A flap motif in human serine hydroxymethyltransferase is important for structural stabilization, ligand binding, and control of product release

Received for publication, April 20, 2019, and in revised form, May 21, 2019. Published, Papers in Press, May 22, 2019. DOI 10.1074/jbc.RA119.007454

Sakunrat Ubonprasert1,*, Juthamas Jaroensuk1,*, Wichai Pornthanakasem1,*, Nuntaporn Kamonsutthipaijit1,*, Peerapong Wongpituk1,*, Pitchayathida Mee-udorn1,*, Thanyada Rungrotmongkol1,*, Penchit Chitnumsub1,*, Ubolsree Leartsakulpanich1,*, Pimchai Chaiyen1,*, and Somchart Maenpuen1

From the 1Department of Biochemistry and Center for Excellence in Protein and Enzyme Technology, Faculty of Science, Mahidol University, Bangkok 10400, Thailand, 2School of Biomolecular Science and Engineering, Vidyasirimedhi Institute of Science and Technology (VISTEC), Rayong 21210, Thailand, 3Biomolecular Analysis and Application Laboratory, National Center for Genetic Engineering and Biotechnology (BIOTEC), and 4National Nanotechnology Center (NANOTEC), National Science and Technology Development Agency, Pathumthani 12120, Thailand, 5Synchrotron Light Research Institute (Public Organization), Nakhon Ratchasima 30000, Thailand, 6††Biocatalyst and Environmental Biotechnology Research Unit, Department of Biochemistry, Faculty of Science, and 7†§†‡‡‡Bioinformatics and Computational Biology Program, Graduate School, Chulalongkorn University, Bangkok 10330, Thailand, and 8†Department of Biochemistry, Faculty of Science, Burapha University, Chonburi 20131, Thailand

Edited by Ruma Banerjee

Human cytosolic serine hydroxymethyltransferase (hcSHMT) is a promising target for anticancer chemotherapy and contains a flexible “flap motif” whose function is yet unknown. Here, using size-exclusion chromatography, analytical ultracentrifugation, small-angle X-ray scattering (SAXS), molecular dynamics (MD) simulations, and ligand-binding and enzyme-kinetic analyses, we studied the functional roles of the flap motif by comparing WT hcSHMT with a flap-deleted variant (hcSHMT/Δflap). We found that deletion of the flap results in a mixture of apo-dimers and holo-tetramers, whereas the WT was mostly in the tetrameric form. MD simulations indicated that the flap stabilizes structural compactness and thereby enhances oligomerization. The hcSHMT/Δflap variant exhibited different catalytic properties in (6S)-tetrahydrofolate (THF)-dependent reactions compared with the WT but had similar activity in THF-independent aldol cleavage of β-hydroxyamino acid. hcSHMT/Δflap was less sensitive to THF inhibition than the WT (K_i of 0.65 and 0.27 mM THF at pH 7.5, respectively), and the THF dissociation constant of the WT was also 3-fold lower than that of hcSHMT/Δflap, indicating that the flap is important for THF binding. hcSHMT/Δflap did not display the burst kinetics observed in the WT. These results indicate that, upon removal of the flap, product release is no longer the rate-limiting step, implying that the flap is important for controlling product release. The findings reported here improve our understanding of the functional roles of the flap motif in hcSHMT and provide fundamental insight into how a flexible loop can be involved in controlling the enzymatic reactions of hcSHMT and other enzymes.

Serine hydroxymethyltransferase (SHMT2; EC 2.1.2.1.) is a pyridoxal 5'-phosphate (PLP)-dependent enzyme in which the PLP cofactor facilitates the reversible transfer of a hydroxymethyl group from L-serine to (6S)-tetrahydrofolate (THF) to yield glycine and 5,10-methylenetetrahydrofolate (5,10-CH2-THF) as products (1–3). SHMT is one of the enzymes in the deoxyxymidylate (dTMP) synthesis cycle in which the other two enzymes, thymidylate synthase (TS) and dihydrofolate reductase (DHFR), also take part in the recycling of folate compounds and production of 5,10-CH2-THF via the SHMT reaction. 5,10-CH2-THF serves as a methyl donor for the TS reaction to convert dUMP to dTMP, which is a requisite precursor for DNA biosynthesis (4). Due to its important role in DNA biosynthesis, cell proliferation, and cell survival, SHMT is one of the attractive targets for antimalarial (5–12) and anticancer chemotherapies (10, 13–16).

This work was supported by Thailand Research Fund Grants RTA5980001 (to P. Chaiyen), MRGS980001 (to S. M.), and RDG6050101 (to P. Chaiyen and S. M.); the Faculty of Science, Mahidol University and School of Biomolecular Science and Engineering, Vidyasirimedhi Institute of Science and Technology (to P. Chaiyen and J. J.); the Faculty of Science, Burapha University (to S. M.); Cluster Program and Management Office, National Science and Technology Development Agency, Thailand Grant CPMP-O-13-00835 (to U. L., W. P., P. Chitnumsub, P. Chaiyen, and S. M.); Thailand Graduate Institute of Science and Technology Scholarship Grant TG-22-14-59-031M (to S. U.); Synchrotron Light Research Institute–Public Organization Grant 1-2561/Project I.D. 3212 (to S. M. and S. U.); and a Science Achievement Scholarship of Thailand (to P. Chaiyen), MRG5980001 (to S. M.), and RDG6050101 (to P. Chaiyen and J. J.); the Faculty of Science, Burapha University (to S. M.); Cluster Program and Management Office, National Science and Technology Development Agency, Thailand Grant CPMP-O-13-00835 (to U. L., W. P., P. Chitnumsub, P. Chaiyen, and S. M.); Thailand Graduate Institute of Science and Technology Scholarship Grant TG-22-14-59-031M (to S. U.); Synchrotron Light Research Institute–Public Organization Grant 1-2561/Project I.D. 3212 (to S. M. and S. U.); and a Science Achievement Scholarship of Thailand (to P. Chaiyen). The authors declare that they have no conflicts of interest with the contents of this article.

1 To whom correspondence should be addressed: Dept. of Biochemistry, Faculty of Science, Burapha University, 169 Long-Hard Bangsaen Rd., Chonburi 20131, Thailand. Tel.: 66-3810-3058 (ext. 29); Fax: 66-3839-3495; E-mail: somchart@go.buu.ac.th.

2 The abbreviations used are: SHMT, serine hydroxymethyltransferase; ADH, alcohol dehydrogenase; AUC, analytical ultracentrifugation; 5,10-CH2-THF, 5,10-methylenetetrahydrofolate; DHFR, dihydrofolate reductase; hcSHMT, human cytosolic SHMT; hcSHMT/Δflap, flap-deleted hcSHMT; hmSHMT, human mitochondrial SHMT; MD, molecular dynamics; MTHFD, 5,10-methylenetetrahydrofolate dehydrogenase; MW, molecular weight; MW_app, apparent MW; PLP, pyridoxal 5'-phosphate; SAXS, small-angle X-ray scattering; SEC, size-exclusion chromatography; THF, (6S)-tetrahydrofolate; T_m, melting temperature; TS, thymidylate synthase; PDB, Protein Data Bank.
In humans, two isoforms of SHMT, cytosolic (hcSHMT or SHMT 1) and mitochondrial (hmSHMT or SHMT 2), have been proposed as targets for anticancer chemotherapy. Studies to explore the potential of SHMT inhibitors in cancer treatment have been performed at the preclinical level (17). hmSHMT is responsible for maintaining intracellular glycine levels, and the enzyme is expressed at a constant level throughout the cell cycle (18–20). However, in glioma tumor cell proliferation, expression of the mitochondrial isoform was demonstrated to be increased significantly, which was reflected by increased glycine accumulation (21, 22). For hcSHMT, its physiological roles in nucleotide biosynthesis and involvement in cancer proliferation have been reported. Its expression level is increased rapidly during cell proliferation, especially in cancer cells (19, 23, 24). Therefore, a better understanding of the structure and mechanism of SHMT is valuable for the development and discovery of better inhibitors or agents to decrease SHMT activity that may be promising candidates for anticancer drugs.

Recent reports revealed that the human cytosolic SHMT displays distinct biochemical and kinetic properties as compared with homologous Plasmodium enzymes (10, 25). These enzymes possess different quaternary structures, homotrimer for hcSHMT and homodimer for Plasmodium SHMT. The turnover number ($k_{cat}$) of the THF-dependent hcSHMT reaction is much faster than that of the Plasmodium enzymes by ~30-fold. The catalytic activity of hcSHMT can be inhibited by high concentrations of THF under low-pH conditions (6.6, 7.1, and 7.5). In contrast, at higher pH values (7.9 and 8.3) THF substrate inhibition was not significant (10, 25). Conversely, substrate inhibition by THF in the reactions of the Plasmodium enzymes is pH-independent and requires much higher concentrations for inhibition (>0.4 mM) (7, 10, 26). Previous studies on the burst kinetics of the reaction indicated that the rate-limiting step of the overall reaction of hcSHMT is glycine product release (10), whereas for the overall reaction of the Plasmodium enzyme, PvSHMT, the rate-limiting step is glycine formation (3).

Structural comparison between human cytosolic (PDB code 1BJ4) (27) and Plasmodium SHMTs (PDB codes 4O6Z and 4PFF) (28, 29) revealed that each subunit of hcSHMT tetramer possesses a unique β-hairpin structure, or “flap motif,” comprising 13 amino acids (PDB code 273VKSVDPKTGKEIL285) located on top of the THF-binding site (Fig. 1). The flap is presented by two flap-connected regions, β-strand (263–272) and α-helix (286–302), linking to the functional sites of the dimer neighboring subunit. This flap motif is conserved only among the mammalian cytosolic SHMTs, such as those from rabbit (PDB code 1LS3) (28, 30), mouse (PDB code 1EJI) (28, 31), and sheep liver enzymes (30, 32). However, the dimeric structures of Plasmodium and bacterial SHMTs do not contain the flap motif (28, 33, 34). Interestingly, alternative splicing in hcSHMT with flap deletion in some cancer cells was reported, but the role has not been extensively investigated (19, 35).

Here, we characterized the functional role of the flap motif in hcSHMT by comparing biochemical and biophysical properties of the WT and the flap-deleted variant (hcSHMTΔflap). Several analytical techniques, namely size-exclusion chromatography (SEC), analytical ultracentrifugation (AUC), small-angle X-ray scattering (SAXS), molecular dynamics (MD) simulations, and ligand-binding and enzyme kinetic studies were employed. Results revealed that the flap motif is important for the binding affinity of THF and crucial for maintaining the dimer integrity for dimer–dimer assembly into tetramer. Taken together, the flap involves entrapping of the glycine product; thereby the product release is the rate-limiting step.
Functional roles of a flexible flap motif of hcSHMT

Results

Expression, purification, and molar absorptivity of hcSHMT/Δflap

The expression and purification of the hcSHMT/Δflap were carried out similarly to procedures described previously for the WT (10, 25) to obtain 220 mg of purified protein/liter of culture with a specific activity of 4.25 units/mg of protein (Fig. S1A and Table S1). The purified hcSHMT/Δflap exhibits a molecular weight of 52 kDa as shown in SDS-PAGE analysis (Fig. S1B) and absorption characteristics typical of a PLP-bound enzyme with a maximum absorption wavelength of 430 nm, similar to the WT (Fig. S1C) (25). The results indicated that removal of the flap motif does not influence PLP absorption. The molar absorption coefficient (ε_{430}) of the hcSHMT/Δflap-bound PLP was determined to be 7.2 ± 0.2 mM^{-1} cm^{-1}, similar to that of the WT (8.0 ± 0.1 mM^{-1} cm^{-1}) (25), suggesting that the molar absorption coefficient of the PLP cofactor residing within the active site of the hcSHMT is not affected by deletion of the flap motif.

The oligomeric states of the hcSHMT/Δflap

To explore whether the flap motif is necessary for stabilization of the oligomeric structure or intersubunit interactions of hcSHMT, three analytical techniques, including SEC, AUC, and SAXS, were employed to determine the oligomeric species and molecular weights (MWs) of hcSHMT/Δflap and compare them with those of WT. The results obtained from these techniques are summarized in Table 1.

The SEC revealed that the purified hcSHMT/Δflap was eluted into two peaks with a similar peak area at 12 and 13.5 ml, corresponding to MWs of 206 and 96 kDa (Figs. 2A and S1D). The results indicated that hcSHMT/Δflap coexists in tetrameric and dimeric forms, which clearly differ from the WT showing almost a single peak at 12 ml of 206 kDa for tetramer and a very small fraction of 96-kDa dimer (Fig. 2B). It is noted that only the 206-kDa peak (tetramer) in both enzymes showed absorption at 430 nm (Fig. 2A, dashed line), suggesting that only the tetrameric form contains bound PLP, whereas the dimeric enzyme from hcSHMT/Δflap exists in an apo form. The melting temperature (T_m) of the apo-dimeric hcSHMT/Δflap form was determined to be 57 °C, lower than those (T_m, 65 °C) of the holo-tetrameric form of both hcSHMT/Δflap and WT. The results suggested that holo-tetramer is more stable than the apo-dimeric form.

The apparent MWs (MW_{app}) of hcSHMT/Δflap (containing both apo-dimeric and holo-tetrameric forms) and WT were further investigated using AUC scans at 280 nm. The plots between continuous size c(s) distribution and sedimentation coefficient (S) using SEDFIT program (36, 37) revealed two distinct ranges of S values, 6–8 and 9–11 (Fig. 2A, inset), corresponding to the observed MW_{app} of 121 and 195 kDa, likely consistent with the dimeric and tetrameric states of the hcSHMT/Δflap, respectively. On the contrary, the WT showed S value of 10–12 corresponding to an observed MW_{app} of 220 kDa (tetrameric form) (Fig. 2B, inset).

For SAXS studies, X-ray scattering data at two different concentrations of the purified hcSHMT/Δflap (4.5 and 9 mg/ml) (Fig. 2C) or WT (9 and 18 mg/ml) (Fig. 2D) were analyzed. Guinier analysis of the X-ray scattering patterns of the hcSHMT/Δflap (Fig. 2C, inset) or WT (Fig. 2D, inset) can be fitted to a linear relationship, indicating no aggregation or interparticle interference. The radius of gyration (R_g) and maximal particle dimensions (D_{max}) obtained from the Guinier slope and pair distribution function (P(r)) analysis (Fig. S2A) were similar between hcSHMT/Δflap (R_g, 40.5–41.2 Å; D_{max}, 127–128 Å) and WT (R_g, 39.4–39.7 Å; D_{max}, 128 Å). Moreover, Kratky analysis suggested similar compactness of the two enzymes (Fig. S2B), suggesting that both enzymes adopt similar compact globular structures. From the above analysis, the size and shape of the two enzymes were similar; however, the calculated MW (see more details under “Experimental procedures”) of the purified hcSHMT/Δflap (156 kDa) was lower than that of the WT (173 kDa). The oligomeric-state analysis of each X-ray scattering curve (Fig. 2E) as fitted by the Oligomer program (38) indicated that the hcSHMT/Δflap (Fig. 2E, red line) contained the mixed forms of dimer and tetramer at 17 and 83%, respectively. The data firmly support that hcSHMT/Δflap exists as mixed oligomers, whereas the WT (Fig. 2E, blue line) showed

| Parameters | hcSHMT/Δflap variant | Wildtype |
|------------|----------------------|---------|
| MWs (kDa)  |                      |         |
| SEC        | Dimer 96*             | ND      |
|           | Tetramer 206*         | 206*    |
| AUC        | Dimer 121*            | ND      |
|           | Tetramer 195*         | 220*    |
| S1X        | Dimer 156*            | ND      |
|           | Tetramer 173*         | 173*    |
| 1-Serine and THF as substrates*<br>\( k_m (s^{-1}) \) | 11.3 ± 0.3 | 19.8 ± 0.8* |
|           | \( K_m (mM) \)       | 0.32 ± 0.05 | 0.22 ± 0.04* |
|           | THF \((x10^{-3})\)   | 19 ± 3 | 18 ± 3 |
|           | 1-Serine \((s^{-1} \text{mM}^{-1})\) | 35 | 90 |
| THF       | 595                  | 1,100* |
| \( K_{THF}^D (mM) \) | 0.65 ± 0.09 | 0.27 ± 0.08* |
| 1-allo-Threonine as a substrate*<br>\( k_m (s^{-1}) \) | 0.42 ± 0.02 | 0.44 ± 0.01 |
|           | \( K_m (mM) \)       | 0.43 ± 0.07 | 0.63 ± 0.02 |
|           | \( k_m/K_m (s^{-1} \text{mM}^{-1}) \) | 0.98 | 0.70 |
| Dissociation constant \((K) (\mu M)^{-1}\) | 108 ± 8 | 35 ± 8 |

* Apoenzyme form obtained from SEC.
* Holoenzyme form obtained from SEC.
* The calculated MW of the purified hcSHMT/Δflap (a mixture of dimer and tetramer at 17 and 83%, respectively, from Oligomer program).
* The calculated MW of wildtype at 100% tetrameric form from Oligomer program.
* The reactions were performed in buffer D.
* Data were obtained from previous report (10).
* \( K_{THF}^D \) value was determined by Equation 3 using the highest concentration of 1-serine at 6.4 and 3.2 mM for the hcSHMT/Δflap and wildtype, respectively.
* The reactions were performed in buffer A.
only a tetramer. Furthermore, the \textit{ab initio} modeling was carried out by two scattering curves of apo-dimeric and holo-tetrameric forms prepared by SEC to give a visual representation of the SAXS data. The \textit{ab initio} modeling was done using the DAMMIF server (39) and DAMAVER (40) with 10 independent bead models without imposed symmetry (P1) of each protein. The models illustrated the mixed forms of apo-dimeric and holo-tetrameric hcSHMT/Δflap (Fig. S3). Only the holo-tetrameric form of hcSHMT/Δflap showed the same symmetrical shape as that of the WT (Fig. S3A). Based on the three analytical techniques demonstrated here, the data clearly support the presence of mixed dimeric and tetrameric forms in the hcSHMT/Δflap, whereas the WT exists mainly in a tetrameric form.

\textbf{Figure 2.} Analysis of oligomeric states of hcSHMT/Δflap compared with WT. The elution profile of purified hcSHMT/Δflap (A) and WT enzymes (B) obtained from SEC are shown. The absorptions at 280 and 430 nm are presented in solid and dashed lines, respectively. The peaks in A at 12 and 13.5 ml at 280 nm correspond to MWs of 206 (tetramer) and 96 (dimer) kDa, respectively. The dimer peak lacks a PLP characteristic peak at 430 nm, indicating the presence of apo-dimer. The insets of A (hcSHMT/Δflap) and B (WT) represent the plots of continuous size (c(s)) distribution versus sedimentation coefficient (S) from AUC scans at 280 nm. The observed MWs are 121 and 195 kDa for hcSHMT/Δflap and 220 kDa for WT. The SAXS X-ray scattering patterns and Guinier analysis (insets) at two different concentrations of purified hcSHMT/Δflap (a mixture of apo-dimer and holo-tetramer) (C) and WT (D) indicate no aggregation. E, X-ray scattering patterns of hcSHMT/Δflap (4.5 mg/ml; black) and WT (9 mg/ml; gray) fitted by the Oligomer program. hcSHMT/Δflap (red) contained a mixture of 17 and 83% dimer and tetramer, whereas the WT (blue) showed only tetramer.
Influence of PLP binding to the shift in equilibrium between the dimer and tetramer of hcSHMT/H9004 flap

Interestingly, the dimeric form of hcSHMT/H9004 flap lost PLP binding, probably influenced by the absence of the flap motif. However, the tetramer was present with bound PLP. We hypothesized that enzyme tetramerization may require PLP binding. Therefore, the shift in dimer–tetramer equilibrium of the hcSHMT/H9004 flap in the presence of excess PLP was monitored by SEC, AUC, and SAXS.

The SEC results showed that addition of 5-fold excess PLP could convert apo-dimers to holo-tetramers (Fig. 3, A and B). Consistently, the equilibrium shift from apo-dimer to holo-tetramer upon addition of PLP was observed in AUC analyses. The holo-tetramer to dimer ratio was 1:2 for apo-dimer and 3:1 for PLP-treated apo-dimer (Fig. 3, A and B, insets). Therefore, PLP binding can promote the formation of the holo-tetramer.

Results from SAXS analysis also indicated that the overall X-ray scattering patterns and Guinier analysis shifted more toward the tetrameric form in the presence of PLP (Fig. 3C). The $R_g$ and $D_{max}$ values of the apo-dimer were determined from Guinier (Fig. 3C, inset) and $P(r)$ analysis (Fig. S2C) as 34.1 and 109–110 Å, respectively. The addition of PLP into the apo-dimer solution altered the scattering patterns (Fig. 3C, green line) to have higher values of $R_g$ of 43.3 Å and $D_{max}$ of 134 Å. These values are similar to the values for the holo-tetramer enzyme separated by SEC ($R_g$ 39.5 Å; $D_{max}$ 129–130 Å). These data were notably different from the values of the apo-dimer. In addition, Kratky analysis of the apo-dimer also showed a more rigid shape than that of the tetrameric enzyme (Fig. S2D). The $ab initio$ model of the PLP-reconstituted enzyme was similar to that of the holo-tetramer fraction and the WT (Fig. S3D). Taken together, these results emphasize that the binding of PLP to apo-dimer can shift equilibrium toward the tetrameric form of the hcSHMT/H9004 flap.

Equilibrium binding of PLP to holo-tetramer of hcSHMT/H9004 flap

With the truncation of the flap motif, the tetrameric form has only been found as a PLP-bound enzyme in hcSHMT/H9004 flap and not as the apo form. To study the effect of the flap motif on PLP binding, we determined the equilibrium binding of PLP to the tetramer via ultrafiltration by collecting the PLP released from the PLP–tetramer at equilibrium. The results showed that only a small amount of released PLP could be detected in the filtrate fraction (Fig. S4). The calculated $K_d$ for the PLP–tetramer was $0.228 \pm 0.005$ μM, whereas the affinity of PLP to the WT ($K_d 0.010 \pm 0.005$ μM) is 23-fold greater than that to the hcSHMT/H9004 flap (25). This indicates that the flap motif has an intrinsic function to promote PLP binding. Furthermore, the binding of PLP reconstitutes Δflap-dimer integrity proper for tetrameric formation.
Use of molecular dynamics simulations to investigate the functional role of the flap motif in protein dynamics

The dimeric hcSHMT/Δflap isolated by SEC was mainly obtained in the apo form, whereas the isolated tetrameric hcSHMT/Δflap was present in the holo form (see results in Fig. 2A). Based on SANS results in Figs. 3C; S2, C and D; and S3, B and C, the X-ray scattering curve, Guinier plot, Kratky, and P(r) analyses as well as the ab initio models, which can differentiate between dimer and tetramer, the data indicated that the isolated apo-dimer mainly existed in that form without any significant formation of the apo-tetramer. Using the same analysis, the isolated holo-tetramer also mainly existed in that form without any significant amount of the holo-dimeric form. Moreover, the experiment in which we probed the effect of PLP on the formation of tetramer confirms the equilibrium between the apo-dimer and holo-tetramer (Figs. 3C and S3D). Therefore, to determine whether the flap motif has a role in controlling protein dynamics, 100-ns MD simulations of hcSHMT/Δflap were carried out for four species, apo-dimer, holo-dimer, apo-tetramer, and holo-tetramer, to assess protein mobility compared with the WT. The output of the simulation is the relationship between the structural integrity and the B-factor, which indicates the thermal motion of the molecule.

The results showed that the obligate tetramer of apo-hcSHMT/Δflap has much greater thermal motion (high B-factor) between dimer–dimer interfaces than the holo forms (Fig. 4). This suggested that loss of the flap motif induced high dynamic disorder or less compact dimers, probably imposed by the high mobility of the flap-connected region (see also Movies S1 and S2), and reduction of tetramer integrity, leading to dissociation into a more thermodynamically stable apo-dimer. Unlike the hcSHMT/Δflap, the MD simulations for the WT showed similar thermal motion for both apo and holo forms of the dimer and tetramer (Fig. S5). Moreover, the B-factors were lower than those of the hcSHMT/Δflap, indicating tightly packed domains in the WT.

MD simulations showed interesting results to explain the structural integrity of hcSHMT/Δflap in that, in the absence of bound PLP, the dynamic disorder of the apo-tetramer is much greater, making it difficult to maintain the structural integrity of the tetramer. High motion between the protein–protein interface would lead to worse tetramer packing and therefore promote release to the more stable apo-dimer form (Fig. 4 and see also Movies S1 and S2). Accordingly, a significant portion of the hcSHMT/Δflap existed in the apo-dimeric form. These results agreed very well with the SEC and AUC results (Fig. 2A), which showed that the oligomeric equilibrium is driven toward the apo-dimer when the supply of PLP is not in excess. However, once PLP is added in excess, the equilibrium is shifted toward the holo-tetramer due to the formation of a more tightly packed dimer induced by stabilization of the structure by the bound PLP (Fig. 4). In contrast, the WT does not require PLP for tetrameric stabilization (Fig. S5). The results support that the human flap motif plays a crucial role in stabilizing the obligate tetrameric structure found naturally for hcSHMT.

In addition to stabilization of the tetrameric structure of hcSHMT, MD simulation showed that the flap motif promotes retention of THF in the binding pocket (see Movies S3 and S4).
Without the flap, THF can be easily released, thereby lowering the binding affinity of THF (see Movie S3). In contrast, in the WT, the flap motif stabilizes THF retention in the pocket (see Movie S4).

Figure 5. Kinetic studies of the hcSHMT/Δflap reactions. A, two-substrate steady-state kinetics of hcSHMT/Δflap (red) and WT enzyme (blue) reactions at pH 7.5. The plot of \( \frac{v}{e} \) versus THF concentrations (0.025–1.6 mM for hcSHMT/Δflap and 0.005–0.16 mM for WT) at fixed L-serine concentrations (0.1–6.4 mM for hcSHMT/Δflap and 0.1–3.2 mM for WT) is shown. Each fit curve represents low to high concentrations of L-serine with relative increasing velocity. Inset a of A, the apparent kinetics of the hcSHMT/Δflap at pH 6.5–8.5 using 0.025–1.6 mM THF and 6.4 mM L-serine. Inset b of A, pH-activity profile (pH 6–10) of hcSHMT/Δflap (red filled circles) and WT enzyme (blue filled squares) at pH 7.0. B, kinetics of glycine product formation of hcSHMT/Δflap (red filled circles) and WT enzyme (blue filled squares) at pH 7.0. C, THF-independent aldol cleavage of L-αllo-threonine (0.1–10 mM) of hcSHMT/Δflap (red filled circles) and WT (blue filled squares) enzymes measured by an SHMT-ADH coupled assay. Error bars represent S.D. (or S.E.).

**Kinetics of the hcSHMT/Δflap reactions**

To identify the catalytic steps for which the flap motif controls, initially, steady-state kinetics of the hcSHMT/Δflap reaction using L-serine (0.1–6.4 mM) and THF (0.025–1.6 mM) as substrates were carried out at pH 7.5, similar to that described previously for the WT (10). The Michaelis–Menten plot revealed that substrate inhibition was apparent when the THF concentration was >0.2 mM (Fig. 5A, red solid lines). This substrate inhibition is less severe than that observed for the WT (>0.08 mM) (Fig. 5A, blue dashed lines) (10). The double-reciprocal plots gave intersecting lines (Fig. S6), indicating that the hcSHMT/Δflap also employs ternary-complex kinetics similar to the reaction catalyzed by the WT (10, 25). Steady-state kinetic parameters were determined and are summarized in Table 1. The \( k_{cat} \) value of hcSHMT/Δflap was 11.3 ± 0.3 s\(^{-1}\), about half of that of the WT (19.8 ± 0.8 s\(^{-1}\)) (10). \( K_m \) values for L-serine and THF for the hcSHMT/Δflap were similar to those of WT. The resulting \( k_{cat}/K_m \) value of the hcSHMT/Δflap was 2–3-fold lower than those of the WT, indicating that hcSHMT/Δflap has lower catalytic efficiency. Therefore, in addition to its effects on protein quaternary structure, structural dynamics, and PLP affinity, deletion of the flap motif also decreases the catalytic efficiency of the enzyme.

Furthermore, the fact that THF inhibition depended on the pH in the WT prompted us to study this effect in hcSHMT/Δflap. We found that THF inhibition in hcSHMT/Δflap showed a tendency similar to the WT in that the inhibition was more pronounced at low pH (Fig. 5A, inset a) (10). The apparent THF inhibition constants (\( K_i^{THF} \)) increased from 0.41 to 1.9 mM upon increasing pH from 6.5 to 8.5. However, the \( K_i^{THF} \) values of the hcSHMT/Δflap are larger than those obtained from the WT (10), suggesting that the THF inhibition is less sensitive in the hcSHMT/Δflap. These data indicate that the flap motif plays a role in THF inhibition, previously hypothesized to be involved in the slow release of product that allows THF binding to form the hcSHMT-Gly-THF dead-end species (10).

The activity of hcSHMT/Δflap investigated at various pH values (6.5–8.5) showed a half-bell-shaped plot, similar to the WT (Fig. 5A, inset b) (25). However, the \( pK_a \) value for the reaction of the hcSHMT/Δflap was 8.8 ± 0.1 (Fig. 5), 1 pH unit higher than that of the WT (7.8 ± 0.2) (25). Shifts in the \( pK_a \) values of ionizable groups in proteins are typically observed upon mutation or truncation (41). The data imply that, upon the removal of flap motif, a key catalytic residue of SHMT that should be protonated to participate in the catalysis has a raised \( pK_a \). In the hcSHMT active site, there are two ionizable residues, Lys-257 and Glu-75. Lys-257 is the active lysine that forms a Schiff base with the PLP cofactor, and it needs to be deprotonated to be active. The \( pK_a \) of Lys-257 is unlikely changed. Glu-75 has been proposed to act as the general acid required to protonate the reaction intermediate to facilitate the product formation upon C–C bond cleavage (2). Therefore, Glu-75 may be a candidate residue responsible for the \( pK_a \) observed in the pH-rate profile and is the residue with the \( pK_a \) shift in the hcSHMT/Δflap. It is known that \( pK_a \) values of Glu in enzyme active sites can vary over a wide range from 2 to 9 (41).
A previous study showed that the glycine release is the rate-determining step of hcSHMT (10). To investigate the implication of flap motif on the glycine release step, we compared pre-steady-state kinetics of hcSHMT/Aflap and WT. The kinetics of glycine product formation for the hcSHMT/Aflap (Fig. 5B, red circles) is clearly different from that of WT (Fig. 5B, blue squares). No burst kinetics was observed for hcSHMT/Aflap reaction, whereas the WT showed burst kinetics, and 41 ± 17 μM glycine formed in the burst phase (0–0.5 s), resulting from the first turnover of the WT reaction. Therefore, in the WT enzyme, the product release is the rate-limiting step as shown previously (10). In contrast, the rate-limiting step of hcSHMT/Aflap is not the product release step. Other processes such as substrate binding or the catalytic steps prior to the product release may be slower than the product release and become the rate-limiting step instead. Taken together, the results indicate that the flap controls the glycine release step.

Removal of the flap motif does not interfere with THF-independent SHMT activity

Because hcSHMT can catalyze aldol cleavage reaction of β-hydroxyamino acid (1, 19, 25), whether the flap interferes with the THF-independent reaction, the conversion of l-allo-threonine to acetaldehyde and glycine, was studied in hcSHMT/Aflap and WT. They showed similar $k_{cat}$ and $K_m$ (Fig. 5C and Table 1), indicating that the flap does not interfere with THF-independent SHMT activity.

Direct evidence showing that the flap motif is involved in THF binding

The above sections suggest the role of the flap motif in controlling glycine release in THF-dependent catalysis by retaining THF binding. Therefore, the dissociation constant of THF was measured to validate the higher affinity of THF in the WT.

THF can bind to both WT and hcSHMT/Aflap (Fig. S7) to form an E-THF binary complex with a maximum absorbance at 490 nm. The $K_d$ values were calculated to be 108 ± 8 and 35 ± 8 μM for the hcSHMT/Aflap and WT, respectively (Fig. 6). This indicated that the flap enhances the binding affinity of THF by 3-fold, agreeing well with the MD results (see Movies S3 and S4).

Functional roles of a flexible flap motif of hcSHMT

Discussion

This study is the first to elucidate the role of the hcSHMT flap motif, a unique β-hairpin structure, that is conserved among mammalian SHMTs. Although the first structure of human SHMT was elucidated almost two decades ago (19, 27), the key observation that the loop is unique only to mammalian SHMTs was only recognized when the structures of SHMTs from *Plasmodium falciparum* and *Plasmodium vivax* were elucidated (28, 29).

The importance of the flap motif in hcSHMT in stabilizing the tetrameric structure and PLP and THF binding has been demonstrated here. These properties are crucial for hcSHMT activity. We showed that the truncation of flap resulted in an apo-dimer, loss of PLP binding, and change of the rate-determining step. Lack of the flap affected the $pK_a$ of the ionizable groups, particularly Glu-75, which may be a general acid in the SHMT reaction. Binding of PLP can shift the equilibrium from apo-dimer to holo-tetramer. This situation is similar to human ornithine δ-aminotransferase in which PLP stabilizes the tetrameric structure (42).

The presence of the flexible loop to control oligomerization, cofactor and substrate binding, and product release is crucial for the protein evolution point of view. Within the SHMT family, the lack of the flap motif results in a dimeric structure for SHMTs in prokaryotes such as *Escherichia coli* (33) and *Bacillus stearothermophilus* (34) and some eukaryotes such as *P. vivax* and *P. falciparum* (3, 7, 26, 28, 29), whereas in higher eukaryotes like human and rabbit the structure is tetrameric with the presence of flap (27, 30). In other enzyme families, a flexible loop/flap region is present in the *Plasmodium* bifunctional DHFR-TS enzyme from a unique insert (so-called Insert 1) and stabilizes the domain attachment between DHFR and TS in the homodimeric structure of the enzyme (43). In other organisms in which the DHFR lacks this flap region, such as human and *E. coli*, DHFR and TS exist as separate enzymes (44). A well-known flap motif is also found in HIV protease, which is well-recognized in its functional roles in pocket formation, ligand binding, and importantly enzyme catalysis in the enzyme homodimer (45).

Our findings about the THF inhibition and binding in hcSHMT/Aflap may explain why hcSHMT mRNA splicing takes place in MCF-7 breast and SH-SY5Y neuroblastoma cancer cells. The alternative mRNA splicing in cancer cells was found in exons 2, 9, and 10 (19, 35), and the splicing of exon 9 leads to production of hcSHMT without the flap motif (residues 273–287). The resultant diminished THF affinity likely raises drug resistance against antifolate chemotherapy. Another example of this is the observed overexpression of alternatively spliced variants of folylpolyglutamate synthase mRNAs in leukemia cells, which causes antifolate anticancer drug resistance in acute lymphoblastic leukemia (46, 47).

In conclusion, these findings provide mechanistic insights into how a flap motif can influence the overall structural architecture and catalytic activity of hcSHMT. Understanding the role of the flap motif paves the way for development of more effective antifolate drugs for anticancer chemotherapy in the future. The results reported here also contribute to the understanding of how a flexi-
Functional roles of a flexible flap motif of hcSHMT

ble loop can take part in stabilization of oligomeric structure and ligand binding in other enzymes in general.

Experimental procedures

Chemicals and reagents

All chemicals and reagents used in this study were analytical grade and of the highest purity commercially available and were prepared as described in previous reports (10, 25). Buffers used throughout this report were (i) buffer A, 10 mM HEPES buffer, pH 7.5, containing 100 mM NaCl, 1 mM DTT, and 0.5 mM EDTA; (ii) buffer B, 10 mM HEPES buffer, pH 7.5, containing 150 mM NaCl; (iii) buffer C, 10 mM HEPES buffer, pH 7.5, containing 100 mM NaCl; and (iv) buffer D, a three-component buffer system (AME buffer, pH 7.5, composed of 0.05M acetic acid, 0.05M MES, and 0.1 M N-ethylmorpholine) containing 1 mM DTT and 0.5 mM EDTA.

Construction of an expression plasmid containing the hcsmt/Δflap gene

An expression plasmid carrying the hcsmt/Δflap gene was generated by site-directed mutagenesis using pET100D/TOPO-hcsmtΔH (25) as a template in a primer of 5′-GATCCTCTACAGGAAAGGAGTGAAATACAACCTG-GAGTCTCTTATCAATTC-3′ and 5′-GAATTGATAAGAGACTCCAGGTGTTATTTCCACTCCTCCGTAGAGG-ATC-3′. The plasmid was sequenced at the 1st Base DNA sequencing service (Malaysia) to verify the deletion of the flap sequence. The expression plasmid was named pET100D/TOPO-hcsmt/Δflap. The sequences encoding 12 amino acids (274KSVDPKTGKEIL285) of the flap motif was deleted to remove the β-hairpin that makes up for the flap motif. Although the overall flap motif includes residues 273–285, we decided to retain residue 273 in the variant to prevent deterioration in protein folding.

Enzyme expression, purification, and assay

The overexpression of the hcsmt/Δflap in E. coli BL21(DE3) was carried out at 16 °C in ZY autoinduction medium system (1% (w/v) peptone, 0.5% (w/v) yeast extract, 5 mM Na2SO4, 2 mM MgSO4, 1X NPS (25 mM Na2HPO4, 25 mM KH2PO4, and 50 mM NH4Cl), and 1X 5052 (0.5% (w/v) glycerol, 0.05% (w/v) D-glucose, and 0.2% (w/v) alpha-lactose) as described previously (10, 25). The purification protocol for the hcsmt/Δflap was performed similarly to that for the WT (10, 25) but with slight modifications. In brief, the hcsmt/Δflap was purified to homogeneity using 1% (w/v) polyethyleneimine and 30–50% (w/v) ammonium sulfate precipitations and diethylaminoethyl (DEAE)-Sepharose and Sephacryl S-200 gel-filtration columns. The buffer system used for enzyme purification and storage was buffer A.

The activity of the hcsmt/Δflap was measured at 25 °C in buffer A and carried out using an SHMT–MTHFD coupled assay method (10, 25) except that 0.2 mM THF and 0.1 μM hcsmt/Δflap were added. The reaction progression of NADPH production was monitored at 375 nm by a diode array spectrophotometer (Hewlett Packard). One unit of SHMT activity is defined as the formation of 1 μmol of NADPH/min at pH 7.5 and 25 °C.

Determination of the molar absorption coefficient of hcsmt/Δflap-bound PLP under denaturing condition

The hcsmt/Δflap was denatured by SDS, and the amount of PLP cofactor liberated was spectroscopically measured and used to determine the molar absorption coefficient of the hcsmt/Δflap. The molar absorption coefficients of PLP and the hcsmt/Δflap-bound PLP were determined in the presence of SDS as described previously (7, 26) except that a final concentration of 2% (w/v) SDS was employed.

Analyses of the MWs, oligomeric states, and quaternary structures

Three analytical techniques, including SEC, AUC, and SAXS, were used to analyze MWs, oligomeric states, and quaternary structures of the hcsmt/Δflap. By the SEC technique, the native MW and oligomeric states of the hcsmt/Δflap (77 μM) were determined using a SuperdexTM 200 Increase 10/300 GL (GE Healthcare) gel-filtration column performed with an AKTA FPLC system (GE Healthcare). The column was calibrated with the known MWs of protein standards (GE Healthcare). All proteins were eluted with buffer B at 25 °C at a constant flow rate of 0.5 ml/min, and the elution volume (Ve) for each protein was measured. The absorbance at 280 nm (A280) was monitored for the proteins, whereas A430 was monitored for bound PLP of the hcsmt/Δflap. The oligomeric form of the hcsmt/Δflap was estimated based on the MW of the subunits of the hcsmt/Δflap for which the amino acid sequence was calculated using the ProtParam program on the ExPaSy Proteomics Server (https://web.expasy.org/protparam/).

For size distribution analysis, peaks corresponding to the apo-dimer, holo-tetramer, and mixed oligomers (as purified form) as separated by Superdex gel filtration and detected by optical detectors (Beckman Coulter, ProteomeLab XL-I) were analyzed by AUC. The hcsmt/Δflap sample (20 μM) was dialyzed overnight and diluted into buffer A. The protein sample (400 μl) and buffer A (425 μl) were injected in each side of a double-sector centerpiece cell. The double-sector cells were placed into an An-60 Ti rotor and then centrifuged at 20,000 rpm at 20 °C for 16 h. The spectrum of each protein sample was obtained continuously at 280 nm using a time interval of 600 s/scan. The sedimentation velocities obtained from multiple scans at 280 nm and at different time intervals were fitted to a continuous c(s) distribution model using the SEDFIT program (36, 37) to obtain the corresponding MW. The partial-specific volume of the protein sample was set as 0.73 ml/g, based on a solvent density of 1.035 g/ml (48).

For SAXS analysis, the experiments were measured using a multipole wiggler source on Beamline 1.3W (BL13W: SAXS) of the Synchrotron Light Research Institute (Public Organization), Thailand. A MarCCD Rayonix SX165 detector with a sample-to-detector distance of 2207 mm was used, and at least two different concentrations (3–20 mg/ml) of each protein sample (apo-dimer, holo-tetramer, the as-purified form of hcsmt/Δflap, and WT in the absence and presence of 5-fold excess PLP) were exposed to X-ray emission (λ = 1.38 Å) at 16 °C for 10 min. This enabled us to capture SAXS data with a q range of 0.022 < q < 0.32 Å−1. The empty sample cell was
measured first followed by the matched reference buffer and the protein sample, respectively. The 2D images were reduced and radially averaged by the program SAXSIT, which was developed by Synchrotron Light Research Institute staff, and BioXTAS RAW (49) to obtain 1D scattering curves. Guinier and Kratky analyses were calculated from the scattering patterns using functions of BioXTAS RAW (49). Pair distribution function ($P(r)$) analysis was calculated from the scattering curves using the GNOM program (50) from ATSAS package version 2.8.2 (51). The MW of each protein was calculated by using a zero extrapolatedintensity

**Functional roles of a flexible flap motif of hcSHMT**

The AMBER ff14SB force field (58) was adopted for the protein, whereas the atom types and the other molecular parameters of PLP were assigned by general AMBER force field version 2 (GAFF2) (59). The protonated model was energy-minimized using two steps consisting of 2500 steps of steepest descents followed by 2500 steps of conjugated gradient. The model was then solvated by a TIP3P water in an octahedron box within 10 Å from the protein surface, neutralized by adding Na$^+$ ions, and minimized with the two steps described above under a restraint on the protein atoms with a weight of 10.0 kcal/molÅ$^2$ followed by minimization on the whole system. The system was then heated to 298 K at 1-atm pressure and subsequently subjected to MD simulation using the isothermal-isobaric (NPT) ensemble for 100 ns as described previously (60). To indicate protein flexibility and mobility, the B-factor was calculated (Equation 1) for the backbone atoms of all minimized structures at the last 20-ns MD trajectories using the CPPTRAJ module and normalized in a relative scale. Herein, $T$ is a set of frames over the last 5-ns trajectories, $r_i(t_j)$ is the position of atom $i$ at time $t_j$, and $r_{ij}$ is the time-averaged position of the same atom $i$. Note that B-factor values are normalized in a range of 0 (blue for low fluctuation) to 1 (red for high fluctuation).

\[
B\text{-factor} = \left[ \frac{1}{T} \sum_{t_j} (r_i(t_j) - \bar{r}_i)^2 \right] \times \frac{8\pi^2}{3} \quad (\text{Eq. 1})
\]

**Steady-state kinetics of two-substrate reactions**

The steady-state kinetics of the hcSHMT/Δflap were studied at pH 7.5 and 25 °C by an SHMT–MTHFD coupled assay (10). The assays were performed in buffer D containing 250 μM NADP$^+$, 10 μM MTHFD, 0.1 μM hcSHMT/Δflap, and various concentrations of L-serine (0.1–6.4 mM) and THF (0.025–1.6 mM) using stopped-flow spectrophotometry in single-mixing mode under anaerobic conditions. The kinetics of NADPH production was monitored at 375 nm, and the initial velocities ($v$) were calculated. Direct plots of initial velocities versus concentrations of either substrate were fitted to Equation 2 where $V_{\text{max}}$ is the maximum rate, $K_A$ and $K_B$ are Michaelis constants for substrates $A$ and $B$, and $K_{\text{ia}}$ is the dissociation constant of substrate $A$. The double-reciprocal plots of initial velocities versus concentrations of either substrate were plotted to identify the steady-state mechanism as described by Dalziel’s equation. Kinetic parameters of the hcSHMT/Δflap were determined without inhibition of THF substrate and compared with those of the WT.

In addition to pH 7.5, the steady-state kinetics at various pH values (6.5–8.5) and THF concentrations (0.025–1.6 mM) at an L-serine concentration 6.4 mM were carried out to determine the magnitude of THF substrate inhibition at different pH values. The inhibition constant ($K_I$) of the THF substrate was determined using Equation 3, which was analyzed by Marquardt–Levenberg algorithms in KaleidaGraph version 4.0 software (Synergy Software).

\[
v = \frac{V_{\text{max}}[A][B]}{K_A[\text{B}] + [A] + [\text{B}]} \quad (\text{Eq. 2})
\]
**Functional roles of a flexible flap motif of hcSHMT**

\[ \nu = \frac{V_{\text{max}} [S]}{K_m + [S]} \left(1 + \frac{[S]}{K'_c}\right) \]  
(Eq. 3)

**pH-activity profile**

The activity of the hcSHMT/Δflap at different pH values was determined by a coupled enzyme assay method as described previously under “Enzyme expression, purification, and assay.” The buffers were as follows: 100 mM HEPES buffer for pH 6–8.5 and 100 mM carbonate buffer for pH 9–10. Initial velocities were plotted as a function of pH and fitted according to Equation 4 to determine the pH value by using Marquardt–Levenberg algorithms in KaleidaGraph version 4.0 program. Here, \( Y \) is the maximum rate, \( C \) is the maximum rate that depends on the pH, and \( pK'_c \) is the dissociation constant of the acid.

\[ Y = \frac{C}{1 + (10^{-pH}/10^{-pK'_c})} \]  
(Eq. 4)

**Analyses of glycine formation kinetics using rapid-quench techniques and HPLC-MS**

All experiments were performed at pH 7.0 similarly to those reported previously using a rapid-acid quench coupled with HPLC-MS techniques (10) except that the final concentration of THF used for the hcSHMT/Δflap and WT was 0.2 and 0.08 mM, respectively, to avoid THF substrate inhibition. The glycine obtained from the rapid-quench technique was monitored and quantified by HPLC-MS (10). The kinetics of glycine product formation of the hcSHMT/Δflap was then analyzed.

**Activity assay of the THF-independent aldol cleavage reaction**

The aldol cleavage kinetics was performed using \( L \)-allo-threonine as a substrate. The formation of acetaldehyde was detected by coupling the reaction with alcohol dehydrogenase (ADH) (26). The assay reaction contained 100 \( \mu \)M NADH; 3 mg/ml ADH; 1 and 0.5 \( \mu \)M hcSHMT for the hcSHMT/Δflap and WT, respectively; and 0.1–10 mM \( L \)-allo-threonine in buffer A. The assay reactions were monitored for an absorbance decrease at 340 nm, and the initial velocities were determined and plotted as a function of \( L \)-allo-threonine concentration. The plot was fitted by Michaelis–Menten equation and analyzed using Marquardt–Levenberg algorithms and KaleidaGraph version 4.0 software to determine the kinetic parameters.

**Measurement of the equilibrium constants for the binding of enzyme with THF**

To measure the \( K_d \) value for the binding of THF to the hcSHMT/Δflap and WT, the binding reactions were measured by spectrophotometry (DH-2000-TGK/CCD-204 spectrophotometer, Ocean Optics, UK) under anaerobic conditions in an anaerobic glovebox (<5 ppm O$_2$; Belle Technology, UK) to avoid oxidation of THF. A 1-ml anaerobic solution of the enzyme ([P]$_{\text{total}}$ 28 \( \mu \)M; \( A_{430} \approx 0.2 \)) in buffer D was added into a quartz cuvette, and the absorption spectrum of the enzyme was recorded as a baseline for spectrum subtraction after each THF addition. Absorption spectra were recorded after adding THF, and the spectrum of the enzyme alone was subtracted to obtain the difference spectrum. The absorbance change values at 490 nm (\( \Delta A \)) for the ligand-bound enzyme were extracted from the difference spectra derived from 16–319 \( \mu \)M THF ([L]$_{\text{total}}$). The \( \Delta A_{\text{max}} \) was determined from the plot between \( \Delta A \) and various [L]$_{\text{total}}$ and input into Equations 5–7 for calculation of the unbound THF concentrations ([L]$_{\text{free}}$), 6–290 \( \mu \)M. The \( K_d \) was determined from the nonlinear least square using Marquardt–Levenberg algorithms in KaleidaGraph version 4.0 software from the plot between the unbound concentrations of THF (6–290 \( \mu \)M) and \( \Delta A \).

\[ \frac{\Delta A}{\Delta A_{\text{max}}} = \frac{[L]_{\text{free}}}{K_d + [L]_{\text{free}}} \]  
(Eq. 5)

\[ [PL] = \frac{\Delta A}{\Delta A_{\text{max}}} [P]_{\text{total}} \]  
(Eq. 6)

\[ [L]_{\text{free}} = [L]_{\text{total}} - [PL] \]  
(Eq. 7)

**Dye-binding thermal shift assay**

The protein sample (40 \( \mu \)M), apo-dimeric, homo-tetrameric, and mixed oligomeric (as-purified) forms of hcSHMT/Δflap, was mixed with a fluorescent SYPRO Orange dye solution (10\( \times \) (Invitrogen). The fluorescence signal was monitored using a real-time PCR machine with a temperature increase from 25 to 95 °C at a constant interval of 1 °C/min (61). The melting curve of temperature versus fluorescence signal was analyzed and used to determine the T$_{m}$ at which half of the total protein is transitioned to an unfolded state.

**Author contributions**—S. U., P. Chitnumsub, U. L., P. Chaiyen, and S. M. conceptualization; S. U., W. P., N. K., P. W., P. M.-u., T. R., O. K., P. Chitnumsub, U. L., P. Chaiyen, and S. M. resources; S. U., J. J., N. K., P. W., P. M.-u., T. R., O. K., P. Chitnumsub, U. L., P. Chaiyen, and S. M. data curation; S. U., N. K., P. W., P. M.-u., T. R., O. K., P. Chitnumsub, U. L., P. Chaiyen, and S. M. formal analysis; S. U., J. J., W. P., N. K., P. W., T. R., O. K., P. Chitnumsub, U. L., P. Chaiyen, and S. M. visualization; S. U., J. J., W. P., N. K., P. W., P. M.-u., T. R., O. K., P. Chitnumsub, U. L., P. Chaiyen, and S. M. investigation; S. U., J. J., N. K., P. W., P. M.-u., T. R., O. K., P. Chitnumsub, U. L., P. Chaiyen, and S. M. methodology; S. U., J. J., W. P., N. K., P. W., P. M.-u., T. R., O. K., P. Chitnumsub, U. L., P. Chaiyen, and S. M. writing–reviewing editing.
References

1. Schirch, L. (1982) Serine hydroxymethyltransferase. Adv. Enzymol. Relat. Areas Mol. Biol. 53, 83–112 CrossRef Medline

2. Schirch, V., and Szebenyi, D. M. (2005) Serine hydroxymethyltransferase revisited. Curr. Opin. Chem. Biol. 9, 482–487 CrossRef Medline

3. Maenpuen, S., Amornwatcharapong, W., Krasatong, P., Sucharitakul, J., Palfey, B. A., Yuthavong, Y., Chitnumsub, P., Leartsakulpanich, U., and Chaiyen, P. (2015) Kinetic mechanism and the rate-limiting step of Plasmodium vivax: Human and mitochondrial isoforms of human serine hydroxymethyltransferase. J. Biol. Chem. 290, 8656–8665 CrossRef Medline

4. Alfadhli, S., and Rathod, P. K. (2000) Gene organization of a Plasmodium falciparum serine hydroxymethyltransferase and its functional expression in Escherichia coli. Mol. Biochem. Parasitol. 110, 283–291 CrossRef Medline

5. Nirmalan, N., Wang, P., Sims, P. F., and Hyde, J. E. (2002) Transcriptional analysis of genes encoding enzymes of the folate pathway in the human malaria parasite Plasmodium falciparum. Mol. Microbiol. 46, 179–190 CrossRef Medline

6. Leartsakulpanich, U., Kongkasriyachai, D., Imwong, M., Chotivanchin, K., and Yuthavong, Y. (2008) Cloning and characterization of Plasmodium vivax serine hydroxymethyltransferase. Parasitol. Int. 57, 223–228 CrossRef Medline

7. Maenpuen, S., Sopithhumkakhun, K., Yuthavong, Y., Chaiyen, P., and Leartsakulpanich, U. (2009) Characterization of Plasmodium falciparum serine hydroxymethyltransferase:A potential antimalarial target. Mol. Biochem. Parasitol. 168, 63–73 CrossRef Medline

8. Pornthanakasem, W., Kongkasriyachai, D., Utaiapibul, C., Yuthavong, Y., and Leartsakulpanich, U. (2012) Plasmodium vivax serine hydroxymethyltransferase: indispensability and display of distinct localization. Malar. J. 11, 387–395 CrossRef Medline

9. Witschel, M. C., Rottmann, M., Schwert, G., Frei, M. S., Witschel, M. C., Rottmann, M., and Chaiyen, P. (2016) Differential 3-bromopyruvate inhibition of cytosolic and mitochondrial human serine hydroxymethyltransferase isoforms, key enzymes in cancer metabolic reprogramming. Biochim. Biophys. Acta 1864, 1506–1517 CrossRef Medline

10. Ducker, G. S., Gherguvich, J. M., Mainolfi, N., Suri, V., Jeong, S. K., Hsin-Jung Li, S., Friedman, A., Manfredi, M. G., Gittai, Z., Kim, H., and Rabinowits, J. D. (2017) Human SHMT inhibitors reveal defective glycine import as a targetable metabolic vulnerability of diffuse large B-cell lymphoma. Proc. Natl. Acad. Sci. U.S.A. 114, 11404–11409 CrossRef Medline

11. Amelio, I., Cutruzzola, F., Antonov, A., Agostini, M., and Melino, G. (2014) Serine and glycine metabolism in cancer. Trends Biochem. Sci. 39, 191–198 CrossRef Medline

12. Stover, P. J., Chen, L. H., Suh, J. R., Stover, D. M., Keyomarsi, K., and Shane, B. (1997) Molecular cloning, characterization, and regulation of the human mitochondrial serine hydroxymethyltransferase gene. J. Biol. Chem. 272, 1842–1848 CrossRef Medline

13. Snell, K., Baumann, U., Byrne, P. C., Chave, K. J., Renwick, S. B., Sanders, P. G., and Whitehouse, S. K. (2000) The genetic organization and protein crystallographic structure of human serine hydroxymethyltransferase. Adv. Enzyme Regul. 40, 353–403 CrossRef Medline

14. Jain, M., Nilsson, R., Sharma, S., Madhusudhan, N., Kitami, T., Souza, A. L., Kaifi, R., Kirschner, M. W., Chish, C. B., and Moorthy, V. K. (2012) Metabolite profiling identifies a key role for glycine in rapid cancer cell proliferation. Science 336, 1040–1044 CrossRef Medline

15. Kim, D., Fiske, B. P., Birsoy, K., Freinkman, E., Kami, K., Possemato, R. L., Chudnovsky, Y., Pacold, M. E., Chen, W. W., Cantor, J. R., Shelton, L. M., Gyi, D. Y., Kwon, M., Ramkissoon, S. H., Ligon, K. L., et al. (2015) SHMT2 drives glioma cell survival in ischaemia but imposes a dependence on glycine clearance. Nature 520, 363–367 CrossRef Medline

16. Wang, B., Wang, W., Zhu, Z., Zhang, X., Tang, F., Wang, D., Liu, X., Yan, X., and Zhuang, H. (2017) Mitochondrial serine hydroxymethyltransferase 2 is a potential diagnostic and prognostic biomarker for human glioma. Clin. Neurol. Neurosurg. 154, 28–33 CrossRef Medline

17. Torndike, J., Pelliniemi, T. T., and Beck, W. S. (1979) Serine hydroxymethyltransferase activity and serine incorporation in leukocytes. Cancer Res. 39, 3435–3440 Medline

18. Rao, N. A., Talwar, R., and Savithri, H. S. (2000) Molecular organization, catalytic mechanism and function of serine hydroxymethyltransferase—a potential target for cancer chemotherapy. Int. J. Biochem. Cell Biol. 32, 405–416 CrossRef Medline

19. Pinthong, C., Maenpuen, S., Amornwatcharapong, W., Yuthavong, Y., Leartsakulpanich, U., and Chaiyen, P. (2014) Distinct biochemical properties of human serine hydroxymethyltransferase compared with the Plasmodium enzyme: implications for selective inhibition. FEBS J. 281, 2570–2583 CrossRef Medline

20. Sopithhumkakhun, K., Maenpuen, S., Yuthavong, Y., Leartsakulpanich, U., and Chaiyen, P. (2009) Serine hydroxymethyltransferase from Plasmodium vivax is different in substrate specificity from its homologues. FEBS J. 276, 4023–4036 CrossRef Medline

21. Renwick, S. B., Snell, K., and Baumann, U. (1998) The crystal structure of human cytosolic serine hydroxymethyltransferase: a target for cancer chemotherapy. Structure 6, 1105–1116 CrossRef Medline

22. Chitnumsub, P., Ittarat, W., Jaruwat, A., Noytanom, K., Amornwatcharapong, W., Pornthanakasem, W., Chaiyen, P., Yuthavong, Y., and Leartsakulpanich, U. (2014) The structure of Plasmodium falciparum serine hydroxymethyltransferase reveals a novel redox switch that regulates its activities. Acta Crystallogr. D Biol. Crystallogr. 70, 1517–1527 CrossRef Medline

23. Chitnumsub, P., Jaruwat, A., Noytanom, K., Amornwatcharapong, W., Pornthanakasem, W., Chaiyen, P., Yuthavong, Y., and Leartsakulpanich, U. (2014) The structure of Plasmodium vivax serine hydroxymethyltransferase: implications for ligand-binding specificity and functional control. Acta Crystallogr. D Biol. Crystallogr. 70, 3177–3186 CrossRef Medline

24. Scarsdale, J. N., Kazanina, G., Radaev, S., Schirch, V., and Wright, H. T. (1999) Crystal structure of rabbit cytosolic serine hydroxymethyl-
Functional roles of a flexible flap motif of hcSHMT

...