HPS-PHI fused enzyme. We cloned the *hps* and *phi* fragments of the fused gene coding for the two-domain HPS-PHI. The His₆-tagged recombinant enzymes named HPS-2 and PHI-2 were expressed in *E. coli* Rosetta cells and purified by one-step metal-chelating chromatography. As estimated by SDS-PAGE, the subunit molecular masses of HPS-2 and PHI-2 were ~20 kDa. Native electrophoresis showed a single band of about 40 kDa for HPS-

2 and a band of 80 kDa for PHI-2 (data not shown), suggesting that the enzymes were dimer and tetramer, respectively. Notably, HPS-2 had four-fold higher synthase activity (120 U/mg protein) and two-fold higher affinity to formaldehyde (apparent $K_m = 0.31 \pm 0.03$ mM) than the HPS-PHI fused protein. Unlike the fused protein, HPS-2 obeyed Hill kinetics ($n = 2.5 \pm 0.6$) rather than Michaelis-Menten kinetics. In contrast to HPS-PHI, the recombinant PHI-2 demonstrated isomerase activity.

Table 2. Properties of hexulosephosphate synthases from microorganisms

Parameter	<i>M. alcaliphilum</i> 20Z		<i>Methylomonas aminofaciens</i> [3]	<i>Methylomonas</i> M15 [2]	<i>Methylococcus capsulatus</i> [1]	<i>Pyrococcus horikoshii</i> [9]
	HPS	HPS-PHI ^a				
A_{max} , U/mg protein	172 ^b	22 ^b	53	66.5	69	526
K_m , mM:						
Formaldehyde	0.98	0.64	0.29	1.1	0.49	6.25
Ru5P	0.13	0.24	0.059	1.6	0.083	nd
k_{cat} , min ⁻¹ :						
Formaldehyde	2749	305	nd	nd	nd	31,560
Ru5P	3528	503	nd	nd	nd	nd
k_{cat}/K_m :						
Formaldehyde	2593	476	nd	nd	nd	5050
Ru5P	27,140	2094	nd	nd	nd	nd
pH-optimum	9.5	8.5	8	7.5-8.0	7	nd
T_{opt} , °C	50	50	nd	nd	nd	80-85
Mr (number of subunits)	40 (2)	70 (2)	45-47 (2)	43 (2)	310 (6)	162 (4)

Notes: nd, not determined.

^a In synthase reaction.

^b A_{max} was determined at 30°C.

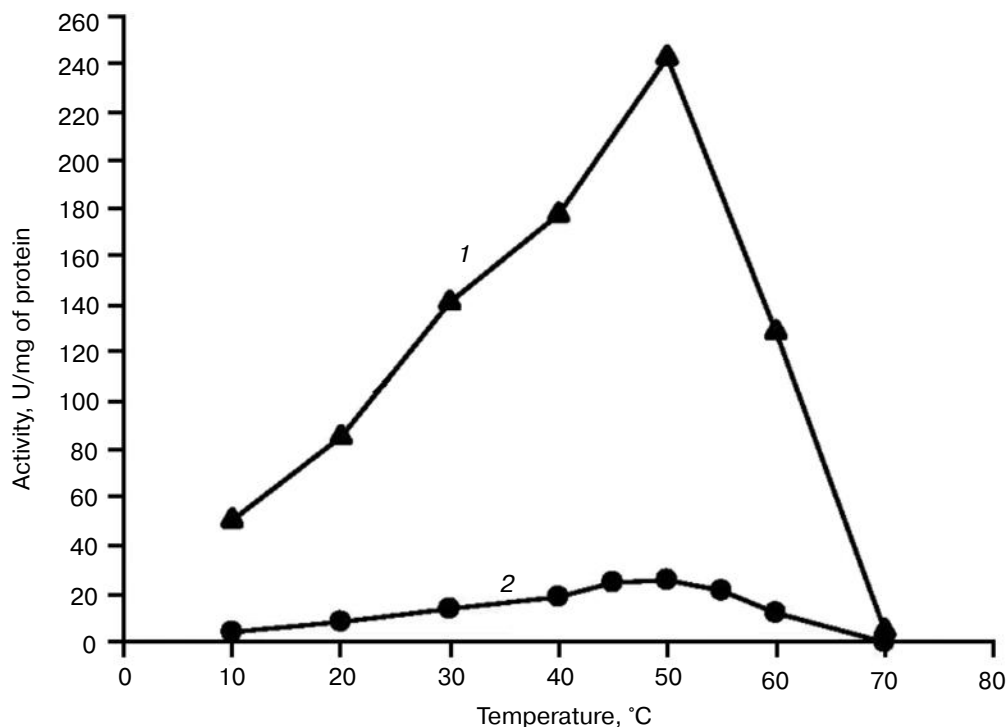


Fig. 3. Dependence of specific activity of recombinant HPS (1) and HPS–PHI fused enzyme (2) on temperature.

Distribution and phylogenetic analysis of HPS and PHI in methylotrophs. BLAST analysis revealed the separate *hps* and *phi* genes in all gammaproteobacterial methanotrophs whose genomes are available in the database; about half have an additional *hps-phi* fused gene, but none of the methanotrophs has a single *hps-phi* fused gene (Table 4). The *hps-phi* fused gene is present in the moderate thermophilic and thermotolerant methanotrophs of the genera *Methylocaldum* and *Methylococcus* and halotolerant and salt-independent species of genera *Methylobacterium*, *Methylosarcina*, and *Methylovulum*. The fused gene is absent in psychrophilic and salt-dependent and -independent *Methylobacter* and *Methylobacterium* representatives, halophilic *Methylohalobius*, thermophilic *Methylothermus*, and all strains of the genus *Methylomonas*. Notably, the *hps-phi* fused gene is absent in all the RuMP pathway methylotrophic bacteria unable to use methane as growth substrate. HPS–PHI from *M. alcaliphilum* 20Z shared 68–99% identity of amino acid sequences with the fused enzymes from other methanotrophs, but only 38–40% identity with the two-domain enzyme from archaea of the *Pyrococcus* and *Thermococcus* genera. Interestingly, *M. alcaliphilum* HPS shared 73% identity with the synthase domain of its own fused enzyme. It should be noted that two or three copies of the *hps* and *phi* genes could be recognized in some representatives of methylotrophs (Table 4).

The phylogenetic analysis revealed several separate branches of HPS (Fig. 4): (1) methanotrophic *Gamma-*

proteobacteria; (2) methylotrophs of *Gamma-* and *Betaproteobacteria* classes; (3) *Flavobacteroides* and *Actinobacteria*; (4) *Epsilonproteobacteria*; (5) *Bacillus* including methylotrophs; (7) archaea. Methanotrophic HPSs are clustered together except for HPS from the halophilic *Methylohalobius crimeensis* and thermophilic *Methylothermus subterraneus* forming a separate branch on the phylogenetic tree.

Effect of *hps-phi* fused gene mutation. To elucidate the further function of the HPS–PHI fused enzyme, we constructed a *hps-phi* insertion mutant via allelic exchange. The growth rate of the *hps-phi* knockout strain was similar to that of the wild type (data not shown), most

Table 3. Influence of some metabolites on activity of HPS–His₆ and HPS–PHI (% of activity without effectors)

Effector (mM)	HPS–His ₆	HPS–PHI
Without effector	100	100
ATP (1)	86	100
ADP (1)	37	11
ADP (5)	0	0
AMP (1)	0	0
PP _i (2)	108	61
P _i (30)	95	107

Table 4. Distribution of *hps*, *phi*, and *hps-phi* fused genes in methanotrophs and methylotrophs

Bacteria	<i>hps</i>	<i>phi</i>	<i>hps/phi</i>
Methanotrophs			
Gammaproteobacteria			
<i>Methylobacter luteus</i> IMV-B-3098T	+	+	—
<i>Methylobacter marinus</i> A45	+	+	—
<i>Methylobacter</i> sp. BBA5.1	+	+	—
<i>Methylobacter tundripaludum</i> SV96	+	+	—
<i>Methylobacter whittenburyi</i> ACM 3310	+	+	—
<i>Methylocaldum szegediense</i> O-12	+	+	+
<i>Methylocaldum</i> sp. 175	+	+	—
<i>Methylogaea oryzae</i> JCM 16910	++	+	+
<i>Methylococcaceae bacterium</i> 73a	++	+	—
<i>Methylococcaceae bacterium</i> Sn10-6	+	+	+
<i>Methylococcus capsulatus</i> Bath	++	++	+
<i>Methylococcus capsulatus</i> Texas	+	+	+
<i>Methyloglobulus morosus</i> KoM1	+	+	+
<i>Methylohalobius crimeensis</i> 10Ki	+	+	—
<i>Methylomarinum vadi</i>	++	+++	—
<i>Methylomicrobium alcaliphilum</i> 20Z	+	+	+
<i>Methylomicrobium agile</i>	+	+	—
<i>Methylomicrobium album</i> BG8	+	+	+
<i>Methylomicrobium buryatense</i> 5G	+++	+++	+
<i>Methylomicrobium kenyense</i> AMO1	++	+	+
<i>Methylomonas denitrificans</i> FJG1	++	+	—
<i>Methylomonas methanica</i> MC09	++	+	—
<i>Methylomonas</i> sp. 11b	++	+	—
<i>Methylomonas</i> sp. 16a	+	+	—
<i>Methylomonas</i> sp. MK1	++	+	—
<i>Methylosarcina fibrata</i> AML-C10	+	+	+
<i>Methylosarcina lacus</i> LW14	+	+	+
<i>Methylothermus subterraneus</i>	+	+	—
<i>Methylovulum miyakonense</i> HT12	+	+	+
Methylobacteria			
Betaproteobacteria			
<i>Methylobacillus flagellatus</i> KT	++	+	—
<i>Methylobacillus glycogenes</i> JCM 2850	++	+	—
<i>Methylobacillus rhizosphaerae</i> Ca-68	++	+	—
<i>Methylotenera</i> sp. 1P/1	++	+	—
<i>Methylotenera</i> sp. 73s	++	+	—
<i>Methylotenera mobilis</i> JLW8	++	+	—
<i>Methylotenera versatilis</i> 301	++	+	—
<i>Methylotenera versatilis</i> 79	++	+	—
<i>Methylophilus methylotrophus</i> DSM 46235	++	+	—
<i>Methylophilus</i> sp. 1	++	+	—
<i>Methylophilus</i> sp. 42	++	+	—
<i>Methylovorus glucosetrophus</i> SIP3-4	++	+	—
<i>Methylovorus</i> sp. MP688	++	+	—
Gammaproteobacteria			
<i>Methylophaga lonarensis</i> MPL	++	+	—
<i>Methylophaga</i> sp. JAM1	++	+	—
<i>Methylophaga</i> sp. JAM7	++	+	—
<i>Methylophaga thiooxydans</i> DMS010	+	+	—
Firmicutes			
<i>Bacillus methanolicus</i> MGA3	+	+	—
Actinobacteria			
<i>Arthrobacter globiformis</i>	++	+	—
<i>Pseudarthrobacter sulfonivorans</i>	+	+	—
<i>Mycobacterium gastris</i>	+	+	—
<i>Amycolatopsis methanolica</i>	+	+	—

Note: The database of www.genoscope.cns.fr, <http://www.ncbi.nlm.nih.gov/>, and <https://img.jgi.doe.gov> were used for analysis. The number of + represents the number of the corresponding genes in the bacterial genome.

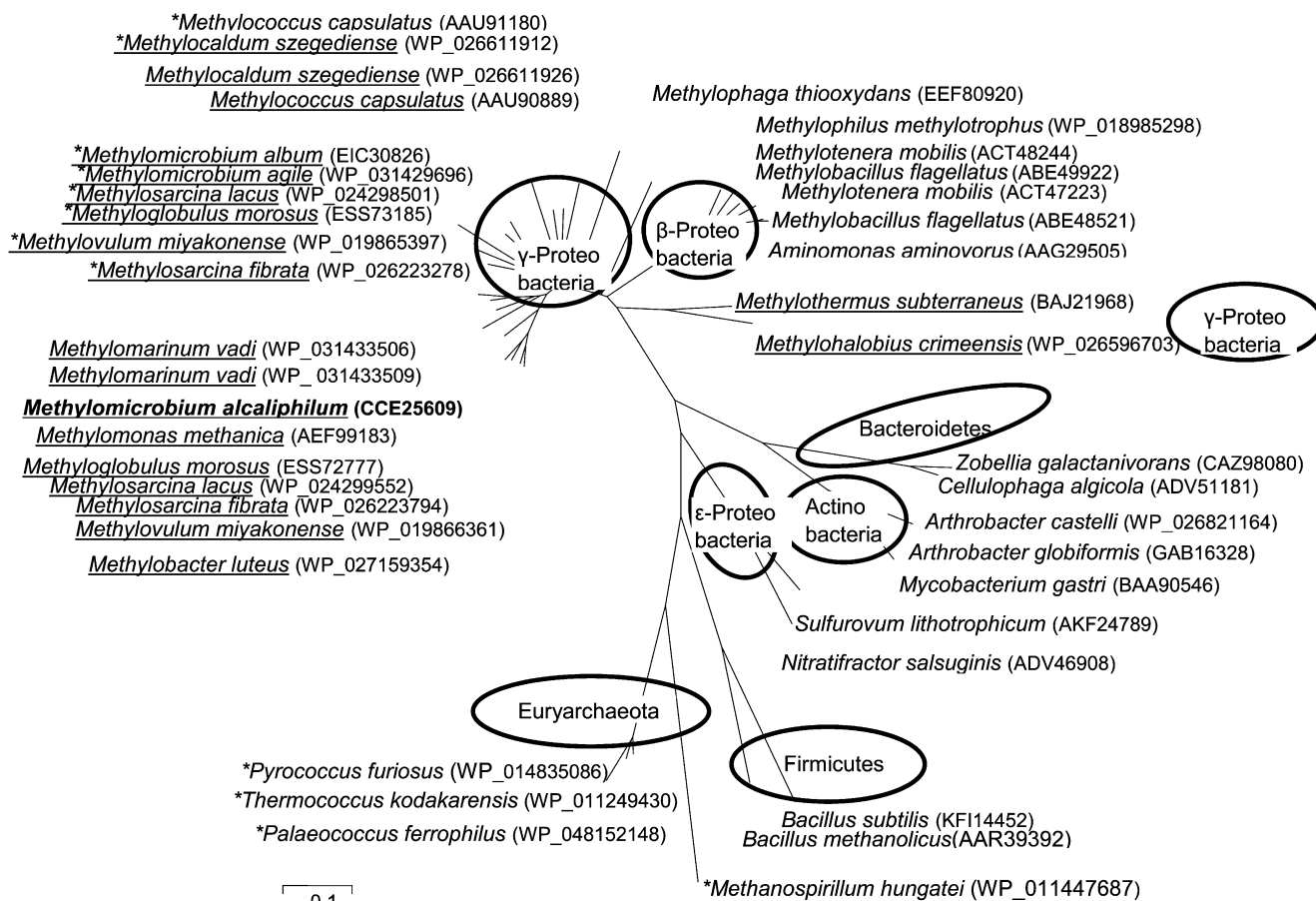


Fig. 4. Phylogenetic tree of amino acid sequences of HPS and HPS-PHI fused proteins. HPS-PHI fused enzymes are marked by asterisks. Methanotrophic bacteria are underlined. Accession numbers of sequences in GenBank are indicated in parentheses.

likely indicating that the HPS-PHI fused enzyme is not essential for primary carbon assimilation. In contrast, we were unable to obtain a mutant deleted in the separately located *hps* or *phi* genes.

DISCUSSION

In this work, we have first characterized two forms of HPS, which is the key enzyme of the RuMP pathway of formaldehyde fixation in aerobic methanotrophs. In *M. alcaliphilum* 20Z, the properties of the HPS-His₆ and HPS-PHI fused enzymes suggested that the enzyme encoded by *hps* is responsible for assimilation of a significant part of methane carbon, whereas the two-domain protein HPS-PHI catalyzed condensation of formaldehyde and Ru5P, but with lower activity and efficiency. Moreover, the strain with an insertion in the *hps-phi* fused gene did not differ in growth rate from the parental strain. In addition, we were unable to obtain a viable strain with the *hps* or *phi* gene deleted.

Although HPS-PHI was inactive as isomerase, both domains became active *in vitro* when expressed separate-

ly, demonstrating corresponding synthase and isomerase activity. Since the active isomerase domain PHI-2 was a tetramer, while HPS-PHI has dimeric structure, we speculate that the absence of isomerase activity of the HPS-PHI fused enzyme is related to its incorrect structure. As shown, all PHI characterized to date are tetramers, and their active site is composed by all monomers of the tetramer [9, 21, 22]. Moreover, the bifunctional enzyme catalyzing the sequential HPS and PHI reactions in the archaeon *Pyrococcus horikoshii* OT3 had tetrameric structure [9]. The importance of quaternary structure for HPS and PHI activities was evidenced by construction of an artificial bifunctional enzyme via expression of fusions from the *hps* and *phi* genes from the methylotrophic bacterium *Mycobacterium gastris* without adding a linker sequence [23]. Only the product of the *hps-phi* fusion expressed in *E. coli* was active, whereas the product of *phi-hps* lacked activity. Since the fused enzyme from *M. alcaliphilum* 20Z had higher affinity to formaldehyde than the separate enzyme (Table 2), we suppose that HPS-PHI is functioning in conditions of low substrate concentrations. Much higher affinity to formaldehyde was demonstrated by the hexameric two-domain protein

from *M. capsulatus* Bath, which also lacked isomerase activity [1].

Finally, our results can be considered in terms of molecular evolution of the RuMP pathway. Comprising separate phylogenetic groups, the sequences of the HPS and HPS domains of the fused protein of the aerobic methanotrophs are very phylogenetically remote from those of anaerobic archaea (Fig. 4). Along with high homology between HPS and the synthase domain of the fused enzyme in the same methanotroph, this could imply rather recent dividing of HPS–PHI into separate enzymes. Absence of the fused enzyme in non-methanotrophic methylotrophs sequenced to date is very intriguing.

Characterization of the key enzymes of the RuMP pathway could be applicable for genetic engineering improvement of methylotrophic producers of useful metabolites (phosphosugars, amino acids), rational fermenting with C₁-compounds as auxiliary substrates, creation of transgenic plants with enhanced capability of taking up formaldehyde from air, and others [24–28]. It should be noted that *Methylomicrobium buryatense* 5G, the closest relative to the strain 20Z but exceeding in growth rate, has three bis-cistronic *hps/phi* operons in its chromosome (Table 4). While the procedure of metabolic modification is simplified when the bifunctional protein is introduced, the properties of the enzymes added demand special attention.

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