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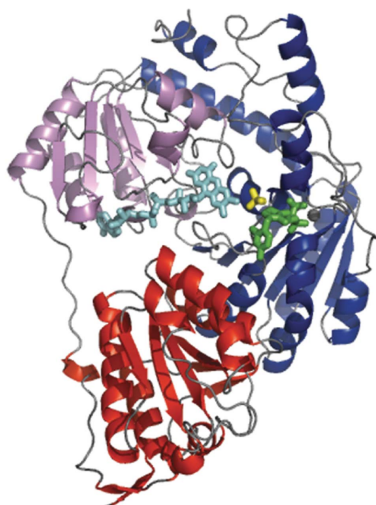
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## The structures of pyruvate oxidase from *Aerococcus viridans* with cofactors and with a reaction intermediate reveal the flexibility of the active-site tunnel for catalysis

The crystal structures of pyruvate oxidase from *Aerococcus viridans* (AvPOX) complexed with flavin adenine dinucleotide (FAD), with FAD and thiamine diphosphate (ThDP) and with FAD and the 2-acetyl-ThDP intermediate (AcThDP) have been determined at 1.6, 1.8 and 1.9 Å resolution, respectively. Each subunit of the homotetrameric AvPOX enzyme consists of three domains, as observed in other ThDP-dependent enzymes. FAD is bound within one subunit in the elongated conformation and with the flavin moiety being planar in the oxidized form, while ThDP is bound in a conserved V-conformation at the subunit–subunit interface. The structures reveal flexible regions in the active-site tunnel which may undergo conformational changes to allow the entrance of the substrates and the exit of the reaction products. Of particular interest is the role of Lys478, the side chain of which may be bent or extended depending on the stage of catalysis. The structures also provide insight into the routes for electron transfer to FAD and the involvement of active-site residues in the catalysis of pyruvate to its products.

### 1. Introduction

Thiamine diphosphate (ThDP) dependent enzymes are present in various organisms and participate in diverse metabolic reactions. On the basis of the reactions they catalyze, ThDP-dependent enzymes have been classified into five families, which have been named after representative enzymes (Duggleby, 2006). The largest family is the pyruvate oxidases (POX) and is comprised of 2-ketoacid decarboxylating enzymes such as pyruvate decarboxylase (PDC), aceto-hydroxyacid synthase (AHAS), acetolactate synthase (ALS), *N*<sup>2</sup>-(2-carboxyethyl)arginine synthase (CEAS), benzaldehyde aldolase (BAL), benzoylformate decarboxylase (BFDC), oxalyl-CoA-decarboxylase (OCDL) and indolepyruvate decarboxylase (IPDC). Recent studies on the POX family have provided valuable information on the mechanism of action of these enzymes and the catalytic role of ThDP. However, important details such as those pertaining to the changes in the active site and the involvement of the intermediates in the reaction pathway remain ambiguous.

POX is a unique enzyme in the sense that it utilizes not only ThDP but also the cofactor flavin adenine dinucleotide (FAD) for the oxidative decarboxylation of pyruvate. In the presence of oxygen and inorganic phosphate, the end products of the reaction are carbon dioxide, hydrogen peroxide and the high-energy metabolite acetyl phosphate. Owing to its attractive substrate and cofactors, POX is employed in a range of biosensors. For example, POX-based biosensors are being developed for the rapid detection of phosphate, which is detrimental to aquatic organisms (Mak *et al.*, 2003), and are also being used in medical diagnosis to detect pyruvate (Zapata-Bacri & Burstein, 1988; Gavalas & Chaniotakis, 2000), the levels of which fluctuate in serum as a result of heart malfunction or poisoning.

At present, the POX-mediated reaction is believed to follow the pathway described by Tittmann and coworkers (Tittmann *et al.*, 2005; Wille *et al.*, 2006). It begins with the abstraction of a proton from the C2 atom of ThDP, which is facilitated by the 4'-amino group of ThDP.

The resulting ylide attacks the carbonyl group of pyruvate to form the tetrahedral intermediate 2-lactyl-ThDP (LThDP). Decarboxylation results in the generation of the  $\alpha$ -carbanion/enamine forms of 2-hydroxyethyl-ThDP (HETHDP). Through electron transfer to FAD, the enamine form is converted to a HETHDP radical and then to an anion-HETHDP adduct or to 2-acetyl-ThDP (AcThDP). The former is formed in the presence of phosphate and is cleaved to acetyl phosphate and ThDP.

Here, we describe the crystal structures of *Aerococcus viridans* pyruvate oxidase (AvPOX) with FAD, with both FAD and ThDP and with FAD and the AcThDP intermediate. Combined with the previously determined structures of *Lactobacillus plantarum* POX (LpPOX) in complex with LThDP, HETHDP and AcThDP (Muller & Schulz, 1993; Muller *et al.*, 1994; Wille *et al.*, 2005, 2006), our structures provide snapshots of different steps in the oxidative decarboxylation of pyruvate. Although most of the interactions of the cofactors and intermediates are similar in AvPOX and LpPOX, some notable changes were found in the present structures. The flavin moiety of FAD is planar in AvPOX, while it is bent in LpPOX. Regions close to the active site are flexible in the former but are rigid in the latter. Such regions may undergo considerable conformational changes to facilitate movement of the substrates and reaction products.

## 2. Materials and methods

### 2.1. Enzyme expression and purification

A plasmid containing the AvPOX gene was introduced into the expression host cell *Escherichia coli* DH1. Cells were grown at 310 K in BHI medium and were collected by centrifugation at 7000g for 10 min. Pellets were suspended in 10 mM phosphate buffer pH 6.5 containing 0.2% lysozyme. The suspension was incubated at 310 K for 2 h with continuous stirring. The supernatant obtained on centrifugation at 7000g for 20 min was loaded onto a DEAE-Sepharose FF column. The enzyme was eluted with a linear gradient of 0–0.5 M KCl in 10 mM phosphate buffer pH 6.5. Ammonium sulfate (13%) was added to the pooled enzyme fractions. The fractions were applied onto a Phenyl-Sepharose column and the enzyme was eluted with a linear gradient of 13–0% ammonium sulfate in 10 mM phosphate buffer pH 6.5. The fractions containing the enzyme were combined and concentrated by membrane filtration (Microcon YM3).

### 2.2. Crystallization

Lyophilized AvPOX sample was dissolved to a concentration of 40 mg ml<sup>-1</sup> in 20 mM sodium phosphate buffer pH 7.0. Crystallizations were performed using the hanging-drop vapour-diffusion method at 293 K. For the AvPOX–FAD crystals, the droplets consisted of equal volumes (5  $\mu$ l) of protein solution and reservoir solution containing 2.0 M ammonium sulfate in 20 mM sodium phosphate buffer pH 7.0. The AvPOX–FAD–ThDP crystals were obtained from similar droplets with the addition of 8 mM ThDP and 20 mM MgCl<sub>2</sub> to the above conditions. The AvPOX–FAD–ThDP–Pyr crystals were prepared by soaking the above AvPOX–FAD–ThDP crystals in a solution containing 1.0 M pyruvate and 2.0 M ammonium sulfate for 3 h. Other crystals were obtained under conditions similar to those used for growing AvPOX–FAD–ThDP but with ammonium phosphate instead of ammonium sulfate. However, when these crystals were soaked in a solution containing 1.0 M pyruvate and 2.0 M ammonium phosphate they immediately dissolved. Other attempts to soak these crystals in solutions containing lower

**Table 1**

Crystal data and statistics of data collection and refinement.

Values in parentheses are for the outer resolution shell.

	AvPOX–FAD	AvPOX–FAD–ThDP	AvPOX–FAD–ThDP–Pyr
Crystal data			
Space group	<i>I</i> 222	<i>I</i> 222	<i>I</i> 222
Unit-cell parameters (Å)			
<i>a</i>	77.6	78.1	78.0
<i>b</i>	105.3	106.1	105.5
<i>c</i>	155.3	155.4	155.9
<i>Z</i> †	1	1	1
Data collection			
Maximum resolution (Å)	1.58	1.80	1.95
Outer shell resolution range (Å)	1.64–1.58	1.86–1.80	2.02–1.95
No. of observed reflections	476332	284643	335885
No. of unique reflections	84186	56020	45577
Completeness (%)	95.5 (77.4)	92.5 (91.8)	98.4 (98.4)
<i>R</i> <sub>merge</sub> ‡ (%)	8.2 (37.9)	5.4 (33.4)	8.1 (28.4)
<i>I</i> σ( <i>I</i> )	21.8 (2.0)	11.0 (1.8)	6.8 (1.9)
Refinement			
Resolution range (Å)	6.0–1.6	10–1.8	10–2.0
Protein atoms	4598	4590	4590
Ligand molecules			
FAD	1	1	1
ThDP	—	1	—
AcThDP	—	—	1
Mg <sup>2+</sup>	—	1	1
SO <sub>4</sub> <sup>2-</sup>	4	4	—
Water	692	563	689
<i>R</i> factor§ (%)	18.3	16.3	19.5
<i>R</i> <sub>free</sub> ¶ (%)	20.6	19.6	24.3

† Number of subunits in the asymmetric unit. ‡  $R_{\text{merge}} = 100 \times \sum_{\mathbf{h}} |I_{\mathbf{h}} - \langle I_{\mathbf{h}} \rangle| / \sum_{\mathbf{h}} I_{\mathbf{h}}$ , where  $I_{\mathbf{h}}$  is the *j*th measurement of the intensity of reflection **h** and  $\langle I_{\mathbf{h}} \rangle$  is its mean value. §  $R$  factor =  $100 \times \sum ||F_o| - |F_c|| / \sum |F_o|$ , where  $|F_o|$  and  $|F_c|$  are the observed and calculated structure-factor amplitudes, respectively. ¶ Calculated using a random set containing 10% of observations that were not included throughout refinement (Brünger, 1992).

concentrations of pyruvate and ammonium phosphate resulted in the same dissolution of the crystals.

### 2.3. X-ray data collection and data processing

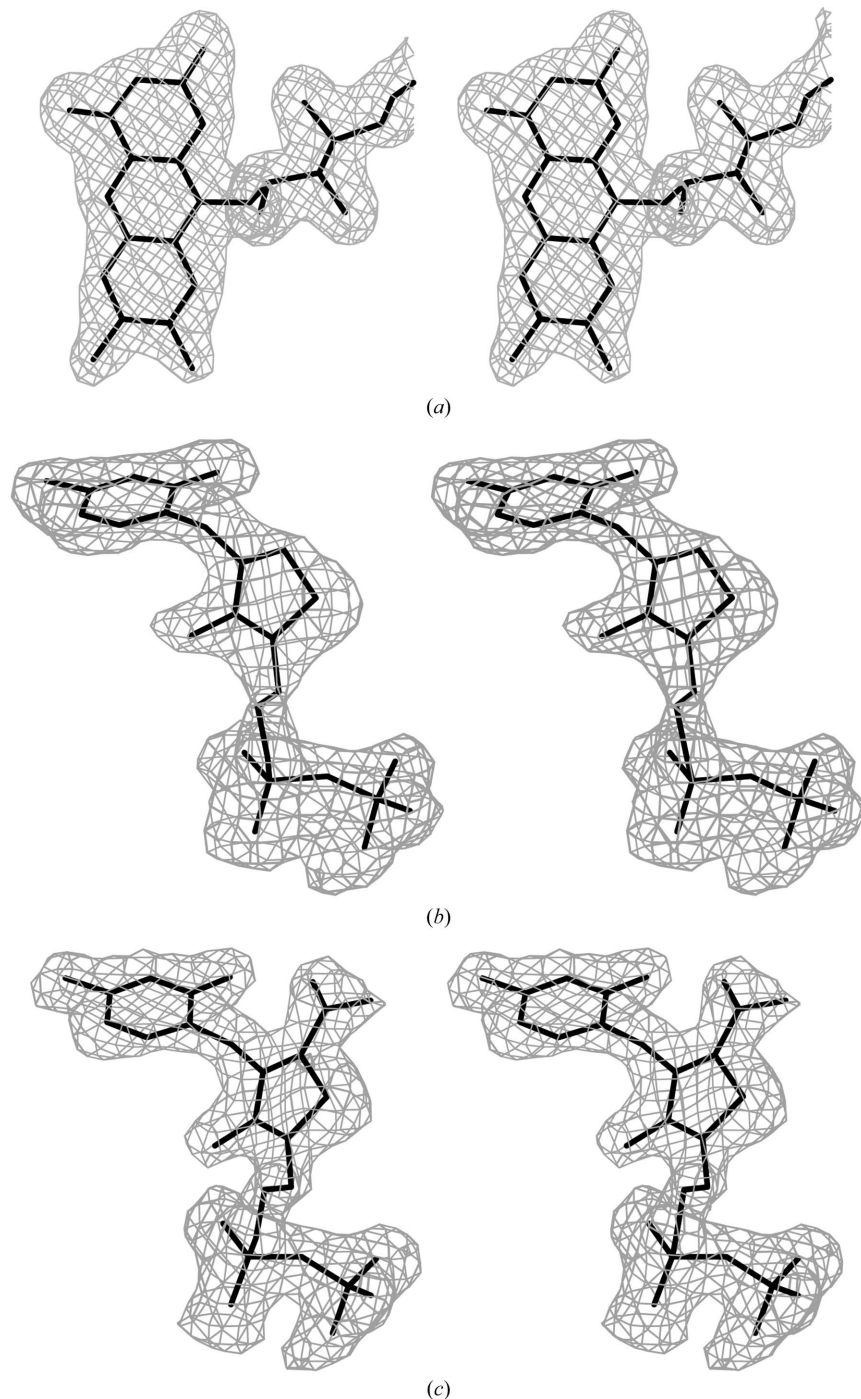
For data collection, all crystals were cryoprotected in solutions containing 2.0 M ammonium sulfate and 30% glycerol in 20 mM sodium phosphate buffer pH 7.0 and flash-frozen in liquid nitrogen. X-ray data were measured at 100 K using synchrotron radiation. For the AvPOX–FAD crystal, X-ray data were obtained at AR-NW12 ( $\lambda = 1.00$  Å) of the Photon Factory (Ibaraki, Japan) and were recorded on an ADSC Quantum 210 CCD detector positioned 140 mm from the crystal. Images were obtained using a 0.3° oscillation range with 5 s exposure. The diffraction data for the AvPOX–FAD–ThDP crystal were measured at BL44XU ( $\lambda = 0.90$  Å) of SPring-8 (Harima, Japan) in 0.5° oscillation frames with 10 s exposure times. The PX210 CCD detector was 300 mm from the crystal. The diffraction patterns of the AvPOX–FAD and AvPOX–FAD–ThDP crystals were indexed, merged and scaled with *HKL-2000* (Otwinowski & Minor, 1997). Conversion of these data to structure-factor amplitudes was performed with *TRUNCATE* from the *CCP4* suite (Collaborative Computational Project, Number 4, 1994). Data collection from the AvPOX–FAD–ThDP–Pyr crystal was carried out at BL18B ( $\lambda = 1.00$  Å) at Photon Factory. X-ray data were measured with an ADSC Quantum 4 CCD detector using 0.5° frames and 60 s exposure and with the detector 160 mm from the crystal. Diffraction patterns were processed with the program *CrystalClear* (Rigaku/MSC; Pflugrath, 1999). Intensity data were put on a relative scale and merged into independent reciprocal space using the programs

*SCALA* and *TRUNCATE* in *CCP4*. The crystal data and statistics of data collection are summarized in Table 1.

## 2.4. Structure determination and refinement

Initial phases were derived by molecular replacement with the program *AMoRe* (Navaza, 1994) using the structure of the wild-type *LpPOX* structure (PDB code 1pox; Muller *et al.*, 1994) as a search model. The structure was constructed on electron-density maps displayed in *QUANTA* (Accelrys Inc.) After rigid-body refinement, the atomic parameters were refined by the restrained maximum-

likelihood least-squares technique in *REFMAC5* from *CCP4* (Murshudov *et al.*, 1997). The structure was revised by interpreting omit maps at every residue. Sulfate and water molecules found in  $|F_o| - |F_c|$  maps ( $>2.5\sigma$ ) were included in the subsequent structure refinements. The final refinements were performed with the program *CNS* (Brünger *et al.*, 1998). Electron density for FAD, ThDP and a ThDP intermediate were observed in the active sites. These ligands were introduced into the models and were included in the structure refinements. In the final refinements, the geometrical restraints imposed on the ligands were completely released to examine their deformability.



**Figure 1**

Stereoviews of omit maps contoured at the  $2.5\sigma$  level for (a) the flavin moiety of FAD, (b) ThDP and (c) AcThDP bound in the *A<sub>v</sub>POX*-FAD, *A<sub>v</sub>POX*-FAD-ThDP and *A<sub>v</sub>POX*-FAD-ThDP-Pyr structures, respectively.

The statistics of refinement for all three crystals are given in Table 1. Fig. 1 shows omit maps of the ligands, which were drawn with the program *O* (Jones *et al.*, 1991). The figures showing the structural details were drawn with *PyMOL* (DeLano Scientific; <http://www.pymol.org>). Structural comparisons of the present structures with those of *Zymomonas mobilis* PDC (ZmPDC; Dobritzsch *et al.*, 1998), yeast AHAS (Pang *et al.*, 2002), *Klebsiella pneumoniae* ALS (Pang *et al.*, 2004), *Streptomyces clavuligerus* CEAS (Caines *et al.*, 2004), *Pseudomonas fluorescens* BAL (Mosbacher *et al.*, 2005), *P. putida* BFDC (Hasson *et al.*, 1998), *Oxalobacter formigenes* OCDC (Berthold *et al.*, 2005) and *Enterobacter cloacae* IPDC (Schütz *et al.*, 2003) were carried out using *Swiss-PdbViewer* v.3.7 (<http://www.expasy.org/spdbv/>).

### 3. Results

#### 3.1. Overall structures

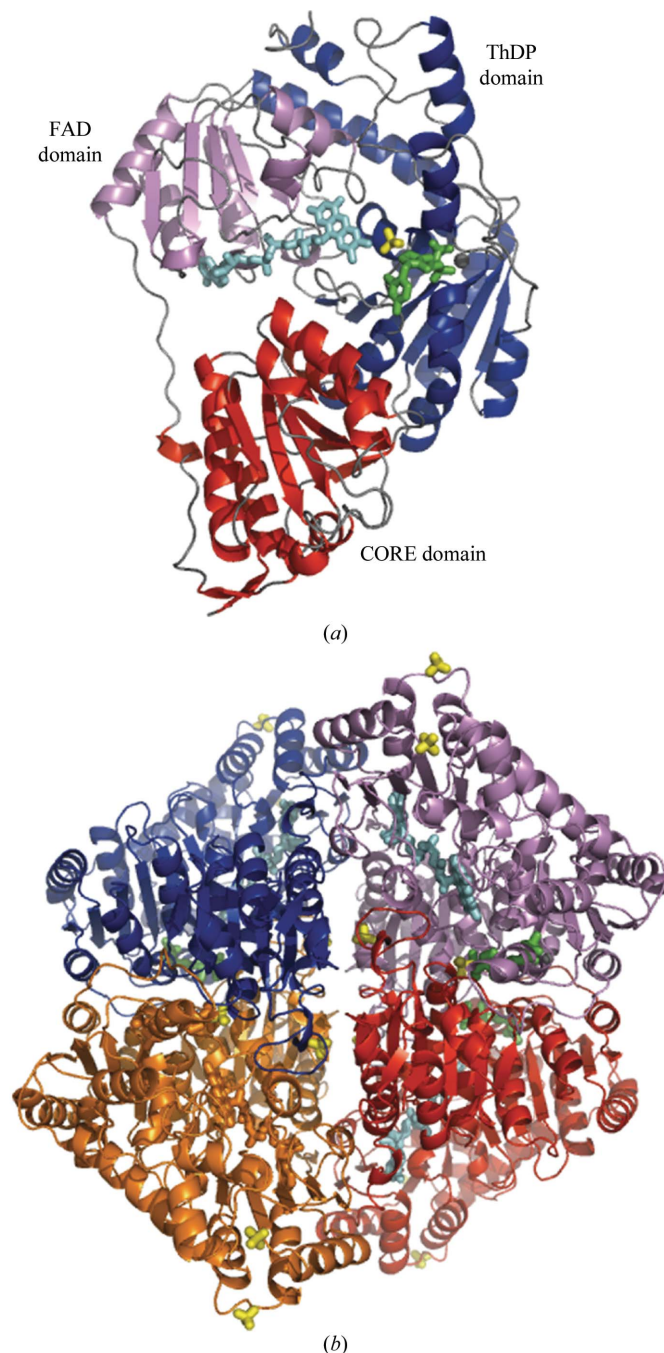
The *Av*POX monomer consists of three regions, denoted CORE (Asn4–Lys186), FAD (Tyr187–Glu337) and ThDP (Ser338–Lys592) domains (Fig. 2*a*). The core scaffold of each subunit is comprised of six parallel  $\beta$ -sheets, which are surrounded by  $\alpha$ -helices. Two subunits of the dimer are related by crystallographic twofold symmetry. The active site is formed by residues from the ThDP domain of one subunit and from the CORE domain of the neighbouring subunit. Another dimer is related by further crystallographic twofold symmetry; thus, the complete tetramer has 222 symmetry (Fig. 2*b*). These features are maintained in all three crystal structures. Superimpositions of corresponding C $\alpha$  atoms among the three structures of *Av*POX gave r.m.s.d. values ranging from 0.26 to 0.39 Å, suggesting that binding of FAD, ThDP and the ThDP intermediate has little effect on the overall protein conformation. Except for the three N-terminal residues, some side chains at the surface of the protein and several residues in a loop region (Asp469–Tyr496) near the active site, the electron densities were well defined over the entire structures of the three *Av*POX structures.

#### 3.2. ThDP binding

In the *Av*POX–FAD–ThDP structure, the ThDP cofactor is bound in a cleft between the ThDP domain of one subunit and the CORE domain of the adjacent subunit, although most of the contacts are made with the ThDP domain (Fig. 3*a*). The diphosphate moiety of ThDP forms hydrogen bonds with the main-chain N atoms of Asn391, Ser392, Gly443, Ala444 and Phe474 and with the side-chain atoms of Ser392, Asp442 and Asn469. The diphosphate terminus is anchored by Mg<sup>2+</sup>, which is octahedrally coordinated by the main-chain O atom of Glu471, the side-chain atoms of Asp442 and Asn469 and a water molecule (Fig. 3*a*). Several of the residues interacting with the diphosphate group are from the <sup>441</sup>GDGX<sub>24</sub>NN<sup>469</sup> motif, which is found in all ThDP-dependent enzymes (Hawkins *et al.*, 1989). In *Av*POX, however, the second-to-last residue of the motif is replaced by serine. The pyrimidine ring forms two hydrogen bonds with the enzyme: N1' to Glu54\* OE2 (where the asterisk denotes a residue in the adjacent subunit) and N4' to Ala415 O. The side chain of Met417 protrudes from the surface, forcing the thiazolium and pyrimidine rings to adopt a V-shaped conformation (the torsional angles  $\varphi_T$  and  $\varphi_P$  are 92° and –62°, respectively). This V-conformation results in the distance between the 4'-amino nitrogen and C2 atoms being 3.2 Å.

#### 3.3. AcThDP binding

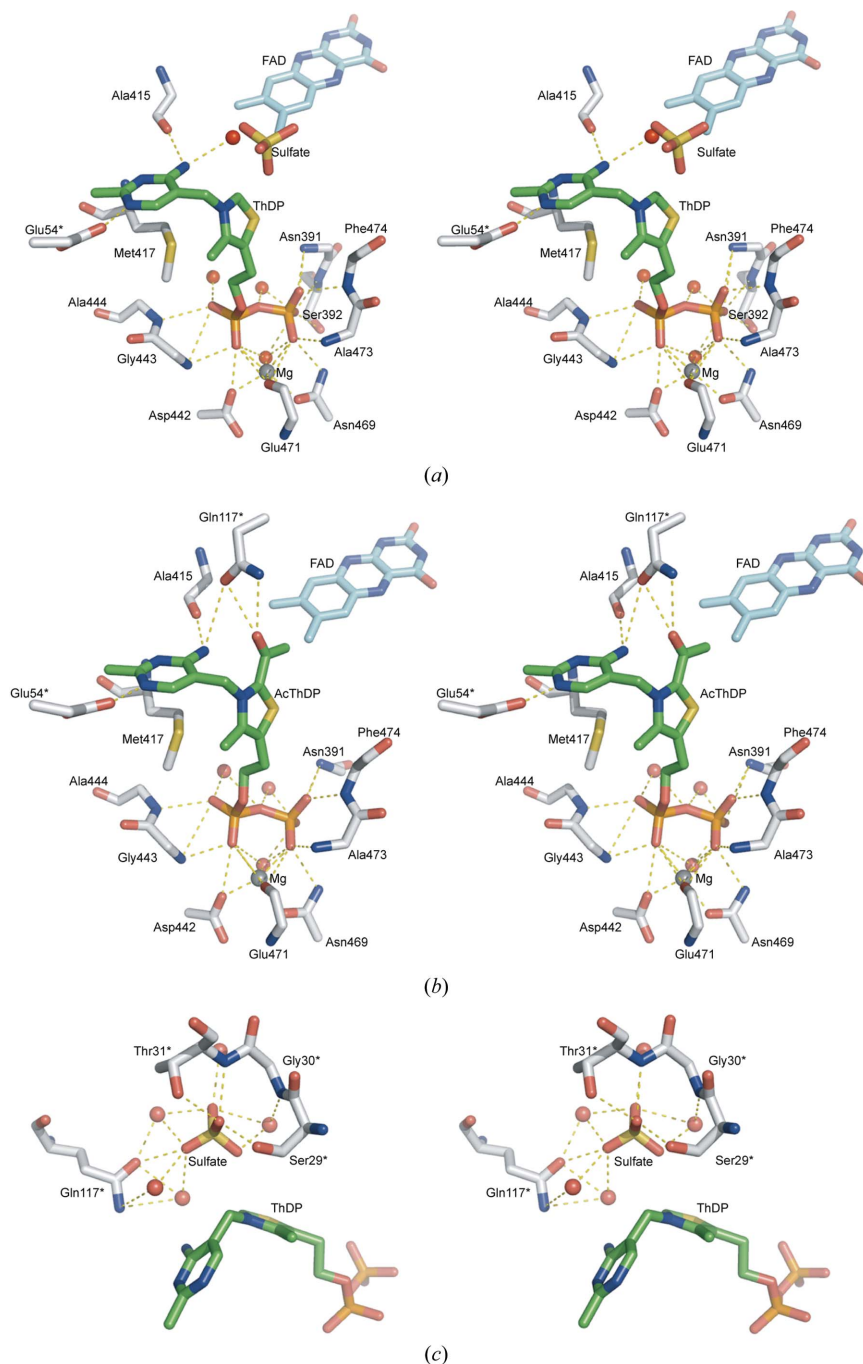
In the initial  $|F_o| - |F_c|$  electron-density map of *Av*POX–FAD–ThDP–Pyr calculated with the phase angles derived from the model of *Av*POX–FAD–ThDP, electron densities were found extending from the C2 atom of ThDP into the active site, indicating the formation of a ThDP intermediate. A well defined density for the cofactor intermediate is also observed in the final omit electron-density map (Fig. 1*c*), which shows that the five atoms of the thia-



**Figure 2**  
The overall structure of *Av*POX–FAD–ThDP. (*a*) The monomeric unit of *Av*POX–FAD–ThDP showing the CORE (red), FAD (violet) and ThDP (blue) domains. (*b*) The tetrahedral *Av*POX–FAD–ThDP homotetramer with 222 symmetry. Subunits A, B, C and D are coloured red, blue, violet and orange, respectively. FAD, ThDP and sulfate ions are shown as cyan, green and yellow stick representations, respectively, while the magnesium ions are shown as grey spheres.

zolium ring, as well as the atoms of the extension, are all in the same plane. The geometrical restraints on this part were released in the final stages of refinement, but the atomic arrangement was not changed, reconfirming the planarity. The intermediate could thus either be the HThDP enamine or AcThDP. We assigned the trapped intermediate as AcThDP. It has been reported that in the absence of phosphate, AcThDP is the predominant intermediate. The crystallization solution for *A<sub>v</sub>*POX–FAD–ThDP–Pyr contained only a very small amount of phosphate, which may have been removed after the long soak in a solution containing no phosphate.

AcThDP is bound to the enzyme in an identical fashion to ThDP and the interactions of the diphosphate moiety with the enzyme and the Mg<sup>2+</sup> ion are similar to those observed in the *A<sub>v</sub>*POX–FAD–ThDP structure (Fig. 3*b*). The interactions with one of the phosphate groups are changed because the side chain of the Ser392 residue is shifted away from the phosphate by about 3 Å. The thiazolium and pyrimidine rings of AcThDP also adopt the V-conformation ( $\varphi_T = 94^\circ$  and  $\varphi_P = -65^\circ$ ). The carbonyl oxygen of the acetyl moiety forms hydrogen bonds to the side-chain atoms of the Gln117\* residue of the neighbouring subunit. The position of the Gln117\* residue is the



**Figure 3** Stereoviews of the active sites, showing the interactions involving (a) ThDP and magnesium in *A<sub>v</sub>*POX–FAD–ThDP, (b) AcThDP and magnesium in *A<sub>v</sub>*POX–FAD–ThDP–Pyr and (c) sulfate in *A<sub>v</sub>*POX–FAD–ThDP. The FAD, ThDP/AcThDP and sulfate molecules are shown as cyan, green and yellow stick representations, respectively. The magnesium ions and the water molecules are shown as grey and red spheres, respectively. All residues marked with an asterisk belong to the symmetry-related subunit. The dashed lines indicate hydrogen-bonding and ionic interactions.

same in the other two *Av*POX structures. Four residues, Ser29\*, Gly30\*, Ser77\* and Gly78\*, interact indirectly with the acetyl group of the intermediate by hydrogen bonding through water molecules. The methyl group of the acetyl moiety makes a van der Waals contact with the C7 methyl group of FAD.

### 3.4. FAD conformation

The cofactor FAD is bound in a cleft surrounded by all three domains in the subunit and adopts the elongated conformation. There are numerous hydrogen-bond formations and van der Waals interactions between the enzyme and FAD. Adenine is bound tightly to the enzyme by hydrogen bonds to the main-chain N atom of Ala321 and the side-chain atom of Asp320. The ribose ring is held in place by hydrogen bonds to the side-chain atoms of Asp301. The phosphate groups of FAD form hydrogen bonds to the main-chain atoms of Ile216 and Asn281 and the side-chain atoms of Thr239, Lys241 and Ser280. The flavin moiety also forms several hydrogen bonds to the enzyme; these contacts are to the main-chain atoms of Thr257, Val260 and Pro412. In the oxidized form of *Lp*POX, the flavin moiety is bent about the N5–N10 axis (Muller & Schulz, 1993; Muller *et al.*, 1994; Wille *et al.*, 2005, 2006). However, in the present oxidized *Av*POX (as evidenced by the bright yellow colour of the crystals), the flavin is planar. This supports the assumption that the conformation of the flavin moiety does not affect its redox potential (Wille *et al.*, 2005).

### 3.5. Sulfate-binding sites

Four sulfate anions are also observed in the *Av*POX–FAD and *Av*POX–FAD–ThDP crystals. In *Av*POX–FAD–ThDP, a sulfate anion is located near the thiazolium ring of ThDP (Fig. 3c). The sulfate anion is bound to the Ser29\*, Thr31\* and Gln117\* residues of a neighbouring subunit. The anion also interacts indirectly with Gly30\* through water mediation. One of the sulfate O atoms interacts with the C2 atom of ThDP. The three other sulfate anions in *Av*POX–FAD–ThDP are located in equivalent positions in *Av*POX–FAD. Two of these common sulfate-binding sites are on the surface of the tetramer, while the remaining common binding site is in a crevice surrounded by three subunits. The sulfate anion present only in *Av*POX–FAD is in a location similar to that of one of the phosphate groups of ThDP in *Av*POX–FAD–ThDP.

**Table 2**

Comparisons of the sequences, structures and flexible regions of the POX family of enzymes.

The *A* subunits from the PDB files were used in all cases.

Protein	Identity (%)	Similarity (%)	R.m.s.d.† (Å)	Region II residues
<i>Av</i> POX (1v5f)	—	—	—	469–496‡
<i>Lp</i> POX (1pox)	47	66	0.96 (568)	474–501
<i>Zm</i> PDC (1zpd)	26	46	1.55 (338)	467–488
AHAS (1jsc)	24	45	1.44 (316)	577–604‡
ALS (1ozf)	23	44	1.77 (322)	474–500
CEAS (1upb)	22	42	1.51 (309)	490–518
BAL (2ag0)	22	40	1.55 (337)	475–503
BFDC (1bfd)	21	40	1.52 (348)	455–482
OCDC (2c31)	21	38	1.63 (351)	479–504
IPDC (1ovm)	21	38	1.62 (293)	462–498

† Calculated by superimposing the corresponding C $\alpha$  atoms. Values in parentheses indicate the number of aligned C $\alpha$  atoms. ‡ Contain disordered regions.

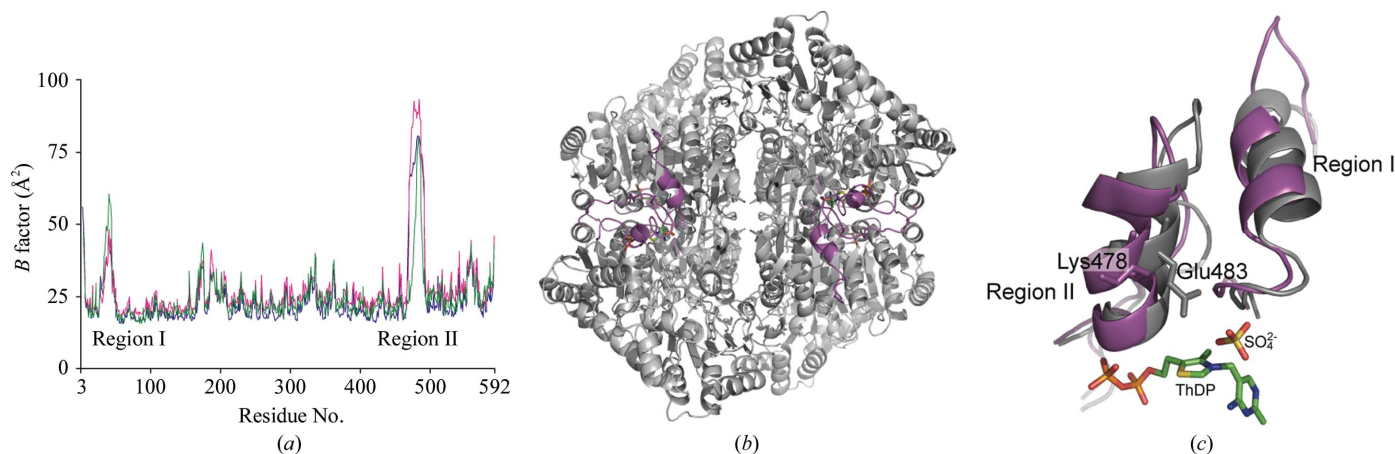
### 3.6. Flexible regions

In general, temperature factors (*B* factors) characterize the conformational flexibility of amino-acid residues. Fig. 4(a) shows plots of the observed *B* factors versus the C $\alpha$  atoms of the residues in the three *Av*POX structures. The graph demonstrates a high degree of flexibility of two regions near the active site. The first region (region I; residues 27–46) is located near the N-terminus, while the second (region II; residues 469–496) includes residues interacting with the diphosphate moiety of ThDP. Examination of the electron densities in these regions revealed that while the residues in region I fit reasonably well into the densities, those in region II are disordered.

## 4. Discussion

### 4.1. Comparison with other POX-family enzymes

Superimpositions of the 570 corresponding C $\alpha$  atoms between the *Lp*POX structure (Muller *et al.*, 1994) and any of the three *Av*POX structures show similarities in the overall main-chain folding, with r.m.s.d. values ranging from 0.82 to 0.90 Å. In the *Av*POX–FAD–ThDP structure, the enzyme is in a state where it is waiting for the substrate pyruvate. The enzyme in this state was compared with related enzymes belonging to the POX family (Table 2). Although AHAS from yeast has a low sequence identity (24%) to *Av*POX, the



**Figure 4**

Flexible regions in *Av*POX. (a) Average *B* factors of the main-chain atoms are shown for *Av*POX–FAD (blue), *Av*POX–FAD–ThDP (magenta) and *Av*POX–FAD–ThDP–Pyr (green). (b) A top view of the tunnels for substrate entry to the active site in *Av*POX–FAD–ThDP. The locations of the flexible regions are coloured in magenta. (c) Side view of the tunnel with the flexible regions in *Av*POX–FAD–ThDP (magenta) superimposed onto the corresponding regions in *Lp*POX (grey; Wille *et al.*, 2006). ThDP, FAD, sulfate, Lys478 and Glu483 are shown as stick representations. Lys478 in *Av*POX corresponds to Glu483 in *Lp*POX.

326 corresponding C $\alpha$  atoms superimposed with an r.m.s.d. of 1.44 Å (Pang *et al.*, 2002). Both POX and AHAS have an essential requirement for the cofactor FAD, although FAD lacks the redox function in the latter enzyme. All other POX-family enzymes showed a good equivalence, with deviations of less than 2 Å for the corresponding C $\alpha$  atoms. The positions of the residues in the FAD domain of AvPOX correlate particularly well with those of the residues in the middle domain (also referred to as the  $\beta$  or R domains) of the other enzymes.

#### 4.2. Role of the flexible regions in catalysis

The two flexible regions occupy the walls of the tunnel leading to the active site (Fig. 4*b*). Both AHAS and OXC were reported to have disordered residues in the same region (Pang *et al.*, 2002; Berthold *et al.*, 2005). In the structures of AHAS in complex with various herbicides, these regions become ordered and close the active site (McCourt *et al.*, 2005), whereas in the OXC structure they are already folded up. In the remaining POX-family enzymes, however, the residues corresponding to these flexible regions have very low *B* factors, indicating stability. Closer examination of the present AvPOX structures and the LpPOX structures revealed several differences in the conformations of the flexible regions (Muller & Schulz, 1993; Muller *et al.*, 1994; Wille *et al.*, 2005, 2006). In the LpPOX structures these regions are more compact compared with those in AvPOX (Fig. 4*c*). This is a consequence of more extensive contacts in the residues occupying the base of the flexible regions in LpPOX; such contacts may lead to their stability. Another stabilizing factor in LpPOX is the additional contact between the Glu483 side chain and the thiazolium moiety of ThDP (or the extension from the thiazolium moiety of the ThDP intermediates). In the LpPOX structures, Glu483 adopts an extended conformation and appears to close the active site. In AvPOX, the main chain of the Lys478 residue occupies a position similar to that of LpPOX Glu483, but its side chain is bent and as such is incapable of forming interactions with ThDP or AcThDP. Lys478 may adopt the ‘open’ or ‘closed’ conformations, acting as a lid to allow the reaction to proceed in the case of the former or to stop in the case of the latter. In a study performed on ZmPDC variants, it was shown that mutation of Glu473, the residue

that corresponds to AvPOX Lys478, slows covalent addition of pyruvate and decarboxylation of LThDP (Tittmann *et al.*, 2003). Further mutation studies on these flexible regions in AvPOX, especially on Lys478, are necessary to determine their roles in pyruvate catalysis.

#### 4.3. Catalytic mechanism

The role of the pyrimidine ring in the reactions of ThDP-dependent enzymes has been widely studied for several decades (Jordan & Mariam, 1978; Nemeria *et al.*, 2007). As shown in Fig. 5, the N1' atom is hydrogen bonded to the carboxyl group of the conserved Glu54\* residue and such an interaction could induce proton release from the N4' amino group. The resulting N4' imino group then forms a hydrogen bond with the carbonyl group of the Ala415 residue. This triggers the transfer of protons from the thiazolium C2 atom to the N4' atom and from the N1' atom to Glu54\*. The C2 atom becomes highly nucleophilic and is favourable for attack by the methoxyl group of pyruvate. This substrate is allowed to enter the active-site tunnel by movements of the flexible regions, particularly of the Lys478 side chain. As a result, the LThDP intermediate is formed, the structure of which was recently reported (Wille *et al.*, 2006). Using the present AvPOX structures, a model of LThDP and several active-site residues in AvPOX was successfully built. The model shows that Gly30\* and Gln117\* may form interactions with LThDP. Lys478, adopting the extended conformation, has its side-chain amide group in a position that is also favourable for interaction with LThDP.

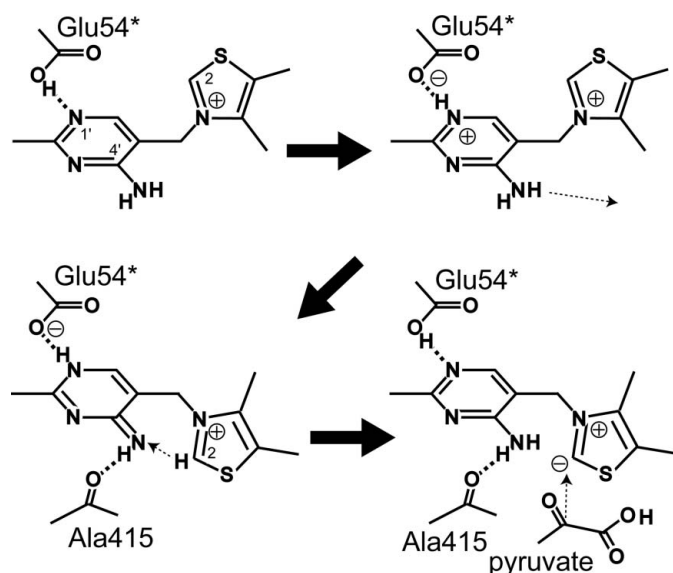
Decarboxylation of LThDP occurs and the resulting CO<sub>2</sub> is expelled from the active site, possibly with Lys478 adopting the open conformation. The  $\alpha$ -carbanion/enamine forms of HETHDP are formed, coupled with the transfer of an electron to FAD. Assuming that the hydroxyethyl moiety of HETHDP occupies the same position as the acetyl moiety of the trapped AcThDP in the AvPOX-FAD-ThDP-Pyr structure, electron transfer could proceed *via* three routes, consistent with previous suggestions (Muller & Schulz, 1993; Muller *et al.*, 1994; Pang *et al.*, 2002). Firstly, the methyl group of the hydroxyethyl moiety of HETHDP may directly transfer an electron to the methyl group on the C7 atom of the flavin. Secondly, an electron may be passed *via* Phe116\* to C7. Thirdly, an electron may be relayed to the methyl group on the C7 of flavin *via* Phe474.

Once again, the active-site lid opens for the phosphate to enter, which may occupy the sulfate-binding site near the thiazolium ring found in the AvPOX-FAD-ThDP structure. The phosphate may also form contacts with Ser29\*, Gly30\*, Thr31\* and Gln117\*, as well as with Lys478 in the extended form. Similar contacts between phosphate and LpPOX have been reported (Wille *et al.*, 2006). The phosphate attacks HETHDP and the anionic phosphate-radical adduct is formed, which subsequently facilitates the transfer of the second electron to FAD. The final products, acetyl phosphate and hydrogen peroxide, are then released to regenerate the ThDP cofactor.

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**Figure 5**  
Proton movements upon ThDP binding to facilitate pyruvate attack.

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