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Structural evidence for a reaction intermediate mimic in the active site of a sulfite dehydrogenase†

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By combining X-ray crystallography, electron paramagnetic resonance techniques and density functional theory-based modelling, we provide evidence for a direct coordination of the product analogue, phosphate, to the molybdenum active site of a sulfite dehydrogenase. This interaction is mimicking the still experimentally uncharacterized reaction intermediate proposed to arise during the catalytic cycle of this class of enzymes. This work opens new perspectives for further deciphering the reaction mechanism of this nearly ubiquitous class of oxidoreductases.

Eukaryotic sulfite oxidases (SOs) and homologous prokaryotic sulfite dehydrogenases (SDHs) are mononuclear molybdoenzymes that catalyze sulfite oxidation by direct oxygen atom transfer from water to the substrate: \( \text{SO}_3^{2-} + \text{H}_2\text{O} \rightarrow \text{SO}_4^{2-} + 2\text{H}^+ + 2\text{e}^- \). They play a key role in sulfur detoxification, notably in the final step of the degradation pathway of sulfur-containing amino acids. In humans, SO deficiency is a genetic disease that causes severe neonatal neurological problems and often early death. In bacteria, SDHs have a central role in converting \( \text{SO}_3^{2-} \) formed during dissimilatory oxidation of reduced sulfur compounds, and are generally linked to energy conservation. In contrast to SOs, SDHs are unable to use \( \text{O}_2 \) as a direct electron acceptor. \( \text{SO}_3^{2-} \) oxidation in these Sulfite Oxidizing Enzymes (SOEs) occurs at their conserved Mo center which cycles between the Mo(IV), Mo(V) and Mo(VI) states (Fig. S1, ESI†). The combined use of X-ray crystallography, extended X-ray absorption fine structure (EXAFS), various electron paramagnetic resonance (EPR) spectroscopic methods and theoretical modelling has been crucial to resolve the geometrical and electronic structure of the Mo cofactor (Moco) at the active site of SOEs.‡ The metal ion is coordinated to five ligand donor atoms in a pseudo-square pyramidal geometric arrangement: three equatorial S donor atoms (one from a conserved cysteine residue and two from a pyranopterin dithiolene cofactor), one axial oxo group and an equatorial OH or \( \text{OH}_2 \) group whose identity varies during the catalytic cycle.

Catalysis in SOEs is a complex multistep process, which involves the synchronization of various chemical events, such as electron and proton transfer and the formation of substrate and product intermediates whose properties are finely tuned by the protein environment. Characterizing them is challenging due to enzyme turnover. For instance, the formation of a covalently bound \( \text{Mo(v)}-\text{O-SO}_3^2 \) reaction intermediate has been proposed following substrate binding (Fig. S1, ESI†), which is supported by recent theoretical investigations. However, no such intermediate could be trapped so far. Three high-resolution structures of SOEs have been solved with either a \( \text{SO}_4^{2-} \) (in chicken SO (cSO) and in \textit{Starkeya novella} SDH (SorAB)), or a \( \text{SO}_3^{2-} \) (in the C185S variant of cSO) molecule close to their active site. However, in all of them, the \( \text{SO}_4^{2-} / \text{SO}_3^{2-} \) oxygen atom closest to the Mo is found at distances \( \geq 3.0 \) Å that preclude direct coordination to the Mo atom. On the other hand, evidences for direct coordination of \( \text{SO}_3^{2-} \) rather than \( \text{SO}_4^{2-} \) to the Mo atom have been provided in plant and human SOs as well as in a variant of SorAB (R55Q) through detailed EPR characterizations of non-catalytic sulfite Mo(v) species (renamed “blocked”).

In this context, structural and functional studies using substrate or product analogues are crucial for providing detailed insights into SOEs’ functioning. So far, direct binding of the product analogues \( \text{PO}_4^{3-} \) and \( \text{AsO}_4^{3-} \) to the Mo(v) ion has been evidenced in vertebrate SOs using EPR and EXAFS spectroscopies, while structural evidence is still lacking.

In this work, we show that SDH from \textit{Thermus thermophilus} (TtSDH) can serve as a model enzyme to remedy this lack of detailed structural information on the binding of the product analogue \( \text{PO}_4^{3-} \) at the Mo active site in SOEs. TtSDH is a
homodimeric enzyme containing only the Mo redox center. It interacts with a diheme cytochrome c550, which acts as physiological electron acceptor. The electrons are then shuttled to O₂ reductases through cytochrome c552, linking SO₄^{2-} oxidation to the aerobic respiration pathway. Here, TtSDH was crystallized in the presence of 1.8 M Na₃PO₄ and its structure solved by X-ray crystallography (Table S1, ESI†). Like other SOEs, the overall structure of TtSDH is organized in two domains: a catalytic domain containing the Moco active site and a dimerization domain (Fig. S2, ESI†). The structure features a well-resolved electron density in the active site of the protein consistent with the presence of the Moco. Interestingly, where the available structures of SOEs normally show an oxo-ligand coordinated to the Mo atom at the equatorial position, an extended positive tetrahedral shaped electron density was observed adjacent to the Mo atom in TtSDH (Fig. 1). This could arise either from SO₄^{2-} present in the culture media or, more likely, from the phosphate present in the crystallization solution at high concentration. As both molecules are indistinguishable based solely on the electron density map, we used EPR techniques to further assess the nature of the anion coordinated to the Mo atom.

The X-band (~9 GHz) CW EPR spectrum of a redox-poised sample of TtSDH containing 1.8 M Na₃PO₄ at pH 5.7 features an intense anisotropic signature with resolved splittings that lead to overlapping EPR lines in the high field region of the spectrum (Fig. 2). X- and Q-band (~34 GHz) (Fig. S3, ESI†) EPR spectra are well simulated using a similar set of parameters corresponding to a single rhombic species with g-values (g₁, g₂, g₃) = (1.9966, 1.9700, 1.9622) and experiencing a strong magnetic interaction with a nucleus with spin value I = 1/2. This signal is assigned to a 4d¹ Mo(V) atom in a low symmetry coordination sphere on the basis of its g-values, its relaxation properties, and the detected lower intensity hyperfine lines from the I = 5/2 ⁹⁵,⁹⁷Mo isotopes (25% natural abundance) that superimpose on the main spectral contribution. Simulations further show that the resolved splittings are well reproduced using a nearly axial hyperfine tensor with principal values (A₁, A₂, A₃) = (22, 21, 29) MHz and assuming collinear g- and hf-main axes (Fig. 2). The identity of the I = 1/2 coupled nucleus was determined using HYSCORE spectroscopy.

The (~−, +) quadrant of the Mo(V) HYSCORE spectrum measured at the g₂ field position is shown in Fig. 3. It features a pair of intense cross-peaks produced by a strongly coupled nucleus with hyperfine coupling constant A satisfying the relation A > 2νₑ (νₑ is the nuclear Zeeman frequency). The correlated nuclear frequencies with maxima around νₑ ≈ (18.5, 6.6) MHz are separated by twice the ³¹P Zeeman frequency (νₑ ≈ 6.0 MHz), and are thus assigned to a ³¹P nucleus. The shape and the position of these cross-peaks are well simulated assuming an axially symmetric ³¹P hyperfine tensor with isotropic and anisotropic components (|Aₙₑ|, |I|) = (26, 3.6) MHz, respectively. These
values are consistent with those inferred from the simulation of the cw EPR spectra (i.e. $|{A_{iso}}|$ [7]) $\approx$ (24, 2.5 MHz). This unambiguously shows that the HYSSCORE cross-peaks analyzed above and the splitting resolved on the X-band cw EPR spectrum originate from a single strongly coupled $^{31}$P nucleus, assigned to a Mo–phosphate adduct. Indeed, the magnitude of the $^{31}$P $A_{iso}$ value is larger$^{7,27–29}$ or comparable to those estimated from previous pulsed EPR experiments in vertebrate SOs and assigned to phosphate coordinated species. Distributed $^{31}$P $A_{iso}$ values ranging from $\sim$0 to $\sim$20 MHz have been estimated in cSO.$^{29}$ The absence of such a wide distribution in TtSDH, comparatively suggests a more rigid environment around the bound phosphate molecule. Eventually, similar spectra were recorded using a TtSDH sample incubated with a lower (i.e. 70 mM) Na$_3$PO$_4$ concentration as typically used in previous SOs studies (Fig. S4, ESI†).$^{21,22,27}$

Upon placement of phosphate in the electron density map and subsequent structure refinement, the product analogue clearly appears to be directly coordinated to the Mo atom, with one of its oxygens being at the position where the equatorial oxo-ligand is normally observed (Fig. 1). This phosphate oxygen is only at 2.16 Å distance from the Mo atom, which is within the known range for Mo–phosphate complexes.$^{30}$

Such a direct coordination is further supported by DFT calculations of EPR parameters performed on large (~670 atoms) molecular models of the TtSDH active site based on atomic coordinates from the phosphate-bound structure (Fig. S5, ESI†). The three geometry-optimized models obtained upon consideration of the different possible protonation states of the coordinated phosphate molecule are referred to as models 1a-$\text{[HPO}_4\text{]}^{2-}$ and 1b-$\text{[HPO}_4\text{]}^{2-}$ (with monoprotonated phosphate) and model 2-$\text{[H}_2\text{PO}_4\text{]}^-\text{[HPO}_4\text{]}^{2-}$ (with diprotonated phosphate). Models 1a and 1b correspond to two energy minima of the bound HPO$_4^{2-}$ placing the phosphate proton with two different orientations (Fig. S6A and B, ESI†). Overall, the three optimized structures are very similar to each other and only subtle structural changes are resolved compared to the original X-ray structure (Table S2, ESI†). Regarding the Mo($v$) $g$-tensor, the phosphate protonation state mainly affects the $g_i$ value which increases significantly when the substrate analogue is deprotonated (Table S3, ESI†). Best agreement between the calculated and experimentally determined $g$-values is achieved for model 1a, suggesting that the phosphate molecule is in the monoprotonated form in the conditions used for EPR analyses (Table S3, ESI†). In addition, $^{31}$P $A_{iso}$ values in the 15–40 MHz range are calculated for the three models, in agreement with the value of 24 MHz measured herein. It is the 15–40 MHz range are calculated for the three models, in agreement with the value of 24 MHz measured herein. It is

The herein characterized HPO$_4^{2-}$-bound complex is likely to mimic the covalently bound Mo($v$)–O–SO$_3$ reaction intermediate proposed to occur in the catalytic cycle of SOs after substrate binding and oxidation, providing new insights into substrate/product recognition in their active site. Indeed, HPO$_4^{2-}$ displays a similar volume and the same charge as SO$_4^{2-}$. Moreover, our modelling studies reveal that the HPO$_4^{2-}$ proton does not form any additional H-bond interaction (Fig. S5, ESI†). On the other hand, our high-resolution structure shows that the phosphate molecule is tightly bound by a dense H-bond network originating from Arg50, Arg107, Asn117, Trp119 and Tyr231 side chains, creating a favorable electrostatic environment for anion binding (Fig. 1). Two water molecules participate in this network by bridging H-bonds to Asn117 main chain and Arg230 and Tyr367 side chains. Besides their involvement in stabilizing the reaction intermediate, these H-bond interactions are predicted to contribute to lower activation barriers.$^{11,32}$

Comparison of the TtSDH structure with those of cSO and SorAB featuring SO$_4^{2-}$ molecules close to the Mo atom reveals significant differences. Indeed, in cSO, the sulfate molecule in the active site is located at 4.6 Å (nearest SO$_4^{2-}$ oxygen) to the Mo atom, which itself still remains coordinated to the equatorial oxygen ligand.$^{12}$ Replacement by Gln of Arg138, proposed as crucial for catalysis, does not affect the binding mode but increases the distance between the SO$_4^{2-}$ molecule and the Mo atom to 5.5 Å.$^{12}$ By contrast, a structure of SorAB with SO$_4^{2-}$ bound at the active site could only be obtained when the equivalent Arg residue (Arg55) was replaced by Met despite high SO$_4^{2-}$ concentration in the crystallization solution (i.e. 2.2 M).$^{13}$ In this R55M variant, the Met residue side chain has been shown to adopt a different orientation compared to the Arg residue, leaving space for the SO$_4^{2-}$ molecule to approach the Mo atom at a distance of 3.0 Å, however, without being coordinated. While no conformational change of active
site residues is observed in the R55M variant of SorAB upon SO$_4^{2-}$ binding, an Arg residue (Arg450) which is solvent exposed in the SO$_4^{2-}$-free structure of the cSO has been shown to point towards the active site in the presence of SO$_4^{2-}$. This Arg residue participates in the H-bond network of what seems to be a leaving SO$_4^{2-}$ molecule and, thus, has been proposed to play a gating role in cSO. In contrast, the side chain of Arg 365 in TtSDH, which occupies the same position as Arg 450 in cSO, is solvent exposed in our phosphate-bound structure, excluding a similar role in TtSDH for this residue. Rather, a further Arg residue in TtSDH (Arg230) is found to be oriented towards the HPO$_4^{2-}$ molecule at 4.1 Å, suggesting a role in directing the substrate/product to or from the active site. The precise role of this arginine therefore deserves to be assessed using site-directed mutagenesis.

In conclusion, by combining X-ray crystallography, CW and pulsed EPR spectroscopy and DFT modelling, we provide clear evidence for direct coordination of HPO$_4^{2-}$ to or from the active site. The precise role of this arginine therefore deserves to be assessed using site-directed mutagenesis.

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**Conflicts of interest**

There are no conflicts to declare.

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