Molecular Basis of the General Base Catalysis of an α/β-Hydrolase Catalytic Triad*

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Background: The α/β-hydrolase MenH uses its Ser-His-Asp triad as a specific general basis with a poorly understood mechanism.

Results: An open-closed conformational change is identified in the enzyme catalysis.

Conclusion: The conformational change controls the formation and reactivity of the catalytic triad.

Significance: The catalytic versatility of the Ser-His-Asp triad is expanded by the open-closed structural change.

The serine-histidine-aspartate triad is well known for its covalent, nucleophilic catalysis in a diverse array of enzymatic transformations. Here we show that its nucleophilicity is shielded and its catalytic role is limited to being a specific general base by an open-closed conformational change in the catalysis of (1R,6R)-2-succinyl-6-hydroxy-2,4-cyclohexadiene-1-carboxylate synthase (or MenH), a typical α/β-hydrolase fold enzyme in the vitamin K biosynthetic pathway. This enzyme is found to adopt an open conformation without a functional triad in its ligand-free form and a closed conformation with a fully functional catalytic triad in the presence of its reaction product. The open-to-closed conformational transition involves movement of half of the α-helical cap domain, which causes extensive structural changes in the α/β-domain and forces the side chain of the triad histidine to adopt an energetically disfavored gauche conformation to form the functional triad. NMR analysis shows that the inactive open conformation without a triad prevails in ligand-free solution and is converted to the closed conformation with a properly formed triad by the reaction product. Mutation of the residues crucial to this open-closed transition either greatly decreases or completely eliminates the enzyme activity, supporting an important catalytic role for the structural change. These findings suggest that the open-closed conformational change tightly couples formation of the catalytic triad to substrate binding to enhance the substrate specificities and simultaneously shield the nucleophilicity of the triad, thus allowing it to expand its catalytic power beyond the nucleophilic catalysis.

The serine (Ser)-histidine (His)-aspartate (Asp) triad is a well known catalytic motif that is utilized in catalysis of a diverse array of chemical transformations by a large number of enzymes. One group of these triad-dependent enzymes includes numerous hydrolytic enzymes in the trypsin and subtilisin families of serine peptidases that cleave C–N bonds in peptide substrates (1). Another group of enzymes utilizing this triad or its variants are members of the α/β-hydrolase fold superfamily that catalyze cleavage of C–O, C–S, C–C, or C–halogen bonds in structurally diverse metabolites or even cofactor-independent haloperoxidation and dioxygenation of electron-rich substrates (2, 3). Consensus with their functional diversity, these enzymes display vast structural diversity in size, structure fold, and three-dimensional architecture. However, they converge to form the conserved triad and the associated oxyanion hole as a stable, integrative catalytic unit from distal residues, which are arranged in one enantiomeric configuration in α/β-hydrolase fold enzymes and in another configuration in members of the trypsin and subtilisin families (3). Besides this structural conservatism, a majority of these enzymes use the triad to catalyze the diverse chemical transformations via a similar nucleophilic mechanism, which is characterized by formation of a covalent enzyme adduct involving nucleophilic addition of the triad serine to the substrate (3, 4).

Besides the traditional nucleophilic mechanism of catalysis, the Ser-His-Asp triad has also been proposed to play the role of a simple general base in the catalysis of some α/β-hydrolases. In the catalysis of cyanohydrin lyases, the triad is postulated to deprotonate hydrogen cyanide for addition to aldehydes and ketones (5, 6) despite the fact that this triad is able to form a catalytically competent complex with trichloroacetaldehyde through nucleophilic addition (6). A similar general base role was also proposed for the triad of C–C bond hydrolases (7–9), which are involved in the degradation pathways of aromatic compounds. However, recent investigations have shown that the C–C bond hydrolase BphD adopts the traditional covalent, nucleophilic mechanism of catalysis because of the finding of an acyl-enzyme intermediate (10, 11). Noticeably, the triad serine in these enzymes has been demonstrated to exhibit nucleophilic reactivity (5, 12). In addition, a nucleophilic mechanism of catalysis...
Recent advances in the understanding of the catalytic mechanism of serine proteases have provided new insights into the catalytic triad of serine proteases. The MenH catalytic triad is indeed capable of new catalytic chemistry. Despite its unique mechanism of catalysis, the MenH triad is not found to be structurally different from that of other triad-utilizing enzymes (Refs. 14, 17, and 18 and Protein Data Bank code 1R3D). In addition, mutational studies found that the MenH catalytic triad most likely catalyzes simple abstraction of a proton from the α-carbon of the succinyl carbonyl group of the substrate to form an enolate intermediate that subsequently undergoes elimination of the enolpyruvyl group in an Elcb mechanism (Fig. 1) (15). This distinctive MenH catalytic mechanism demonstrates that the classical Ser-His-Asp triad is indeed capable of new catalytic chemistry.

The proposed general basis catalysis of 2,5-elimination of pyruvate by the Ser-His-Asp triad of the vitamin K biosynthetic enzyme MenH.

FIGURE 1.
acquired consisting of 128,000 data points with a spectral width of 35 ppm at 4°C. The concentration of MenH or the H232A mutant was fixed at 0.33 mM, and SHHC was titrated into the MenH solution to a concentration of 0.165, 0.33, and 0.66 mM. As a control, one-dimension proton NMR was collected for 0.36 mM SHHC alone. To determine which of the protons at the side chain of His232, a mixture of 0.40 mM 15N-labeled MenH and 0.80 mM SHHC was used for analysis by heteronuclear single quantum coherence spectroscopy. However, no cross-peaks were found probably because of the weak 15N signals.

**Fluorometric Determination of Binding Affinity**—The dissociation constant \(K_D\) for binding of the SHHC ligand by MenH or its H232A mutant was determined by the ligand-dependent quenching of tryptophan fluorescence with a PerkinElmer Life Sciences Model LS-55 luminescence spectrometer. Specifically, the SHHC ligand was titrated into the solution of MenH or the H232A mutant at 2.0 \(\mu\)M in the 25 mM Tris-HCl buffer, pH 8.0. The tryptophan fluorophore was excited at 280 nm, and its emission \(F\) at 340 nm was recorded. The dissociation constant was determined by the least square fitting of the fluorescence-\([SHHC]\) curves according to the equation \(F/F_0 = K_D/(K_D + [SHHC])\) where \(F_0\) is the tryptophan fluorescence of the protein in the absence of any ligands.

**Stopped-flow Kinetics**—Rapid kinetic measurements of either UV-visible absorption or fluorescence emission were performed on an Applied Photophysics SX.18MV-R stopped-flow reaction analyzer mounted with an Applied Photophysics spectral kinetic monochromator. In pre-steady state kinetic measurements, 50-\(\mu\)l aliquots of MenH at 2.0 \(\mu\)M were mixed with an equal volume of SEPHHC solution at concentrations varying from 5.0 to 240 \(\mu\)M. The release of the SHHC product was monitored at 290 nm for a total of 200 ms with a 1-ms dead time. In the kinetic measurement of SHHC binding, 50-\(\mu\)l aliquots of MenH at 1.0 \(\mu\)M were mixed with an equal volume of SHHC at concentrations varying from 0 to 360 \(\mu\)M. Quenching of the tryptophan fluorescence emission at 340 nm was determined over a period of 200 ms as a change of fluorescence \(\Delta F\) at 340 nm relative to the \(F_0\) value.

**Crystallization**—Initial screening of crystallization conditions for ligand-free MenH and MenH in complex with SHHC was performed using hanging drop vapor diffusion with a range of screen kits (Hampton Research) at 293 K. The concentrations of MenH and SHHC were 10 mg/ml and 2 mM, respectively. The protein solution was mixed with the reservoir solution in a 1:1 ratio in these screening trials. Small ellipsoid and rhombohedral crystals were obtained within 1 week under the same conditions for both the complex and the ligand-free protein. After several rounds of optimization, the longest dimension of the crystals reached about 100 \(\mu\)M. However, neither the complex nor the ligand-free protein crystals diffracted to a high resolution on an in-house x-ray diffractometer. After optimization via additive screens using the Crystal Screen kit and Crystal Screen II kit (Hampton Research), cuboid-shaped crystals of ligand-free MenH were observed in solution containing 0.2 M Li2SO4, 0.1 M Tris buffer, pH 9.0, 30% (v/v) glycerol, and 17% PEG 3350 with a dimension of 300 \(\times\) 300 \(\times\) 200 \(\mu\)M. Large, rod-shaped crystals (500 \(\mu\)M for the longest dimension) of the MenH complex were obtained when the initial screens were rechecked 2 months later in solution containing 1.6 mM sodium/potassium phosphate, pH 6.9. The ternary complex was crystallized under the same conditions as for the binary complex with the addition of 2 mM pyruvate. The crystals were mounted and soaked in cryoprotectant containing 20% glycerol in the mother liquor and then flash frozen in liquid N2. For data collection, a cryo-streamer was used with an ADSC Quantum 315R charge-coupled device detector. Diffraction images were indexed, integrated, and scaled using HKL2000 (24). The ligand-free MenH structure was solved by molecular replacement with Phaser (25) in the CCP4 suite (26) using the structure of Vibrio cholerae MenH (Protein Data Bank code 1R3D) as the search model. The initial electron density map indicated that one MenH polypeptide chain was correctly located in the asymmetric unit in space group \(P_{3}1_2_1_2_{1}\). The model was then extended via automatic model building using the program ARP/wARP (27) and then further built manually using Coot (28) followed by refinement using REFMAC (29) and PHENIX (30). The overall quality of the structural model was assessed by PROCHECK (31) and MolProbity (32). Data collection and refinement statistics are summarized in Table 1.

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Based on the diffraction patterns, the MenH-SHHC complex crystal is in the \(P_{3}1_2_1_{2}_{1}\) space group. The crystal structure solved by molecular replacement with Phaser using the ligand-free MenH structure as the search model contained a high solvent content of 66.1% with one-third of the crystal volume occupied by non-interpretable electron densities. Although the unit cell contains MenH molecules with very well defined electron densities, this structure did not appear to be a correct solution. The diffraction data set was then reindexed to the \(P_{3}1_{2}_{1}_{2}_{1}\) space group for structural determination by molecular replacement using Phaser and BALBES (33). The resulting structure contained two well defined MenH molecules in the asymmetric unit, which also contained uninterpreted electron densities in one-third of its volume. A third MenH molecule was successfully generated in the asymmetric unit after model building by OASIS (34) albeit with poor and disconnected electron densities. By applying the merohedral twin law \((h, -h-k, -l)\), the model was eventually refined by Phenix.refine to a high resolution with \(R_{work}/R_{free} = 0.13/0.15\). However, the actual \(R_{work}/R_{free}\) was found to be 0.23/0.25 in the validation process, indicative of the absence of twinning in the crystal. Examination of the reflection images and the self-rotation function found that the crystal likely had an order-disorder structure (35, 36). The diffused electron densities of the third MenH chain in the middle of the asymmetric unit in the \(P_{3}1_{2}_{1}_{2}_{1}\) space group were then interpreted as two overlapping MenH chains in a 1:1 ratio that were related by a 2-fold axis. A “statistical” \(2_{1}\) axis is generated by the two overlapping MenH chains so that the symmetry of the crystal is \(P_{3}1_{2}_{1}_{2}_{1}\). The structure was finally solved with \(R_{work}/R_{free}\) = 0.13/0.15.
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$R_{free} = 0.13/0.17$. Restraints of SHCHC were generated and refined using eLBOW (37). The ternary complex crystal also had an order-disorder structure and was similarly solved. Data collection and refinement statistics are summarized in Table 1.

Structural Analysis and Sequence Alignment—Sequence was analyzed with ClustalW2.0 (38). PyMOL was used in the structural analysis and generation of all graphics (39). PISA was used to analyze the protein interfaces (40), and CASTp was used to identify the binding cavities in the protein structures and to calculate the volume of the active site cavity (41).

RESULTS

Overall Structure—The ligand-free MenH, MenH-SHCHC, and MenH-SHCHC-pyruvate crystal structures were determined and refined to a resolution of 1.45, 1.37, and 1.42 Å, respectively. The asymmetric unit of the ligand-free MenH crystal contains one molecule that binds one sulfate ion and one chloride ion in close proximity in the active site. In comparison, the asymmetric unit of the binary or the ternary complex crystal structure contains three protein molecules of which two bind one or both small molecule ligands and the third in the middle is present in two C2-symmetric conformations in a 1:1 ratio without any ligands. The electron densities of the SHCHC and pyruvate ligands are very well defined with 100% occupancy for all atoms in the MenH-SHCHC and MenH-SHCHC-pyruvate complexes. The MenH molecules in the complexes are essentially identical with a root mean square difference (r.m.s.d.) of <0.25 Å over all 252 Ca atoms but are different from the molecule in the ligand-free protein crystal with a 12.2 Å r.m.s.d.

All the MenH molecules are composed of an α-helical cap domain and a typical α/β-hydrolase fold domain (Fig. 2A). The α/β-domain contains seven β-strands (β2 to β8) in the central β-pleated sheet flanked by helices (αA to αC) on both sides; it is missing the β1 strand of the canonical α/β-hydrolase fold (42). It also contains an additional 3₁₀-helix (residues 51–53) in the β4-to-αB loop and an additional short helix (αF; residues 239–251) at the αF N terminus, both of which are absent in the canonical α/β-hydrolase fold. The α-helical cap domain is inserted in the loop connecting β6 to αD and consists of four long α-helices (α1 to α4) and two short 3₁₀-helices (α2' and α4') connected to the C termini of α2 and α4, respectively.

Open and Closed Conformations—Analysis of the electrostatic potential surface with APBS (43) revealed a major difference between the structures of ligand-free MenH and MenH in complex with one or both products. As shown in Fig. 2B, the positively charged crevice corresponds to the active site cavity formed between the helical cap domain and the α/β-domain and is significantly larger and wider in the ligand-free MenH structure than in all three molecules in the asymmetric units of the complex structures containing SHCHC (shown in Fig. 2B) or both SHCHC and pyruvate (not shown), including the central molecule without any ligands. Using CASTp for analysis (41), the volume of the cavity was calculated to be 1662 Å³ in ligand-free MenH and 1013 Å³ in the MenH-SHCHC complex, indicating a reduction of ~650 Å³ in size upon binding of the product. This drastic change shows that MenH is in an open conformation in the ligand-free protein structure and is in a closed conformation when complexed with the ligands.

The catalytic triad is not formed in the open conformation. As shown in Fig. 2C, the side chains of the three triad residues Ser²³⁶, His²³², and Asp²¹⁰ are well separated without direct hydrogen bonding connections. The side chain of His²³² exists in two rotamers, RT1 and RT2, of which the former is present with ~30% of the electron density in an energetically disfavored gauche conformation, and the latter possesses ~70% of the electron densities in an energetically favored trans conformation. The two nitrogen atoms in the RT1 imidazole side chain were assigned to avoid clash with the bound water molecule, which would occur after a 180° flip of the imidazole ring. Neither RT1 nor RT2 of His²³² takes a position to interact directly with either Ser²³⁶ or Asp²¹⁰. Consequently, the open conformation of MenH is inactive without a functional catalytic triad.

In contrast, a functional catalytic triad is readily found in the closed MenH conformation with strong hydrogen bonds directly connecting Ser²³⁶, His²³², and Asp²¹⁰ (Fig. 2C), demonstrating that the closed state of MenH is active. This functional triad is found in all molecules with or without the product ligands in the asymmetric units of both the binary and ternary complex structures. It is also found in the three reported MenH structures without any product ligands (Refs. 16 and 17 and Protein Data Bank code 1R3D), indicating that they are also in the active closed conformation. Indeed, the ligand-free MenH structure from E. coli (Protein Data Bank code 4GDM) is superimposable with the closed structure in the MenH-SHCHC complex with an r.m.s.d. of 0.35 Å over all 252 Ca atoms. It is clear from this comparison that E. coli MenH exists in two structurally distinct forms in the absence of ligands of which one is the “open” structure determined in this study and the other is the previously determined “closed” conformation (18).

Open-to-closed Conformational Change—The open-closed structural difference at the MenH active site is a result of global conformational change. From a superposition of the open and closed conformations (Fig. 3A), a significant change is identified in the reorientation of about one-half of the cap domain consisting of α2, α2', α3, and the connecting loops (Fig. 3B). This orientation change is responsible for the significant shrinkage of the active site cavity in the closed conformation (Fig. 2B) because the reoriented structure constitutes a large part of the active site. It also enables extensive interaction of the protein with the SHCHC ligand, including the hydrogen bonds between the SHCHC succinyl carboxylate and Arg¹⁶⁸ (mediated by water), Trp¹⁴⁷, and Tyr¹⁴⁸ and the hydrophobic contacts between the ligand and Trp¹⁴⁷, Val¹⁵², and Phe¹⁵³ (Fig. 3D). These new interactions significantly stabilize the closed conformation and may even induce its formation.

The hinged movement in the cap domain leads to a downward shift of the 3₁₀-helix α2' by 2.3 Å toward the α/β-domain. As a result, the hydrophobic side chain of Val¹⁵² in the α-helix, which packs against the imidazole ring of His²³² RT1 rotamer in the open conformation, is dislocated to pack against the His²³² imidazole ring of the functional catalytic triad in the closed conformation, which corresponds to the position of His²³² RT2 rotamer in the open conformation. In the meantime, the bulky side chain of the adjacent Phe¹⁵³ is moved into the position of the imidazole ring in the RT1 rotamer of His²³² in the open conformation, forcing the triad residue to adopt the energeti-
cally disfavored gauche conformation in the closed conformation (Fig. 3E). To accommodate these changes, Asp\(^{210}\) and His\(^{232}\) undergo a subtle positional movement of about 0.7 and 0.9 Å, respectively, in the closed conformation relative to the open conformation to finally form the functional catalytic triad (Fig. 3E). The positional change of Asp\(^{210}\) and His\(^{232}\) is a result of backbone shift in a large portion of the \(\alpha/\beta\)-domain after the cap domain insertion, including \(\alpha D, \alpha E, \alpha F, \beta 7,\) and \(\beta 8\) (Fig. 3C).

The \(\alpha/\beta\)-domain change causes abrupt and significant backbone change with a maximum displacement of more than 2.4 Å in the loop connecting \(\alpha E\) to \(\beta 8\) (Fig. 3C), which includes residues 221–224 in the open conformation and residues 220–222 in the closed conformation. As a result, \(\alpha E\) is shorter by one residue and \(\beta 8\) is longer by two residues in the closed conformation in comparison with the open conformation. In addition, Cys\(^{206}\) (in \(\beta 7\)) and Cys\(^{224}\) are suitably positioned to form a disulfide bond in the open conformation but are too far apart to form a similar disulfide bond in the closed conformation. The disulfide bond would lock the enzyme in the inactive conformation and could be utilized to regulate MenH activity through a redox-responsive mechanism. However, the two cysteine res-
idues are not conserved among MenH orthologues. Their potential involvement in regulating the enzyme activity needs further investigation.

**Enzyme-Ligand Interactions**—The SHCHC product is bound firmly at identical position and orientation at the active sites of the binary and ternary complexes with a markedly lower B factor in comparison with other parts of the structures. Its average B factor is 12.8 Å² in the binary complex and 9.9 Å² in the ternary complex compared with an average B factor of 28.0 and 24.9 Å² for the protein (Table 1), respectively. The extensive interactions of this ligand with the enzyme include hydrophobic interactions with Trp147, Val152, and Phe153 (Fig. 3D); hydrogen bonding interactions with Tyr148, Ser86, Arg90, Arg124, Trp147, Tyr148, Arg168, and Asn223; and salt bridge interaction with Arg90 (Fig. 4). These interactions are consistent with findings in a previous mutational study (15), although the interaction with Arg168 is a water-mediated hydrogen bond rather than a salt bridge interaction. In addition, the product is also stabilized by two short hydrogen bonds provided by the backbone amides of Phe23 and Leu87 to the SHCHC succinyl keto group where an oxyanionic enolate intermediate is expected after H-proton abstraction in the MenH-catalyzed reaction. This finding strongly supports that the backbone amides of these residues or their equivalents in MenH orthologues form the oxyanion hole as proposed on the basis of modeling results (17, 18).

In the ternary complex, the pyruvate product is bound at a position close to the opening of the active site cavity to the bulk solvent. It is stabilized by only one hydrogen bond with the side-chain hydroxyl of Ser86 (Fig. 4C) and is close to the side-chain guanidinium group of Arg124 without direct interaction. This weak interaction suggests that pyruvate occupies a transient binding site on its exit from the active site after it is cleaved.
from the SEPHCHC substrate. Its close positioning to the Arg₁₂⁴ side chain is consistent with the suggestion that this residue is involved in the SEPHCHC substrate binding by forming a salt bridge with the pyruvyl carboxylate (15).

Site-directed Mutagenesis—In the closed conformation, Trp₁⁴⁷ and Tyr₁⁴⁸ strongly interact with the SHCHC ligand through both hydrogen bonding and hydrophobic interaction, whereas Val₁⁵² and Phe₁⁵³ directly interact with the catalytic triad and force its formation. All these residues are highly conserved among MenH homologues (15, 18). To assess their catalytic contribution, the mutants V₁⁵₂A and F₁⁵₃A, were created and characterized in previous work (15). In addition, two double mutants, W₁⁴⁷A/Y₁⁴⁸A and W₁⁴⁷F, were created and characterized in previous work (15). In addition, two double mutants, W₁⁴⁷A/Y₁⁴⁸A and W₁⁴⁷F, were created and characterized in previous work (15). In addition, two double mutants, W₁⁴⁷A/Y₁⁴⁸A and W₁⁴⁷F, were created and characterized in previous work (15). In addition, two double mutants, W₁⁴⁷A/Y₁⁴⁸A and W₁⁴⁷F, were created and characterized in previous work (15).

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The catalytic efficiency of V₁⁵₂A and F₁⁵₃A was reduced by 1230- and 11-fold (Table 2), respectively. When both residues were mutated, the catalytic activity was completely eliminated. To understand the meaning of these kinetic data, it is worth noting that Phe₁⁵³ is not limited to their contribution to the binding of substrate through hydrophobic interactions observed in the MenH-SHCHC complex (Fig. 3D); rather, their role in stabilizing and inducing the active closed conformation from the inactive conformation is likely a much more important catalytic contribution. Thus, the kinetic results of these mutants strongly support an essential role of the open-to-closed conformational change in MenH catalysis.

Both Trp₁⁴⁷ and Tyr₁⁴⁸ play an important role in the open-closed structural change by strongly interacting with the ligand through hydrogen bonding and hydrophobic interaction. This role is consistent with the kinetic properties of the proteins with a mutation at either position. As reported earlier (15), the catalytic efficiency of the W₁⁴⁷A and W₁⁴⁷F mutants is reduced by 5.9 × 10⁻³ and 62-fold, respectively. Similarly, mutation at the adjacent Tyr₁⁴⁸ also caused a significant decrease in the second order rate constant (Table 2).

NMR Detection of the Catalytic Triad—It is well known that the hydrogen bond between the central histidine and the acidic residue (Asp or Glu) is strongly deshielded in the catalytic triads of serine proteases and α/β-hydrolase enzymes, affording a proton nuclear resonance signal in the downfield region (usually 12–16 ppm). When these enzymes interact with their mechanism-based inhibitors, the imidazole ring in the side chain of the central histidine is protonated, and its hydrogen bond with the acidic residue is greatly strengthened, leading to a significant downfield shift of the corresponding proton signal into the 16–19.5-ppm range (45–47). Based on these facts, we used NMR spectroscopy to detect the proton in the strong hydrogen bond between the side chains of Asp₁⁵⁰ and His₁⁵² in the catalytic triad of ¹⁵N-labeled MenH. As shown in Fig. 5, there is no signal in the 13–22-ppm range in the absence of a
**FIGURE 4. Enzyme-ligand interactions.**

A, stereodiagram of the active site of MenH in the binary MenH-SHCHC complex. The active site residues and the SHCHC are represented in sticks with green and yellow carbon atoms, respectively. Electron density \(2\sigma_F - \sigma_C\) of SHCHC is displayed in blue mesh and is contoured at \(1.5\sigma\). B, LIGPLOT of enzyme-ligand interactions in the binary MenH-SHCHC complex. C, LIGPLOT of enzyme-ligand interactions in the ternary MenH-SHCHC-pyruvate complex. Dashed lines indicate hydrogen bonds with a distance less than 3.5 Å. Water molecules (W1–W3) are shown as red spheres. The plots were generated with LIGPLOT V.4.5.3 (44).

### TABLE 2

Kinetic constants of the E. coli MenH mutants

| Protein          | \(K_m\) \(\mu M\) | \(k_{cat}\) \(min^{-1}\) | \(k_{cat}/K_m\) \(min^{-1} \mu M^{-1}\) |
|------------------|------------------|-----------------|---------------------------------|
| Wild type        | \((1.3 \pm 0.3) \times 10^3\) | \((5.1 \pm 0.2) \times 10^2\) | \((3.2 \pm 0.5) \times 10^7\) |
| F153A            | \((7 \pm 1) \times 10^3\) | \((2.0 \pm 0.1) \times 10^2\) | \((2.9 \pm 0.1) \times 10^6\) |
| V152G            | \((1.6 \pm 0.2) \times 10^2\) | \((1.2 \pm 0.1) \times 10^1\) | \((7.3 \pm 0.8) \times 10^4\) |
| V152A            | \((3.0 \pm 0.7) \times 10^2\) | \(7.7 \pm 0.9\) | \((2.6 \pm 0.2) \times 10^4\) |
| Y148F            | \((5.0 \pm 0.3) \times 10^3\) | \(7.5 \pm 0.8\) | \((1.6 \pm 0.3) \times 10^5\) |
| W147A/Y148A      | ND\(^a\) | ND | ND |
| V152G/F153A      | ND | ND | ND |

\(^a\) ND, not detectable.
The crystal structures of the closed conformation were determined in the absence of ligands in previous investigations (Refs. 17 and 18 and Protein Data Bank code 1R3D) might be due to the better crystallizability of the closed structure in comparison with the open structure.

**Fast Kinetics of Enzyme-Ligand Interactions**—Fast kinetics of enzyme-ligand interactions were determined to understand the mechanism of MenH recognition of its substrate or other ligands. First, the stopped-flow technique was used to determine the pre-steady state kinetics of the MenH catalysis by monitoring formation of the SHCHC product, which was measured by the ultraviolet absorbance at 290 nm. No lag phase was detected in the product formation curves, and the calculated reaction rates were consistent with those obtained from the steady state measurements at various substrate concentrations. The absence of the lag phase strongly suggests that the substrate-induced open-closed structural change is much faster than the catalytic steps and is beyond the detection limit of the stopped-flow technique. Next, the MenH-SHCHC binding kinetics were determined by monitoring the quenching of tryptophan fluorescence at 340 nm, which is indicative of hydrogen bonding of the Trp147 side chain to SHCHC in the enzyme-ligand complex (Fig. 3D). As shown in Fig. 7, the observed rate of the approach to binding equilibrium $k_{obs}$ defined as the fluorescence quenching rate, increases hyperbolically with increasing ligand concentration. This positive hyperbolic $k_{obs}$-[$SHCHC$] relationship is predicted by an induced fit mechanism for the enzyme-ligand recognition under the rapid equilibrium approximation (51, 52). However, it is not unambiguous evidence for the induced fit mechanism. Actually, a conformational selection mechanism is also able to afford a similar rate-ligand concentration curve (53, 54), which has been demonstrated for several enzyme-ligand systems (53–56). Thus, the observed $k_{obs}$-[$SHCHC$] curve does not exclude a conformational selection mechanism for the MenH-ligand interaction.

**DISCUSSION**

The discovery of the inactive open conformation of MenH in its ligand-free form demonstrates the conformational flexibility of the $\alpha/\beta$-hydrolase fold enzyme. Together with the active closed conformation determined previously (Ref. 17 and Protein Data Bank code 1R3D), this finding shows that the vitamin K biosynthetic enzyme is present in aqueous solution in one or both of the open and closed conformations. NMR evidence shows that MenH predominantly exists in the inactive open conformation in its ligand-free form, and it has to undergo a ligand-dependent open-closed conformational change in the catalytic process, similar to the structural change caused by the product ligand SHCHC. This open-closed structural change could occur via an induced fit mechanism (57) or a conformational selection mechanism (58). Unfortunately, neither the pre-steady state kinetics nor fast kinetic measurement of the MenH-SHCHC binding process provides an unambiguous distinction between these two possibilities. Nevertheless, the fact that the both open and closed conformations without a ligand have been readily crystallized indicates an equilibrium between them in solutions and strongly suggests that the conforma-
tional change likely occurs via a conformational selection mechanism.

Results from the site-directed mutagenesis are consistent with an important catalytic role for the ligand-dependent open-closed conformational change (Table 2). This conformational change suggests a novel mechanism for modulating the intrinsic nucleophilic reactivity of the catalytic triad (Fig. 8). In the absence of a cognate ligand, the enzyme is in its dominant open conformation without a functional catalytic triad. The catalytic triad is formed and activated only when the substrate binds to the enzyme active site and stabilizes it in the active form. Such a triad may still retain its nucleophilicity but is unable to exhibit this reactivity because of the lack of a suitable electrophile in the enzyme-substrate complex. However, its basicity can be utilized to carry out the needed \( \alpha \)-proton abstraction in the catalysis because the \( \text{O}_2 \) of \( \text{Ser}^{\text{Asp}} \) is optimally positioned for abstraction of a proton from \( \text{C}_2 \) of the \( \text{SEPHCHC} \) substrate as suggested by its close distance of 2.3 Å from the \( \text{C}_2 \) carbon of the \( \text{SHCHC} \) ligand and results from modeling of the substrate into the triad-containing structures of MenH (17, 18). In this way, the open-closed conformational change tightly couples formation of the catalytic triad to binding of the \( \text{SEPHCHC} \) substrate, thereby completely shielding the nucleophilicity of the triad and limiting its catalytic role to a specific general base.
This ligand-dependent triad formation also places a stringent constraint on the structure of the activating ligand and consequently greatly increases the substrate specificity of the enzyme, providing a rationale for its lack of hydrolytic activity toward nonnative esters or thioesters (14, 15).

With the modulation of the triad activity by the open-closed conformational change, a new catalytic mechanism is proposed for the MenH-catalyzed reaction as shown in Fig. 6. In this mechanism, MenH is stabilized in the active conformation after the negative SEPHCHC substrate is bound to the positively charged active site cavity through interactions with the conserved active site residues identified previously and in the current study. Subsequently, an oxyanion is generated via deprotonation of the Ser86 side-chain hydroxyl group by the Asp-His dyad to serve as a general base for abstraction of the C2 proton from the substrate. The resulting dienolate intermediate is stabilized by the oxyanion hole comprising the backbone amides of Phe23 and Leu87. Next, the dienolate intermediate undergoes an E1cb elimination of the pyruvate to form the SHCHC product as proposed previously (15) using either water or the protonated side chain of Lys214 as the general acid to protonate the pyruvyl enolate. Finally, both products dissociate from the enzyme, and the protonated Asp-His dyad is deprotonated to transiently form the functional triad again. However, the triad then disintegrates after dissociation of the SHCHC product to complete a catalytic cycle.

The ligand-dependent conformational change is one of the most important features of the MenH catalysis. This type of conformational change, which usually involves local conformational change and no change of the catalytic triad, is well known to increase the substrate specificities and allosterically regulate the activities of serine proteases (4, 59, 60). In addition, conformational change is also known to play a critical role in the “interfacial activation” of lipases within the α/β-hydrolase superfamily that involves a low activity closed state in homogenous solution and a highly active open state when exposed to the substrate in an oil droplet. This structural change involves the movement of a long surface loop covering the active site in most lipases (61, 62) but involves an open-closed conformational change in the α-helical cap domain in other lipases (63, 64) that is similar to the changes observed for MenH. However, the interfacial activation of the lipases does not involve changes in the catalytic triad or the α/β-domain. In addition, the closed conformation of the lipases is inactive, which is opposite to what was observed for MenH. Thus, the open-closed conformational change of MenH is a novel, catalytically important structural change, which contributes to the structural and functional diversity of the α/β-hydrolase superfamily.

The ligand-dependent formation of the Ser-His-Asp triad is another important feature of the MenH catalysis that is rarely seen among triad-utilizing enzymes. Besides MenH, the serine protease Factor D in the alternative complement system is the only other enzyme that forms a functional triad after induction by substrate binding. In its resting state, Factor D also has a nonfunctional triad with the central histidine taking the energetically more favorable trans conformation and rotated out of the catalytic position due to the presence of a self-inhibitory loop (65). When its native substrate, a complex of the complement Factor C3b and Factor B, is bound, Factor D undergoes extensive conformational change to move the inhibitory loop and form the functional catalytic triad (16). Despite these similarities, the substrate induces conformation change to remove an obstacle to form the triad in Factor D from the three conserved residues that are thermodynamically favored to form the
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triad, whereas the ligand-dependent conformational change in MenH forces the formation of the triad from three separated residues that are thermodynamically disfavored to form the catalytic motif. In addition, the Factor D catalytic triad still catalyzes the hydrolysis of C3b via the covalent, nucleophilic mechanism of catalysis like other serine proteases, whereas its reactivity is restricted to that of a general base in the MenH catalysis. Therefore, the ligand-dependent triad formation in MenH is a novel allosteric mechanism to modulate the reactivity of the catalytic triad, allowing the triad to expand its catalytic power beyond the covalent, nucleophilic catalysis.

In summary, we have found an open-closed conformational change critical to the catalysis of the α/β-hydrolase fold enzyme MenH that involves the formation of its Ser-His-Asp catalytic triad in the closed conformation and strictly subjects its enzymatic activity to control by the binding of a cognate ligand such as the substrate or the product. The coupling of the triad formation to the substrate binding shields the nucleophilicity of the traditional catalytic triad and effectively limits the triad to the role of a simple general base in the catalysis. Moreover, this control of the triad formation greatly increases the substrate specificities of the enzyme. These findings not only provide new insights into the catalytic mechanism of the typical α/β-hydrolase fold enzyme but also demonstrate that the traditional triad is capable of new catalytic chemistry through appropriate modulation.

Acknowledgments—We thank Nathaniel Echols, Pavel Afonine, Paul Adams, and Tom Terwilliger of the PHENIX team for help in solving the order-disorder structures.

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