Interdomain Conformational Changes Provide Allosteric Regulation en Route to Chorismate**

Ali Reza Nazmi‡, Eric J. M. Lang‡, Yu Bai‡, Timothy M. Allison‡, Mohamad H. Othman‡, Santosh Panjikar‡, Vickery L. Arcus**, and Emily J. Parker**‡

From the ‡Biomolecular Interaction Centre and Department of Chemistry, University of Canterbury, P. O. Box 4800, Christchurch 8140, New Zealand, the §Department of Chemistry, University of Oxford, South Parks Road, Oxford OX1 3QY, United Kingdom, the †Australian Synchrotron, Clayton, Melbourne, Victoria 3168, Australia, the ‡Department of Biochemistry and Molecular Biology, Monash University, Clayton Campus, Melbourne, Victoria 3800, Australia, the **School of Science, University of Waikato, Hamilton 3240, New Zealand, and the ‡‡Maurice Wilkins Centre, Biomolecular Interaction Centre and Department of Chemistry, University of Canterbury, P. O. Box 4800, Christchurch 8140, New Zealand

Multifunctional proteins play a variety of roles in metabolism. Here, we examine the catalytic function of the combined 3-deoxy-D-arabino-heptulosonate-7-phosphate synthase (DAH7PS) and chorismate mutase (CM) from Geobacillus sp. DAH7PS operates at the start of the biosynthetic pathway for aromatic metabolites, whereas CM operates in a dedicated branch of the pathway for the biosynthesis of amino acids tyrosine and phenylalanine. In line with sequence predictions, the two catalytic functions are located in distinct domains, and these two activities can be separated and retain functionality. For the full-length protein, prephenate, the product of the CM reaction, acts as an allosteric inhibitor for the DAH7PS. The crystal structure of the full-length protein with prephenate bound and the accompanying small angle x-ray scattering data reveal the molecular mechanism of the allostery. Prephenate binding results in the tighter association between the dimeric CM domains and the tetrameric DAH7PS, occluding the active site and therefore disrupting DAH7PS function. Acquisition of a physical gating mechanism to control catalytic function through gene fusion appears to be a general mechanism for providing allosterity for this enzyme.

Allostericity is the biochemical process whereby binding a remote ligand triggers protein functional changes. Allosteric regulation of key metabolic enzymes is an important control system for many metabolic pathways. The shikimate pathway, which is found in plants and microorganisms, is the biosynthetic route for the formation of the aromatic amino acids and other important aromatic metabolites (1). The first committed step of the pathway, catalyzed by 3-deoxy-D-arabino-heptulosonate-7-phosphate synthase (DAH7PS), is generally controlled by allosteric inhibition arising from the remote binding of end products or intermediates of the pathway (2).

DAH7PS catalyzes the metal ion-dependent condensation reaction of phosphoenolpyruvate (PEP) and erythrose 4-phosphate (E4P) to generate 3-deoxy-D-arabino-heptulosonate-7-phosphate (DAH7P, Fig. 1). The pathway has six further steps to the common intermediate chorismate. Chorismate is substrate for a variety of enzymes, including those that catalyze steps that ultimately lead to the aromatic amino acids Phe, Tyr, or Trp. Allosteric regulation is observed both at the pathway entry via DAH7PS and at the branch points from chorismate. All known DAH7PS enzymes share a (βα)8 barrel catalytic domain, which houses the active site. However, there are distinct variations between DAH7PS enzymes from different organisms in the structural extensions to the barrel that support allosteric.

DAH7PS enzymes are divided into two distinct types, which differ in sequence, subunit size, and quaternary structure. Type I enzymes are generally smaller than their type II counterparts and have been shown to display allosteric control from a single allosteric effector or to display no allosterity. Type I enzymes are further divided into two subtypes. The type Iα subgroup contains the well characterized Escherichia coli, Saccharomyces cerevisiae, and Neisseria meningitidis enzymes (3–5). These enzymes have an allosteric site formed by an N-terminal and loop extension and are sensitive to one of the aromatic amino acids. Type Iβ enzymes are more varied and either comprise just the core catalytic barrel and show no allosteric regulation or have a discrete domain found at either the N or C termini of the barrel (6). This addition can be either an ACT domain (7, 8) (named after the enzymes in which it was originally identified, aspartokinase, chorismate mutase, and TyrA) or a chorismate mutase (CM) enzyme (9), which catalyzes a branch point step of the pathway (Fig. 1). For the latter, previously characterized DAH7PS enzymes have dual catalytic functionality (both DAH7PS and CM activity), and the DAH7PS activity is inhib-
In accordance with the different extensions to the core catalytic barrel, the molecular mechanisms by which allostery is achieved vary between the DAH7PS types. Whereas the type Iβ DAH7PS from *Thermotoga maritima* undergoes significant ACT domain movement associated with the binding of inhibitor Tyr (7), for other enzymes, such as the type II *M. tuberculosis* DAH7PS (13) and the type Iα *N. meningitidis* DAH7PS (14), allosteric inhibition is conferred by more subtle structural changes and alteration in protein dynamics. The variation in allosteric machinery and mechanism for this enzyme is intriguing both from a mechanistic and evolutionary point of view.

Here, we examine in detail the mechanism of allosteric for the DAH7PS from *Geobacillus* sp. (strain Y412MC61), a thermophilic soil bacterium. From sequence inspection, this DAH7PS is predicted to have an N-terminal CM domain and be a member of the type Iβ group. The independence of the individual catalytic activities of the DAH7PS and CM domains has already been demonstrated for the *Bacillus subtilis* enzyme (15), and a structure of an uninhibited form of an enzyme of this type has been previously determined for the DAH7PS from *Listeria monocytogenes* (*LmoDAH7PS*) (16). Here, we explore the functional properties of this dual activity protein and demonstrate directly the structural changes associated with the allosteric response induced by prephenate binding. Based on our new x-ray crystal structure and small angle x-ray scattering experiments, we reveal that prephenate binding to the CM domain stabilizes a more compact conformation of the protein, limiting the catalytic functionality of the DAH7PS domain active site.

**Results**

**DAH7PS and CM Catalytic Activities of Wild-type GspDAH7PS Can Be Separated into Two Functional Enzymes—**

The DAH7PS protein complex from *Geobacillus* sp. putatively includes two functional units, represented by a DAH7PS and a CM domain. To test this prediction and to investigate the dependence of the catalytic activity of each functional unit of the GspDAH7PS complex, the full-length protein (*GspDAH7PSWT*) was expressed, and two truncated forms were generated consisting of either only the predicted DAH7PS (*GspDAH7PSD*) or CM (*GspDAH7PSC*) domains (Fig. 2). The full-length protein indeed displayed both DAH7PS and CM activity (Table 1), and as for other characterized DAH7PS enzymes, the DAH7PS activity of *GspDAH7PSWT* was found to be dependent on the presence of a divalent metal ion (Fig. 3A). Consistent with the thermophilic nature of *Geobacillus* sp., maximum DAH7PS and CM activity was observed at 70 °C (Fig. 3, B and C). The DAH7PS from *Geobacillus* sp. is therefore a bifunctional thermophilic enzyme, which possesses both DAH7PS and CM activity.

Consistent with compartmentalization of activities into the respective domains, in the truncated variants only metal-dependent DAH7PS activity was observed for *GspDAH7PSD*, and only CM activity was detected for *GspDAH7PSC*. Intriguingly, whereas the DAH7PS activity of the separated DAH7PS domain was greater than that of the wild-type protein, the efficiency of catalysis of the CM reaction was reduced for the isolated CM domain (Table 1). This change was principally caused by a 5-fold increase in the chisomerase *K*<sub>c</sub> value for the shorter *GspDAH7PSD* protein, yet the catalytic turnover, represented by the *k*<sub>cat</sub> was 2-fold higher. The activities of both separated domains were also found to be slightly more sensitive to temperature than was found for the full-length protein (Fig. 3). Thus, the properties of the two functional domains comprising the full DAH7PS-CM protein are affected by its formation, suggesting the possibility of inter-domain communication.

**Inhibition of DAH7PS Activity by Prephenate—**

Structural extensions to DAH7PS core catalytic barrels is a common

---

**FIGURE 1. Shikimate pathway for aromatic amino acid biosynthesis.** The first step of the pathway is catalyzed by the enzyme DAH7PS. The pathway branches at chorismate, and chorismate mutase catalyzes the formation of prephenate from chorismate, which in turn will form Tyr and Phe.
A mechanism for elaborating feedback inhibition on DAH7PS activity (4, 7, 17). The combination of DAH7PS and CM domains may represent another case of this phenomenon. To test this, we assessed whether the product of CM, prephenate, was an inhibitor of the DAH7PS activity, as this would be consistent with the fusion of the CM domain to DAH7PS providing feedback regulation to the pathway. Notably, prephenate inhibited the DAH7PS activity of the full-length GspDAH7PSWT protein but not of separated GspDAH7PSD (Fig. 4). The DAH7PS activity of the full-length enzyme was reduced to \( \frac{1}{2} \) at 20 \( \mu M \) prephenate and to less than 2% at higher prephenate concentrations compared with the activity in the absence of prephenate. This observation suggests that the inhibitory mechanism directly involves the CM domain, which may provide an allosteric binding site for prephenate. Due to the high CM activity of GspDAH7PS WT, the same inhibition response was observed with chorismate rather than prephenate.

Small Angle X-ray Scattering Shows a Conformational Change in the Presence of Prephenate—To examine the effect on the structure of the complex in the presence of prephenate, experimental SAXS data were obtained for GspDAH7PS in the absence and presence of prephenate to shed light on the conformational changes associated with inhibition (Fig. 5). Comparison of the SAXS profiles of GspDAH7PS in the absence and presence of prephenate shows clear differences, consistent with a conformational change resulting from the binding of prephenate (Fig. 5A). This change is associated with an increased compactness of the enzyme in the presence of prephenate by the smaller radius of gyration, derived from the Guinier plot (\( R_g = 34.8 \pm 0.6 \) Å with prephenate compared with \( R_g = 36.5 \pm 0.4 \) Å without) (Fig. 5A).

Notably, the bell-shaped curve observed in the Kratky plot (Fig. 5B) shows that in both the presence and absence of prephenate the protein exhibits a typical folded shape, without random coils. This verifies that both domains remain folded even in the absence of prephenate, although the slightly more flattened curve for the prephenate-free sample may indicate an increased flexibility of the protein. The Porod-Debye plots (Fig. 5C) present two different plateaus in these two cases, suggesting that the protein experiences a discrete conformational change upon ligand binding without any radical change in its

### TABLE 1

| Protein     | DAH7PS activity* | CM activity |
|-------------|------------------|-------------|
|             | \( K_m \) PEP  | \( k_m \) | \( k_m/K_m \) | \( k_m/K_m \) |
| GspDAH7PSWT | 87 ± 9 \( \mu M \) | 95 ± 8 \( \mu M \) | 45 ± 4 | 0.52 | 0.47 |
| GspDAH7PSD  | 98 ± 11  | 62 ± 5  | 63 ± 5 | 0.64 | 1.0 |
| GspDAH7PSC  | 450 ± 40 | 5.5 ± 0.5 | 1.2 × 10^{-2} |

*The Michaelis constants for the DAH7PS activity are apparent values determined with a fixed concentration of the invariant substrate (as described under Experimental Procedures).
intrinsic flexibility (18, 19). This conclusion is supported by the $q^3 I(q)$ versus $q^3$ plot (Fig. 5D) that displays a distinct decrease, rather than a plateau as one would expect in the case of a highly flexible enzyme.

Prephenate Binds to the CM Domain at the Active Site Located within a Helical Bundle—To further probe the intriguing conformational changes in the presence of prephenate and to identify the prephenate-binding site, we determined the crystal structure of GspDAH7PS in complex with the CM product and DAH7PS inhibitor prephenate. GspDAH7PS was crystallized in the presence of Mn$^{2+}$ and chorismate, and its structure was determined at a resolution of 2.75 Å in the space group P6$_2$. The asymmetric unit contains two adjacent chains, with each comprising a catalytic $\beta/\alpha$$_8$ barrel common to all DAH7PS enzymes (residues 106–362) and a CM domain (residues 1–86) connected by a short linker region (residues 86–105) (Fig. 6A). The two catalytic barrels from each chain in the asymmetric unit are highly similar (C$_{\text{r.m.s.d.}}$ 0.11 Å), whereas the CM domains are slightly less similar (C$_{\text{r.m.s.d.}}$ 0.23 Å), with one of the chains having more unresolved residues than the other in the linker region connecting the CM domain with the catalytic barrel. However, when taken as a whole, the two chains present a C$_{\text{r.m.s.d.}}$ of 3.93 Å. This large difference arises from the difference in location of the two CM domains relative to the barrels in each chain (Fig. 6A).

A 2-fold symmetry operation reveals the homotetrameric biological unit of GspDAH7PS (Fig. 6B). The DAH7PS barrel subunits are arranged in a similar fashion to the tetrameric assemblies observed for other structurally characterized type Iβ DAH7PS enzymes (6), with neighboring protomers placed so that adjacent barrels face in opposite directions around the tetramer plane. A manganese ion is modeled at the C-terminal end of each of the barrels and is coordinated to residues Cys-

FIGURE 3. A, activation of the DAH7PS activity of GspDAH7PSWT (green) and GspDAH7PSD (blue) by different divalent metal ions (100 μM) in comparison with the activity with Cd$^{2+}$. Also shown is the activity of both enzymes in the presence of the metal chelator EDTA. B, dependence of reaction rates of DAH7PS activity (B) and CM activity (C) on temperature. GspDAH7PSWT activity is shown in green; GspDAH7PSD activity is shown in blue, and GspDAH7PSC activity is shown in pink. Error bars represent the standard deviation from triplicate measurements.

FIGURE 4. Response of GspDAH7PSWT (blue) and GspDAH7PSD (red) DAH7PS activity to increasing concentrations of prephenate at 60 °C. Error bars represent the standard deviation from triplicate measurements.

FIGURE 5. Analysis of the SAXS data for GspDAH7PS in the absence (blue) and presence (red) of prephenate. A, SAXS profiles (log $I(q)$ versus $q$) with Guinier plots inset. B, Kratky plot ($q^2 I(q)$ versus $q$), C, Porod-Debye plot ($q^4 I(q)$ versus $q^4$), and D, SIBYLS plot ($q^3 I(q)$ versus $q^3$), limited to the range of the SAXS data for which the Guinier linearity is observed.
Bifunctional DAH7PS from Geobacillus

FIGURE 6. Crystal structure of GspDAH7PS. A, two chains of the asymmetric unit, chain A (left) and chain B (right). The catalytic TIM barrel is shown in blue, the CM domain in green, and the linker region in purple. Allosteric inhibitor prephenate (Pre) is represented with gray sticks, and the active site is identified with the metal ion, represented with an orange sphere. B, tetrameric assembly with the components of one asymmetric unit colored (TIM barrel is shown in blue, the CM domain in green, and the linker region in purple) and the symmetry-related dimer is colored gray. C and D, binding sites of prephenate in the two unique subunits of the asymmetric unit. Residues important for the binding of prephenate and the catalytic activity of the CM domain are shown with sticks. One CM domain is shown in green, and the adjacent chain forming the catalytically active CM is shown in blue. Prephenate is represented with gray sticks. The χA-weighted Fo – Fc, omit maps for prephenate are represented with a blue mesh and were generated by deleting the prephenate molecules from the n-1 round of refinement. Electron density is contoured at 3σ (0.1457e/Å³).

126, His-296, Glu-322, and Asp-333, identifying the locality of the DAH7PS active site. These metal-binding residues are conserved in all structurally characterized DAH7PS enzymes. The helical CM domains of two diagonally opposed chains associate in an interlaced dimer that resembles a type II or AroQ helical CM (20).

Difference electron density was observed in the area corresponding to the catalytic site of the CM domain. Because of the observed CM activity of GspDAH7PS, prephenate is generated from the added chorismate, and prephenate presence was confirmed by difference density at the CM domain active site (Fig. 6, C and D). The prephenate-binding site reveals that the residues involved in its binding correspond to a subset of the residues responsible for the catalytic activity of the CM domain (Fig. 6, C and D). More specifically, the position of Arg-10, Arg-27, Lys-38, Asp-47, and Glu-51 suggests they form hydrogen bonds and other electrostatic interactions with prephenate, whereas Val-31, Ile-34, Met-54, Phe-79, and Leu-83 create a hydrophobic shell surrounding the ligand and the aliphatic side chain of Arg-50 contributing to these hydrophobic interactions.

Despite low sequence identity (35%, Fig. 2) the CM domains of GspDAH7PS align well with the structure of E. coli AroQ CM (20) (EcoCM, r.m.s.d. = 2.23 Å) (PDB code 1ECM; Fig. 7A). The GspDAH7PS CM helices show more curvature than is observed in the EcoCM structure, especially noticeable at the end of the first helix (Fig. 7A), which may be the result of the interactions between the CM domains and the DAH7PS barrels. Moreover, the second and third helices of the GspCM domains are shorter than their EcoCM counterparts, with the end of the third α-helix of EcoCM corresponding to the beginning of the linker region, which connects the GspDAH7PS CM domain to the catalytic barrel. The well-characterized catalytic site of EcoCM (20–23) exhibits an excellent structural alignment with the active site of the CM domain of GspDAH7PS (Fig. 7B), with the catalytically important charged and polar residues shared by both proteins. Hydrophobic residues Ala-32, Val-35, Ile-81, and Val-85 of EcoCM are not conserved in GspDAH7PS but instead are replaced by alternative hydrophobic residues Val-31, Ile-34, Phe-79, and Leu-83, respectively. In addition, Ser-84, which had been postulated as being important in EcoCM (21), is not conserved and replaced by Ala in GspDAH7PS. These alterations may account for the relatively low CM activity of the GspDAH7PSWT and the truncated GspDAH7PS protein.

For the structures of LmoDAH7PS, which have no ligand bound in the CM active site, the CM domains are found further from the catalytic DAH7PS barrels in an “open” position when compared with the “closed” position they adopt in the inhibited prephenate-bound GspDAH7PS structure. LmoDAH7PS shares sequence identity of 75.1% over 361 residues (91.1% similarity, Fig. 2) with GspDAH7PS, and structural alignment of
tetrameric GspDAH7PS with LmoDAH7PS yields r.m.s.d. values of 0.82 Å (PDB code 3NVT) and 1.08 Å (PDB code 3TFC).

Prephenate Binding Is Associated with Closer Association between the Catalytic and Regulatory Domains—To estimate the likely structural changes of GspDAH7PS on the binding of prephenate, a full model of GspDAH7PS in its active open form, referred to hereafter as the “open model,” based on the two known prephenate-free LmoDAH7PS structures was constructed by homology using MODELLER (24). In addition, missing residues and side chains in the crystal structure of GspDAH7PS were reconstructed, and their position optimized, with MODELLER (referred to as the “closed model”).

Comparison of these open and closed models sheds light on the structural changes likely associated with the binding of prephenate and mechanism of the allosteric inhibition of GspDAH7PS (Fig. 8). In the absence of prephenate, there is little contact between the CM domain and the DAH7PS barrels (7.6% solvent-accessible surface area (SASA) of the tetrameric DAH7PS barrels buried by the CM domains), and the structure is extended (Fig. 8A), with the entrance to the active sites of the DAH7PS domains accessible for substrate binding (Fig. 8C). In comparison, the closed model with prephenate bound is more compact (Fig. 8B), with the CM domains slightly bent to partly follow the shape of the DAH7PS barrels (15.6% of the SASA of the tetrameric DAH7PS barrels buried by the CM domains). In addition, a number of hydrogen bonds are formed between the DAH7PS and CM domains, in particular between Asp-326 and Arg-52, Tyr-46 and Glu-129, and Arg-175 and Gly-42, occluding in part the entrance to the active sites, suggesting an explanation for the inhibitory effect of prephenate against the DAH7PS activity (Fig. 8C).

Portions of CM domains that are disordered in the open model, namely the inter-helical loops, the N-terminal region, and the beginning of the linker region, are found to be ordered in the closed model and lead to extended CM helices (Fig. 8, A and B). These parts of the CM domains include residues identified as being responsible for the catalytic activity of the CM domains and for the binding of prephenate, in particular, Ile-34, Lys-38, Asp-47, Arg-50, and Glu-51. We can therefore postulate that the binding of chorismate or prephenate is accompanied by a reorganization of the secondary structure of the CM domains, viz. the formation of longer helices. This reorganization, together with the binding of the allosteric ligand itself, significantly rigidifies the CM domains, favoring a conformation in which these domains lean on the DAH7PS catalytic barrels.

SAXS Data Confirm That the Conformational Compression upon Prephenate Binding Observed Crystallographically Also Occurs in Solution—To confirm the conformational changes upon prephenate binding observed in the crystal structure of GspDAH7PS and to gain insights into the possible allosteric regulation mechanism adopted by this enzyme, the SAXS data recorded in the absence and presence of prephenate were compared with the open and closed (prephenate-bound) models of the protein (Fig. 9). The theoretical scattering curves calculated for each model present a good fit with the corresponding experimental curves ($\chi^2 = 0.70$ and 0.37 for the closed and open forms, respectively). In contrast, the comparisons of the theoretical curve of the open structure to the data for prephenate-bound GspDAH7PS, and of the closed form to the apoenzyme data, give poor fits ($\chi^2 = 8.02$ and 3.13, respectively).

The experimental SAXS data were further fitted using a mixture of open and closed models of GspDAH7PS. In the absence of prephenate, the analysis shows a better fit of the theoretical scattering from a combination of both open and closed GspDAH7PS models ($\chi^2 = 0.22$) to the experimental data obtained in the absence of prephenate. This combination predicts that $\approx73\%$ of the protein adopts the open conformation, and $\approx27\%$ of the protein adopts the closed form conformation (Fig. 9A). On this basis, a conformational equilibrium may be established between the open and closed forms in the absence of ligand binding to the CM domains, and intermediate states between the open and closed forms may be present. Nevertheless, the open conformation or open-like structures are predominant in the absence of prephenate, suggesting that the closed form is less stable or more difficult to access. In the presence of prephenate, the best fit to the experimental profile ($\chi^2 = 0.37$) is obtained with that of the closed GspDAH7PS model alone, predicting that the closed structure, or a set of
conformationally similar structures, is the sole form adopted by the prephenate-bound protein (Fig. 9B). These data are consistent with the full loss of DAH7PS activity of GspDAH7PS by prephenate binding observed experimentally.

Discussion

Multifunctional enzymes are observed in many living systems, and they play important roles in substrate channeling, enzyme compartmentalization, and metabolic control (25). In the thermophilic soil bacterium Geobacillus sp., a CM domain is found fused to the N terminus of the first enzyme dedicated to aromatic amino acid biosynthesis, DAH7PS. We have shown here that this fusion gives rise to a bifunctional enzyme presenting both DAH7PS and CM catalytic activity. Additionally, we have shown that the smaller enzyme, CM, can bind its product prephenate, and this inhibits the activity of the fusion partner DAH7PS.

The helical AroQ form of CM is frequently observed as a fusion partner to enzymes involved in Tyr and Phe biosynthesis. For example, not only is it attached to the DAH7PS enzyme studied herein, it is found fused to both prephenate dehydratase and prephenate dehydrogenase enzymes (26–30). Geobacillus sp. has both the AroQ form (fused) and the other form of CM, AroH, encoded by its genome. Sequence inspection reveals that this AroH variant does not seem to be fused to another protein. For the B. subtilis DAH7PS enzyme, the CM catalytic activity of the AroQ in the fusion protein was shown to be extremely low, and it was predicted that the AroH protein was the major contributor to CM activity, and the primary role of the CM domain of the equivalent CM-DAH7PS fusion was for allosteric control (15). In contrast, the AroQ CM domain of the hyperthermophilic Geobacillus sp. fusion protein has moderate activity, suggesting it may play a dual role in both providing catalytic function in tandem with the AroH protein as well as providing a feedback mechanism to control entry into the pathway. Notably, the CM catalytic efficiency is enhanced by its association with DAH7PS, principally due to its significantly lower Michaelis constant.

Through a combination of enzyme kinetics, x-ray crystallography, SAXS, and modeling, we can now build a picture of the molecular details of how the feedback inhibition of DAH7PS is achieved in this class of DAH7PS enzymes. In the absence of the allosteric ligand prephenate, the CM domains remain at a distance from the DAH7PS catalytic barrels, permitting the substrates to enter the DAH7PS active site. The binding of prephenate to the CM domain is then accompanied by a conformational change, evident from SAXS data and comparison of our prephenate-bound crystal structure with the homologous unbound structure of LmoDAH7PS. The latter structure shows a far more extended structure in the absence of ligands within the CM domain binding site. In the presence of prephenate, the protein adopts a more compact and more ordered form, with the extension of helices in the CM domain, which bear residues forming binding contacts with prephenate. Furthermore, the SAXS data suggest that the CM domains remain close to the DAH7PS catalytic barrels in the presence of prephenate, because of the very good fit obtained for the data in the presence of inhibitor, using the closed form model. The compactness of the structure impacts on substrate access to the DAH7PS active
site and thus inhibits the enzyme. Conformational transitions and the occasional sampling of conformations resembling the closed form may account for the observed slightly higher activity of the truncated DAH7PS mutant lacking the CM domain. This mechanism of allostery involving physical gating of the active site has been employed for other DAH7PS enzymes, albeit with structurally distinct regulatory domains. The *T. maritima* DAH7PS bears an ACT regulatory domain, which on inhibitor Tyr binding also restricts access to the active site (7). These two quite distinct allosteric solutions suggest that adopting this structural approach for the control DAH7PS represents a general mechanism, albeit that both of these allosteric control mechanisms require the same basic tetrameric structure for the DAH7PS modules to operate. Other DAH7PS enzymes have been shown to adopt allosteric regulation mechanisms based on changes in protein dynamics (13, 14) rather than using a physical gating mechanism, which reinforces the versatility of both the protein fold and the approaches to allosteric control.

As noted above, the acquisition of the CM domain by DAH7PS comes with a slight burden to its catalytic function, as evidenced by the moderate increase in DAH7PS catalytic efficiency observed for the truncated protein of GspDAH7PS<sup>S</sup>. In contrast, the CM activity benefits from its association with its DAH7PS partner, with its independent activity following excision being reduced. This reduction suggests that the transient contacts the CM dimer forms with its DAH7PS partner in the full-length protein are important for its catalytic function as well as for the delivery of allosterically. Although it is difficult to unravel the individual contributions to binding and catalysis from the steady-state kinetic parameters, it is notable that the CM activity of the full-length protein has a lower chorismate *K*<sub>m</sub> value (by a factor of almost 5) and a lower *k*<sub>cat</sub> value (almost a 3-fold decrease). These changes reveal the impact domain interaction observed in the x-ray structure has on CM catalytic function. The decrease in *K*<sub>m</sub> value for the fused CM may reflect the contribution that conformational restriction affords by the interaction with DAH7PS, whereas the lower *k*<sub>cat</sub> value may reflect the tailoring of the CM active site for prephenate binding and the provision of allosterically for the DAH7PS activity, rather than for catalyzing the CM reaction.

The connected and overlapping functions between enzymatic activities demonstrate the complexity of cellular mechanisms for both providing allosteric control and delivering cellular biosynthetic functionality. DAH7PS, through its associations and varied mechanisms for providing allosteric control, provides insight into the evolutionary processes that provide solutions to biosynthetic control.

### Experimental Procedures

**Construction of Plasmids**—The gene sequence encoding GspDAH7PS<sup>WT</sup> (strain Y412MC61) was codon-optimized for expression in *E. coli*, synthesized by Geneart (Life Technologies, Inc.), and ligated into the expression vector pET-28b between the Ncol and Xhol sites to give a protein with a non-cleavable C-terminal His<sub>6</sub> tag. This plasmid was used as the template for the preparation of constructs for expressing GspDAH7PS<sup>S</sup> (the DAH7PS portion of GspDAH7PS<sup>WT</sup>) and GspDAH7PS<sup>C</sup> (the CM portion of GspDAH7PS<sup>WT</sup>) variants, created by truncating between Lys-92 and Ala-93. Genes

---

**TABLE 2**

| Primers for generation of constructs | GspDAH7PS<sup>S</sup> primers (round 1) | Reverse primer | GspDAH7PS<sup>C</sup> primers (round 1) | Reverse primer | Generic primers (round 2) | Reverse primer |
|------------------------------------|----------------------------------------|----------------|----------------------------------------|----------------|--------------------------|----------------|
| Forward primer | 5′-GCGACCCGGCTGATTTTTCTAGCAAGAGGCTCCGAA | 5′-GCGACCCGGCTGATTTTTCTAGCAAGAGGCTCCGAA | Forward primer | 5′-GCGACCCGGCTGATTTTTCTAGCAAGAGGCTCCGAA | Forward primer | 5′-GCGACCCGGCTGATTTTTCTAGCAAGAGGCTCCGAA |
| Reverse primer | 5′-GGCGGACCCGGCTGATTTTTCTAGCAAGAGGCTCCGAA | 5′-GGCGGACCCGGCTGATTTTTCTAGCAAGAGGCTCCGAA | Reverse primer | 5′-GGCGGACCCGGCTGATTTTTCTAGCAAGAGGCTCCGAA | Reverse primer | 5′-GGCGGACCCGGCTGATTTTTCTAGCAAGAGGCTCCGAA |
encoding the truncated proteins were amplified using gene-specific primers (Table 2) for cloning using the Gateway system (Life Technologies, Inc.). The products from the first round of PCR were extended using primers that incorporated the attB1 site and attB2 site, and the PCR products were gel-purified. The purified products were cloned into the donor vector pDONR-221 using BP clonase enzyme mix (Life Technologies, Inc.), and the product was subsequently transformed into *E. coli* One Shot® TOP10 cells (Life Technologies, Inc.). Purified plasmids from transformants were sequence-verified. LR clonase enzyme mix (Life Technologies, Inc.) was used to sub-clone from transformants were sequence-verified. LR clonase enzyme mix (Life Technologies, Inc.) was used to sub-clone.

### Purification of GspDAH7PS WT, GspDAH7PSD, and GspDAH7PS C—

- The vectors bearing the genes of GspDAH7PS WT and GspDAH7PS C were transformed into *E. coli* BL21 (DE3) cells, whereas the vector bearing the gene coding for GspDAH7PS D was transformed into *E. coli* BL21 (DE3) pBB540/pBB542 cells for expression. Cultures were grown at 37 °C in 1 liter of Luria-Bertani (LB) medium supplemented with 100 μg/ml kanamycin for GspDAH7PS WT, 100 μg/ml ampicillin for GspDAH7PS C; and a combination of 100 μg/ml ampicillin, 25 μg/ml chloramphenicol, and 10 μg/ml spectinomycin for GspDAH7PS D. Expression of target protein was induced through the addition of 0.5 mM isopropyl β-D-thiogalactopyranoside, and the cultures were grown for a further 4 h at 37 °C (for expression of GspDAH7PS WT) or overnight at 23 °C (for the expression of GspDAH7PS D and GspDAH7PS C). Cells were harvested by centrifugation and resuspended in lysis buffer (50 mM 1,3-bis(tris(hydroxymethyl) sodium phosphate, 500 mM NaCl, 50 mM imidazole, pH 7.4). The column was washed with binding buffer, before overnight treatment with TEV protease at a 1% mol/mol concentration. TEV-treated enzyme was subsequently run through a second GSTrap HP column to separate protease and GST tag from cleaved enzyme. Untagged enzyme eluted in the flow-through of the second column, and fractions were pooled for SEC.

### Determination of Enzymatic Activity—Assays were performed in 50 mM BTP buffer, pH 6.8, at the stated temperature in a total volume of 1 ml. Initial rates were fitted to the Michaelis-Menten equation to determine kinetic parameters using GraFit (Erich tus software).

**DAH7PS Assay**—A modified continuous assay of Schoner and Herrmann (31) was used as described previously by Schoefield et al. (32) for use at elevated temperatures. Enzyme activity was monitored by following the loss of absorbance at 232 nm due to consumption of PEP at 60 °C in the presence of CdCl2 (final concentration of 100 μM) following the addition of E4P to the assay solution containing PEP and enzyme (2.8—3.5 × 10−2 μM). For the determination of kinetic parameters, initial rates of reaction were determined by keeping the concentration of one substrate fixed (PEP concentration was fixed at 338 μM and E4P concentration was fixed at 329 μM) while varying the concentration of the other substrate. The metal activation profiles of GspDAH7PS WT and GspDAH7PS D were determined using standard assays with 195 μM PEP and 197 μM E4P at 60 °C in the presence of 100 μM CdCl2, MnSO4, MgSO4, Zn(CO2CH3)2, NiCl2, CoCl2, EDTA, dipicolinic acid, or no chelator or metal ion (control) and enzyme (2.8 × 10−2 μM).

**CM Assay**—Steady-state kinetics were measured using a continuous assay monitoring the disappearance of chorismate at 274 nm at 50 °C (extinction coefficient of 2,630 M−1 cm−1 at 50 °C and pH 6.8). Reactions were initiated with the addition of chorismate (Sigma) at a desired concentration to a buffer solution containing enzyme (0.94 μM).

### Temperature Dependence of Enzyme Activity—DAH7PS activity was measured at 325 μM PEP, 475 μM E4P for GspDAH7PS WT and 337 μM PEP, 356 μM E4P for GspDAH7PS D and 100 μM Cd2+ in 50 mM BTP, pH 6.8. Enzyme (2.5 × 10−2 μM for GspDAH7PS WT and 10−2 μM for GspDAH7PS D) was added for 5 min prior to the initiation of activity by the addition of E4P. CM activity was measured at 70 μM chorismate for GspDAH7PS WT and 80 μM chorismate for GspDAH7PS C in 50 mM BTP, pH 6.8. Enzyme (0.9 μM for GspDAH7PS WT and 1.25 μM for GspDAH7PS D) was added 5 min prior to initiation of the reaction with chorismate.
Bifunctional DAH7PS from Geobacillus

Inhibition by Prephenate—The inhibition profiles of DAH7PS activity for GspDAH7PS<sup>WT</sup> and GspDAH7PS<sup>D</sup> were determined by varying the concentration of prephenate (Sigma) in buffer with CdCl<sub>2</sub> (100 μM), PEP (121 μM), and enzyme (2.5 × 10<sup>-2</sup> μM) incubated at 60 °C for 10 min. The reactions were initiated by the addition of E4P (123 μM).

Crystallization and Structure Solution—Crystals of GspDAH7PS<sup>WT</sup> were grown by hanging-drop vapor diffusion. A protein solution (11 mg ml<sup>-1</sup> in 20 mM BTP, 40 mM KCl, 200 μM PEP, pH 7.4) was mixed 1:1 (v/v) with a reservoir solution containing 0.2 M sodium citrate, 0.1 M BTP, pH 6.5, 20% w/v polyethylene glycol (PEG) 3350, and 0.2 mM chorismic acid. The drop sizes were 2 μl, and the reservoir volume was 500 μl. The crystallization trays were incubated at 20 °C until immediately before loading the crystals into the loop for flash cooling because of the weak electron density in these regions. These residues, as well as some unmodded side chains, were restructured using MODELLER (24) version 9.13. First, the tetrameric form was generated from the asymmetric dimeric unit in PyMOL (40), and the topologies and parameters for prephenate and Mn<sup>2+</sup>, obtained from the SwissParam server (41) and the MM3 force field (42–44), respectively, were added to the MODELLER library. Second, 100 models of GspDAH7PS were generated using the “automodel” routine of MODELLER tuned to improve model accuracy. More specifically, the variable target function method was set to slow, and the maximum number of conjugate gradient iterations was set to 300, although the level of molecular dynamics refinement was adjusted to slow and the complete optimization-refinement process was repeated twice. The “env.io.hetatm” instruction was defined as true to include the metal ion and prephenate in the model. During the model generation, only the missing residues and residues with missing atoms were allowed to move, and the chorismate mutase domains were restrained to keep their secondary structure intact, and a symmetry restraint was applied on the α-carbons of the symmetric chains of the tetramer. The best model was selected among the models presenting the lowest MODELLER objective function value and the lowest global DOPE score, following visual inspection and residue-by-residue DOPE score analysis. The C-terminal tail of this model was further refined using the “loopmodel” class of MODELLER (45) using the same protocol as described above, and the best of the 100 models generated was assessed using the same evaluation procedure.

Homology Model Construction of GspDAH7PS in Its Open Form—Homology modeling of GspDAH7PS in its open, active form was performed using the two previously determined structures of DAH7PS from L. monocytogenes EGD-e (PDB codes 3NTV and 3TFC) as templates. For each structure of LmoDAH7PS, the tetrameric form was generated in PyMOL, and the topology and parameters of PEP obtained with SwissParam as well as those of Mn<sup>2+</sup> were included into MODELLER libraries. The missing residues and side chains were then added with MODELLER using automodel to create 25 models using the same parameters and the same assessment procedure as described above. The best model for each of the structures was aligned, and their sequences were aligned against the full sequence of GspDAH7PS using T-COFFEE (46). The homology model of GspDAH7PS in its open form was performed with MODELLER using the automodel routine as before with the env.io.hetatm instruction set to true to include PEP and Mn<sup>2+</sup> in the model. Secondary structure restraints were used to place the CM domains in the difference Fourier map using phased molecular replacement in MOLREP (37). Further structural refinement was performed using REFMAC5, and electron density maps were analyzed using Coot (38). Prephenate was modeled after the penultimate round of refinement. Structures were validated using the MolProbity server (39) and the tools of Coot.
were applied to the chorismate mutase domains, and symmetry restraints were used for the symmetric chains of the tetramer. A total of 100 models were generated, as above, and the best model was selected among the models presenting the lowest MODELLER objective function value and the lowest global DOPE score, following visual inspection and residue-by-residue DOPE score analysis.

SAXS—Measurements were performed at the Australian Synchrotron. SAXS/WAXS beamline was equipped with a Pilatus detector (1M, 170 × 170 mm, effective pixel size, 172 × 172 μm). The wavelength of the x-rays was 1.0332 Å. The sample detector distance was 2.6 m, which provided an s range of 0.01–0.35 Å⁻¹ (where q is the magnitude of the scattering vector, which is related to the scattering angle 2θ and the wavelength (λ) as follows: s = (4λ/π)sin(θ). Data were collected from samples in a 1.5-mm thin-walled glass capillary at 25 °C at 2-s intervals.

Scattering data were collected from GspDAH7PSWT following elution from an SEC column (Superdex 200 5/150), pre-equilibrated with SEC buffer (10 mM BTP, pH 7.4, 150 mM NaCl, 200 μM PEP) or SEC buffer with prephenate (10 mM BTP, pH 7.4, 150 mM NaCl, 200 μM PEP, 400 μM prephenate).

Raw scattering data were processed with Scatterbrain developed at the Australian Synchrotron. Scattered intensity (I) was plotted versus s using Primus (47) and Origin 8. All samples were devoid of an intensity increase at low s (indicative of aggregation). Guinier plots were linear for sRg <1.3. The data sets for structural analysis were recorded with 447 data points over the range 0.005 ≤ s ≤ 0.35 Å⁻¹. Theoretical scattering curves were generated from the models of open form (active) and closed form (inactive) GspDAH7PS generated as described above and compared with experimental scattering curves using OLGOMER (47).

Theoretical scattering data of two distinct GspDAH7PS forms were further mixed and fitted to the individual experimental scattering profile in the presence or absence of prephenate with OLGOMER to investigate the volume fraction of each form in the solution with and without ligand (47).

Additional Computational Methods—SASA for the open and closed models were calculated in VMD (48) using the “measure sasa” command and using a solvent probe radius of 1.4 Å. Sequence identities were determined using BLASTN (49) using the BLOSUM50 scoring matrix with an opening gap penalty of −12 and an extending gap penalty of −2. Sequence alignments were generated with T-Coffee (50) and rendered with ESPript 3 (51). Structural figures were generated with PyMOL.

Author Contributions—A. R. N., M. H. O., and Y. B. carried out the experiments. E. J. M. L., Y. B., and M. H. O. analyzed the data. S. P., V. L. M., and T. M. A. assisted with the x-ray structure solution. E. J. P. conceived and coordinated the study. A. R. N., E. J. M. L., T. M. A., Y. B., and E. J. P. wrote the paper. All authors reviewed the results and approved the final version of the manuscript.

Acknowledgments—Data were collected at the MX and SAXS beamlines of the Australian Synchrotron with access provided by the New Zealand Synchrotron Group.

References

1. Knaggs, A. R. (2003) The biosynthesis of shikimate metabolites. Nat. Prod. Rep. 20, 119–136
2. Light, S. H., and Anderson, W. F. (2013) The diversity of allosteric controls at the gateway to aromatic amino acid biosynthesis. Protein Sci. 22, 395–404
3. König, V., Pfeil, A., Braus, G. H., and Schneider, T. R. (2004) Substrate and metal complexes of 3-deoxy-D-arabino-heptulosonate-7-phosphate synthase from Saccharomyces cerevisiae provide new insights into the catalytic mechanism. J. Mol. Biol. 337, 675–690
4. Shumilin, I. A., Zhao, C., Bauerle, R., and Krettsinger, R. H. (2002) Allosteric inhibition of 3-deoxy-D-arabino-heptulosonate-7-phosphate synthase alters the coordination of both substrates. J. Mol. Biol. 320, 1147–1156
5. Cross, P. J., Pietersma, A. L., Allison, T. M., Wilson-Couts, S. M., Cochrane, F. C., and Parker, E. J. (2013) Neisseria meningitidis expresses a single 3-deoxy-D-arabino-heptulosonate 7-phosphate synthase that is inhibited primarily by phenylalanine. Protein Sci. 22, 1087–1099
6. Schofield, L. R., Anderson, B. F., Patchett, M. L., Norris, G. E., Jameson, G. B., and Parker, E. J. (2005) Substrate ambiguity and crystal structure of Pyrococcus furiosus 3-deoxy-D-arabino-heptulosonate-7-phosphate synthase: an ancestral 3-deoxyald-2-ulosonate-phosphate synthase? Biochemistry 44, 11950–11962
7. Cross, P. J., Dobson, R. C., Patchett, M. L., and Parker, E. J. (2011) Tyrosine latching of a regulatory gate affords allosteric control of aromatic amino acid biosynthesis. J. Biol. Chem. 286, 10216–10224
8. Lang, E. J., Cross, P. J., Mittelstädt, G., Jameson, G. B., and Parker, E. J. (2014) Allosteric ACTion: the varied ACT domains regulating enzymes of amino acid metabolism. Curr. Opin. Struct. Biol. 29, 102–111
9. Wu, J., Shehban, G. Y., and Woodard, R. W. (2005) Bacillus subtilis 3-deoxy-D-arabino-heptulosonate 7-phosphate synthase revisited: resolution of two long-standing enigmas. Biochem. J. 390, 583–590
10. Webbey, C. J., Baker, H. M., Lott, J. S., Baker, E. N., and Parker, E. J. (2005) The structure of 3-deoxy-D-arabino-heptulosonate 7-phosphate synthase from Mycobacterium tuberculosis reveals a common catalytic scaffold and ancestry for type I and type II enzymes. J. Mol. Biol. 354, 927–939
11. Blackmore, N. J., Reichau, S., Jiao, W., Hutton, R. D., Baker, E. N., Jameson, G. B., and Parker, E. J. (2013) Three sites and you are out: ternary synergistic allosteric controls aromatic amino acid biosynthesis in Mycobacterium tuberculosis. J. Mol. Biol. 425, 1582–1592
12. Sasso, S., Okvist, M., Roderer, K., Gamper, M., Codoni, G., Krengel, U., and Kast, P. (2009) Structure and function of a complex between chorismate mutase and DAHP synthase: efficiency boost for the junior partner. EMBO J. 28, 2128–2142
13. Jiao, W., Hutton, R. D., Cross, P. J., Jameson, G. B., and Parker, E. J. (2012) Dynamic cross-talk among remote binding sites: the molecular basis for unusual synergistic alloster, J. Mol. Biol. 415, 716–726
14. Lang, E. J., Heyes, L. C., Jameson, G. B., and Parker, E. J. (2016) Calculated pKa variations expose dynamic allosteric communication networks. J. Am. Chem. Soc. 138, 2036–2045
15. Wu, J., and Woodard, R. W. (2006) New insights into the evolutionary links relating to the 3-deoxy-D-arabino-heptulosonate 7-phosphate synthase subfamilies. J. Biol. Chem. 281, 4042–4048
16. Light, S. H., Halavaty, A. S., Minasov, G., Shuvalova, L., and Anderson, W. F. (2012) Structural analysis of a 3-deoxy-D-arabino-heptulosonate 7-phosphate synthase with an N-terminal chorismate mutase-like regulatory domain. Protein Sci. 21, 887–895
17. Webbey, C. J., Jiao, W., Hutton, R. D., Blackmore, N. J., Baker, H. M., Baker, E. N., Jameson, G. B., and Parker, E. J. (2010) Synergistic alloster, a sophisticated regulatory network for the control of aromatic amino acid biosynthesis. J. Biol. Chem. 285, 30567–30576
18. Rambo, R. P., and Tainer, J. A. (2011) Characterizing flexible and intrinsically unstructured biological macromolecules by SAS using the Porod-Debye law. Biopolymers 95, 559–571
19. Hammel, M. (2012) Validation of macromolecular flexibility in solution by small-angle x-ray scattering (SAXS). Eur. Biophys. J. 41, 789–799
Bifunctional DAH7PS from Geobacillus

20. Lee, A. Y., Karplus, P. A., Ganem, B., and Clardy, J. (1995) Atomic structure of the buried catalytic pocket of Escherichia coli chorismate mutase. J. Am. Chem. Soc. 117, 3627–3628

21. Hur, S., and Bruce, T. C. (2002) The mechanism of catalysis of the chorismate to prephenate reaction by the Escherichia coli mutase enzyme. Proc. Natl. Acad. Sci. U.S.A. 99, 1176–1181

22. Liu, D. R., Cloud, S. T., Pastor, R. M., and Schultz, P. G. (1996) Analysis of active site residues in Escherichia coli chorismate mutase by site-directed mutagenesis. J. Am. Chem. Soc. 118, 1789–1790

23. Lassila, J. K., Keeffe, J. R., Kast, P., and Mayo, S. L. (2007) Exhaustive mutagenesis of six secondary active-site residues in Escherichia coli chorismate shows the importance of hydrophobic side chains and a helix N-capping position for stability and catalysis. Biochemistry 46, 6883–6891

24. Sali, A., and Blundell, T. L. (1993) Comparative protein modelling by satisfaction of spatial restraints. J. Mol. Biol. 234, 779–815

25. Elleuch, S. (2015) Bringing functions together with fusion enzymes—from nature’s inventions to biotechnological applications. Appl. Microbiol. Biotechnol. 99, 1545–1556

26. Tzin, V., Malitsky, S., Aharoni, A., and Galili, G. (2009) Expression of a bacterial bi-functional chorismate mutase/prephenate dehydratase modules primary and secondary metabolism associated with aromatic amino acids in Arabidopsis. Plant J. 60, 156–167

27. Lütke-Eversloh, T., and Stephanopoulos, G. (2005) Feedback inhibition of chorismate mutase/prephenate dehydratase from Escherichia coli: generation and characterization of tyrosine-insensitive mutants. J. Biol. Chem. 280, 2782–2789

28. Vincent, S., Chen, S., Wilson, D. B., and Ganem, B. (2002) Probing the overlap of chorismate mutase and prephenate dehydrogenase sites in the Escherichia coli T-protein: a dehydrogenase-selective inhibitor. Bioorg. Med. Chem. Lett. 12, 929–931

29. Zhang, S., Pohnert, G., Kongsaeer, P., Wilson, D. B., Clardy, J., and Ganem, B. (1998) Chorismate mutase-prephenate dehydratase from Escherichia coli—study of catalytic and regulatory domains using genetically engineered proteins. J. Biol. Chem. 273, 6248–6253

30. Turnbull, J., Cleland, W. W., and Morrison, J. F. (1990) Chorismate mutase prephenate dehydrogenase from Escherichia coli. 1. Kinetic characterization of the dehydrogenase reaction by use of alternative substrates. Biochemistry 29, 10245–10254

31. Schoner, R., and Herrmann, K. M. (1976) 3-Deoxy-D-arabino-heptulosonate 7-phosphate synthase. Purification, properties, and kinetics of the tyrosine-sensitive isoenzyme from Escherichia coli. J. Biol. Chem. 251, 5440–5447

32. Schofield, L. R., Patchett, M. L., and Parker, E. J. (2004) Expression, purification, and characterization of 3-deoxy-D-arabino-heptulosonate 7-phosphate synthase from Pyrococcus furiosus. Protein Expr. Purif. 34, 17–27

33. Kabsch, W. (2010) XDS. Acta Crystallogr. D Biol. Crystallogr. 66, 125–132

34. Evans, P. R. (2011) An introduction to data reduction: space-group determination, scaling and intensity statistics. Acta Crystallogr. D Biol. Crystallogr. 67, 282–292

35. McCoy, A. J., Grosse-Kunstleve, R. W., Adams, P. D., Winn, M. D., Storoni, L. C., and Read, R. J. (2007) Phaser crystallographic software. J. Appl. Crystallogr. 40, 658–674

36. Murshudov, G. N., Skubak, P., Lebedev, A. A., Pannu, N. S., Steiner, R. A., Nicholls, R. A., Winn, M. D., Long, F., and Vagin, A. A. (2011) REFMAC5 for the refinement of macromolecular crystal structures. Acta Crystallogr. D Biol. Crystallogr. 67, 355–367

37. Vagin, A., and Teplyakov, A. (2010) Molecular replacement with MOLREP. Acta Crystallogr. D Biol. Crystallogr. 66, 22–25

38. Emsley, P., Lohkamp, B., Scott, W. G., and Cowtan, K. (2010) Features and development of Coot. Acta Crystallogr. D Biol. Crystallogr. 66, 486–501

39. Chen, V. B., Arendall, W. B., 3rd., Headd J. J., Keedy, D. A., Immormino, R. M., Kapral, G. J., Murray, I. W., Richardson, J. S., and Richardson, D. C. (2010) MolProbity: all-atom structure validation for macromolecular crystallography. Acta Crystallogr. D Biol. Crystallogr. 66, 12–21

40. Delano, W. L. (2010) The PyMOL Molecular Graphics System, Version 1.3ri, Schrodinger, LLC, New York

41. Zoete, V., Cucu, M. A., Grosdidier, A., and Michielin, O. (2011) Swiss-Param: a fast force field generation tool for small organic molecules. J. Comput. Chem. 32, 2359–2368

42. Allinger, N. L., Yuh, Y. H., and Lii, J. H. (1989) Molecular mechanics. The MM3 force field for hydrocarbons. 1. J. Am. Chem. Soc. 111, 8551–8566

43. Lii, J. H., and Allinger, N. L. (1989) Molecular mechanics. The MM3 force field for hydrocarbons. 2. Vibrational frequencies and thermodynamics. J. Am. Chem. Soc. 111, 8566–8575

44. Lii, J. H., and Allinger, N. L. (1989) Molecular mechanics. The MM3 force field for hydrocarbons. 3. The van der Waals’ potentials and crystal data for aliphatic and aromatic hydrocarbons. J. Am. Chem. Soc. 111, 8576–8582

45. Fiser, A., Do, R. K., and Sali, A. (2000) Modeling of loops in protein structures. Protein Sci. 9, 1753–1773

46. Notredame, C., Higgins, D. G., and Heringa, J. (2000) T-Coffee: A novel method for fast and accurate multiple sequence alignment. J. Mol. Biol. 302, 205–217

47. Konarev, P. V., Volkov, V. V., Sokolova, A. V., Koch, M. H. J., and Svergun, D. I. (2003) PRIMUS: a Windows PC-based system for small-angle scattering data analysis. J. Appl. Crystallogr. 36, 1277–1282

48. Humphrey, W., Dalke, A., and Schulten, K. (1996) VMD: Visualization of molecular dynamics. J. Mol. Graph. 14, 33–38

49. Huang, X. C., and Miller, W. (1991) A time-efficient, linear-space local similarity algorithm. Adv. Appl. Math. 12, 337–357

50. Di Tommaso, P., Moretti, S., Xenarios, I., Orbritig, M., Montanyola, A., Chang, J. M., Taly, J. F., and Notredame, C. (2011) T-Coffee: a web server for the multiple sequence alignment of protein and RNA sequences using structural information and homology extension. Nucleic Acids Res. 39, W13–W17

51. Robert, X., and Gouet, P. (2014) Deciphering key features in protein structures with the new ENDscript server. Nucleic Acids Res. 42, W320–W324
Interdomain Conformational Changes Provide Allosteric Regulation en Route to Chorismate
Ali Reza Nazmi, Eric J. M. Lang, Yu Bai, Timothy M. Allison, Mohamad H. Othman, Santosh Panjikar, Vickery L. Arcus and Emily J. Parker

J. Biol. Chem. 2016, 291:21836-21847. doi: 10.1074/jbc.M116.741637 originally published online August 8, 2016

Access the most updated version of this article at doi: 10.1074/jbc.M116.741637

Alerts:
- When this article is cited
- When a correction for this article is posted

Click here to choose from all of JBC's e-mail alerts

This article cites 50 references, 9 of which can be accessed free at http://www.jbc.org/content/291/42/21836.full.html#ref-list-1