

Determination of the Substrate Binding Mode to the Active Site Iron of (*S*)-2-Hydroxypropylphosphonic Acid Epoxidase Using ¹⁷O-Enriched Substrates and Substrate Analogues[†]

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ABSTRACT: (*S*)-2-Hydroxypropylphosphonic acid epoxidase (HppE) is an O₂-dependent, nonheme Fe(II)-containing oxidase that converts (*S*)-2-hydroxypropylphosphonic acid ((*S*)-HPP) to the regio- and enantiomerically specific epoxide, fosfomycin. Use of (*R*)-2-hydroxypropylphosphonic acid ((*R*)-HPP) yields the 2-keto-adduct rather than the epoxide. Here we report the chemical synthesis of a range of HPP analogues designed to probe the basis for this specificity. In past studies, NO has been used as an O₂ surrogate to provide an EPR probe of the Fe(II) environment. These studies suggest that O₂ binds to the iron, and substrates bind in a single orientation that strongly perturbs the iron environment. Recently, the X-ray crystal structure showed direct binding of the substrate to the iron, but both monodentate (via the phosphonate) and chelated (via the hydroxyl and phosphonate) orientations were observed. In the current study, hyperfine broadening of the homogeneous *S* = 3/2 EPR spectrum of the HppE-NO-HPP complex was observed when either the hydroxyl or the phosphonate group of HPP was enriched with ¹⁷O (*I* = 5/2). These results indicate that both functional groups of HPP bind to Fe(II) ion at the same time as NO, suggesting that the chelated substrate binding mode dominates in solution. (*R*)- and (*S*)-analogue compounds that maintained the core structure of HPP but added bulky terminal groups were turned over to give products analogous to those from (*R*)- and (*S*)-HPP, respectively. In contrast, substrate analogues lacking either the phosphonate or hydroxyl group were not turned over. Elongation of the carbon chain between the hydroxyl and phosphonate allowed binding to the iron in a variety of orientations to give keto and diol products at positions determined by the hydroxyl substituent, but no stable epoxide was formed. These studies show the importance of the Fe(II)-substrate chelate structure to active antibiotic formation. This fixed orientation may align the substrate next to the iron-bound activated oxygen species thought to mediate hydrogen atom abstraction from the nearest substrate carbon.

The epoxide ring is a common structural element found in many natural products. The inherent strain energy associated with the small ring renders it reactive with enzyme nucleophiles. This has allowed many epoxide-containing compounds to be used as enzyme inhibitors and therapeutic agents. A representative example is fosfomycin ((1*R*,2*S*)-epoxypropylphosphonic acid, **1**), an antibiotic used to treat lower urinary tract infections (*1*, *2*). The biological target of

fosfomycin has been identified as UDP- α -D-GlcNAc-*O*-enolpyruvyl¹ transferase (*3*), which catalyzes the attachment of phosphoenolpyruvate to UDP- α -D-GlcNAc, a key step in the assembly of the peptidoglycan layer within the bacterial cell wall.

In most cases, the oxirane moiety of epoxide-containing natural products is biosynthesized through oxidation of the corresponding alkenes by monooxygenases with incorporation of an oxygen atom originated from dioxygen (*4*, *5*). The synthesis of fosfomycin (**1**) is unusual in that the secondary hydroxyl oxygen of the immediate precursor, (*S*)-2-hydroxypropylphosphonic acid ((*S*)-HPP, **2-S**), is retained in the

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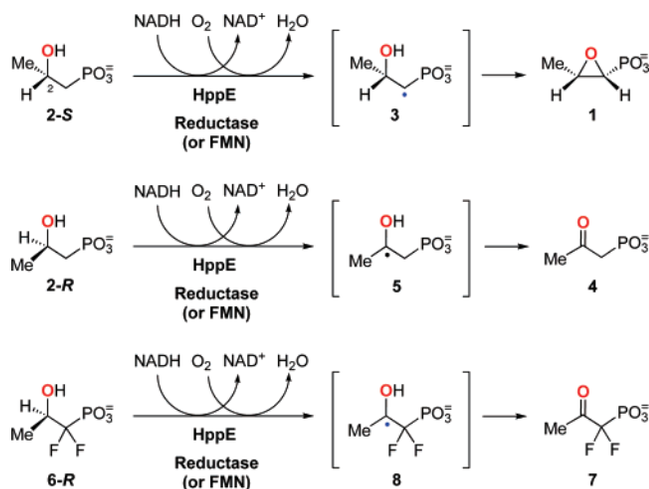
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¹ Abbreviations: ACCO, 1-aminocyclopropane-1-carboxylate oxidase; CS, clavaminase synthase; DTT, dithiothreitol; EDTA, ethylenediaminetetraacetate; EPR, electron paramagnetic resonance; GlcNAc, *N*-acetyl- α -D-glucosamine; HBA, 2-hydroxybutyric acid; HBP, 3-hydroxybutylphosphonic acid; HPEP, 2-hydroxy-2-phenylethylphosphonic acid; HPP, 2-hydroxypropylphosphonic acid; HppE, (*S*)-2-hydroxypropylphosphonic acid epoxidase; IPNS, isopenicillin N synthase; NO, nitric oxide; UDP, uridine 5'-diphosphate.

Scheme 1



fosfomycin product (6–9). Thus, the transformation catalyzed by (*S*)-2-hydroxypropylphosphonic acid epoxidase (HppE) is effectively a dehydrogenation reaction as shown in Scheme 1. The overall epoxidation reaction is a four-electron redox process involving the oxidation of substrate and NAD(P)H and the reduction of dioxygen to water.

We have expressed the HppE gene from *Streptomyces wedmorensis* and purified the encoded protein (8, 9). The purified HppE, after reconstitution, contains a mononuclear nonheme iron that is essential and cannot be replaced by other divalent ions (10, 11). The *in vitro* activity of HppE also requires NAD(P)H, dioxygen, and an auxiliary electron mediator to transfer electrons from NAD(P)H to the enzyme active site (8, 9, 11). While this mediator's role can be fulfilled by small molecule electron carriers, such as FMN, FAD, riboflavin, and benzyl viologen (11), the physiological mediator is expected to be a reductase, which has not yet been identified. Since the only reductase gene in the three known fosfomycin biosynthetic gene clusters of *S. wedmorensis* (12), *Pseudomonas syringae* (13), and *S. fradiae* (14, 15), the electron transfer in the HppE reaction may rely on a promiscuous reductase within the cell.

The detailed mechanism of HppE is unknown, but results from several experimental approaches are consistent with formation of a radical intermediate through the action of an iron-bound activated oxygen species. For example, the natural substrate, 2-S, is converted to 1 by HppE, while its epimer, (*R*)-HPP (2-R), is converted to a keto product, 2-oxopropylphosphonic acid (4, Scheme 1) (16). The distinct fates of these two epimers can be explained on the basis of regiospecific, radical-initiated, hydrogen atom abstraction by HppE, and this was tested by fluorine substitution at C1 of 1. It was found that (*S*)-[1-F₂]HPP (6-S) is an inhibitor for HppE, whereas the epimeric (*R*)-[1-F₂]HPP (6-R, Scheme 1) can still undergo turnover to the corresponding 2-keto product 7 by HppE (16). These results are most easily explained by initial radical formation at C-1 (via 3) in the *S* isomers and at C-2 (via 5 or 8) in the *R* isomers (16).

Not only does HppE exhibit highly regiospecific hydrogen atom abstraction, which is governed by the C-2 stereochemistry of HPP, it can also distinguish the two diastereotopic hydrogens at C-1 of HPP during turnover. The removal of a hydrogen atom from C-1 of 2-S by HppE had been

determined to be pro-*R* stereospecific (17). The strict regio- as well as stereospecificity observed for HppE reaction must stem from a specific substrate binding orientation relative to the reactive iron–oxygen species, which is responsible for the hydrogen abstraction step.

Recent crystallographic characterizations of HppE confirmed that the mononuclear metal ion in the HppE active site is coordinated by a 2-His-1-carboxylate (D/E) core as implicated by sequence alignment and mutagenesis studies (10, 18, 19). Surprisingly, the substrate, 2-S, was found to bind to the metal ion in two different configurations in the enzyme–substrate binary complex (18). As illustrated in Figure 1A, 2-S could bind to the metal ion in a monodentate mode via a phosphonate–oxygen linkage. It could also form a bidentate coordination to the metal ion using the 2-OH group and an oxygen atom from the phosphonate group as ligands (Figure 1B). The presence of two different enzyme–substrate binding modes is unusual. It is not clear whether both binding modes are catalytically relevant. It is also not known whether a similar situation applies for 2-R when it is present as the substrate.

It is important for mechanistic considerations to determine whether one or both of the substrate binding modes detected by X-ray crystallography applies to the functioning enzyme. The substrate chelate mode would suggest that the stereo- and regiospecific reaction is enforced through the use of multiple ligand sites on the iron, whereas these roles would shift to the overall active site structure if the monodentate binding mode applies. To learn more about the substrate–enzyme complex and to investigate the catalytic relevance of the mono- and bidentate coordination, we prepared a series of substrate analogues and [¹⁷O]-labeled HPPs, and studied their binding to HppE and the HppE–NO complexes. The results of this work provide significant insight into how the enzyme and substrate interact and how this interaction determines the regio- and stereospecificity of the enzyme.

EXPERIMENTAL PROCEDURES

General. HppE from *S. wedmorensis* was obtained by expressing the plasmid pPL1001, which contains the corresponding gene (*fom4*), in *E. coli* BL21(DE3)/pPL1001 (9). The overproduced HppE was purified to near homogeneity as previously described (9). All reagents and solvents were purchased from commercial sources and were used without further purification unless otherwise noted. Biochemicals, including fosfomycin (1) and (*S*)-2-hydroxybutyric acid ((*S*)-HBA, 33), were acquired from Aldrich-Sigma (St. Louis, MO). Culture medium ingredients were products of Difco (Detroit, MI). Isotope-labeled water (36.8% ¹⁷O and 48.8% ¹⁸O) was obtained from Isotec Inc. (Miamisburg, OH).

Preparation of Structural Analogues of (*S*)-HPP (2-S). (*R*)-HPP (2-R) and (*S*)- and (*R*)-1,1-difluoro-HPP ([1-F₂]HPP) (6-S, 6-R) were chemically synthesized as previously reported (16).

Synthesis of Isobutylphosphonic Acid (IBP, 9) (Scheme 2). A mixture of isobutyl bromide (10, 1.37 g, 10 mmol), triethylphosphine (P(OEt)₃, 4.2 mL, 20 mmol), and tetrabutylammonium iodide (Bu₄NI, 50 mg, 0.14 mmol) was heated at 150 °C for 3 h and then dried under reduced pressure. The crude products were purified by flash silica gel column chromatography with hexanes/ethyl acetate (10:1 to 1:1) to

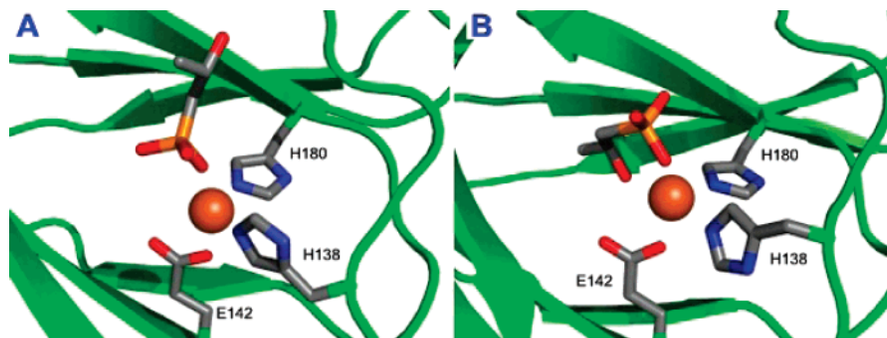
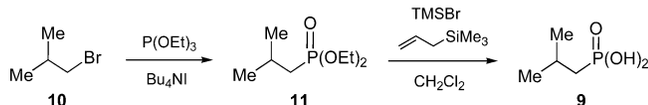


FIGURE 1: X-ray Structures of the modeled HppE active site bound with Fe(II) and 2-S in monodentate (1A) and bidentate (1B) binding modes. Models were generated on the basis of the published crystal coordinates for the *S. wendomorensis* enzyme (PDB entries 1ZZ7 and 1ZZ8).

Scheme 2



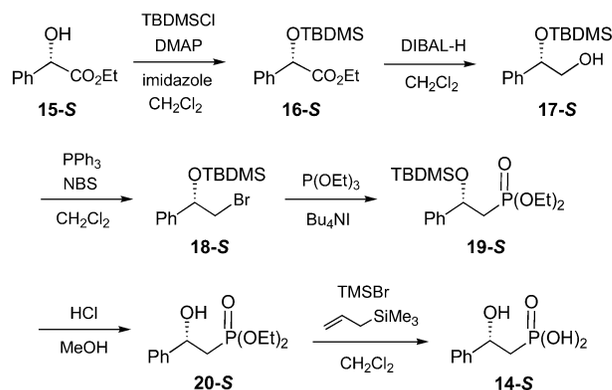
give diethyl isobutylphosphonate (**11**) in 90% yield. A portion of **11** (0.5 g, 2.6 mmol) was then incubated with trimethylsilyl bromide (TMSBr, 0.5 mL, 3.8 mmol) and allyltrimethylsilane (0.25 mL, 1.6 mmol) for 24 h at room temperature. The reaction mixture was neutralized with $\text{NH}_4\text{-HCO}_3$ and extracted with water, and the aqueous fractions were pooled and lyophilized to give IBP (**9**) in white powder form. The overall yield was 95%. ^1H NMR (300 MHz, CDCl_3) δ 2.05 (1H, m), 1.66 (2H, dd, $J = 6.9$, 18), 1.04 (6H, d, $J = 6.9$). ^{13}C NMR (75 MHz, CDCl_3) δ 31.1, 21.9, 21.5 (d, $J = 136.2$). ^{31}P NMR (121 MHz, CDCl_3) δ 32.7.

Preparation of ^{17}O -Labeled HPPs. (*R*)-HPP with an ^{17}O -enriched hydroxyl group (**12-R**) were synthesized by the established procedure (9), except for the replacement of THF with *p*-toluenesulfonic acid (*p*TSA) and H_2^{17}O in the first step. Preparation of the (*S*)-isomer (**12-S**) was also attempted. Unfortunately, the isotope enrichment of this sample was lower than that of (*R*)- ^{17}O HPP. Thus, only the **12-R** sample was subjected to EPR study. The (*R*)-HPP with ^{17}O enriched in the phosphonate group (**13-R**) was prepared by dissolving chemically synthesized (*R*)-HPP (0.04 mmol) in 0.2 mL of ^{17}O H_2O . The solution was incubated at room temperature for 2 weeks. The samples were lyophilized and the ^{17}O isotope enrichment in the products was 20%, as estimated by mass spectrometry performed by the MS facility of the Department of Chemistry and Biochemistry at the University of Texas at Austin.

Synthesis of (*S*)- and (*R*)-2-Hydroxy-2-phenylethylphosphonic Acids ((*S*)- and (*R*)-HPEP, **14-S and **14-R**) (Scheme 3).**

(i) **Ethyl (*S*)-(tert-Butyldimethylsilyloxy)phenylacetate (**16-S**).** To a solution of ethyl (*S*)-hydroxyphenylacetate (**15-S**, 5.0 g, 27.7 mmol), 4-dimethylaminopyridine (DMAP, 0.3 g, 2.5 mmol), and imidazole (4.0 g, 56 mmol) in 25 mL of dry CH_2Cl_2 was added portionwise a solution of *tert*-butyldimethylsilyl chloride (TBDMSCl, 4.6 g, 30 mmol) in 25 mL of CH_2Cl_2 at 0 °C. After being stirred at room temperature for 15 h, the mixture was diluted with CH_2Cl_2 , washed with brine, and dried over anhydrous Na_2SO_4 . The solvents were evaporated under reduced pressure, and the residue was purified by flash silica gel column chromatog-

Scheme 3



raphy (hexanes). The yield of **16-S** was 95%. ^1H NMR (300 MHz, CDCl_3) δ 7.48 (2H, m), 7.31 (3H, m), 5.24 (1H, s), 4.15 (2H, q, $J = 7.2$), 1.22 (3H, t, $J = 7.2$), 0.94 (9H, s), 0.13 (3H, s), 0.06 (3H, s). ^{13}C NMR (75 MHz, CDCl_3) δ 172.2, 139.3, 128.4, 128.1, 126.4, 74.5, 61.1, 25.8, 18.4, 14.2, -4.9, -5.1.

(ii) **(*S*)-(tert-Butyldimethylsilyloxy)-2-phenylethanol (**17-S**).** To a solution of ethyl (*S*)-(tert-butyldimethylsilyloxy)phenylacetate (**16-S**, 7.8 g, 26.5 mmol) in 100 mL of dry CH_2Cl_2 was added dropwise 53 mL of diisobutylaluminum hydride (DIBAL-H) solution (1.5 M in toluene) at -78 °C. After being stirred at room temperature for 4 h, the reaction was cooled to -20 °C, and 14 mL of 1:1 mixture of AcOH/ H_2O was added slowly. The resulting mixture was stirred at -20 °C for 30 min, poured into a saturated NaHCO_3 solution, and extracted with CH_2Cl_2 . The combined organic extracts were dried over anhydrous Na_2SO_4 , and the solvents were removed under reduced pressure. The residue was purified by flash silica gel column chromatography (hexanes:ethyl acetate, 5:1) to give **17-S** in 67% yield. ^1H NMR (300 MHz, CDCl_3) δ 7.30 (5H, m), 4.76 (1H, dd, $J = 6.9$, 5.1), 3.59 (2H, m), 0.93 (9H, s), 0.08 (3H, s), -0.08 (3H, s). ^{13}C NMR (75 MHz, CDCl_3) δ 141.5, 128.3, 127.7, 126.3, 76.0, 69.0, 25.9, 18.3, -4.4, -4.9.

(iii) **(*S*)-2-Bromo-1-phenylethoxy-tert-butyldimethylsilane (**18-S**).** To a solution of (*S*)-(tert-butyldimethylsilyloxy)-2-phenylethanol (**17-S**, 2.53 g, 10 mmol) and PPh_3 (2.90 g, 11 mmol) in 20 mL of CH_2Cl_2 was added *N*-bromosuccinimide (NBS, 2.0 g, 11 mmol) at -78 °C. After being stirred at the same temperature for 10 min and at room temperature for 3 h, the mixture was diluted with ether, washed with brine, and dried over anhydrous Na_2SO_4 . The solvents were removed under reduced pressure, and the residue was purified

by flash silica gel column chromatography (hexanes) to give **18-S** in 78% yield. ^1H NMR (300 MHz, CDCl_3) δ 7.36 (5H, m), 4.88 (1H, dd, $J = 7.8, 5.1$), 3.48 (2H, m), 0.93 (9H, s), 0.14 (3H, s), -0.06 (3H, s). ^{13}C NMR (75 MHz, CDCl_3) δ 142.3, 128.5, 128.1, 126.3, 75.4, 39.7, 25.9, 18.4, -4.6 , -4.8 .

(iv) Diethyl (*S*)-2-Phenyl-2-(*tert*-butyldimethylsilyloxy)ethylphosphonate (**19-S**). A mixture of (*S*)-2-bromo-1-phenylethoxy-*tert*-butyldimethylsilane (**18-S**, 1.09 g, 3.4 mmol), $\text{P}(\text{OEt})_3$ (2.0 mL, 13.6 mmol), and Bu_4NI (40 mg, 0.11 mmol) was heated at 120 °C for 23 h and then dried under reduced pressure. The residue was loaded onto a silica gel column, and the product was eluted with hexanes/ethyl acetate (10:1 to 1:1). Compound **19-S** was isolated in 45% yield. ^1H NMR (300 MHz, CDCl_3) δ 7.29 (5H, m), 5.03 (1H, m), 3.92 (4H, m), 2.32 (1H, ddd, $J = 16.8, 15.3, 6.9$), 2.12 (1H, ddd, $J = 17.4, 15.3, 5.7$), 1.23 (3H, t, $J = 7.2$), 1.16 (3H, t, $J = 7.2$), 0.82 (9H, s), 0.03 (3H, s), -0.22 (3H, s). ^{13}C NMR (75 MHz, CDCl_3) δ 144.8 (d, $J = 9.6$), 128.3, 127.6, 126.2, 70.8, 61.3 (d, $J = 4.1$), 61.2 (d, $J = 4.1$), 37.8 (d, $J = 137.0$), 25.8, 18.1, 16.4, 16.3, -4.6 , -4.9 . ^{31}P NMR (121 MHz, CDCl_3) δ 27.9.

(v) Diethyl (*S*)-2-Hydroxy-2-phenylethylphosphonate (**20-S**). A mixture of diethyl (*S*)-2-phenyl-2-(*tert*-butyldimethylsilyloxy)ethylphosphonate (**19-S**, 0.48 g, 1.2 mmol) and concentrated HCl (0.5 mL) in 5 mL of methanol was stirred for 1 h at room temperature. The solvents were removed *in vacuo*, and the residue was purified by flash silica gel column chromatography (hexanes:ethyl acetate, 2:1 to 1:3). The yield of **20-S** was 87%. ^1H NMR (300 MHz, CDCl_3) δ 7.34 (5H, m), 5.09 (1H, m), 4.09 (4H, m), 3.96 (1H, d, $J = 3.0$), 2.19 (2H, m), 1.33 (3H, t, $J = 7.5$), 1.28 (3H, t, $J = 6.9$). ^{13}C NMR (75 MHz, CDCl_3) δ 143.5 (d, $J = 16.1$), 128.6, 127.8, 125.6, 68.9 (d, $J = 4.3$), 62.2 (d, $J = 6.0$), 62.0 (d, $J = 7.0$), 36.0 (d, $J = 135.8$), 16.5 (d, $J = 6.0$), 16.4 (d, $J = 6.6$). ^{31}P NMR (121 MHz, CDCl_3) δ 29.9.

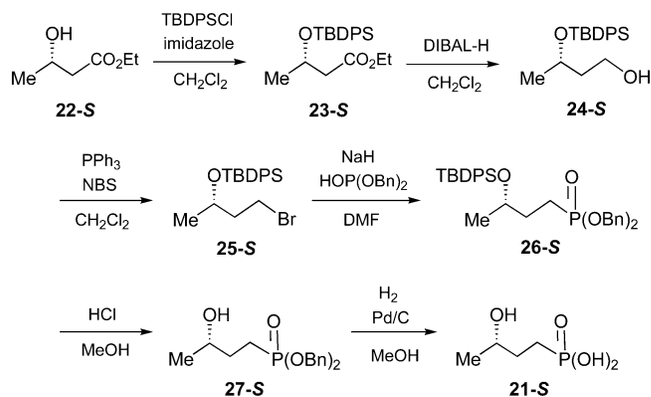
(vi) (*S*)-2-Hydroxy-2-phenylethylphosphonic Acid ((*S*)-HPEP, **14-S**). A solution of diethyl (*S*)-2-hydroxy-2-phenylethylphosphonate (**20-S**, 0.24 g, 0.9 mmol), TMSBr (0.4 mL, 3.0 mmol) and allyltrimethylsilane (0.2 mL, 1.3 mmol) in 4 mL of CH_2Cl_2 was stirred for 25 h at room temperature. The solvents were removed *in vacuo*, and the residue was vigorously stirred with 5 mL of water for 10 min. The aqueous solution was neutralized with NH_4HCO_3 , washed with CHCl_3 , and then lyophilized to give product **14-S** as a white powder in 96% yield. ^1H NMR (300 MHz, D_2O) δ 7.26 (5H, m), 4.76 (1H, m), 1.75 (2H, m). ^{13}C NMR (75 MHz, D_2O) δ 143.9 (d, $J = 13.2$), 128.6, 127.8, 126.1, 70.6 (d, $J = 29.0$), 37.5 (d, $J = 126.3$). ^{31}P NMR (121 MHz, D_2O) δ 20.9.

(vii) (*R*)-2-Hydroxy-2-phenylethylphosphonic Acid ((*R*)-HPEP, **14-R**). Preparation of this compound followed an identical procedure developed for the synthesis of (*S*)-2-hydroxy-2-phenylethylphosphonic acid (**14-S**) using ethyl (*R*)-(*tert*-butyldimethylsilyloxy)phenylacetate as the starting material.

Synthesis of (*S*)- and (*R*)-3-Hydroxybutylphosphonic Acid ((*S*)- and (*R*)-HBP, **21-S** and **21-R**) (Scheme 4).

(i) Ethyl (*S*)-3-(*tert*-Butyldiphenylsilyloxy)butyrate (**23-S**). To a solution of ethyl (*S*)-3-hydroxybutyrate (**22-S**, 2.0 g, 16 mmol) and imidazole (1.3 g, 19 mmol) in 100 mL of dry CH_2Cl_2 was added dropwise *tert*-butyldiphenylsilyl chloride (TBDPSCI, 5.0 g, ~ 18 mmol) at 0 °C. After being stirred

Scheme 4



overnight at room temperature, the mixture was diluted with CH_2Cl_2 , washed with brine, and dried over anhydrous Na_2SO_4 . The solvents were then removed under reduced pressure, and the residue was purified by flash silica gel column chromatography (hexanes). The yield of **23-S** was 92%. ^1H NMR (300 MHz, CDCl_3) δ 7.70 (4H, m), 7.42 (6H, m), 4.33 (1H, m), 4.15 (2H, q, $J = 7.2$), 2.59 (1H, dd, $J = 14.7, 7.2$), 2.41 (1H, dd, $J = 14.7, 5.7$), 1.13 (3H, t, $J = 7.2$), 1.06 (9H, s). ^{13}C NMR (75 MHz, CDCl_3) δ 171.9, 136.0, 135.9, 134.4, 133.9, 129.7, 129.6, 127.7, 127.6, 67.0, 51.5, 44.5, 27.0, 23.7, 19.3.

(ii) (*S*)-3-(*tert*-Butyldiphenylsilyl)butane-1,3-diol (**24-S**). To a solution of ethyl (*S*)-3-(*tert*-butyldiphenylsilyloxy)butyrate (**23-S**, 5.4 g, 15 mmol) in 100 mL of dry CH_2Cl_2 was added dropwise DIBAL-H (34 mL, 1.5 M solution in toluene) at -78 °C. After being stirred at room temperature for 2 h, the mixture was cooled to -20 °C, quenched by the addition of 10 mL of saturated NH_4Cl solution, poured into saturated NaHCO_3 solution, and extracted with CH_2Cl_2 . The organic extracts were combined and dried over anhydrous Na_2SO_4 . The solvents were removed under reduced pressure, and the residue was purified by flash silica gel column chromatography (hexanes:ethyl acetate, 10:1 to 3:1). Compound **24-S** was isolated in 55% yield. ^1H NMR (300 MHz, CDCl_3) δ 7.73 (4H, m), 7.36 (6H, m), 4.13 (1H, m), 3.86 (1H, m), 3.71 (1H, m), 2.24 (1H, br s), 1.84 (1H, m), 1.67 (1H, m), 1.11 (3H, d, $J = 6.3$), 1.08 (9H, s). ^{13}C NMR (75 MHz, CDCl_3) δ 135.9, 135.8, 134.2, 133.7, 129.8, 129.7, 127.7, 127.6, 68.8, 60.0, 40.7, 27.0, 23.0, 19.2.

(iii) (*S*)-1-Bromo-3-(*tert*-butyldiphenylsilyloxy)butane (**25-S**). To a solution of (*S*)-3-(*tert*-butyldiphenylsilyl)butane-1,3-diol (**24-S**, 3.3 g, 10.1 mmol) and triphenylphosphine (PPh_3 , 3.2 g, 12.3 mmol) in 25 mL of CH_2Cl_2 was added *N*-bromosuccinimide (2.0 g, 11.2 mmol) at -78 °C. After being stirred for 2 h at room temperature, the mixture was diluted with ether, washed with brine, and dried over anhydrous Na_2SO_4 . The solvents were removed under reduced pressure, and the residue was purified by flash silica gel column chromatography (hexanes) to give **25-S** in 56% yield. ^1H NMR (300 MHz, CDCl_3) δ 7.72 (4H, m), 7.41 (6H, m), 4.04 (1H, m), 3.49 (2H, m), 2.10 (1H, m), 1.95 (1H, m), 1.08 (12H, m). ^{13}C NMR (75 MHz, CDCl_3) δ 135.9, 134.5, 133.8, 129.7, 129.6, 127.7, 127.5, 68.1, 42.6, 30.2, 27.1, 23.3, 19.4.

(iv) Dibenzyldiphenylsilyloxy (*S*)-3-(*tert*-Butyldiphenylsilyloxy)butylphosphonate (**26-S**). To a solution of dibenzyl phosphite (0.66 g, 2.5 mmol) in dry DMF (3 mL) was added NaH (2.5

mmol) at room temperature. After the mixture was stirred for 10 min, (*S*)-1-bromo-3-(*tert*-butyldiphenylsilyloxy)butane (**25-S**, 0.50 g, 1.3 mmol) in dry DMF (2 mL) was added. The resulting mixture was stirred overnight, diluted with CH₂-Cl₂, washed with water, and dried over anhydrous Na₂SO₄. The solvents were evaporated under reduced pressure, and the residue was flash chromatographed on silica gel (hexanes: ethyl acetate, 3:1 to 1:1) to give **26-S** in 82% yield. ¹H NMR (300 MHz, CDCl₃) δ 7.65 (4H, m), 7.32 (16H, m), 5.00 (4H, m), 3.85 (1H, m), 1.99–1.67 (4H, m), 1.04 (9H, s), 1.00 (3H, d, *J* = 6.3). ¹³C NMR (75 MHz, CDCl₃) δ 136.6, 136.5, 135.9, 135.8, 134.4, 134.0, 129.7, 129.6, 128.6, 128.3, 127.9, 127.8, 127.6, 127.5, 69.0 (d, *J* = 19.6), 67.0 (d, *J* = 6.3), 31.6 (d, *J* = 5.1), 27.0, 22.6, 21.6 (d, *J* = 141.9), 19.3. ³¹P NMR (121 MHz, CDCl₃) δ 35.1.

(v) *Dibenzyl (S)-3-Hydroxybutylphosphonate (27-S)*. A mixture of dibenzyl (*S*)-3-(*tert*-butyldiphenylsilyloxy)butylphosphonate (**26-S**, 0.31 g, 0.5 mmol) and concentrated HCl (0.35 mL) in 3 mL of methanol was stirred for 13 h at room temperature. The solvents were removed *in vacuo*, and the residue was purified by flash silica gel column chromatography (hexanes:ethyl acetate, 2:1 to 1:2). The yield of **27-S** was 72%. ¹H NMR (300 MHz, CD₃OD) δ 7.34 (10H, m), 4.99 (4H, m), 3.65 (1H, m), 2.02–1.52 (4H, m), 1.08 (3H, d, *J* = 5.7). ¹³C NMR (75 MHz, CD₃OD) δ 136.3, 128.4, 128.3, 127.9, 67.4, 66.9, 66.7, 31.1, 21.9, 21.5 (d, *J* = 138.2). ³¹P NMR (121 MHz, CD₃OD) δ 35.6.

(vi) *(S)-3-Hydroxybutylphosphonic Acid ((S)-HBP, 21-S)*. A mixture of 10% Pd/C (80 mg) and dibenzyl (*S*)-3-hydroxybutylphosphonate (**27-S**, 140 mg, 0.42 mmol) in 10 mL of methanol was vigorously stirred for 30 min under a hydrogen atmosphere at room temperature. The mixture was filtered through a short celite column. The solvents were then removed under reduced pressure, and the residue was dissolved in water and neutralized with NH₄HCO₃. The aqueous solution was lyophilized to give **21-S** as a white powder in 96% yield. ¹H NMR (300 MHz, D₂O) δ 3.58 (1H, m), 1.34 (4H, m), 0.92 (3H, d, *J* = 6.0). ¹³C NMR (75 MHz, D₂O) δ 68.3 (d, *J* = 17.9), 32.1 (d, *J* = 4.0), 24.2 (d, *J* = 134.6), 21.3. ³¹P NMR (121 MHz, D₂O) δ 26.2.

(*R*)-3-Hydroxybutylphosphonic Acid ((*R*)-HBP, **21-R**). This compound was synthesized by the same procedure as described for 3-(*S*)-hydroxybutylphosphonic acid (**21-S**) using ethyl (*R*)-3-(*tert*-butyldiphenylsilyloxy)butyrate as the starting material in the first step.

Characterization of Turnover Products Generated in the Incubation of Substrate Analogues with HppE. Chemically synthesized substrate analogues were individually examined for their competence as substrates by the *in vitro* HppE assay (9). The products generated in the incubation mixture were purified by a P-2 gel filtration column with H₂O as the eluent. The desired fractions were pooled and lyophilized, and the purified products were dissolved in D₂O for NMR analysis. For 1,2-oxy-2-phenyl-ethylphosphonic acid (**28**) derived from **14-S**: ¹H NMR (300 MHz, D₂O) δ 7.15 (5H, m), 3.78 (1H, m), 2.81 (1H, dd, *J* = 19.2, 3); ¹³C NMR (75 MHz, D₂O) δ 136.4, 128.5, 128.4, 125.7, 58.9 (d, *J* = 170.0), 57.1; ³¹P NMR (121 MHz, D₂O) δ 9.9. For 2-oxo-2-phenyl-ethylphosphonic acid (**29**) derived from **14-R**: ¹H NMR (300 MHz, D₂O) δ 7.20–7.91 (2H, m), 7.48–7.52 (1H, m), 7.36–7.40 (2H, m), 3.30 (2H, d, *J* = 20.5); ³¹P NMR (121 MHz, D₂O) δ 10.8. For (3*R*)-3,4-dihydroxybutylphosphonic acid (**31**)

derived from **21-S**: ¹H NMR (500 MHz, D₂O) δ 3.44–3.49 (1H, m), 3.39 (1H, dd, *J* = 11.5, 3.5), 3.29 (1H, dd, *J* = 11.5, 6.5), 1.41–1.52 (1H, m), 1.30–1.40 (2H, m), 1.16–1.28 (1H, m); ¹³C NMR (75 MHz, D₂O) δ 70.3, 68.2, 33.1 (d, *J* = 4.0), 24.3 (d, *J* = 134.6); ³¹P NMR (121 MHz, D₂O) δ 23.3. For 3-oxo-butylphosphonic acid (**32**) derived from **21-R**: ¹H NMR (500 MHz, D₂O) δ 2.57 (2H, t, *J* = 7.5), 2.08 (3H, s), 1.42–1.48 (2H, m); ¹³C NMR (75 MHz, D₂O) δ 215.6, 38.3, 28.9, 22.6 (d, *J* = 133.0); ³¹P NMR (121 MHz, D₂O) δ 22.5.

Preparation of NO Samples of Fe^{II}-HppE Bound with Substrate or Substrate Analogues. Double distilled H₂O (ddH₂O) and Tris·HCl (50 mM, pH 7.5) buffer were made anaerobic by repetitive vacuum and bubbling with argon that had been passed over a column of copper catalyst (BASF Inc.) heated at 150 °C for approximately 3 h. The Fe(II) stock solution was prepared by adding 1 mL of the above ddH₂O through a gastight syringe into a serum vial containing preweighted Fe(NH₄)₂(SO₄)₂·6H₂O (99.997%, Aldrich) under anaerobic conditions. Anaerobic solutions of metal-free apoprotein, substrate, and substrate analogues were prepared in reaction vials by repeated cycles of vacuum and flushing with argon and then transferred to the EPR sample tubes under an argon atmosphere. Nitric oxide (NO) was introduced through a gastight syringe to the headspace of the quartz EPR tubes or UV-vis cuvettes containing the ferrous enzymes prepared under anaerobic conditions. An argon flush was maintained above samples to protect them from oxidation by O₂ and to minimize an anomalous EPR signal near *g* = 2 which derives from NO.

The enzyme–substrate nitrosyl ternary complexes were made by adding substrate (or substrate analogues) and then NO (or *vice versa*) to the enzyme in its reduced state. For the isotope-labeling experiments, ¹⁷OH-labeled substrates were first added to the enzyme, followed by NO gas. Control samples were prepared in parallel using natural-abundance substrate. EPR samples were frozen by slow immersion in liquid N₂ for latter use.

EPR Spectroscopy. The ¹⁷O-EPR spectra were recorded at 9.6 GHz frequency using a Bruker Elexsys E500 spectrometer. The EPR data of various substrate analogues were acquired using a Bruker EMX spectrometer and a high sensitivity cavity at 9.38 GHz. Both spectrometers were equipped with an Oxford Instruments liquid helium cryostat. Temperature was controlled by a digitalized Oxford temperature controller. A Bruker ER035M Gauss Meter was used during measurements to assist the *g*-value determination. EPR spectra of the *S* = 3/2 complexes were analyzed according to the spin Hamiltonian:

$$\hat{\mathcal{H}} = g\beta_e B_0 S + D[\hat{S}_z^2 - 5/4 + E/D(S_x^2 - S_y^2)]$$

where *B*₀ is the magnetic field, *D* is the axial zero-field splitting parameter, and *E/D* indicates the degree of rhombic distortion in the electronic environment. Changes in the *E/D* term are diagnostic of changes in the environment of the iron. Computer simulation of experimental EPR data was performed using SimFonia (Bruker).

RESULTS

Electronic Absorption Spectra of Fe(II)-HppE Nitrosyl Complexes. As a result of using DTT and EDTA in the

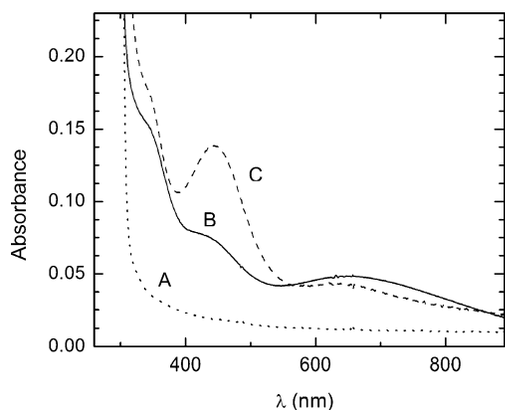


FIGURE 2: Electronic absorption spectra of Fe(II)–HppE nitrosyl complexes. (A) As-isolated HppE (200 μ M) (metal-free); (B) HppE (180 μ M) reconstituted with Fe(II) and exposed to NO; (C) to sample B was added (S)-HPP (4 mM).

protein purification procedure, the as-purified HppE is essentially an apoprotein (9), and it has little UV–vis absorbance above 300 nm (Figure 2, curve A). Upon anaerobic reconstitution with a stoichiometric amount of Fe-(NH₄)₂(SO₄)₂, the resultant Fe(II)–HppE remained colorless above 300 nm (same as curve A in Figure 2). However, the addition of NO to the ferrous enzyme yielded a pale yellow complex with two absorption maxima at 438 nm ($\epsilon \approx 750$ M⁻¹ cm⁻¹ per iron) and 640 nm ($\epsilon \approx 250$ M⁻¹ cm⁻¹ per iron) (Figure 2, curve B). These two bands can be assigned, based on previous studies of the ferrous-nitrosyl complexes of various enzymes and model compounds, to the electronic absorption of the NO⁻ → Fe³⁺ charge transfer and the Fe³⁺ ligand field transition, respectively (20). This observation provided initial evidence for binding of NO to the metal center. The visible spectrum also has a shoulder at 354 nm that was observed to vary in intensity depending on the concentration of NO. A similar spectral feature at 340 nm was previously noted in the spectrum of the nitrosyl complex of isopenicillin N synthase (IPNS) and was ascribed to nonspecific interactions of NO with protein (21).

When the Fe(II)–HppE nitrosyl binary complex was subjected to repeated cycles of vacuum and flushing with argon, the pale-yellow color gradually faded due to loss of NO, and the enzyme eventually returned to the colorless state. Interestingly, the addition of NO to Fe(II)–HppE in the presence of **2-S** led to a ternary complex with an optical spectrum exhibiting maxima at 438 ($\epsilon \approx 400$ M⁻¹ cm⁻¹ per iron) and 680 nm ($\epsilon \approx 300$ M⁻¹ cm⁻¹ per iron) (Figure 2, curve C). The large change in absorptivity of the 438 nm Fe-nitrosyl band indicates significant changes in the metal coordination environment upon substrate binding.

EPR Spectra of Fe(II)–HppE Nitrosyl Complexes. Previously EPR studies showed that the spectrum of the nitrosyl complex of Fe(II)-reconstituted HppE is composed of two superimposed $S = 3/2$ species (Table 1) with distinct relaxation properties (9). Upon addition of **2-S**, most of the heterogeneity disappears and a single species ($S = 3/2$) with increased rhombicity, $E/D = 0.066$, having principal g -values of 4.42, 3.63, and 1.97, prevails (Figure 3) (9). The observed spectral change is independent of the addition order of **2-S** and NO to the EPR sample and is consistent with the change of the electronic spectrum observed upon substrate binding to the Fe(II)–HppE nitrosyl complex (Figure 2) (9). These

Table 1: EPR Properties of Fe(II)–HppE Nitrosyl Complexes

complex	compd	$g_{ }$	E/D	rel %
HppE-Fe(II)•(S)-HPP•NO	2-S	4.42, 3.63	0.066	>95
HppE-Fe(II)•(R)-HPP•NO	2-R	4.42, 3.63	0.066	>95
HppE-Fe(II)•(S)-HFPP•NO	6-S	4.42, 3.63	0.066	>95
HppE-Fe(II)•(R)-HFPP•NO	6-R	4.42, 3.63	0.066	>95
HppE-Fe(II)•HBA•NO	33	4.26, 3.80	0.038	93
HppE-Fe(II)•IBP•NO	9	4.45, 3.61	0.070	63
		4.17, 3.89	0.024	33
HppE-Fe(II)•fosfomycin•NO	1	4.47, 3.60	0.073	54
		4.23, 3.84	0.033	40
HppE-Fe(II)•(S)-HPEP•NO	14-S	4.37, 3.69	0.056	43
		4.21, 3.88	0.028	52
HppE-Fe(II)•(R)-HPEP•NO	14-R	4.53, 3.51	0.085	29
		4.37, 3.70	0.056	26
		4.21, 3.87	0.028	26
HppE-Fe(II)•(S)-HBP•NO	21-S	4.48, 3.57	0.075	26
		4.31, 3.76	0.046	46
		4.17, 3.89	0.023	23
HppE-Fe(II)•(R)-HBP•NO	21-R	4.40, 3.66	0.061	63
		4.18, 3.89	0.024	34

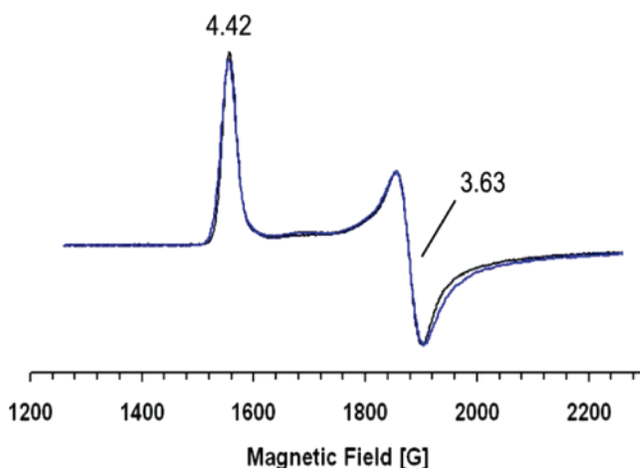


FIGURE 3: EPR spectra of Fe(II)–HppE•HPP nitrosyl complexes. Blue line: Fe(II)–HppE (250 μ M) mixed with **2-S** (2.5 mM) and exposed to NO; Black line: Fe(II)–HppE (250 μ M) mixed with **2-R** (2.5 mM) and exposed to NO. Instrumental parameters: temperature 2 K, microwave frequency 9.6 GHz, microwave power 0.6 mW, modulation amplitude 5 G, time constant 0.02 s, and a sweep rate of 50 G/s.

results suggest that **2-S** binds near or directly to the active site Fe(II) leading to reorganization of the metal coordination core. The sensitivity of the EPR spectrum to changes in the structure or environment of the Fe-NO complex increases as the rhombicity increases in the range of 0 to ~ 0.2 and then decreases as E/D approaches the limiting value of 0.33. Thus, the observation of an EPR spectrum from only one species with an intermediate size E/D value shows that the iron center has become quite homogeneous.

EPR Spectra of Fe(II)–HppE Complexed with Nitric Oxide and Stereoisomers of HPP and [1-F₂]HPP. To gain more insight into substrate binding to HppE in the presence of NO, the EPR spectra of Fe(II)–HppE nitrosyl complex coordinated with **2-R**, and the fluorinated substrate analogues **6-S** and **6-R** were studied. It was found that the EPR spectrum of **2-R**-bound Fe(II)–HppE nitrosyl complex is nearly superimposable on that of **2-S**-bound complex (Figure 3). This suggests that **2-S** and **2-R** bind to the HppE active site in a similar manner. Very similar spectra were also obtained when **6-S** and **6-R** were bound (Table 1). Appar-

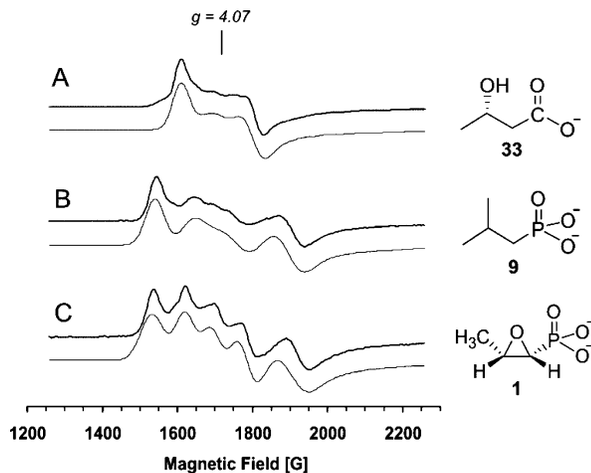


FIGURE 4: EPR spectra of Fe(II)–HppE complexed with NO and different substrate analogues. (A) (*S*)-HBA (**33**); (B) IBP (**9**); (C) fosfomycin (**1**). Instrumental parameters are the same as shown in Figure 3. Gray lines represent simulated spectra.

ently, the stereochemistry at C-2 and the fluorine substitution at C-1 have little effect on the electronic symmetry of the enzyme–substrate–nitrosyl complex, suggesting that the same functional groups are bound in the same ligand sites.

EPR Spectra of Fe(II)–HppE Complexed with Nitric Oxide and HPP Analogues Lacking 2-OH or Phosphonate Group. To probe the role of the hydroxyl and phosphonate groups of HPP in binding to the metal center, two substrate analogues, (*S*)-2-hydroxybutyric acid ((*S*)-HBA, **33**, which has the phosphonate group replaced by a carboxylic group, and isobutylphosphonic acid (IBP, **9**) which carries a methyl group instead of a hydroxyl group at C-2, were prepared. Neither compound is a substrate for HppE based on the *in vitro* activity assay (9). Each of them was added to the nitrosyl complex of Fe(II)–HppE for EPR analysis, and the resulting EPR spectra are shown in Figure 4. A minor resonance at $g = 4.07$ was present in all spectra and is assigned to a small amount of denatured enzyme.²

The spectrum of the nitrosyl complex of **33**–HppE (Figure 4A) exhibits an E/D value for the dominant species of 0.038 that is much smaller than those detected for the **2-S** and **2-R** complexes, indicating that the iron environment is not the same as observed for true substrates (Table 1). Nevertheless, the (*S*)-HBA spectrum is dominated by one species like those of the **2-S** and **2-R** complexes, suggesting that this molecule binds predominantly in one binding orientation. Thus, the carboxylate group appears to be an effective substitute for phosphonate group in this role. In contrast, the spectrum of the nitrosyl complex of the enzyme with **9** is composed of two major EPR species (Figure 4B): one ($g = 4.45, 3.61$; $E/D = 0.070$) has a slightly larger E/D value than the corresponding spectrum of the **2-S** complex, while the other ($g = 4.17, 3.89$; $E/D = 0.024$) has a much smaller E/D value (Table 1). Since **9** is an achiral molecule, the two sets EPR signals cannot be an artifact due to the presence of two stereoisomers in the active site. Instead, they are likely to be the result of binding of **9** to Fe(II) in two different

² The intensity of the $g = 4.07$ signal was dependent on NO concentration. When excess NO (3 psi) was introduced to Fe(II)–HppE, the enzyme became denatured. Gentle shaking caused the enzyme to redissolve but the EPR spectrum exhibited the $g = 4.07$ signal, which is not observed for the nitrosyl complex of the active enzyme.

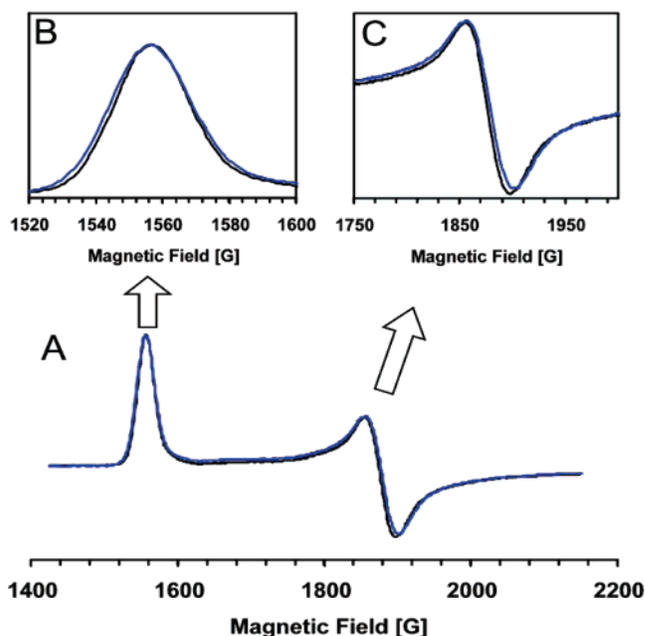


FIGURE 5: EPR spectra of Fe(II)–HppE nitrosyl complex with (*R*)-[¹⁷O]HPP (**12-R**). Panel A: full low field region of the spectrum; Panel B: detailed view of the $g = 4.42$ region; Panel C: detailed view of the $g = 3.63$ region. Black lines: (*R*)-[¹⁶O]HPP; Blue lines: (*R*)-[¹⁷O]HPP. Instrumental parameters are the same as shown in Figure 3.

configurations. This is made more likely by the fact that **9** only has one potential metal binding group to provide orientation, whereas each of the other binding molecules discussed thus far has two. The fact that **9** binds to HppE in two different configurations when the 2-OH group is absent is suggestive of an active role of 2-OH in metal binding. Interestingly, the EPR spectrum of **1** (which also has only one binding determinant) bound to the nitrosyl complex of Fe(II)–HppE (Figure 4C, Table 1) resembles that of **9**. A free hydroxyl group at C-2 of the substrate is apparently important to metal coordination.

EPR Spectra of Fe(II)–HppE Complexed with Nitric Oxide and ¹⁷O-Labeled HPP. In order to fully characterize substrate coordination to the iron center of HppE, ¹⁷O-labeled HPP was chemically synthesized and used in sample preparation. Figure 5 shows the EPR spectra of Fe(II)–HppE·HPP nitrosyl complexes prepared using (*R*)-HPP with ¹⁷O enriched in the 2-OH group ((*R*)-[¹⁷O]HPP, **12-R**). The spectrum is superimposed with a reference spectrum obtained using unlabeled **2-R** for comparison. The small, but clearly discernible, line width broadening due to hyperfine interactions between ¹⁷O nuclear spin ($I = 5/2$) and {Fe–NO}⁷ electronic spin ($S = 3/2$) is obtained. As shown in Figures 5B and 5C, the ¹⁷O labeling causes broadening of 2.3 and 2.5 G of the spectral features near $g = 4.42$ and 3.63, respectively. Considering the low enrichment (~20%) of ¹⁷O in the sample, this change is comparable to those reported for other ¹⁷O-labeled-substrate–enzyme complexes and thus demonstrates direct coordination of the hydroxyl oxygen of HPP to the iron center (22–25).

(*R*)-HPP with ¹⁷O enriched in the phosphonate group was prepared by incubating unlabeled **2-R** in [¹⁷O]-enriched water for several weeks. Binding of this (*R*)-[P¹⁷O₃]HPP (**13-R**, 36.8% enrichment of ¹⁷O) to the Fe(II)–HppE nitrosyl complex led to the apparent hyperfine broadening in each

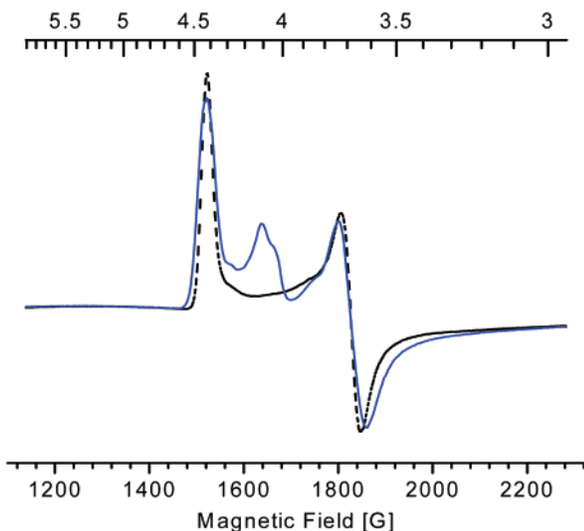
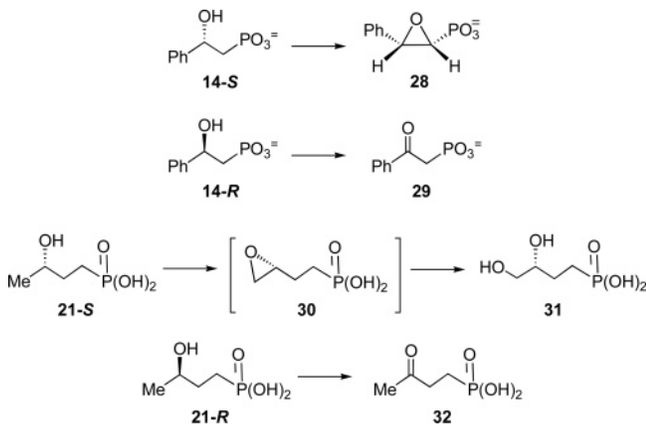


FIGURE 6: EPR spectra of Fe(II)-HppE-nitrosyl complex with (*R*)-[P¹⁷O₃]HPP (**13-R**). Black dotted line: (*R*)-[P¹⁶O₃]HPP; Blue solid line: (*R*)-[P¹⁷O₃]HPP. Instrumental parameters are the same as shown in Figure 5. Spectra are aligned to give approximately the same number of integrated spins of the *E/D* = 0.066 species. The minor resonance at *g* = 4.07 in the spectrum of the (*R*)-[P¹⁷O₃]HPP (**13-R**) containing sample is due to a minor amount of denatured enzyme.²

Scheme 5



spectral resonance (Figure 6). On the basis of the change in width at half-height of the resonance near *g* = 4.42, the broadening is approximately 11 G, the largest reported for interactions of this type. This demonstrates the direct coordination between the phosphonate oxygen and the iron center. Since the EPR spectrum of the nitrosyl complex of the enzyme-substrate complexes shows that there is only one species present, the hyperfine broadening observed when either the OH or the phosphonate substituent is labeled with ¹⁷O shows that these groups simultaneously bind to the iron. Moreover, the presence of NO as a ligand shows that three ligand sites are simultaneously occupied by exogenous ligands in the complex frozen from solution.

Activity Assay and EPR Analysis of Other Substrate Analogues. Four other HPP analogues were synthesized and used to study substrate binding in the HppE active site (Scheme 5). The phenyl-substituted (*S*)- and (*R*)-HPEP (**14-S** and **14-R**) are both substrates for HppE. They could be converted to the corresponding epoxide (**28**) and ketone (**29**) product, albeit at much reduced rates. Their addition to the nitrosyl complex of Fe(II)-HppE led to complex EPR

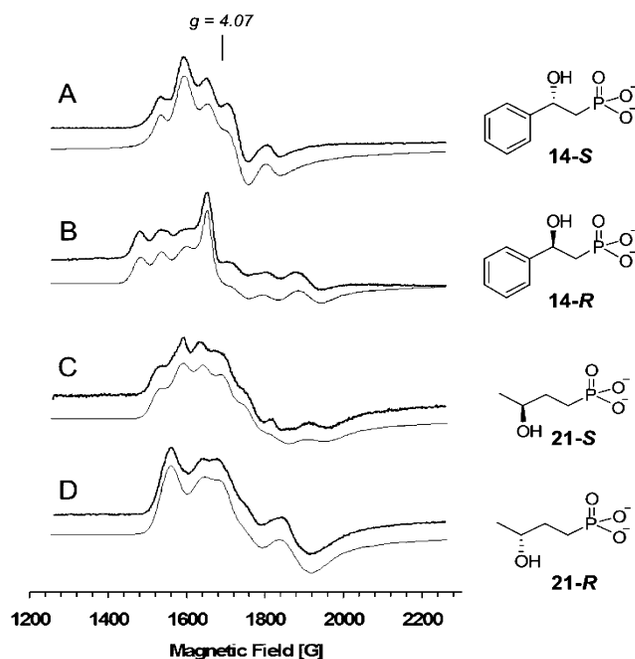


FIGURE 7: EPR spectra of Fe(II)-HppE-nitrosyl complexes with different substrate analogues. (A) (*S*)-HPEP (**14-S**); (B) (*R*)-HPEP (**14-R**); (C) (*S*)-HBP (**21-S**); (D) (*R*)-HBP (**21-R**). Instrumental parameters are the same as shown in Figure 3. Gray lines represent simulated spectra.

spectra with multiple overlapped signals in the *g* = 4 region (Figure 7). Table 1 summarizes the relative spin quantitation of each of the EPR signals based on simulations. In contrast to the nearly identical spectra arising from the nitrosyl complexes of HppE bound with **2S** and **2R**, those from the complexes with (*S*)-HPEP (**14-S**) and (*R*)-HPEP (**14-R**) are quite different (Figure 7A and 7B). It appears that many binding orientations are possible for each isomer, likely stemming from the bulk of the phenyl group.

(*S*)- and (*R*)-HBP (**21-S** and **21-R**) which contain an extra methylene unit between the hydroxyl and phosphonate groups were also prepared. The increased distance between these two iron-binding groups is expected to have significant impact on the regio- as well as stereoselectivity of turnover. Reaction of HBP with HppE showed that both isomers could serve as substrates. The product of (*S*)-HBP (**21-S**) is a diol derivative (**31**), while that of (*R*)-HBP (**21-R**) is a ketophosphonate (**32**). The former may derive from hydrolysis of an epoxide precursor (**30**, Scheme 5). In both cases, the oxidation occurred at C-4 or C-3, instead of at C-1 or C-2. These results imply a more crucial role of the hydroxyl group in determining the regiochemistry of the reaction. The EPR spectra of Fe(II)-HppE nitrosyl complexes with **21-S** or **21-R** (Figures 7C and 7D) are different from those with **2-S** or **2-R**, consistent with the different products that are generated.

DISCUSSION

Substrate Binding to HppE in the Presence of NO. Nitric oxide has been used extensively as a probe for the oxygen-binding site in nonheme iron enzymes because of its similarity to O₂ in size and binding mode as well as its ability to bind to ferrous ions to form EPR-visible *S* = 1/2 (26–29) or *S* = 3/2 (21, 23, 30–36){FeNO}⁷ complexes (37). Using this probe, we have conducted optical and EPR

Although the details of the catalytic mechanism remain to be elucidated, it is generally believed to involve an enzyme-bound substrate radical that is generated via hydrogen abstraction by an activated iron–oxygen species. As depicted in Scheme 7, the initial steps involve substrate binding and activation of dioxygen by the Fe(II) center, which in the presence of exogenous electron(s) from NADH, can produce various oxidative iron–oxygen intermediates, such as Fe(III)–superoxo (**35**), Fe(III)–hydroperoxo (**36**), or Fe(IV)-oxo (**37**). One of these intermediates is expected to be responsible for the abstraction of a C1 hydrogen atom from enzyme-bound **2-S** (in **34**) to generate a substrate radical intermediate (**38**, **39**, or **40**). Radical induced homolytic cleavage of the Fe–O bond in this intermediate will produce **1** and also regenerate the iron core.

The direct coordination between active site iron of HppE and the hydroxyl oxygen of **2-S**, or **2-R**, is likely important to HppE catalysis by facilitating a fast electron transfer from the substrate radical intermediate to the iron center. Given that epoxide **1** and ketone **4** are exclusively produced from **2-S** and **2-R**, respectively (**16**), a rapid electron-transfer process may be necessary to shorten the lifetime of the radical intermediates and thus prevent potential radical rearrangements. Moreover, a fast electron transfer through the bond connecting the iron center and the hydroxyl oxygen of **2-S** may be the key in determining the reaction course toward epoxidation. In the absence of the above Fe–O bond between enzyme and substrate, reaction may proceed to give hydroxylation product, since epoxidation in this case would require the electron be transferred through space, a much slower process.

Conclusion. This study has established a clear picture for the binding of substrates and substrate analogues to HppE in the presence of dioxygen analogue, nitric oxide. By using isotope-labeled substrate, the bidentate mode of substrate binding to the active site iron in the solution state has been fully established. The further characterization of HppE activity using chemically synthesized mechanistic probes has also shown that the binding of the hydroxyl group of the substrates determines the site of oxidation. The combined bioorganic and EPR spectroscopic characterizations have resulted in more detailed understanding of the enzyme–substrate interactions and insight into the stereospecificity of the enzyme action. This study has also provided support for the previously proposed catalytic mechanism of HppE.

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