

Reprogramming the Chemodiversity of Terpenoid Cyclization by Remolding the Active Site Contour of *epi*-Isozizaene Synthase

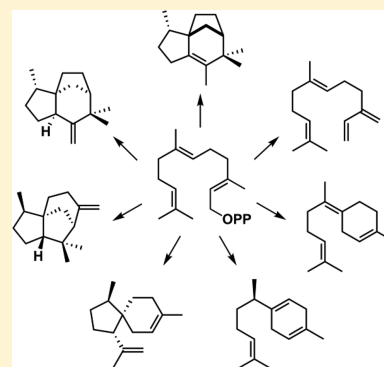
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ABSTRACT: The class I terpenoid cyclase *epi*-isozizaene synthase (EIZS) utilizes the universal achiral isoprenoid substrate, farnesyl diphosphate, to generate *epi*-isozizaene as the predominant sesquiterpene cyclization product and at least five minor sesquiterpene products, making EIZS an ideal platform for the exploration of fidelity and promiscuity in a terpenoid cyclization reaction. The hydrophobic active site contour of EIZS serves as a template that enforces a single substrate conformation, and chaperones subsequently formed carbocation intermediates through a well-defined mechanistic sequence. Here, we have used the crystal structure of EIZS as a guide to systematically remodel the hydrophobic active site contour in a library of 26 site-specific mutants. Remolded cyclization templates reprogram the reaction cascade not only by reproportioning products generated by the wild-type enzyme but also by generating completely new products of diverse structure. Specifically, we have tripled the overall number of characterized products generated by EIZS. Moreover, we have converted EIZS into six different sesquiterpene synthases: F96A EIZS is an (*E*)- β -farnesene synthase, F96W EIZS is a zizaene synthase, F95H EIZS is a β -cucumene synthase, F95M EIZS is a β -acoradiene synthase, F198L EIZS is a β -cedrene synthase, and F96V EIZS and W203F EIZS are (*Z*)- γ -bisabolene synthases. Active site aromatic residues appear to be hot spots for reprogramming the cyclization cascade by manipulating the stability and conformation of critical carbocation intermediates. A majority of mutant enzymes exhibit only relatively modest 2–100-fold losses of catalytic activity, suggesting that residues responsible for triggering substrate ionization readily tolerate mutations deeper in the active site cavity.



The family of terpenoid natural products, which includes steroids and carotenoids, contains more than 70000 unique compounds that have been identified to date [Dictionary of Natural Products (<http://dnp.chemnetbase.com>)]. This vast chemodiversity belies relatively simple roots in just two fundamental isoprenoid building blocks, the C₅ compounds dimethylallyl diphosphate and isopentenyl diphosphate.^{1,2} These simple universal precursors can be coupled in head-to-tail fashion to generate longer acyclic isoprenoids such as geranyl diphosphate (C₁₀), farnesyl diphosphate (C₁₅), and geranylgeranyl diphosphate (C₂₀),^{3,4} which in turn are substrates for terpenoid cyclases. While the specific mechanism of isoprenoid diphosphate cyclization varies from one cyclase to another, common mechanistic features include initial ionization of the allylic diphosphate substrate followed by a complex reaction sequence that proceeds through electrophilic attack on the internal double bonds involving multiple carbocation intermediates.^{5–9}

Product chemodiversity for an individual terpenoid cyclase is a measure of its function as a template and chaperone. A terpenoid cyclase such as aristolochene synthase from *Aspergillus terreus* can be a strict chaperone and generate a single product with high fidelity.¹⁰ Alternatively, a terpenoid

cyclase can serve as a more permissive chaperone and allow for multiple cyclization cascades leading to complex product mixtures. For example, γ -humulene synthase from *Abies grandis* generates 52 different sesquiterpene products from the substrate farnesyl diphosphate.¹¹ Thus, the catalytic activity of a terpenoid cyclase is described not only by its reaction kinetics but also by the number and structural diversity of the reaction products it generates.

The structural basis of fidelity and promiscuity in terpenoid cyclase reactions, particularly with regard to the code that relates the three-dimensional contour of an active site template with its cyclization product array, is a topic of intense interest in the field of enzyme structure and function. Recently, we determined the X-ray crystal structure of the sesquiterpene cyclase *epi*-isozizaene synthase (EIZS) from *Streptomyces coelicolor* at 1.6 Å resolution.^{12–14} EIZS is classified as a class I terpenoid cyclase because it utilizes a trinuclear magnesium cluster liganded by conserved DDXXD and NDXSXXXE motifs¹⁵ to trigger the formation of the initial substrate carbocation and inorganic pyrophosphate (PP_i). Because

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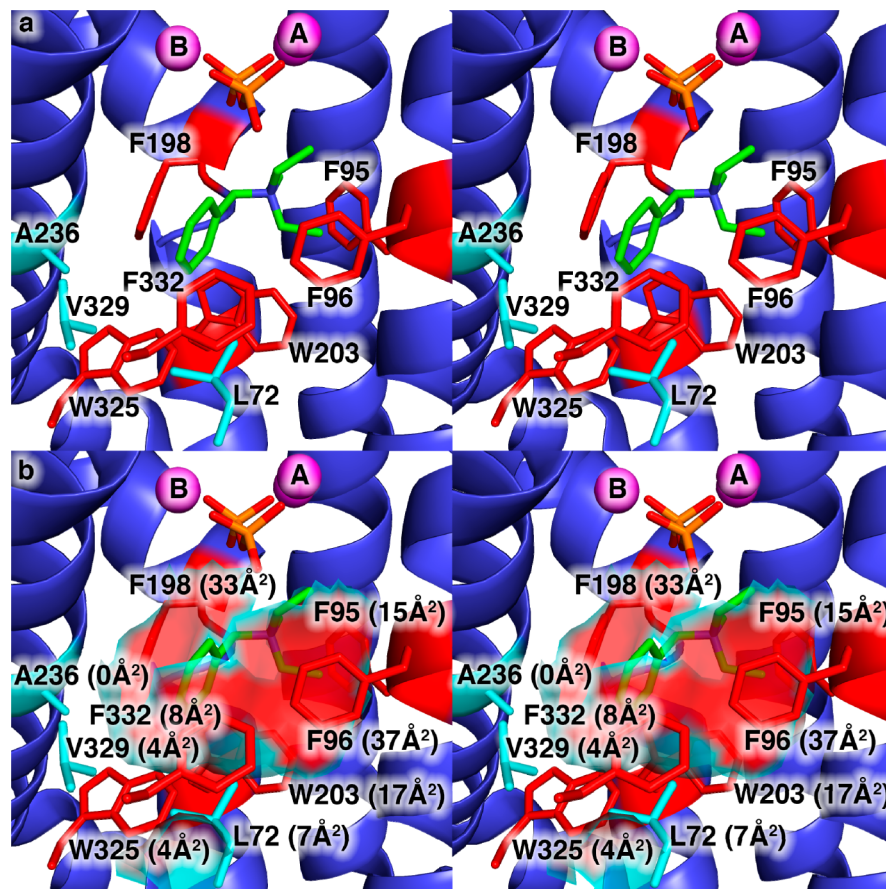


Figure 1. (a) View of the hydrophobic EIZS active site in the wild-type EIZS–Mg²⁺–PP_i–BTAC complex showing the locations of aromatic residues (red) and aliphatic residues (cyan) that largely define the active site contour shown in panel b. Portions of the protein structure in the foreground, including the α -helix containing W325, V329, and F332, are cut away to allow the active site to be viewed. Mg²⁺ ions are shown as magenta spheres. P and O atoms of the PP_i anion are colored orange and red, respectively. C and N atoms of the BTAC cation are colored green and blue, respectively. The solvent-accessible surface area contributed by each residue is given in parentheses. Note that although A236 does not contribute to the solvent-accessible surface area in the wild-type enzyme, it is sufficiently close to the surface that substitution of larger side chains will impact the surface contour.

EIZS is moderately promiscuous, generating at least five minor sesquiterpene products in addition to 79% *epi*-isozizaene from the universal sesquiterpene substrate farnesyl diphosphate (FPP),^{13,14} it is an ideal platform for the exploration of fidelity and promiscuity in a terpenoid cyclization reaction. Minor products generated by an enzyme may signal potential toeholds for the introduction of new activities in nature and in the laboratory,¹⁶ and this perspective applies to the terpenoid cyclases. For example, limited mutations in the active site contour of γ -humulene synthase can minimize the formation of several major products and maximize the formation of a variety of normally minor products.¹⁷ Importantly, residues beyond those that define the immediate active site contour can also significantly influence the product ratios of a terpenoid cyclase.¹⁸

Here, we describe the preparation, assay, and comparison of 26 single-site mutants of EIZS (three of which have been previously described¹³) in which several key aromatic and aliphatic amino acids comprising the hydrophobic active site contour have been systematically varied (Figure 1). We reasoned that by leaving intact the residues responsible for triggering initial substrate ionization (those residues that coordinate to divalent metal ions and residues that hydrogen bond with the diphosphate group of FPP), each mutant enzyme

would be able to generate an initial allylic cation from substrate FPP that might possibly be chaperoned through an alternative reaction sequence because of the remolded active site contour. We focused on three types of residues: (1) the aromatic triad of F95, F96, and F198 [the aromatic ring faces of these residues encircle the active site cavity and engage in cation– π interactions with a quaternary ammonium analogue, the benzyltriethylammonium cation (BTAC)];¹³ (2) other aromatic residues that define the active site contour, W203, W325, and F332; and (3) aliphatic residues that contribute to the active site contour (L72 and V329) or lie just beneath the solvent-accessible surface of the contour (A236).

X-ray crystal structure determinations of selected mutants confirm the structure of the remolded active site contour and illuminate the binding mode of the analogue BTAC. Mutants retain varying levels of catalytic activity and exhibit altered product profiles; indeed, some mutants essentially lose the capacity to generate *epi*-isozizaene and instead generate completely new cyclization products. Specifically, we have tripled the overall product profile, represented by the number of characterized major and minor products generated from substrate FPP. Additionally, we have converted EIZS into six nominally different sesquiterpene synthases, each of which

Table 1. Mutagenic Primers Utilized To Generate EIZS Mutants^a

mutation	F or R ^b	5'-3' sequence
L72A	F	GTC TGT GCT ACA CGG ACg ctA TGG CGG GCT ACT ACC TG
	R	CAG GTA GTA GCC CGC CAT agc GTC CGT GTA GCA CAG AC
L72V	F	CTG TGC TAC ACG GAC gtg ATG GCG GGC TAC TAC
	R	GTA GTA GCC CGC CAT cac GTC CGT GTA GCA CAG
L72M	F	GTC TGT GCT ACA CGG ACa tgA TGG CGG GCT ACT ACC TG
	R	CAG GTA GTA GCC CGC CAT cat GTC CGT GTA GCA CAG AC
F95A	F	GAC TAC AGC GCG TGG gca TTC GTC TGG GAC GAC
	R	GTC GTC CCA GAC GAA tgc CCA CGC GCT GTA GTC
F95V	F	GAT AGC GGA CTA CAG CGC GTG Ggt tTT TGT CTG GGA CGA CCG TCA C
	R	GTG ACG GTC GTC CCA GAC AAA aac CCA CGC GCT GTA GTC CGC TAT C
F95M	F	GAT AGC GGA CTA CAG CGC GTG Gat gTT TGT CTG GGA CGA CCG TCA C
	R	GTG ACG GTC GTC CCA GAC AAA cat CCA CGC GCT GTA GTC CGC TAT C
F95H	F	GAT AGC GGA CTA CAG CGC GTG Gca tTT TGT CTG GGA CGA CCG TCA C
	R	GTG ACG GTC GTC CCA GAC AAA atg CCA CGC GCT GTA GTC CGC TAT C
F96V	F	GAT AGC GGA CTA CAG CGC GTG gtt CGT TGT CTG GGA CGA CCG TCA C
	R	GTG ACG GTC GTC CCA GAC AAC GAA cca CGC GCT GTA GTC CGC TAT C
F96L	F	GAC TAC AGC GCG TGG TTC tta GTC TGG GAC GAC CGT CAC
	R	GTG ACG GTC GTC CCA GAC taa GAA CCA CGC GCT GTA GTC
F96Y	F	GAT AGC GGA CTA CAG CGC GTG GTT Cta tGT CTG GGA CGA CCG TCA C
	R	GTG ACG GTC GTC CCA GAC ata GAA CCA CGC GCT GTA GTC CGC TAT C
F96W	F	GAC TAC AGC GCG TGG TTC tgg GTC TGG GAC GAC CGT CAC
	R	GTG ACG GTC GTC CCA GAC cca GAA CCA CGC GCT GTA GTC
F198V	F	GAA CTG CGC CGG CTC ACG gtt GCT CAC TGG ATC TGG AC
	R	GTC CAG ATC CAG TGA GCa acC GTG AGC CGG CGC AGT TC
F198L	F	CTC GAA CTG CGC CGG CTC ACG tta GCG CAC TGG ATC TGG ACC GAC
	R	GTC GGT CCA GAT CCA GTG CGC taa CGT GAG CCG CGC CAG TTC GAG
F198Y	F	GTA CCT CGA ACT GCG CCG GCT CAC Gta tGC ACA CTG GAT CTG GAC CGA C
	R	GTC GGT CCA GAT CCA GTG TGC ata CGT GAG CCG GCG CAG TTC GAG GTA C
A236G	F	GTC AGG AAT TCg gcG CCT GGT ACA AC
	R	GTT GTC CCA GGC gcc GAA TTC CTG AC
A236F	F	GCG TTG CTG AGT CAG GAA TTC ttG GCC TGG TAC AAC GAC CTC TGC
	R	GCA GAG GTC GTT GTC CCA GGC aaa GAA TTC CTG ACT CAG CAA CGC
A236M	F	GCG TTG CTG AGT CAG GAA TTC atg GCC TGG TAC AAC GAC CTC TGC
	R	GCA GAG GTC GTT GTC CCA GGC cat GAA TTC CTG ACT CAG CAA CGC
W325F	F	GGC AAT ATG CGG AAC TTC ttc AGT TCC GTC TAC TG
	R	CAG TAG ACG GAA CTG AAg aaG TTC CGC ATA TTG CC
V329A	F	CTG GTT CAG TTC Cgg tTA CTG GTT CCA CCA CGA GTC
	R	GGT GGA ACC AGT Acg cGG AAC TGA ACC AG
V329G	F	CTG GTT CAG TTC Cgg tTA CTG GTT CCA CCA CGA GTC
	R	GAC TCG TGG TGG AAC CAG TAa ccG GAA CTG AAC CAG
V329I	F	CTG GTT CAG TTC Cat tTA CTG GTT CCA CCA CGA GTC
	R	GAC TCG TGG TGG AAC CAG TAa atG GAA CTG AAC CAG
V329M	F	CTG GTT CAG TTC Cat gTA CTG GTT CCA CCA CGA GTC
	R	GAC TCG TGG TGG AAC CAG TAc atG GAA CTG AAC CAG
F332A	F	CAG TTC CGT CTA CTG Ggc aCA CCA CGA GTC CGG C
	R	GCC GGA CTC GTG tgc CCA GTA GAC GGA ACT G

^aLowercase letters represent the introduced mutant codon. ^bF, forward primer; R, reverse primer.

generates an alternative sesquiterpene as the predominant product.

MATERIALS AND METHODS

Mutagenesis and Kinetic Assays. A total of 23 new single-site mutants of EIZS were prepared by polymerase chain reaction (PCR) mutagenesis. Mutations were introduced into the wild-type EIZS plasmid using forward and reverse primers for each mutant as listed in Table 1. Two optimal conditions for PCR amplification of the insert were applied: (1) 100 ng of each forward and reverse primer, 3 μ L of 10 mM dNTP mix, 100 ng of template plasmid, 5 μ L of Pfu turbo polymerase

buffer, and 1 unit of Pfu turbo polymerase diluted with water to a final volume of 50 μ L and (2) 150 ng of each forward and reverse primer, 3 μ L of 10 mM dNTP mix, 150 ng of template plasmid, 5 μ L of 10 \times Pfu Ultra High-Fidelity DNA polymerase buffer, and 1 unit of Pfu Ultra High-Fidelity DNA polymerase diluted with water to a final volume of 50 μ L. Optimal PCR conditions required initial denaturation of the reaction mixture at 95 °C for 5 min, addition of polymerase followed by 30 cycles (denaturation for 1 min at 95 °C, annealing for 1 min at primer T_m – 5 or 10 °C, and extension for 7 min at 72 °C), and a final 10 min extension at 72 °C followed by a final hold at 4 °C. One microliter of DpnI was added to the PCR mixture and

Table 2. X-ray Crystal Structure Data Collection and Refinement Statistics

	unliganded wild type	L72V-Mg ²⁺ -PP _i -BTAC complex	F95M-Mg ²⁺ -PP _i -BTAC complex	F95H-Mg ²⁺ -PP _i -BTAC complex	A236G-Mg ²⁺ -PP _i -BTAC complex	W325F-Mg ²⁺ -PP _i complex	V329A-Mg ²⁺ -PP _i -BTAC complex
Data Collection							
wavelength (Å)	1.075	1.008	1.542	1.075	0.9795	0.9795	1.008
resolution (Å)	50–2.40	50–2.10	50–2.45	50–2.10	50–1.76	50–2.46	50–1.95
no. of unique reflections	13825	22011	13884	23067	37050	13755	26839
completeness (%) ^a	95.8 (94.5)	99.9 (99.2)	99.9 (100.0)	100.0 (100.0)	99.2 (98.2)	99.9 (99.9)	97.3 (97.4)
redundancy	3.1 (2.9)	3.6 (3.5)	4.4 (4.2)	5.4 (5.4)	3.3 (3.1)	3.6 (3.4)	4.2 (3.7)
I/σ^a	11.9 (2.4)	12.2 (3.5)	10.9 (3.5)	11.9 (4.9)	13.7 (2.7)	7.6 (3.0)	13.0 (3.0)
$R_{\text{merge}}^{a,b}$	0.091 (0.335)	0.104 (0.301)	0.119 (0.400)	0.136 (0.323)	0.071 (0.339)	0.148 (0.427)	0.116 (0.432)
Refinement							
$R_{\text{cryst}}/R_{\text{free}}^c$	0.191/0.250	0.173/0.206	0.185/0.224	0.160/0.204	0.164/0.198	0.182/0.231	0.152/0.201
rmsd							
bonds (Å)	0.004	0.002	0.002	0.003	0.007	0.002	0.007
angles (deg)	0.8	0.7	0.7	0.7	1.1	0.6	1.0
dihedral angles (deg)	17	13	15	14	15	12	17
no. of protein atoms	2577	2789	2778	2795	2804	2770	2812
no. of solvent atoms	53	241	80	268	283	47	376
no. of Mg ²⁺ cations		3	3	3	3	3	3
no. of PP _i anions		1	1	1	1	1	1
no. of BTAC cations		1	1	1	1		1
no. of glycerols			1	4			1
no. of SO ₄ ²⁻ anions	2	1	1		1	1	2
no. of HPO ₄ ²⁻ anions				1			
Ramachandran plot (%) ^d							
most favored	91.0	94.7	94.7	94.7	95.7	94.7	94.0
additional allowed	8.6	5.3	5.3	5.3	4.3	5.3	6.0
generously allowed	0.4						
Protein Data Bank entry	4LTV	4LXW	4LTZ	4LZ3	4LZO	4LZC	4LUU

^aValues in parentheses refer to data for the highest-resolution shell. ^b $R_{\text{merge}} = \sum |I_h - \langle I_h \rangle| / \sum I_h$, where $\langle I_h \rangle$ is the average intensity calculated from replicate reflections. ^c $R_{\text{cryst}} = \sum ||F_o|| - ||F_c|| / \sum |F_o|$ for reflections contained in the test set held aside during refinement. $|F_o|$ and $|F_c|$ are the observed and calculated structure factor amplitudes, respectively. ^dCalculated with PROCHECK version 3.4.4.

incubated at 37 °C for 1.5 h to digest the template. PCR products were transformed into XL1-Blue cells for DNA isolation and sequencing. DNA was purified (Qiagen mini-prep kit) from cultures from single colonies, and DNA sequencing (DNA Sequencing Facility, University of Pennsylvania) confirmed incorporation of the mutations.

Mutant proteins were expressed and purified using procedures identical to those previously reported.¹³ Steady-state kinetic parameters for the generation of hydrocarbon products by EIZS mutants were measured as previously described.¹³ Briefly, enzymes were solubilized in 50 mM piperazine-*N,N'*-bis(2-ethanesulfonic acid) (PIPES) (pH 6.5), 20% glycerol, 100 mM NaCl, 10 mM MgCl₂, and 5 mM β-mercaptoethanol (BME). Each series of assays was performed two or three times using 0.025–50 μM [¹H]FPP (100 mCi/mmol). The optimal concentration of each EIZS mutant was

determined to be that at which the dependence of product formation on enzyme concentration was linear and the level of substrate turnover was <10%. A 1 mL reaction mixture in a test tube was overlaid with 1 mL of hexane immediately after addition of substrate, covered with aluminum foil, and incubated for 12–15 min at 30 °C. The reaction was quenched by the addition of 75 μL of 500 mM EDTA (pH 8.0), and the mixture was vortexed for 20 s. The hexane extract was passed through a silica gel column (1 cm of silica gel in a Pasteur pipet) directly into a scintillation vial containing 5 mL of scintillation fluid. The aqueous phase was extracted with an additional 2 × 1 mL of hexane and passed through the same silica gel column. Finally, the column was washed with an additional 1 mL of diethyl ether. A Beckman scintillation counter was used to measure total product formation, and the substrate concentration versus rate of product formation data

were fit by nonlinear regression using Prism to determine steady-state kinetic parameters, k_{cat} and K_M , based on the known total enzyme concentration.

Analysis of Sesquiterpene Product Arrays. FPP (80 μM) was incubated with 0.8 μM EIZS wild type or each mutant in 6 mL of buffer [50 mM PIPES (pH 6.5), 15 mM MgCl₂, 100 mM NaCl, 20% glycerol, and 5 mM BME] and overlaid with 3 mL of HPLC grade *n*-pentane in a glass test tube at 30 °C for 18 h. The reaction mixture was extracted three times with *n*-pentane; pentane extracts were dried with anhydrous MgSO₄, filtered, and concentrated on an ice/water mixture under reduced pressure until the volume reached 100 μL . Extracts were analyzed using an Agilent 5890 Series II mass spectrometer with a 30 m × 0.25 mm HPMSMS capillary column in EI positive mode using a temperature program with an initial hold at 60 °C for 2 min, a 20 °C/min temperature gradient from 60 to 280 °C, a final hold at 280 °C for 2 min, and a solvent delay of 3 min. Analysis of the organic extracts resulting from the incubation of FPP with the mutant cyclases by gas chromatography and mass spectrometry (GC–MS) revealed the formation of mixtures of sesquiterpene hydrocarbons at *m/z* 204. Compounds were identified by comparison of their individual mass spectra and chromatographic retention indices with those of authentic compounds using the MassFinder 4.0 Program and Database.¹⁹

X-ray Crystal Structure Determinations. The L72V, A236G, W325F, and V329A EIZS mutants were crystallized by the hanging drop vapor diffusion method using the same conditions employed for crystallization of the wild-type enzyme.¹³ Briefly, a 4 μL drop of a protein solution [8 mg/mL EIZS mutant, 20 mM Tris-HCl (pH 7.5), 300 mM NaCl, 10 mM MgCl₂, 10% glycerol, 1 mM TCEP, 2 mM sodium pyrophosphate, and 2 mM BTAC] was added to a 4 μL drop of precipitant solution [100 mM Bis-Tris (pH 5.5), 25–28% polyethylene glycol 3350, and 0.2 M (NH₄)₂SO₄] and equilibrated against a 1 mL reservoir of precipitant solution at room temperature. Crystals of each mutant were isomorphous with those of the wild-type enzyme (space group P2₁, $a = 53.2 \text{ \AA}$, $b = 47.4 \text{ \AA}$, $c = 75.4 \text{ \AA}$, $\beta = 95.5^\circ$)¹³ and diffracted to 1.76–2.30 \AA resolution at the National Synchrotron Light Source (NSLS), beamline X-29, or the Advanced Photon Source (APS), NE-CAT beamline 24-ID-C. Data collection statistics are listed in Table 2.

Crystals of unliganded wild-type EIZS were prepared by soaking crystals of the wild-type EIZS–Mg²⁺–BTAC complex¹³ in chelation buffer [5 mM 1,4,7,10-tetraazacyclododecane-1,4,7,10-tetraacetic acid, 15 mM tetrasodium ethylenediaminetetraacetic acid, 100 mM Bis-Tris (pH 5.5), 30% polyethylene glycol 3350, 200 mM (NH₄)₂SO₄, and 15% glycerol] for 7 days. Crystals diffracted X-rays to 2.4 \AA resolution at NSLS beamline X-29 and were isomorphous with those of the liganded enzyme.¹³ Data collection statistics are listed in Table 2.

F95M EIZS was crystallized by the sitting drop vapor diffusion method. A 0.6 μL drop of protein solution [15 mg/mL F95M EIZS, 20 mM Tris-HCl (pH 7.5), 300 mM NaCl, 10 mM MgCl₂, 10% glycerol, 2 mM TCEP, 2 mM sodium pyrophosphate, and 2 mM BTAC] was added to 0.6 μL of precipitant solution [100 mM Bis-Tris (pH 5.5), 200 mM (NH₄)₂SO₄, and 25% polyethylene glycol 3350] and equilibrated against a 110 μL well reservoir of precipitant solution. Crystals appeared within 1 week. X-ray diffraction data (2.45 \AA resolution) were collected on a Rigaku IV⁺⁺ image plate

area detector mounted on a Rigaku RU200HB rotating anode X-ray generator (8.048 keV and 1.5406 \AA); crystals were isomorphous with those of the wild-type enzyme.¹³ Data collection statistics are listed in Table 2.

F95H EIZS was crystallized by the sitting drop vapor diffusion method. A 0.6 μL drop of protein solution [10 mg/mL F95H EIZS, 20 mM Tris-HCl (pH 7.5), 300 mM NaCl, 10 mM MgCl₂, 10% glycerol, 2 mM TCEP, 2 mM sodium pyrophosphate, and 2 mM BTAC] was added to 0.6 μL of precipitant solution (40 mM KH₂PO₄, 16% polyethylene glycol 8000, and 20% glycerol) and equilibrated against a 110 μL well reservoir of precipitant solution. Crystals appeared in 1 week and diffracted X-rays to 2.1 \AA resolution at NSLS beamline X-29. Crystals belonged to space group P2₁2₁2₁ with the following unit cell parameters: $a = 46.7 \text{ \AA}$, $b = 75.6 \text{ \AA}$, and $c = 107.6 \text{ \AA}$ (one monomer in the asymmetric unit).

Crystal structures were determined by molecular replacement using PHASER,^{20,21} the atomic coordinates of wild-type EIZS less ligand and solvent molecules were used as a search probe for rotation and translation function calculations. Initial electron density maps of each mutant clearly revealed the substituted amino acid side chain, which was modeled into density and refined. The initial electron density map of unliganded EIZS revealed an empty active site. Iterative cycles of refinement and manual model building were performed with PHENIX and COOT, respectively.^{22,23} Ligands and water molecules were included in later cycles of refinement. Individual atomic B factors were utilized in refinement. Data collection and refinement statistics for each structure are listed in Table 2. The rmsds of C α atoms between wild-type and mutant EIZS complexes were calculated with COOT.

RESULTS

Catalytic Activity of EIZS Mutants. Steady-state kinetic parameters for all mutants reflect the generation of ³H-labeled hexane-extractable products and are reported in Table 3; data for the F96A, F198A, and W203F mutants, which were previously reported by Aaron and colleagues,¹³ are included for comparison. In general, K_M values for mutants of aromatic and aliphatic active site residues are within 1 order of magnitude of those measured for wild-type EIZS. The most significant difference in K_M is observed for V329M EIZS, which exhibits a 20-fold higher K_M value compared with that of the wild-type enzyme. More significant changes are observed for the k_{cat} values of various mutants; overall, changes in k_{cat} tend to be greater for replacement of aromatic residues than for substitution of aliphatic residues. In particular, substitution of F95, F96, and F198 with smaller aliphatic side chains results in 11–730-fold decreases in k_{cat} . This aromatic triad encircles the active site (Figure 1) and is proposed to stabilize carbocationic transition states and intermediates through cation–π interactions.¹³

Two tryptophan residues, W203 and W325, although only marginally exposed to the central active site cavity compared to F95 and F96, were substituted with phenylalanine to probe the importance of their aromatic indole side chains. Whereas W203F EIZS exhibits a 250-fold reduction in k_{cat} compared to that of wild-type EIZS, W325F EIZS shows only a modest 4-fold reduction in k_{cat} . Thus, W203 has a stronger influence on catalysis, probably because W203 contributes more surface area to the active site contour than W325 (Figure 1). The F332A mutant exhibits an 800-fold reduction in k_{cat} , consistent with

Table 3. Kinetic Parameters for Wild-Type and Mutant EIZS Enzymes

enzyme	k_{cat} (ms ⁻¹)	K_M (nM)	k_{cat}/K_M (M ⁻¹ s ⁻¹)
WT	84 ± 1	330 ± 20	260000 ± 10000
Aromatic Mutants			
F95A	1.8 ± 0.2	672 ± 2	2700 ± 300
F95V	7.4 ± 0.4	1700 ± 300	4400 ± 600
F95M	3.16 ± 0.09	275 ± 4	11500 ± 500
F95H	1.72 ± 0.02	650 ± 30	2600 ± 100
F96A ^a	0.24 ± 0.02	800 ± 100	310 ± 60
F96V	1.15 ± 0.02	580 ± 30	199 ± 7
F96L	2.57 ± 0.02	820 ± 60	3100 ± 200
F96Y	0.19 ± 0.01	1800 ± 100	107 ± 2
F96W	0.35 ± 0.02	310 ± 70	1200 ± 400
F198A ^a	0.30 ± 0.02	1200 ± 200	250 ± 50
F198V	1.66 ± 0.09	2800 ± 400	600 ± 60
F198L	4.75 ± 0.05	4200 ± 400	1100 ± 100
F198Y	15.04 ± 0.06	4100 ± 100	3600 ± 100
W203F ^a	0.34 ± 0.03	1500 ± 200	230 ± 40
W325F	22 ± 2	160 ± 60	14000 ± 5000
F332A	0.105 ± 0.008	360 ± 90	290 ± 80
Aliphatic Mutants			
L72A	8.81 ± 0.09	3000 ± 100	2900 ± 100
L72V	44 ± 3	600 ± 100	70000 ± 20000
L72M	11 ± 2	3200 ± 900	3400 ± 400
A236G	66 ± 2	450 ± 50	150000 ± 20000
A236M	—	—	—
A236F	0.0424 ± 0.0007	230 ± 50	190 ± 40
V329A	126 ± 7	470 ± 80	270000 ± 50000
V329G	19.8 ± 0.1	1590 ± 60	12500 ± 500
V329I	22.8 ± 0.3	770 ± 40	29000 ± 1000
V329M	0.640 ± 0.004	6500 ± 300	98 ± 3

^aMutants previously reported (ref 13).

the fact that the edge of this aromatic residue also defines part of the active site contour.

Substitution of aliphatic residues L72, V329, and A236 with either larger or smaller aliphatic side chains appears to be better tolerated, with k_{cat} values comparable to or within 1 order of magnitude of that measured for the wild-type enzyme ($k_{\text{cat}} = 0.084 \text{ s}^{-1}$), except for V329M EIZS, for which k_{cat} is reduced more than 100-fold to $6.4 \times 10^{-4} \text{ s}^{-1}$. Interestingly, A236M EIZS yields no detectable products, while A236F EIZS exhibits a nearly 2000-fold reduction in k_{cat} , suggesting that a small side chain at residue 236 is mandatory for efficient catalysis. Accordingly, A236G EIZS exhibits nearly the same steady-state kinetic parameters as wild-type EIZS.

Sesquiterpene Product Arrays Generated by EIZS Mutants. Sesquiterpene hydrocarbon mixtures generated by each of the EIZS mutants were subjected to analysis and quantification by gas chromatography–mass spectrometry (GC–MS) (Table 4). Sesquiterpene products identified in these mixtures are shown in Figure 2. In general, replacement of F95, F96, or F198 in the aromatic triad compromises the generation of *epi*-isozaene (18), generally more so for F96 and F198, which each contribute more solvent-accessible surface area to the active site contour than does F95 (Figure 1). It appears that the fidelity of *epi*-isozaene formation is most sensitive to substitution of F96, particularly with smaller aliphatic residues; the F96A, F96V, and F96L mutants generate <10% *epi*-isozaene. Only F96Y EIZS, representing a nearly isosteric substitution of the aromatic ring of residue 96, retains

the fidelity of *epi*-isozaene generation, in spite of a substantial 2400-fold reduction in catalytic efficiency (k_{cat}/K_M). It is nonetheless notable that the product distribution can be significantly reprogrammed by only a single amino acid substitution for each member of the aromatic triad, often with retention of significant catalytic activity. For example, the F95M mutation converts EIZS into a predominant β -acoradiene synthase, generating a mixture consisting of 68% β -acoradiene (13) and only 32% *epi*-isozaene (18), with only a 23-fold reduction in catalytic efficiency. F95M EIZS is thus directed to make an entirely different predominant cyclization product with relatively good catalytic efficiency. Analysis of the biosynthetic manifold (Figure 3) suggests that the observed change in product distribution is a consequence of diversion of the normal cyclization of the homobisabolyl cation intermediate.

By contrast, replacement of the phenyl ring with an imidazole group in F95H EIZS results in the generation of a mixture of 50% β -curcumene (6) and 44% *epi*-isozaene (18), accompanied by only 2% β -acoradiene (13), along with an additional 4% of two products of unassigned structure. Although the overall catalytic efficiency is compromised 100-fold in this mutant, the rigidity and perhaps the polarity of the substituted imidazole side chain apparently favor early termination of the cyclization cascade by direct deprotonation of the homobisabolyl cation. This suggests that F95 plays a key role in stabilizing and directing the chemistry of this monocyclic intermediate (Figure 3).

Similarly, substitution of F198 with the progressively smaller aliphatic residues leucine, valine, and alanine reduces the proportion of *epi*-isozaene to 24, 10, and 0% of the total sesquiterpene products, respectively. Notably, F198L EIZS generates instead 61% β -cedrene (12), of which 80% is (−)- β -cedrene and 20% is (+)- β -cedrene based on GC–MS comparison with authentic standards; F198L EIZS also generates 24% *epi*-isozaene (18), 14% zizaene (17), and 1% sesquisabinene-A (7). While the catalytic efficiency of this mutant is 236-fold lower than that of wild-type EIZS, it is notable that a single amino acid substitution is sufficient to convert EIZS predominantly into a β -cedrene synthase. Importantly, the generation of (+)- β -cedrene indicates that synthetic sequences proceeding through the alternative (7R)-homobisabolyl enantiomer, rather than the normal (7S)-enantiomer leading to (−)- β -cedrene and *epi*-isozaene formation, are feasible. The absolute configurations of the remaining aberrant sesquiterpene cyclization products shown in Figure 3 have not been determined and are arbitrarily illustrated.

Interestingly, F96V EIZS and W203F EIZS generate 44 and 47% (Z)- γ -bisabolene (4), respectively, as part of more complex product mixtures containing significantly reduced proportions of *epi*-isozaene (6 and 14%, respectively). Early termination of the cyclization sequence by elimination of a proton from the (4R)-bisabolyl cation intermediate to yield 4 suggests that the steric bulk of each of these residues, as well as presumed cation–π interactions with F96, plays a key role in directing the biosynthetic fate of the (4R)-bisabolyl cation.

Finally, the F96W substitution generates a mixture of 65% zizaene (17), 9% (3Z,6E)- α -farnesene (2), and only 3% *epi*-isozaene (18), along with 23% of three unidentified sesquiterpene hydrocarbon products. An increase in steric bulk at residue 96 thus redirects the cyclization of the (7S)-homobisabolyl cation to favor attack on the opposite face of the

Table 4. Sesquiterpene Product Arrays Generated by EIZS Mutants

enzyme	relative average percentage of sesquiterpene products (product numbering shown in Figure 2)																		
	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18	N/A ^a
WT ^b	5						2	1		2							9	79	
WT ^c		1	1														5	93	
WT ^d																	99	1	
L72A							6			6						13	67	8 (2)	
L72V							3			2						14	43	38	
L72M										7		2				10	80		
F95A			4			3					35						1	56 (3)	
F95V											26	2					66	6 (2)	
F95M											68						32		
F95H					50						2						44	4 (2)	
F96A ^b	70		7			9											8	7 (1)	
F96V		44									14	5					6	31 (1)	
F96L		2		4	10	8		3		16			2	30	14	9	2 (1)		
F96Y										3		1		18	78				
F96W	9														65	3	23 (3)		
F198A ^b	5		24	6		20	13				12							20 (3)	
F198V			3			3		2		48	6		3	4	13	10	8 (3)		
F198L						1				61					14	24			
F198Y																	89	5 (2)	
W203F ^b	6		47			7										7	14	21 (4)	
A236G																2	70	28 (4)	
A236M	—	—	—	—	—	—	—	—	—	—	—	—	—	—	—	—	—	—	
A236F											8					15	77		
W325F																45	50	5 (3)	
V329A						4										3	85	8 (1)	
V329G		2		8	6	2					7						75		
V329I																2	98		
V329M																33	62	5 (1)	
F332A						1										37	56	6 (3)	

^aN/A, not assigned; value in parentheses indicates the number of unknown products detected by GC–MS. ^bProduct array reported in ref 13. ^cAssay conducted at 20 °C. ^dAssay conducted at 4 °C.

side chain trisubstituted double bond. By contrast, as previously reported,¹³ F96A EIZS, bearing a much smaller, nonaromatic methyl side chain, generates a mixture containing 70% of the acyclic sesquiterpene (*E*)- β -farnesene (**1**); replacement of this key aromatic side chain with a methyl group thus results in the quenching of the initially formed farnesyl cation. While catalytic efficiency is reduced >800-fold in this mutant relative to that of wild-type EIZS, the catalytic efficiency for enzymatic formation of (*E*)- β -farnesene by F96A EIZS ($k_{\text{cat}}/K_M = 310 \text{ M}^{-1} \text{ s}^{-1}$) is nonetheless only ~14-fold lower than that reported for the wild-type wormwood (*E*)- β -farnesene synthase ($k_{\text{cat}}/K_M = 4500 \text{ M}^{-1} \text{ s}^{-1}$).²⁴

Substitutions for the aliphatic residues L72, A236, and V329 mostly result in less dramatic changes in the sesquiterpene product spectrum compared to the changes seen upon replacement of aromatic residues. All such mutants generate mixtures containing at least 60% *epi*-isoizaene, with the exception of L72V EIZS, which generates a mixture of 43% *epi*-isoizaene (**18**), 14% zizaene (**17**), 3% sesquisabinene-A (**7**), 2% β -cedrene (**12**), and 38% of an as yet unidentified sesquiterpene hydrocarbon product. The majority of these aliphatic mutants generate zizaene in addition to *epi*-isoizaene, representing the alternative cyclization stereochemistry for the common (7*S*)-homobisabolyl cation intermediate (Figure 3). Because the side chains of these aliphatic amino acids make smaller contributions to the active site contour of wild-type EIZS, the effects of their replacement may be limited. The

overall catalytic activity and the cyclization fidelity both strongly depend on the presence of a small residue at position 236, which is normally buried just beneath the solvent-accessible surface of the active site contour. Thus, A236G EIZS generates a mixture of 70% *epi*-isoizaene (**18**), 2% zizaene (**17**), and 28% of a mixture of four unidentified sesquiterpenes with a near-normal catalytic rate, whereas the corresponding A236F EIZS mutant generates a mixture of 77% *epi*-isoizaene (**18**), 15% zizaene (**17**), and 8% β -cedrene (**12**) with a nearly 2000-fold reduction in k_{cat} . By contrast, A236M EIZS is completely devoid of activity.

Crystal Structures of Unliganded EIZS and Selected Mutants.

X-ray crystal structures of unliganded EIZS and selected mutants were determined at resolutions of 1.76–2.46 Å. The structure of unliganded EIZS, the first to be determined of the unliganded wild-type enzyme, reveals an open active site conformation that is essentially identical to that previously observed in the corresponding unliganded D99N mutant,¹³ with an rmsd of 0.51 Å for 308 C α atoms between the two structures. Interestingly, active site aromatic residue F96 adopts a different conformation in the unliganded enzymes compared with the structures of all other wild-type or mutant EIZS enzymes complexed with three Mg $^{2+}$ ions and PP $_i$, regardless of whether the carbocation mimic, benzyltriethylammonium cation (BTAC), is also bound.¹³ The change in the conformation of F96 triggered by the binding of three Mg $^{2+}$ ions and PP $_i$ and consequent full active site closure corresponds

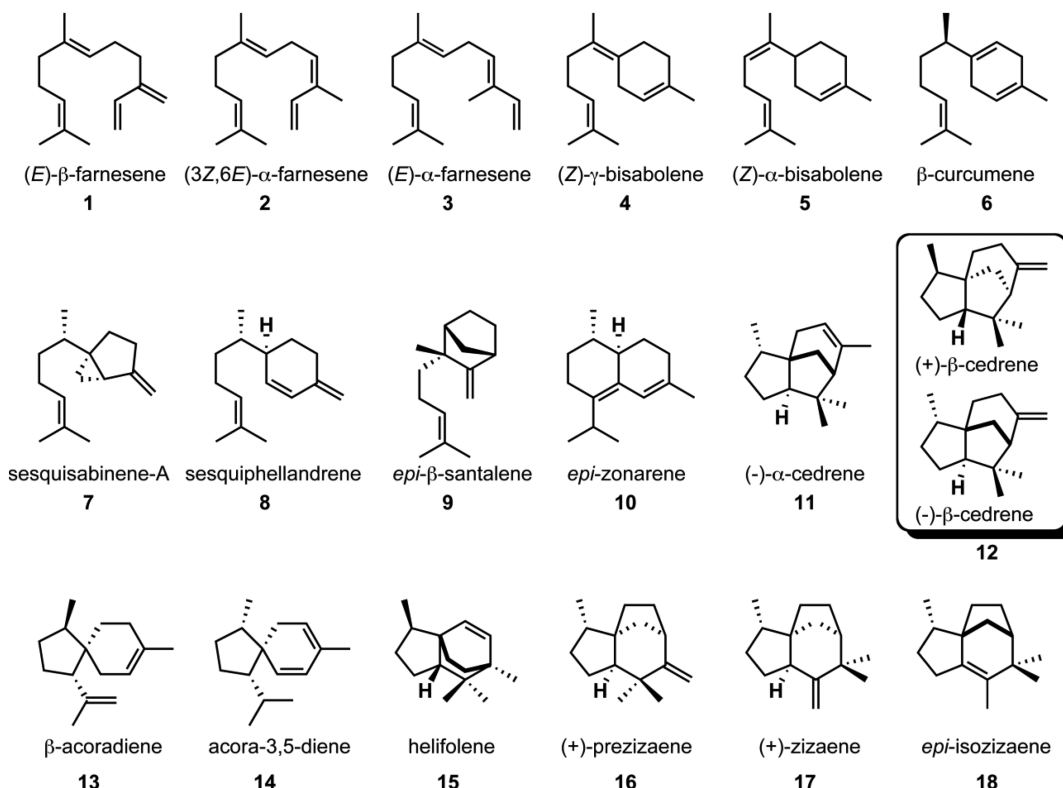


Figure 2. Sesquiterpene products generated by EIZS mutants as detected by GC–MS analysis and identified by comparison of their individual mass spectra and chromatographic retention indices with those of authentic compounds in the MassFinder 4.0 Database.

to rotations of $\sim 105^\circ$ and $\sim 10^\circ$ about side chain torsion angles χ_1 and χ_2 , respectively (Figure 4). Such induced-fit conformational changes in the active site contour of a terpenoid cyclase are unusual, but not unprecedented. For example, F578 of bornyl diphosphate synthase undergoes a significant conformational change upon ligand binding.²⁵ Comparisons of high-resolution unliganded and liganded contours of other terpenoid cyclases, however, generally reveal conformational changes that reflect rigid-body movements rather than significant modification of side chain conformations.

X-ray crystal structures of $Mg^{2+}_3\text{--PP}_i\text{--BTAC}$ complexes were determined for the L72V, F95M, F95H, A236G, and V329A mutants of EIZS, while the structure of the complex with three Mg^{2+} ions and PP_i was determined for the W325F mutant. Each of these mutants was selected because it exhibits significant residual catalytic activity [indeed, V329A exhibits enhanced catalytic activity (see Table 3)] and/or interesting differences in the array of resulting sesquiterpene products compared to the wild-type enzyme (Table 4). All of these mutants, except for F95H EIZS, were readily crystallized under the same conditions as the wild-type protein. In general, each single-amino acid substitution causes minimal structural perturbations, with the following rmsds from the structure of the wild-type EIZS– $Mg^{2+}_3\text{--PP}_i$ complex: L72V, 0.13 Å (340 C α atoms); F95M, 0.17 Å (340 C α atoms); F95H, 0.22 Å (340 C α atoms); A236G, 0.16 Å (340 C α atoms); V329A, 0.08 Å (340 C α atoms); W325F, 0.18 Å (338 C α atoms). The binding mode of the $Mg^{2+}_3\text{--PP}_i$ cluster is generally unchanged in each mutant, except that the PP_i anion rotates slightly in F95M EIZS such that the bridging oxygen atom moves ~ 0.6 Å, and the Mg^{2+}_C ion moves $\sim 1.0\text{--}1.5$ Å in the F95M and W325F mutants (Figure 5). Thus, site-specific mutations can be made in the hydrophobic region of the active site without significantly

perturbing neighboring residues, including those that play an important role in triggering substrate ionization.

The position and orientation of BTAC are unchanged in A236G EIZS and V329A EIZS, which still generate *epi*-isozizaene as the major sesquiterpene cyclization product [70 and 85%, respectively (see Table 4)]. However, BTAC rotates by $\sim 20^\circ$ and moves ~ 0.5 Å deeper into the active site of L72V EIZS as a consequence of the smaller side chain at this position (Figure 5). The proportion of *epi*-isozizaene generated is reduced to 43% in this mutant. Interestingly, BTAC appears to be absent from the active site of W325F EIZS, although some uninterpretable electron density peaks are observed that could correspond to disordered BTAC; the level of formation of *epi*-isozizaene generated by this mutant is reduced to 50%. It thus appears that the movement or disorder of BTAC in crystalline complexes with EIZS mutants correlates with a higher potential for alternative cyclization sequences. Possibly, more authentic mimics of carbocation intermediates in the EIZS-catalyzed cyclization mechanism, such as (4*R*)-azabisabolene,²⁶ which mimics the (4*R*)-bisabolyl cation intermediate, would provide an even more sensitive indicator of the potential for redirected cyclization cascades in EIZS mutants.

■ DISCUSSION

Active Site Mutations Dramatically Enhance Product Chemodiversity. We have demonstrated that the hydrophobic active site contour of EIZS can be remolded with single amino acid substitutions to adopt a predictable, alternative shape based on analysis of three-dimensional structures of wild-type and mutant enzymes. Furthermore, in a library of 26 site-specific mutants, we have tripled the overall number of major and minor characterized products, and we have converted EIZS

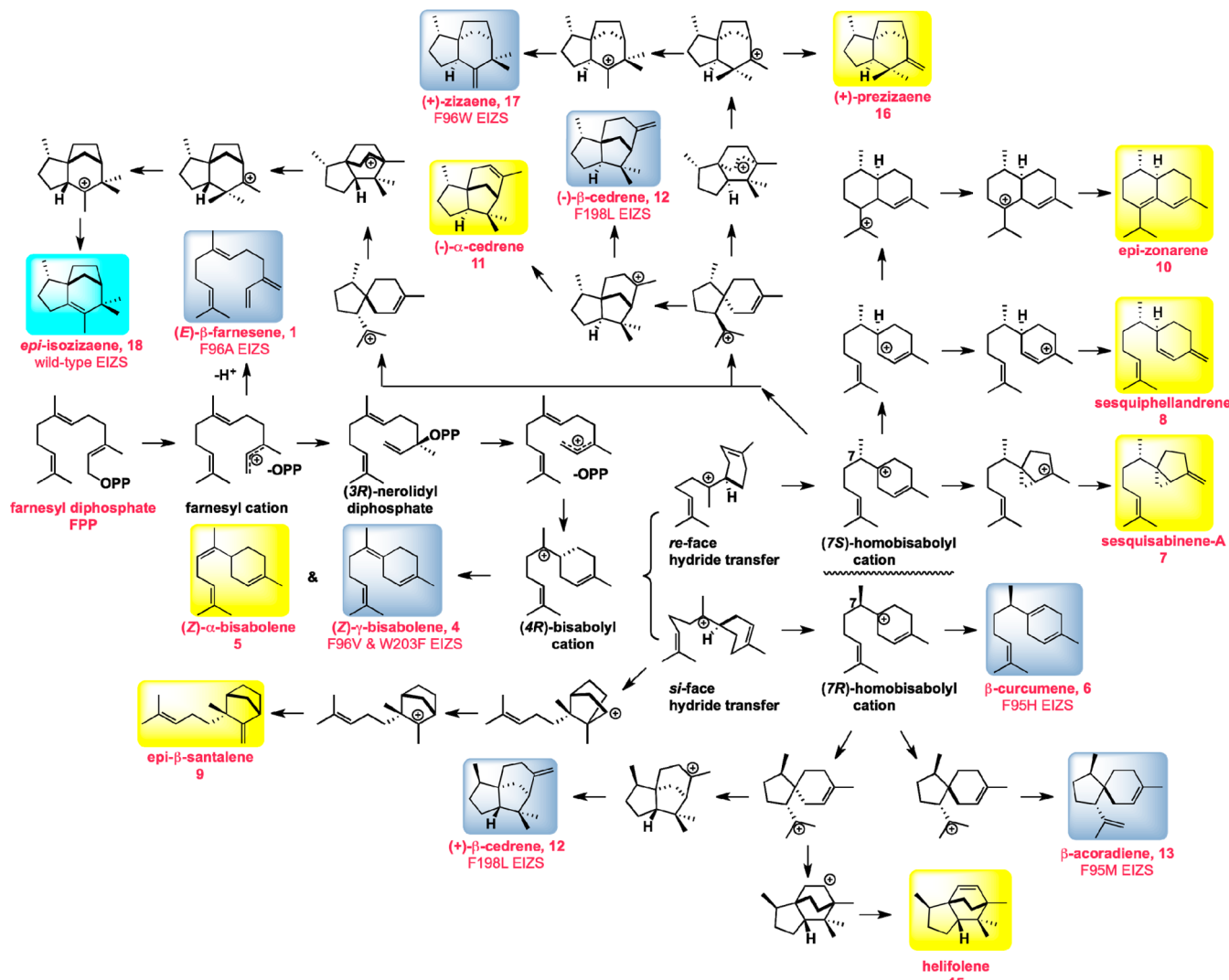


Figure 3. Biosynthetic manifold illustrating the proposed mechanism of FPP cyclization reactions catalyzed by EIZS and its site-specific mutants. The product numbering is identical to that in Figure 2. The predominant products of wild-type EIZS and certain EIZS mutants are highlighted in cyan and blue boxes, respectively, and side products are highlighted in yellow boxes; product percentages are listed in Table 4. For the sake of clarity, farnesene isomers 2 and 3 generated at low levels (1%) by wild-type EIZS are not shown, nor is acora-3,5-diene (14) generated at low levels (2%) by F95V EIZS. With regard to product stereochemistry, it should be noted that the determination of the absolute configuration of the (+)-epi-isozizaene produced by EIZS has been previously reported.¹² Although the absolute configurations of (+)-prezizaene (16) and (+)-zizaene (17) produced by EIZS mutants have not been directly determined,¹⁴ they can be provisionally assigned from the natural cyclization mechanism, because they represent simple late-stage diversions of the pathway. By contrast, generation of both enantiomers of β-cedrene (12) has been directly determined for F198L EIZS by direct chiral GC–MS comparison with authentic standards, as described in the text. The formation of both (−)- and (+)-12 requires that EIZS or its mutants be capable of generating both the (7S)- and (7R)-homobisabolyl cations, as shown. The absolute configurations of the remaining aberrant sesquiterpene cyclization products have not been determined and are arbitrarily illustrated for (Z)-α-bisabolene (5), β-curcumene (6), sesquisabinene A (7), sesquiphellandrene (8), epi-β-santalene (9), epi-zonarene (10), (−)-α-cedrene (11), β-acoradiene (13), and helifolene (15). Formation of the enantiomers of any of these compounds can readily be accommodated by simple variations of the mechanistic scheme.

into six different sesquiterpene synthases that each generate an alternative predominant cyclization product or, in one case, an elimination product. Thus, F96W EIZS is a zizaene synthase, F95H EIZS a β-curcumene synthase, F95M EIZS a β-acoradiene synthase, F198L EIZS a β-cedrene synthase, and F96V EIZS and W203F EIZS are (Z)-γ-bisabolene synthases. F96A EIZS is a simple (E)-β-farnesene synthase (Table 4). New cyclization products include simple abortive derailment products resulting from direct quenching of the (R)-bisabolyl cation to yield (Z)-α-bisabolene and (Z)-γ-bisabolene, while other new cyclization products are, in effect, shunt metabolites

resulting from diversion of the natural cyclization cascade and subsequent quenching of the derived carbocations, such as the alternative reaction sequences leading to the generation of epi-zonarene or β-acoradiene.

It is also noteworthy that the experimentally observed product array of wild-type EIZS can be reconfigured by single-site mutagenesis to convert a minor product into a major product, and vice versa, in accord with previous studies of terpenoid cyclases. For example, we previously reported that F96A EIZS is an (E)-β-farnesene synthase, because (E)-β-farnesene is the predominant sesquiterpene product (70%); wild-type EIZS generates only 5% (E)-β-farnesene.¹³ By

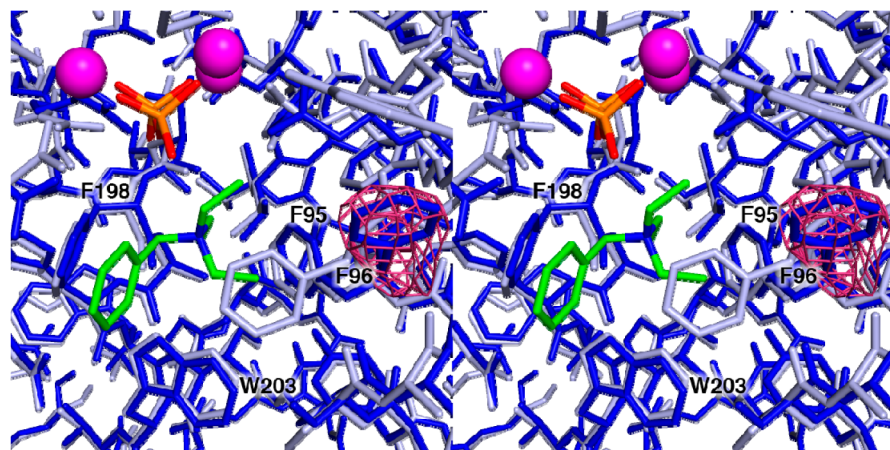


Figure 4. Omit electron density map contoured at 2.2σ showing the conformation of F96 in the unliganded enzyme (dark blue), compared with the conformation of this residue in EIZS–ligand complexes as represented by the wild-type EIZS– Mg^{2+} – PP_i –BTAC complex (light blue). Mg^{2+} ions are shown as magenta spheres. P and O atoms of the PP_i anion are colored orange and red, respectively. C and N atoms of the BTAC cation are colored green and blue, respectively. Selected active site residues are labeled. The aromatic side chain of F96 undergoes a significant induced-fit conformational change as the terpenoid cyclase active site undergoes the transition from a fully open to a fully closed conformation.

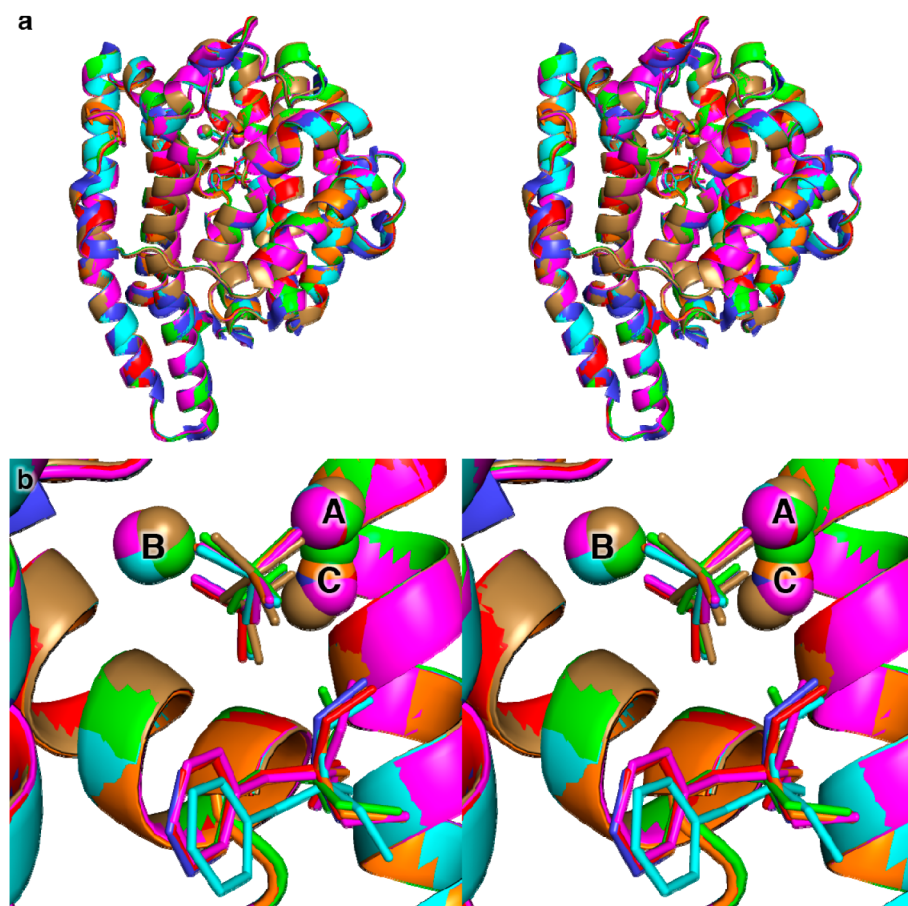


Figure 5. Superposition of wild-type (blue), L72V (cyan), F95M (green), F95H (magenta), A236G (orange), and V329A (red) EIZS complexed with three Mg^{2+} ions, PP_i , and BTAC, and W325F EIZS complexed with three Mg^{2+} ions and PP_i (tan).

contrast, we now show that F96W EIZS is a zizaene synthase, because zizaene is the predominant sesquiterpene product (65%) (Table 4); wild-type EIZS generates only 9% zizaene. In the absence of an experimentally determined crystal structure of γ -humulene synthase, Keasling and colleagues used homology modeling to guide the mutagenesis of a limited number of active site hydrophobic residues, similarly remolding the active

site to reconfigure the product array.¹⁷ Noel and colleagues demonstrated that terpene product ratios can also be reconfigured by mutagenesis of residues remote from the active site, presumably by long-range structural effects that propagate through the protein scaffolding.¹⁸

Aromatic Residues Are Hot Spots for Reprogramming or Reconfiguring the Cyclization Cascade. Whether the

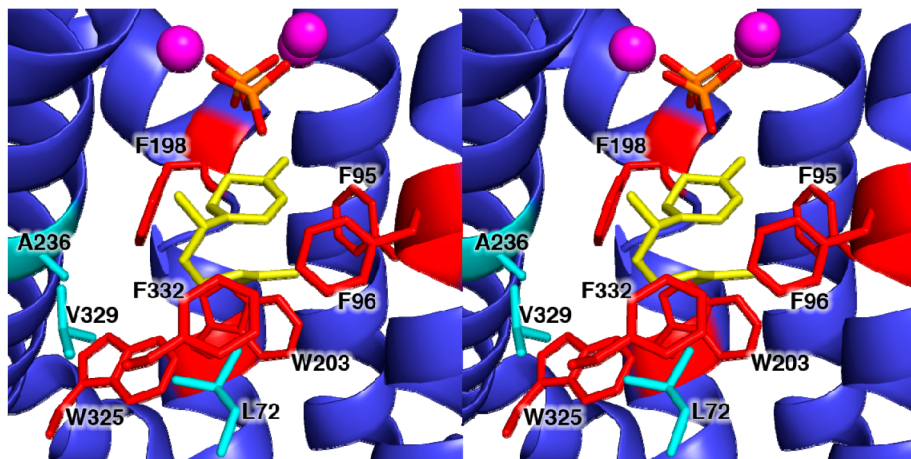


Figure 6. Model of the (7S)-homobisabolyl cation bound in the active site of wild-type EIZS, showing how the L72/W325/V329/F332 cluster could chaperone the precise conformation of the prenyl side chain that would generate the acorenyl cation leading to *epi*-isozaene formation as summarized in Figure 3.

cyclization cascade of EIZS is reprogrammed to generate entirely new products or whether the proportions of major and minor products are reconfigured, the most striking results are obtained when aromatic residues are mutated. Active site aromatic residues can stabilize carbocation intermediates through cation–π interactions, as suggested for the side chains of the aromatic triad of F95, F96, and F198.¹³ Additionally, aromatic residues provide significant steric bulk and hydrophobic surface area as they define the contour of the active site, which is evident in Figure 1.

Interestingly, certain product distributions in Table 4 can be mapped to specific aromatic residues with regard to their influence on a specific carbocation intermediate. For example, consider F96A EIZS. The steric bulk and weakly polar nature of the F96 side chain presumably play a role in stabilizing the farnesyl cation resulting from the initial ionization of FPP, because F96A EIZS generates (E)-β-farnesene as an immediate derailment product. Inspection of Figure 1 reveals that the side chain of F96 is located ~8 Å beneath the plane defined by Mg²⁺_A, Mg²⁺_B, and Mg²⁺_C. Furthermore, the plane formed by Mg²⁺_A, Mg²⁺_C, and F96 is approximately perpendicular to that formed by the three metal ions. We propose that F96 is ideally positioned to interact with the initial farnesyl cation intermediate through both its steric bulk and its capacity for cation–π interactions. The diminished progress along the reaction sequence as the product arrays of F96Y, F96L, F96V, and F96A are compared suggests that steric bulk at position 96 is functionally important, presumably for ensuring the close contact between the C1 and C6 atoms allowing (R)-bisabolyl cation formation.

In another more subtle example, the partitioning of the (7S)-homobisabolyl cation intermediate that leads either to normal formation of *epi*-isozaene or to aberrant formation of zizaene is a function of the folding of the side chain of this intermediate. The conformation of the (7S)-homobisabolyl cation directs the electrophilic attack on one face or the other of the trisubstituted double bond of the side chain, and this conformation is presumably enforced by the three-dimensional contour of the active site. Mutants that exhibit significant increases in the level of zizaene formation cluster in the lower region of the active site where they decrease steric bulk: L72V, W325F, V329M, and F332A. Although the methionine side chain in V329M EIZS is slightly longer than the wild-type valine side chain, this

substitution reduces the steric bulk around the Cβ atom of the parent branched valine side chain. Alternatively, both F96Y EIZS and F96W EIZS also produce significantly elevated proportions of zizaene. Both mutants retain the original aromatic character of the wild-type phenylalanine side chain, and their larger side chains are located on the opposite side of the active site from the L72/W325/V329/F332 cluster. Thus, the partitioning of the (7S)-homobisabolyl cation toward zizaene formation (Figure 3) can be favored either by pulling the intermediate into the void created by smaller amino acid mutations in the L72/W325/V329/F332 cluster or by pushing the intermediate toward this cluster by introducing larger amino acids on the opposite side of the active site. This suggests that the L72/W325/V329/F332 cluster may chaperone the conformation of the prenyl side chain of the (7S)-homobisabolyl cation to generate the acorenyl cation leading to *epi*-isozaene formation. A model of the (7S)-homobisabolyl cation bound in the active site of EIZS shows how the L72/W325/V329/F332 cluster may achieve this chaperone function (Figure 6).

Finally, the side chains of F95 and F198 are located in the rear of the active site as viewed in Figure 1. Substitution of these residues with smaller side chains other than alanine may disfavor the 1,2-hydride shift that generates the natural (7S)-homobisabolyl cation intermediate, diverting the reaction instead to the enantiomeric (7R)-homobisabolyl cation, which in turn serves as a precursor of β-acoradiene and β-cedrene. Indeed, of the 61% β-cedrene generated by F198L EIZS, 20% is (+)-β-cedrene, which is derived from the (7R)-homobisabolyl cation intermediate (Figure 3). As a further variation of this pathway branching, F95H EIZS generates predominantly β-curcumene by direct deprotonation of the homobisabolyl cation. Whether the (7S)- or (7R)-homobisabolyl cation is formed is presumably a consequence of the precise folding of the preceding (4R)-bisabolyl cation enforced by the active site contour. Thus, bulky side chains in the rear of the active site (as viewed in Figure 1) ensure the formation of the natural (7S)-homobisabolyl cation intermediate that ultimately leads to *epi*-isozaene. Smaller side chains in this region may allow the formation of the aberrant (7R)-homobisabolyl cation intermediate, as observed for F198L EIZS.

Product Chemodiversity and Biotechnology Applications. While numerous studies have been published to date

describing mutagenesis of class I terpenoid cyclases and resultant changes in products generated, many of these studies have focused on residues outside of the hydrophobic active site pocket, such as residues coordinating to catalytically obligatory metal ions,^{10,27–29} residues that hydrogen bond with the substrate diphosphate group,^{30,31} or residues thought to function in general acid–general base or electrostatic catalysis.^{10,29,32} Relatively few studies have focused on site-specific mutants specifically prepared to remodel the hydrophobic active site contour of a class I terpenoid cyclase, and fewer still have been explicitly guided or accompanied by experimentally determined crystal structures.^{17,18,33–36} Accordingly, we suggest that our work with EIZS represents the first small step in a journey that will ultimately allow us to correlate the three-dimensional structures of experimentally verified remodeled active site contours with reprogrammed isoprenoid cyclization cascades.

Structure-based reprogramming of terpenoid cyclase activity should have myriad applications in chemistry, biology, and biotechnology. For example, given that plant (*E*)- β -farnesene synthase has been commercialized for the large-scale production of (*E*)- β -farnesene in the development of an aviation biofuel,^{37–39} it is possible that a bacterial or fungal sesquiterpene synthase, such as F96A EIZS¹³ or the Y92A mutant of aristolochene synthase from *Penicillium roqueforti*,⁴⁰ could be similarly adapted for large-scale biofuel generation. The terpenoid cyclase step is typically the bottleneck in microbial biosynthetic engineering experiments,^{41,42} and plant sesquiterpene synthases can be difficult to express in microbial fermentation systems.⁴³ Thus, the availability of a microbial sesquiterpene synthase engineered to generate a desired sesquiterpene product may provide a useful alternative for the microbial production of advanced biofuels.

It is possible, too, that the use of an engineered bacterial terpenoid cyclase could enhance the flux of sesquiterpene generation by virtue of enhanced catalytic activity relative to that of a naturally occurring plant enzyme. For example, bisabolane has recently been identified as a new biosynthetic alternative to D2 diesel fuel, which is utilized in compression ignition engines.⁴³ Bisabolane is generated by hydrogenation of (*Z*)- α -bisabolene, which in turn is generated by the three-domain plant sesquiterpene cyclase, α -bisabolene synthase from *Abies grandis*,⁴⁴ in a microbial expression system (Figure 7). The reported catalytic efficiency (k_{cat}/K_M) of this α -bisabolene synthase is $38 \text{ M}^{-1} \text{ s}^{-1}$. However, bisabolane can also be generated by the hydrogenation of β -curcumene (Figure 7), which is generated by F95H EIZS with a k_{cat}/K_M of $2600 \text{ M}^{-1} \text{ s}^{-1}$ (Table 3). Thus, the generation of the bisabolane precursor β -curcumene by F95H EIZS is 68-fold more efficient than the generation of the bisabolane precursor (*Z*)- α -bisabolene by wild-type α -bisabolene synthase. Because no naturally occurring β -curcumene synthase has been characterized (apart from a grape sesquiterpene cyclase that generates a mixture of sesquiterpene products, including 22% β -curcumene⁴⁵), the reprogrammed catalytic activity of F95H EIZS may allow a more efficient approach for economical biofuel generation.

Concluding Remarks. In closing, we note that the grand challenge in the engineering of a terpenoid cyclase is predictability: can a terpenoid cyclase be reprogrammed in true *a priori* fashion using structure-based design? The reprogrammed cyclization cascades of EIZS mutants result from divergent mechanistic sequences that branch from a handful of key carbocation intermediates, such as the farnesyl

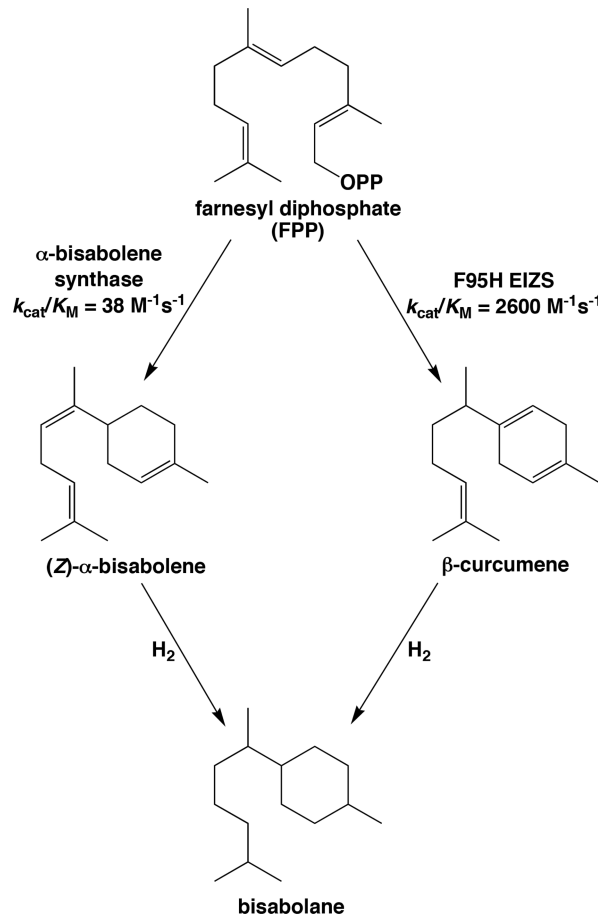


Figure 7. D2 diesel fuel substitute bisabolane can be generated by the hydrogenation of (*Z*)- α -bisabolene or β -curcumene. F95H EIZS exhibits 68-fold enhanced catalytic efficiency in the generation of β -curcumene compared with the generation of (*Z*)- α -bisabolene by α -bisabolene synthase, so it is possible that F95H EIZS could serve as a superior terpenoid synthase for microbial biofuel generation.

cation, the (4*R*)-bisabolyl cation, or the (7*S*)- or (7*R*)-homobisabolyl cation. The next step will be to study how the binding of stable analogues of these common intermediates is affected by active site mutations. Such an approach will provide the required foundation for predicting how a carbocation intermediate can be managed and manipulated through alternative mechanistic sequences. Future work in this regard will be reported in due course.

ASSOCIATED CONTENT

Accession Codes

The atomic coordinates and structure factors of unliganded wild-type EIZS, the L72V EIZS–Mg²⁺–PP_i–BTAC complex, the F95M EIZS–Mg²⁺–PP_i–BTAC complex, the F95H EIZS–Mg²⁺–PP_i–BTAC complex, the A236G EIZS–Mg²⁺–PP_i–BTAC complex, the W325F EIZS–Mg²⁺–PP_i complex, and the V329A EIZS–Mg²⁺–PP_i–BTAC complex have been deposited in the Protein Data Bank as entries 4LTV, 4LXW, 4LTZ, 4LZ3, 4LZ0, 4LZC, and 4LUU, respectively.

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Notes

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ABBREVIATIONS

BME, β -mercaptoethanol; BTAC, benzyltriethylammonium cation; EIZS, *epi*-isozaene synthase; FPP, farnesyl diphosphate; GC, gas chromatography; MS, mass spectrometry; NSLS, National Synchrotron Light Source; PEG, polyethylene glycol; PIPES, piperazine-*N,N'*-bis(2-ethanesulfonic acid); PP_i, inorganic pyrophosphate; rmsd, root-mean-square deviation; TCEP, tris(2-carboxyethyl)phosphine; WT, wild-type.

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