OPTICS

X-ray free-electron laser studies reveal correlated motion during isopenicillin *N* synthase catalysis

Patrick Rabe¹, Jos J. A. G. Kamps^{1,2,3}, Kyle D. Sutherlin⁴, James D. S. Linyard¹, Pierre Aller^{2,3}, Cindy C. Pham⁴, Hiroki Makita⁴, Ian Clifton¹, Michael A. McDonough¹, Thomas M. Leissing¹, Denis Shutin¹, Pauline A. Lang¹, Agata Butryn^{2,3}, Jürgen Brem¹, Sheraz Gul⁴, Franklin D. Fuller⁵, In-Sik Kim⁴, Mun Hon Cheah⁶, Thomas Fransson⁷, Asmit Bhowmick⁴, Iris D. Young^{4,8}, Lee O'Riordan⁴, Aaron S. Brewster⁴, Ilaria Pettinati¹, Margaret Doyle⁴, Yasumasa Joti^{9,10}, Shigeki Owada^{9,10}, Kensuke Tono^{9,10}, Alexander Batyuk⁵, Mark S. Hunter⁵, Roberto Alonso-Mori⁵, Uwe Bergmann^{11,12}, Robin L. Owen², Nicholas K. Sauter⁴, Timothy D. W. Claridge¹, Carol V. Robinson¹, Vittal K. Yachandra⁴, Junko Yano⁴, Jan F. Kern⁴*, Allen M. Orville^{2,3}*, Christopher J. Schofield¹*

Isopenicillin *N* synthase (IPNS) catalyzes the unique reaction of L- δ -(α -aminoadipoyl)-L-cysteinyl-D-valine (ACV) with dioxygen giving isopenicillin *N* (IPN), the precursor of all natural penicillins and cephalosporins. X-ray free-electron laser studies including time-resolved crystallography and emission spectroscopy reveal how reaction of IPNS:Fe(II):ACV with dioxygen to yield an Fe(III) superoxide causes differences in active site volume and unexpected conformational changes that propagate to structurally remote regions. Combined with solution studies, the results reveal the importance of protein dynamics in regulating intermediate conformations during conversion of ACV to IPN. The results have implications for catalysis by multiple IPNS-related oxygenases, including those involved in the human hypoxic response, and highlight the power of serial femtosecond crystallography to provide insight into long-range enzyme dynamics during reactions presently impossible for nonprotein catalysts.

INTRODUCTION

Following pioneering studies demonstrating L- δ -(α -aminoadipoyl)-L-cysteinyl-D-valine (ACV) is the precursor of all natural penicillins, isopenicillin *N* synthase (IPNS) was shown to catalyze formation of both penicillin β -lactam and thiazolidine rings, in an iron- and dioxygen-dependent reaction without synthetic precedent (*1*–*3*). IPNS is a member of the Fe(II) and 2-oxoglutarate (2OG) oxygenase structural superfamily (*4*); such enzymes are widespread in nature and have important roles, including in collagen biosynthesis, lipid metabolism, nucleic acid repair, and signaling (*5*, *6*). 2OG oxygenases play key roles in the human hypoxic response via hydroxylation of the hypoxia inducible factors in a manner regulated by dioxygen availability (*7*, *8*). Understanding how dioxygen interacts with the 2OG oxygenases superfamily is of interest from catalytic and physiological perspectives.

2OG oxygenases typically catalyze hydroxylation reactions that are coupled to 2OG oxidation giving succinate and CO_2 (4, 5). IPNS catalyzes a unique four-electron oxidation involving two

challenging C—H bond cleavages during conversion of the inactive peptide ACV to the conformationally strained antibiotic isopenicillin N (IPN) (Fig. 1 and fig. S1). Substrate analog (2, 9), kinetic (10), spectroscopic (11, 12), and modeling (10, 11, 13–17) studies imply binding of dioxygen to the IPNS:Fe(II):ACV complex that yields an Fe-linked superoxide, which abstracts a C-3 hydrogen from the ACV cysteine giving a thioaldehyde, which undergoes 4-*exo*-tricyclization to produce a β -lactam linked via S to an Fe(IV)=O species. Thiazolidine formation occurs via valine hydrogen abstraction followed by reductive elimination giving the bicyclic IPN ring system (fig. S1) (4, 6, 10). There is a lack of knowledge of how the IPNS protein mediates the conformational changes required for penicillin formation, and structural knowledge of the intermediate formed by reaction of IPNS:Fe(II):ACV with dioxygen has been unavailable.

Redox labile intermediates are not amenable to synchrotron analysis because they react via x-ray-induced photoelectric effects (18). X-ray free-electron lasers (XFELs) provide intense femtosecondlong x-ray pulses enabling diffraction data from thousands of micron-size crystals, limiting radiation-induced chemistry (serial femtosecond crystallography, SFX) (19). To enable time-resolved (tr) SFX studies (20–22) on IPNS catalysis, we exposed anaerobic IPNS:Fe(II):ACV microcrystals to dioxygen using acoustic droplet ejection (ADE) tape drive methods (23) with simultaneous monitoring of the Fe oxidation state by x-ray emission spectroscopy (XES) (24, 25). We subsequently used ¹⁹F NMR (nuclear magnetic resonance) to probe the solution relevance of the dynamics revealed by tr-SFX.

RESULTS

To focus tr-SFX analyses on events associated with O₂ reaction rather than ACV binding and/or IPN release, we co-crystallized the anaerobic IPNS:Fe(II):ACV complex under anaerobic conditions to produce a dense microcrystal slurry (~3 μ m × 3 μ m × 60 μ m, needle

Copyright © 2021 The Authors, some rights reserved; exclusive licensee American Association for the Advancement of Science. No claim to original U.S. Government Works. Distributed under a Creative Commons Attribution License 4.0 (CC BY).

¹Chemistry Research Laboratory, Department of Chemistry and the Ineos Oxford Institute for Antimicrobial Research, University of Oxford, 12 Mansfield Road, Oxford OX1 3TA, UK. ²Diamond Light Source, Diamond House, Harwell Science and Innovation Campus, Didcot OX11 0DE, UK. ³Research Complex at Harwell, Rutherford Appleton Laboratory, Didcot, Oxfordshire OX11 0FA, UK. ⁴Molecular Biophysics and Integrated Bioimaging Division, Lawrence Berkeley National Laboratory, 1 Cyclotron Road, Berkeley, CA 94720, USA. ⁵Linac Coherent Light Source, SLAC National Accelerator Laboratory, Menlo Park, CA 94025, USA. ⁶Department of Chemistry - Ångström, Molecular Biomimetics, Uppsala University, SE 751 20 Uppsala, Sweden. ⁷Interdisciplinary Center for Scientific Computing, University of Heidelberg, 69120 Heidelberg, Germany. ⁸Department of Bioengineering and Therapeutic Sciences, University of California, San Francisco, 600 16th Street, San Francisco, CA 94158, USA. ⁹Japan Synchrotron Radiation Research Institute, 1-1-1 Kouto, Sayo-cho, Sayo-gun, Hyogo 679-5198, Japan. ¹⁰RIKEN SPring-8 Center, 1-1-1 Kouto, Sayo-cho, Sayo-gun, Hyogo 679-5148, Japan. ¹¹Stanford PULSE Institute, SLAC National Accelerator Laboratory, Menlo Park, CA 94025, USA. ¹²Department of Physics, University of Wisconsin–Madison, 1150 University Avenue, Madison, WI 53706, USA. *Corresponding author. Email: christopher.schofield@chem.ox.ac.uk (C.J.S.); allen.orville@diamond.ac.uk (A.M.O.); jfkern@lbl.gov (J.F.K.)



Fig. 1. IPNS catalysis is amenable to tr-SFX analysis. (A) Protein fold of anaerobic IPNS:Fe(II):ACV (conformation A, PDB: 1BLZ) (26); Fe, orange sphere; ACV, yellow; α -helices, purple; β strands, green; except elements involved in ACV binding/flexible regions due to O_2 binding; red. (B) Composite SFX $2mF_o$ -DF_c omit electron density map for anaerobic IPNS:Fe:ACV (PDB: 6ZAE, 1.0 σ contour level, 1.40-Å resolution; fig. S2), showing key ACV interactions (R87, Y91, S183, Y189, L223, N230, R279, and S281) highlighting S281 (red), which is important in dynamics. (C) Composite SFX $2mF_o$ -DF_c omit map for IPNS:Fe:IPN obtained by ~30-min O_2 exposure of IPNS:Fe:ACV micro-crystals (PDB: 6ZAQ, 1.0 σ contour level, 1.60-Å resolution; fig. S2). (D) Key intermediates with the superoxide (ACV conformations B to D, further addressed in Fig. 3 due to ACV rearrangement after O_2 binding) in a blue dashed box. Note: The exact localization of the electrons in the Fe superoxo complex is unknown. To reflect this, we have used the nomenclature Fe- O_2^{3e-} .

morphology, $P2_12_12_1$ space group). The microcrystals that diffract to high resolution (tables S1 and S2) are amenable to efficient reaction initiation by exposure to O₂ and trap IPN at the active site due to lattice constraints (fig. S2), ensuring single turnover conditions.

The anaerobicity of our ADE tape drive sample delivery system and suitability of IPNS microcrystals for tr-SFX analysis was validated by determination of an SFX structure of anaerobic IPNS:Fe:ACV (Fig. 1 and fig. S2), which is similar to reported synchrotron structures under cryo- and room temperature conditions [C- α RMSDs: 0.338 Å/Protein Data Bank (PDB) 1BK0 (*26*) and 0.234 Å/6Y00 (*27*)]. The active site iron refines to full occupancy and exhibits square pyramidal coordination with ligation by His²¹⁴, Asp²¹⁶, His²⁷⁰, a water, the ACV thiolate, with an open coordination site trans to Asp²¹⁶. ACV interactions relevant to the tr-SFX analyses include a salt bridge of its α -aminoadipoyl (Aad) carboxylate with Arg⁸⁷ (which hydrogen bonds with Thr³³¹ on the C-terminal helix α 10), hydrogen bonds between Tyr⁹¹ and the Aad amino group, and of the ACV valine carboxylate with Tyr¹⁸⁹ and Ser²⁸¹ (fig. S1).

Exposure of room temperature IPNS:Fe:ACV microcrystals to air for \geq 30 min resulted in high-resolution structures and IPN refined to full occupancy, with diffraction data collected at cryogenic or room temperatures, at both synchrotrons and XFELs, implying a single, O₂-dependent ACV turnover (SFX rt.: PDB: 6ZAQ; MX cryo: PDB: 6ZAO; SSX fixed target: PDB: 6Y0P; Fig. 1C and fig. S2). In accord with prior work (28), exposure of large single IPNS:Fe(II):ACV crystals to O₂ resulted in incomplete conversion to IPN (26), even with high pressure/prolonged reaction, implying more efficient O₂ diffusion in microcrystals. For the IPN product complexes reported here, we refined a five-coordinate distorted square pyramidal Fe site, ligated by His²¹⁴, Asp²¹⁶, His²⁷⁰, a water, and the IPN sulfur (Fe-S: 2.98 Å), differing from the square pyramidal coordination in IPNS:Fe:ACV and modeling of the proposed IPN complex (10). In our IPN product complexes derived from microcrystals, the Fe refined to convergence with low occupancy (40 to 60%). However, solution studies using NMR and non-denaturing mass spectrometry (MS) provide no evidence for IPN-promoted Fe displacement (fig. S2). Thus, the apparent loss of Fe on formation of IPN in crystallo likely reflects tight IPN binding due to lattice packing interactions, which promote release of Fe rather than IPN.

We investigated the reaction of IPNS:Fe(II):ACV with O_2 by exposing microdroplets of the anaerobic microcrystal slurry to a 100% O_2 atmosphere, regulating the reaction time by changing the ADE tape drive speed (23). On the basis of the high ratios of indexed crystal lattices to images collected, multiple datasets (for 400-, 500-, 800-, 1600-, and 3000-ms O_2 exposure times) were collected from the same microcrystal batch (table S1) enabling robust dataset comparison.

Diffusion of O_2 into microcrystals slightly alters the unit cell parameters, suggesting changes that alter fold and/or lattice packing. No clear structural differences at 400, 500, or 800 ms were apparent (fig. S3); however, analysis of the ACV B-factors at 500 ms and longer times reveals substantial increases (22 Å²/anaerobic to 36 Å²/1600 ms; Fig. 2 and fig. S3). The ACV value B-factors increased first (500 ms), followed by those of the entire ACV (\geq 800 ms), including the Aad side chain. The latter was unexpected, because all ACV/IPN atoms are fully ordered in both, the anaerobic ACV substrate and IPN product complexes (Fig. 1 and figs. S2 and S3). These observations suggest that O_2 binding induces motions beyond those directly involved in covalent reaction at the active site. We observed that addition of glycerol (to minimize dehydration) reduces the reaction rate, consistent with O_2 diffusion being slowed by increased viscosity in the presence of glycerol (fig. S4) (29).

Positive electron density appeared at the Fe coordination site trans to Asp²¹⁶ in the 1600- and 3000-ms SFX datasets (Fig. 3 and fig. S5). The 1600-ms dataset electron density was initially fit and



Fig. 2. O_2 binding to IPNS:Fe:ACV causes dynamics throughout the protein. Color gradient B-factor analysis of tr-SFX data from blue (24 Å²), to white, to red (36 Å²) for (**A**) Fe, O_2 , substrate/intermediate/product and (**B**) the protein including α 3 (amino acids 47 to 64) and the C-terminal region (α 10; amino acids 315 to 331). Note: (i) α 3 B-factors rise on O_2 exposure and decrease on IPN formation, (ii) ACV valine–derived atoms become disordered on O_2 exposure but are ordered in the IPN complex, (iii) α 10 is involved in ACV binding/IPN product release and remains disordered in IPNS:Fe:IPN complex since IPN is trapped due to lattice packing constraints.

refined with a coordinated water, but difference features remained and this model did not adequately reflect the scattering. The best refined fit to the 1600-ms dataset includes an O_2 bound "end-on" to the Fe (figs. S5 and S6), consistent with modeling (fig. S7) (11, 16).

In refinements for the 1600- and 3000-ms datasets, we omitted those ACV atoms with very weak or lacking $2mF_o$ -DF_c density, which did not generate mF_o -DF_c difference features when removed. These atoms are present but are disordered in intermediates rendering them invisible to x-ray diffraction (XRD), despite the use of 35-fs x-ray pulses. By contrast, before reaction initiation and following its completion, all nonhydrogen ACV or IPN atoms are visible. Thus, as O₂-initiated reaction progresses, the ACV becomes dynamic, with the extent of disorder varying with the substrate region.

The results imply that O_2 binding causes the ACV Val isopropyl group to rotate away from the Fe to avoid a steric clash; similar rotation is observed in synchrotron cryo-structures for the IPNS:Fe:ACV:NO complex, wherein NO is an O_2 surrogate (Fig. 3 and fig. S7). By contrast with the IPNS:Fe:ACV:NO complex, in the 1600-ms dataset, electron density for the ACV Val is weak and the Val isopropyl methyls were deleted in the refinement models [ACV conformation (conf.) A, 20% occupancy (– O_2) and conf. B, 80% occupancy (+ O_2); Fig. 3 and fig. S5].

Similar considerations apply to the 3000-ms dataset, where we fit and refined the ACV with the entire Val isopropyl deleted. In this dataset, we also observed decreased electron density, consistent with partial disorder, for both the valine carboxylate and Aad side chain (conf. C; Fig. 3 and fig. S5). In the 3000-ms dataset, an additional conformation (conf. D) was proposed in which the ACV cysteinyl methylene rotates such that the pro- $SC_{Cys,\beta}$ —H bond is close to the distal O of the Fe-O₂ (2.4 Å compared to 3.1 Å in conf. C). This produces a complex aligned for stereospecific cysteinyl 3—H bond cleavage (*10*) leading to a thioaldehyde, which is preorganized for stereoelectronically favored β -lactam formation (Fig. 3 and figs. S5 and S6). Thus, the tr-SFX results reveal that, at this stage of

the reaction, the ACV Cys/Val atoms involved in thioaldehyde and β -lactam formation appear to be more ordered than those Aad and Val atoms that are not directly involved in these steps of the reaction cycle.

The tape drive setup enables simultaneous collection of tr-SFX and tr-XES data (23), allowing assignment of the Fe oxidation states (30). XES analysis of the anaerobic IPNS:Fe:ACV microcrystal complexes supports the Fe(II) oxidation state (Fig. 4A and 4B, and fig. S6). In the O₂-bound datasets (1600 and 3000 ms and 500 ms without glycerol; figs. S4 and S6), changes are observed in the XES and difference spectra (Fig. 4A), especially in the full width at half maximum (FWHM) (25) of the K α_1 peak (Fig. 4B). These changes are similar in magnitude to those on going from Fe(II) to Fe(III) in Mn/Fe containing R2a ribonucleotide reductase (23), in the diiron center of methane monooxygenase (31), and in small-molecule Fe complexes (32), supporting formation of an IPNS:Fe(III):ACV:O₂⁻ intermediate consistent with solution studies (10, 11, 17) and our tr-SFX data.

The tr-SFX datasets reveal clear changes of average B-factors in two structurally separate regions: (i) the C-terminal region including $\alpha 10$ [which is important for productive catalysis (3, 33)] and the loop connecting $\alpha 10$ with the protein core and (ii) $\alpha 3$, for which no function is ascribed and which is located on the exterior of the double stranded β helix core (DSBH) fold (Fig. 2 and figs. S3 and S4). Two other exterior regions also refine with increased B-factors (amino acids 105 to 120, including α 5) and an exterior loop (amino acids 194 to 206), although these do not exhibit changes correlating well with increasing O_2 exposure time. The $\alpha 10$ B-factors increase with time and remain high in the IPNS:Fe:IPN complex. Those for a3 increase with time, but in the IPNS:Fe:IPN complex (Fig. 2 and fig. S3), they refine back to values close to those for anaerobic IPNS:Fe:ACV. Comparison of the F_{obs} – F_{obs} isomorphous difference maps for the 400- and 3000-ms datasets reveals clear evidence for dynamic changes in the $\alpha 3/\beta 11$ conformations, consistent with the calculated mFo-DFc maps (fig. S3).

SCIENCE ADVANCES | RESEARCH ARTICLE



Fig. 3. tr-SFX reveals mode of O₂ binding to IPNS:Fe:ACV. ACV-derived conformations (**A** to **D**) were fit and refined to the 1600-ms (ACV conformation A and B) and 3000-ms (ACV conformation C and D) tr-SFX datasets (PDB: 6ZAI and 6ZAJ). $2mF_o-DF_c$ omit maps (1.0 σ contour level, 1.53 and 1.55 Å resolution, respectively) are shown. Conf. A: The same as the anaerobic ACV complex (Fig. 1). Conf. B: The major 1600-ms IPNS:Fe:ACV + O₂ conformation (B, 80%) refined with the ACV Val isopropyl methyls deleted (teal). Left inset: Overlay of IPNS:Fe:ACV (A, 20%, details in Fig. 1B) and IPNS:Fe:ACV:O₂ (B, 80%) 1600-ms models. Conf. C: (3000 ms), where all Val isopropyl atoms are disordered. Central inset: The anaerobic IPNS:Fe:ACV:NO complex (PDB: 6ZAN, cryo MX, 1.39-Å resolution). Conf. D: New proposed conformation in the 3000-ms dataset with the Cys side-chain rotated. Right inset: Superimposition of confs. C and D, showing how the ACV cysteinyl methylene rotation positions the pro-S C_{Cys,β}—H bond close to the distal O of the Fe-O₂ for stereospecific C—H cleavage (fig. S6).

B-factor analyses of the tr-SFX O₂–exposed datasets reveals increased dynamics of α 3; however, we were unable to confidently fit and refine discrete new conformations because the electron density maps indicated that these are present at low occupancy (10 to 15%). Therefore, we attempted to trap thermodynamically unstable conformations by exposing single IPNS:Fe:ACV crystals to O₂ and then plunging them into liquid nitrogen for cryo-synchrotron analysis. Analysis of ~100 IPNS:Fe:ACV crystals exposed to O₂ for times ranging from 30 to 600 s shows evidence for electron density corresponding to O₂ binding trans to Asp²¹⁶ [although photoelectric reduction likely occurs (18)] with a trend to disorder of the ACV valine, along with increased α 10 and α 3 B-factors. Notably, under the cryo-conditions, along with the starting conformations as observed previously (I), we unequivocally observed the formation of discrete new conformations for α 3 and β 11 (II) (Fig. 5 and fig. S7).

Similar changes in value disorder, increased B-factors and the new $\alpha 3/\beta 11$ conformations, were observed on formation of the IPNS:Fe:ACV:NO complex in single crystals (Fig. 5 and fig. S7). A structure of the IPNS:Fe:ACV:NO complex (PDB: 1BLZ) (26) has been reported; alternative confs. of $\alpha 3$ and $\beta 11$ due to NO binding

were not refined, possibly due to low occupancies of the flexible regions. As for the tr-SFX-observed O₂ binding and as reported in a synchrotron derived structure (26), NO binds trans to Asp²¹⁶, but with a slightly different orientation to O₂ (Fig. 5D). These NO or O₂-exposed structures were refined with both α 3/ β 11 conformations (I and II)—the IPNS:Fe:ACV:NO complex and a representative IPNS:Fe:ACV:O₂ complex with 1:1 occupancy of conformations I and II were selected for deposition. The difference between the cryo-condition work, where we observed the discrete new conformations for α 3/ β 11, and the tr-SFX studies, where we did not accrue unequivocal evidence for them, likely in part reflects higher mobility under the latter room temperature conditions.

The combined tr studies enable experimentally based proposals for the underlying interactions involved in dynamics during IPNS catalysis. Although motions other than those involving $\alpha 3/\beta 11$, including the C-terminal region in substate binding and product release, are likely involved, our studies reveal a critical role for $\alpha 3/\beta 11$ dynamics in β -lactam ring formation. O₂ binding induces mobility of the ACV valine weakening its interaction with Ser²⁸¹_{$\beta 11$} causing $\beta 11$ to move away from ACV (Fig. 5, fig. S7, and movie S1). This



Fig. 4. Tr-XES analysis reveals superoxide formation. (**A**) Fe K $\alpha_{1,2}$ XES of anaerobic IPNS:Fe:ACV microcrystals (red), after 1600-ms O₂ exposure (black); a difference spectrum between the two (blue, 5× enlarged) and a difference spectrum between 10 mM solutions of Fe(III)Cl₃ and Fe(II)Cl₂ (green, 2× enlarged). a.u., arbitrary units. (**B**) Full width at half maximum (FWHM) of the Fe K $_{\alpha 1}$ XES peak following O₂ exposure. Early times (\leq 800 ms) show only small changes in the FWHM (+/- 0.03 eV); the 1600- and 3000-ms times show a shift by ~0.1 eV, indicating Fe oxidation. (**C**) 2mF $_0$ -DF $_c$ map of the IPNS:Fe³⁺:ACV:O₂^{1o-} complex (PDB: 6ZAI).



Fig. 5. O₂ **binding to IPNS:Fe:ACV causes unexpected conformational changes.** (**A**) Refined model for a cryo-cooled IPNS:Fe:ACV:O₂ complex after O₂ exposure (60 s) reveals Fe-O₂ binding and induced rearrangements from confs. I to II of α 3/ β 11 [PDB: 6ZAP, cryo MX, 1.36-Å resolution, yellow: conf. I ($-O_2$); teal: conf. II ($+O_2$)]. Ser²⁸¹, which is involved in ACV value carboxylate binding shifts by 1.2 Å. Movements are indicated by red arrows; hydrophobic interactions ($L^{282}_{\beta 11}/L^{54}_{\alpha 3}$) by dashed lines (fig. S7). (**B**) The IPNS:Fe:ACV:NO (PDB: 6ZAN, 1.39-Å resolution) complex refines to two equal α 3/ β 11 confs. I/II (fig. S7). (**C**) Hydrogen bond network involved in O₂ binding induced α 3/ β 11 rearrangements. (fig. S7). (**D**) Superimposition of both complexes reveals different NO and O₂ binding modes [Fe-O₁-O₂ (135°) and Fe-N-O (130°); fig. S7]. (**E** and **F**) Polder omit maps (3.0 σ contour level) of residues involved in O₂ binding–induced rearrangement carved around β 11/ α 3, omitting (E) conf. I and (F) conf. II, showing confs. I and II of β 11/ α 3, respectively.

increases the active site volume by ~13 Å³, so enabling productive movement of the CV unit of ACV to initially form a thioaldehyde and then a monocyclic β -lactam (Fig. 6).

As a consequence of the movement of Ser²⁸¹_{$\beta 11$} and other $\beta 11$ residues, conformational changes occur throughout $\alpha 3$ on the protein surface (amino acids 47 to 64, $\alpha 3$ 51 to 64). The correlated

motions involving $\alpha 3/\beta 11$ are enabled by interactions between $Gln^{280}{}_{\beta 11}$ and $Ser^{55}{}_{\alpha 3}$, $Leu^{282}{}_{\beta 11}$ and $Leu^{54}{}_{\alpha 3}$, $Tyr^{224}{}_{\alpha 6}$ and $Ser^{55}{}_{\alpha 3}$ [Tyr²²⁴ is conserved and adjacent to Leu^{223} , which is part of a hydrophobic pocket binding the ACV valine isopropyl (26, 33)] and $His^{62}{}_{\alpha 3}$ and $Tyr^{90}{}_{\alpha 3}$ (Fig. 5E and fig. S7). Tyr⁹⁰ is adjacent to Tyr^{91} , which binds the ACV Aad amino group via a hydrogen bond



Fig. 6. Comparison of active site volume before and after O₂ binding-induced rearrangement of \beta11 and \alpha3. Calculations of the active site volume were performed with Pyvol (*61***). (A**) Active site volume presentation of the IPNS:Fe:ACV complex (yellow). The alternative conf. II obtained after O₂ binding is in teal with high transparency. (**B**) Active site volume superimposition of the atomic model obtained from the single-crystal IPNS:Fe:ACV:O₂ (PDB: 6ZAP) complex by cryo-cooled synchrotron data collection highlighting the two different conformations of β 11 and α 3 triggered by O₂ binding. The two panels show the active site volume in an open (left) and closed (right) view. To perform active site volume calculations, either conf. A (ground state, yellow) or B (O₂ bound, teal) was removed from the model. The active site volume after O₂ binding (teal, 1374 Å³) appears to increase by 13 Å³ compared to conf. A, as obtained in the anaerobic ground state (yellow, 1361 Å³). Note that these changes appear solely in the region where the bond-breaking and bond-forming steps for penicillin formation are happening. (**C**) Active site volume returns back to its ground state after turnover. These observations suggest a controlled active site volume expansion during penicillin ring formation. The alternative conf. II obtained after O₂ binding is shown in teal with high transparency.

(Fig. 5E). The Aad side chain, including its amino group, becomes disordered following O_2 exposure but is ordered in the IPN product complex. α 3 thus plays a central role in conformational changes involved in the conversion of ACV to IPN via dynamic interactions with both the ends of ACV/intermediates, i.e., with the Aad amino group and the Val carboxylate.

We then used ¹⁹F-NMR to investigate whether the conformational changes observed for α 3 in crystals occur in solution, by monitoring changes of IPNS ¹⁹F-labeled (*34*) on α 3 (IPNS*; Fig. 7 and fig. S8). The results reveal distinct signals for the apo, Fe(II), and Fe(II):ACV complexes. Addition of NO to IPNS*:Fe:ACV, but not IPNS*:Fe or IPNS*:Cd:ACV (Fig. 7 and fig. S8) complexes, resulted in movement of the IPNS* ¹⁹F signal (Fig. 7 and fig. S8), supporting the proposal that O₂/NO binding induces α 3 dynamics in presence of ACV, since NO should not bind to the Cd(II) complex.

DISCUSSION

Our studies provide new insights how correlated motions induced by O_2 binding enable IPNS to catalyze β -lactam formation, including by transiently altering the active site volume and arranging the substrate conformation in preparation for ring formation. Sequence comparisons reveal that $\alpha 3/\beta 11$ and key residues involved in the associated correlated motions are conserved in IPNS ranging from prokaryotes to an animal (fig. S9A) (35). β11 (which extends one β sheet of the core DSBH fold) and α 3 (one of the two conserved exterior N-terminal helices) are present in multiple 2OG oxygenases, ranging from those of cephalosporin biosynthesis which are closely related to IPNS to those involved in DNA/RNA repair, histone demethylation, lipid metabolism, and the human hypoxic response (fig. S9B) (5, 6). Thus, although there are likely variations, and the extent of protein dynamics during catalysis is likely underestimated in crystallographic studies, the types of correlated motions observed by tr-SFX during IPNS catalysis are likely of widespread relevance. Detailed knowledge of how oxygenases work should enlighten research concerning their biological roles as sensors, on engineering them to alter the course of biosynthesis and on modulating their activity for therapeutic benefit.

The roles of dynamics in tuning catalysis of individual steps are of general interest in catalysis, particularly for enzymes that catalyze chemically challenging reactions involving substantial conformational changes (36). tr-SFX is a powerful method for uncovering

SCIENCE ADVANCES | RESEARCH ARTICLE



Fig. 7. Spectroscopic analyses in solution support structural dynamics as a consequence of O₂ binding. (A) ¹⁹F NMR spectra from anaerobic solutions of IPNS* \pm Fe²⁺ \pm ACV \pm NO support α 3 dynamics after NO binding. (i) IPNS* (¹⁹F-labeled IPNS); (ii) IPNS* + Fe²⁺; (iii) IPNS* + Fe²⁺ + ACV; (iv) IPNS* + Fe²⁺ + ACV; (iv) IPNS* + Cd²⁺ \pm ACV \pm NO; and (v) IPNS* + Fe²⁺ + ACV; (iii) IPNS* \pm Cd²⁺ \pm ACV \pm NO; (ii) IPNS* \pm Cd²⁺ \pm ACV \pm NO; (ii) IPNS* \pm Cd²⁺ \pm ACV \pm NO; (iii) IPNS* \pm Cd²⁺ \pm ACV \pm NO; (iii) IPNS* \pm Cd²⁺ \pm ACV; (iii) IPNS* \pm Cd²⁺ \pm ACV \pm NO; (iii) IPNS* \pm Cd²⁺ \pm ACV \pm O(Fig. S9). (**C**) Comparison of the structural elements of α 3/ β 11 in IPNS:Fe:ACV:O₂ (PDB: 6ZAP) and IPNS*:Fe:ACV (cryo-cooled single-crystal analysis, PDB: 6ZAM, 1.55-Å resolution, used for ¹⁹F NMR, showing keto and gem diol forms of fluorine label). (**D**) Overview of the IPNS* complex with α 3 and β 11 in red and presence of the fluorine label on α 3. (**E**) View of the active site and α 3 of the IPNS*:Fe:ACV complex, showing the distance from the ACV valinyl carboxylate to the alkylated Cys⁵⁵ (16.5 Å). Changes in the ACV valinyl carboxylate conformation as a function of O₂ binding are proposed to induce structural rearrangement of α 3/ β 11. Cys⁵⁵ was alkylated with a CH₂COCF₃ group to study conformational changes in solution by ¹⁹F NMR. Tryptic digestion confirmed the presence of alkylated Cys⁵⁵ (fig. S10). (**F**) Views of the active site of the aerobically grown IPNS:Cd:ACV complex (PDB: 6ZW8, 1.22-Å resolution) obtained by single-crystal cryogenic data collection. A Polder omit map of the active site of IPNS:Cd:ACV after omitting Cd and ACV (3.0 σ contour level, green, modeled occupancy of 90% for Cd). Unmodeled electron density was obse

these, including, as shown here, motions involved in steps after substrate binding and before product release. We hope increased knowledge of how IPNS works will promote the use of tr-SFX to study synthetically challenging reactions and inspire the discovery of new types of nonprotein catalysts making densely functionalized and conformationally strained ring systems. However, our results demonstrating that motions outside the active site are important in catalyzing conversion of a "simple" peptide to the penicillin nucleus suggest that such biomimetic catalysts may need to be macromolecular.

MATERIALS AND METHODS

Chemicals for preparation of buffers and crystallization screens were from commercial suppliers and were used without further purification. ACV was synthesized by solid phase peptide synthesis and purified using a Shimadzu high-performance liquid chromatography (HPLC) system, equipped with a SunFire semiprep column (C18, 5 μ m, 150-mm length, 10-mm diameter). The mass of ACV was confirmed by LC-MS; Agilent Technologies 1260 Infinity Series,

equipped with a 6120 quadrupole mass spectrometer using a Merck Chromolith Performance C18 (100 mm by 4.2 mm) HPLC column. ACV purity was confirmed by NMR (Bruker AVIII HD 600 equipped with a BB-F/H N₂ Prodigy CryoProbe). Recombinant wt-IPNS (pJB703) from *Aspergillus nidulans* in NM554 *Escherichia coli* was produced by the reported procedure (*37*).

Cloning of pCOLD_IPNS

DNA (codon-optimized for expression in *E. coli*) of IPNS (GeneArt; Thermo Fisher Scientific, UK) was inserted into the pCOLD I vector (Addgene, UK) using Sal I and Not I restriction sites. This vector enabled production of IPNS with an *N*-terminal hexa-histidine tag (6xHis) with an *N*-terminal C human rhinovirus protease cleavage site as reported (38).

Mutation of pCOLD_IPNS S55C

A Ser⁵⁵ to Cys variant of codon-optimized IPNS produced with the pCOLD vector was obtained using the Q5 site–directed mutagenesis kit (New England Biolabs, USA) followed by DpnI digestion at 32°C

for 5 hours. Primers used are as follows: IPNS_S55C_fwd: cattaatgtgcagcgtctgtgccagaaaaccaaagaatt and IPNS_S55C_rev: aattctttggttttctggcacagacgctgcacattaatg. The polymerase chain reaction product was transformed into XL10 Gold ultracompetent cells (Agilent Technologies, USA) by heat shock and grown overnight on 2YT agar plates containing ampicillin (50 μ g ml⁻¹). A single colony was picked and cultured in liquid 2YT media while shaking (150 rpm) overnight at 37°C. Plasmid DNA was isolated using the GeneJET Plasmid Miniprep kit (Thermo Fisher Scientific, USA). The presence of the desired mutation was confirmed by DNA sequencing (Eurofins, Germany).

Expression and purification of IPNS_S55C and labeling of IPNS S55C with BTFA (34, 39)

The pCOLD_IPNS_S55C plasmid was transformed into E. coli BL21 cells and grown in 2YT agar plates containing ampicillin (50 μ g ml⁻¹). Single colonies were picked and cultured in 2YT media (100 ml), supplemented with ampicillin (50 µg ml⁻¹), overnight with shaking (150 rpm, 37°C). A starter culture (6 ml; 1:100, v/v) was used to inoculate a large-scale growth in 2YT media (600 ml) supplemented with ampicillin (50 μ g ml⁻¹) with shaking (150 rpm, 37°C) until an appropriate OD_{600} (optical density at 600 nm) = 0.6 was reached. Expression was induced by β-D-1-thiogalactopyranoside (1 mM final concentration) addition; cultures were incubated overnight at 15°C. Cells were harvested by centrifugation (11,000g, 10 min, 4°C) and stored at -80°C.

A cell pellet (25 g) was resuspended (1:4 w:v) in buffer G [50 mM tris (pH 7.5), 200 mM NaCl, and 5 mM imidazole] supplemented with deoxyribonuclease I (10 µg ml⁻¹), phenylmethylsulfonyl fluoride (10 μ g ml⁻¹), and lysozyme (0.2 mg ml⁻¹) by stirring at 4°C and for 30 min. Cells were lysed by sonication (9 s on: 9 s off, 60% amplitude, 12-min total time, 4°C; Sonics Vibra-Cell), cell debris was removed by centrifugation (58,000g, 30 min, 4°C), and the supernatant was filtered (0.45 µm). The filtrate was loaded onto a nickel affinity column (5 ml HisTrap HP, GE Healthcare, USA), pre-equilibrated with buffer G (20 CV), and eluted with a gradient from buffer G to buffer H [50 mM tris (pH 7.5), 200 mM NaCl, 500 mM imidazole, and 20 CV]. Fractions containing IPNS were identified by SDSpolyacrylamide gel electrophoresis (PAGE) and concentrated by centrifugation [10,000 molecular weight cut-off (MWCO), 3000g, 4°C; Merck Millipore, USA], before purification by Superdex 75 (300 ml, GE Healthcare, USA) chromatography using a column pre-equilibrated with buffer I [25 mM tris (pH 7.5), 500 mM NaCl, and 1 CV]. Fractions containing protein were identified by ultraviolet analysis and analyzed by SDS-PAGE. Fractions containing purified IPNS S55C were concentrated using a centrifugation tube (10,000 MWCO, 3000g, 4°C; Amicon Ultra). Removal of the N-terminal hexa-histidine affinity tag was achieved by incubation with 3C protease (1:100, w/w) overnight at 4°C. Successful cleavage was confirmed by protein MS (Quattro Premier XE, Waters, USA). Untagged IPNS_S55C was subsequently purified using a nickel affinity column (5 ml HisTrap HP, GE Healthcare, USA) using buffer G (5 CV). A solution of IPNS_S55C (350 µM) was incubated with tris(2-carboxyethyl) phosphine [350 µM, 1 equivalent (eq.)] in buffer J [25 mM tris (pH 8.5) and 100 mM NaCl] for 5 min at 4°C. 3-Bromo-1,1,1-trifluoro acetone (BTFA; 10.5 mM final concentration, 30 eq.) was added, and the solution was incubated (90 min) at 4°C. Alkylation of IPNS_S55C was confirmed by MS (Quattro Premier XE, Waters, USA); the observed mass for IPNS_S55C (37,768 Da) demonstrated an increase

by 128 Da when alkylated to give IPNS* (37,896 Da, BTFA-labeled IPNS_S55C). Note that the predicted mass difference based on the calculated mass of CH₂C(O)CF₃ (110 Da) was higher by 18 Da than observed, suggesting that the ketone exists mainly in its hydrated form, as observed for a different protein (34, 40, 41), and consistent with our crystallization analyses (Fig. 7). The reaction was stopped by buffer exchange using PD-10 columns, pre-equilibrated with buffer J. The BTFA-labeled IPNS_S55C (IPNS*) variant was concentrated to a volume < 2 ml and further purified by size exclusion chromatography using a Superdex 75 column (300 ml, GE Healthcare, USA) pre-equilibrated with buffer J. Protein containing fractions were subsequently dialyzed as described above. The apo-IPNS* was concentrated to 50 mg ml⁻¹, aliquoted, and stored at -80°C. Trypsin digestion was performed to confirm the labeling (fig. S10).

Tryptic digestion

Tryptic digestion was performed according to a reported procedure (42). Labeled IPNS* (15μ l, 1 mg ml⁻¹) was treated with dithiothreitol (DTT; 2 µl, 85 mM), in ammonium bicarbonate buffer (10 mM), for 40 min at 56°C. The protein was then alkylated with iodoacetamide $(7 \mu l, 55 \text{ mM})$ for 30 min and then treated with DTT $(3 \mu l, 85 \text{ mM})$ for 10 min at room temperature (rt) in the dark. Trypsin (3 µl; 1:20, w/w) was added and reacted for 16 hours at 37°C. The sample was then diluted with acetonitrile (120 µl; 1:100, v/v) and allowed to react for 3 hours at 37°C. Digestion was stopped by the addition of formic acid (7.5 µl), and the mixture was then vacuum centrifuged (Eppendorf, Germany) to dryness. The residue was redissolved in (Eppendort, Germany) to dryness. The residue was redissolved in $H_2O(18 \ \mu)$ supplemented with aqueous formic acid (0.1% v/v). After pre-equilibrating a ZipTip column (Merck Millipore, USA) with aqueous acetonitrile (98:2, 0.1% formic acid), the sample was aspirated into the ZipTip column. The fragments were eluted from the ZipTip column using aqueous acetonitrile (40:60, 0.1% aqueous formic acid), vacuum centrifuged to dryness and redissolved in aqueous acetonitrile (98:2, 0.1% formic acid), and analyzed by nanoLC (Thermo Elite, Thermo Fisher Scientific, USA), and data were analyzed using BSI PEAKS studio 8.5. The results from the trypsin digestion are shown in fig. S10. **Nitric oxide exposure of IPNS crystals and solutions**Experiments involving nitric oxide (NO) were carried out by two researchers using a fume hood. A two-chambered, four-valve glass apparatus (fig. S11) was used for controlled NO exposure, minimizing O₂ and user exposure. Anaerobic protein/crystal samples, sealed in J Young valve NMR tubes (5 mm, Norell, USA), were attached to the NMR valve without opening the NMR tube (fig. S11). The system was first purged with argon (10 min) with the gas outlet and separating valves open. The gas outlet valves were closed, under a slight overpressure of argon. Chamber 2 was sealed by closing the armsting when we have a sing the product of the NME tube of the product H_2O (18 µl) supplemented with aqueous formic acid (0.1% v/v).

slight overpressure of argon. Chamber 2 was sealed by closing the separating valve, and vacuum was applied by opening the chamber 2 outlet valve. To create a mild vacuum in the NMR tube, the outlet valve was closed before opening the NMR valve. After equilibration (~2 min), the NMR valve was closed and the chamber 2 outlet valve was reopened. This process was repeated twice. Chamber 1 was purged with NO [1000 parts per million (ppm) in nitrogen, (0.1% NO and 99.9% N₂), 10 min; BOC, UK]. After purging chamber 1, the outlet valve was closed while the system was under a positive pressure, and the balloon was partially filled with NO. To expose the sample to NO, the separating and NMR valves were opened. Samples were incubated with NO for 10 to 30 min to yield a pale pink solution (12) in the NMR tube. The NMR valve was closed and samples was removed for analysis. To remove NO from the system, all valves were opened and flushed with argon for several minutes. Note that, for safety reasons, no vacuum was applied once the system was filled with NO.

NMR studies

NMR experiments on IPNS* [the ¹⁹F-BTFA-labeled IPNS_S55C mutant isoform (trifluoroacetonyl labeled); Fig. 7 and fig. S8] were recorded using a Bruker AVIII HD 600 equipped with a BB-F/H ProdigyN2 CryoProbe, in 5-mm regular or J Young valve NMR tube (5 mm; Norell, USA), at 298 K, unless stated otherwise. ¹⁹F NMR spectra were referenced to CF₃CO₂H (100 μ M, at $\delta_F = -76.55$ ppm) and processed with 3- or 30-Hz Lorentzian line broadening using MestReNova 14.1 (MestReLabs, Spain; www.mestrelab.com) and TopSpin 3.6.1 (Bruker, Germany; www.bruker.com). Sample preparation was performed under anaerobic conditions (<2-ppm O₂) in an anaerobic chamber (Belle Technology, UK). Solutions [in 25 mM tris-d₁₁ buffer (pH 7.5)] were deoxygenated by argon purging (30 min) before placing in an anaerobic chamber. Solids (FeSO₄ and ACV) and NMR tubes were transferred into the glovebox and left to equilibrate (16 hours). Solutions of apo-IPNS* (50 mg ml⁻¹, 1.35 mM, and 50 µl) were transferred to the anaerobic chamber immediately before use. Appropriate stock solutions of ACV (10 mM), FeSO₄ (100 mM), and CF₃CO₂H (10 mM) were prepared in a glovebox. The total volume of a typical sample was 450 µl and contained 10% (v/v) D_2O . After equilibration, the sample was transferred from an Eppendorf tube (1.5 ml) into a 5-mm J Young valve NMR tube (Norell, USA) for analysis.

For the addition of subsequent solutions to the sample, a J Young valve NMR tube was transferred into the glovebox and equilibrated for 5 to 10 min before opening. The appropriate solution was added, and the tube was closed and inverted to equilibrate. For exposing a sample inside a J Young valve NMR tube to nitric oxide (NO), the tube was connected to an appropriate glass apparatus (fig. S11), and the sample was treated as described above. The J Young valve was closed before disconnecting the tube and inverted carefully several times to allow mixing of the gas in the headspace with the solution.

Preparation of salt free IPNS and nondenaturing MS

Purified IPNS was used for nondenaturing electrospray ionization MS after buffer exchange into ammonium acetate (1 M). Buffer exchange was performed using Zeba Micro Spin desalting columns (Thermo Fisher Scientific) followed by 3-hour dialysis with ammonium acetate (1 M). Samples were mixed with the different (co-) substrates dissolved in Milli-Q after different incubation times. Nondenaturing MS data were acquired using an Orbitrap extended mass range prototype (43) machine. Data were recorded in positive ion mode, from a static nanospray source, using a gold-plated capillary prepared in house. Nitrogen was used as a collision gas [the pressure of around 1×10^{-9} mbar recorded in the Orbitrap, no HCD voltage (0 V) applied]. The capillary temperature was set to 30°C, and a spray voltage of 1.4 kV was applied. Acquired data were processed and analyzed using Thermo Xcalibur 4.13 (Thermo Fisher Scientific, UK) (43).

Optimization of microcrystals

Crystallization was conducted within an anaerobic chamber maintained at 2 ppm or less O_2 (Belle Technologies, UK) with plates, solutions, and other equipment used for crystallization deoxygenated within the chamber for at least 24 hours. The IPNS solution that was used to grow crystals was deoxygenated in the anaerobic chamber for 1 to 2 hours before use.

IPNS microcrystals were prepared as reported (27). For setting up a batch plate of microcrystals, IPNS (200 μ l, $c = 52 \text{ mg ml}^{-1}$), ACV (2.2 mg), FeSO₄·7H₂O (1 eq., 100 mM stock in H₂O, 2.78 µl) and seed stock solution (12 µl) were transferred into an anaerobic chamber (Belle Technologies, UK). Using a 96-well plate (Corning, USA), solutions for the batch setup were prepared as follows. For each well, Li_2SO_4 (66.5 µl of 2.7 M in H_2O), H_2O (13.5 µl), tris buffer (10 µl of 1.0 M in H₂O, pH 8.5), and IPNS mixture (6.5 µl) were required. For one aliquot of IPNS solution (200 μ l, 50 mg ml⁻¹, 1.4 mM), a solution for 30 wells comprising Li₂SO₄ (1.995 ml, 2.7 M in Milli-Q), Milli-Q (405 µl), and tris [300 µl, 1.0 M in Milli-Q (pH 8.5)] was prepared. Before addition of IPNS to the crystallization solution, a freshly prepared FeSO₄ solution (2.8 µl, 1.0 equivalent, 100 mM in Milli-Q water) was added to the IPNS solution; the mixture was equilibrated for 10 min. ACV (2.2 mg) was dissolved in tris buffer [40 μ], 25 mM in Milli-Q (pH 8.5)] and added in 4 \times 10 μ l portions carefully to the protein solution; precipitation was removed by centrifugation. The clear protein solution (containing ACV and FeSO₄, ~220 µl) was added to the crystallization solution. The seed stock (12 µl) was then added and the mixture was pipetted into the wells (100 µl). The plate was carefully sealed (Polyolefin StarSeal, Starlab, UK) and shaken (700 rpm) using a microplate shaker (SciQuip, UK) for 16 to 30 hours depending on the temperature and the relative humidity in the anaerobic chamber. The appropriate shaking speed enabled production of needle morphology microcrystals (1 ml, $c = 2 \times 10^7$ microcrystals/ml) of homogeneous size typically with $3 \mu m \times 3 \mu m \times (40 \text{ to } 60) \mu m$ size range (fig. S12).

Single-crystal preparation methods for MX: IPNS*:Fe:ACV and IPNS:Cd:ACV

Purified recombinant apo-IPNS* or apo-IPNS was used for crystallization. IPNS crystals were grown in 24-well hanging drop VDX plates (Hampton Research, USA), anaerobically for the IPNS*:Fe:ACV and aerobically for the IPNS:Cd(II):ACV complex. The IPNS crystallization solution was prepared by mixing freshly prepared FeSO4 or CdCl₂ (4 µl, 100 mM) with IPNS [in 25 mM tris (pH 8.0), 80 µl of 50 to 52 mg ml⁻¹], followed by a subsequent addition of ACV $[4 \times 5 \mu$ l, 2.1 mg in 20 μ l of 25 mM tris (pH 8.5)]. A screen, varying the pH [0.1 M tris (pH 8.1) to 8.7 in steps of 0.2, vertical axis] and the salt concentration (Li₂SO₄ 1.5 to 2.0 M in steps of 0.1, horizontal axis), was carried out (28). Crystals were prepared using the hanging drop method by combining the reservoir solutions (3 µl) and protein solutions (3 µl). Crystals (60 to 200 µm) formed after 24 to 36 hours were harvested and cryo-cooled by rapid plunging into liquid N2 before data collection. Single IPNS crystals were cryoprotected by transferring to a solution of mother liquor [1.7 M Li₂SO₄ and 0.1 M tris (pH 8.5)] supplemented with 20% (v/v) glycerol before being cryo-cooled in liquid N2. Data for the single crystals were collected at 100 K using synchrotron radiation at the Diamond Light Source beamlines I03, I04, I04-1, and I24 and processed using the Xia2 pipelines (table S1) (44).

Note that formation of IPNS:Cd:ACV crystals under aerobic conditions was only possible when the starting enzyme was completely metal ion free, possibly due to ACV turnover by a small amount of Fe(II)-bound enzyme in solution under aerobic conditions. Needle-shaped crystals were used to prepare seeds using the PTFE Seed Bead Kit as described by the manufacturer (Hampton Research, USA). These crystals were used in batch methods (as described above) to obtain more IPNS:Cd:ACV crystals for NO exposure experiments. Note that while all complexes crystallized with Fe(II) show clear turnover in crystallo and in solution (followed by NMR and mass spectroscopy) (33, 45) after removal from the anaerobic chamber and exposure to O₂ from air, the Cd(II) complex crystallized under aerobic conditions and showed no turnover of ACV to IPN.

Setup and sample injection for crystallographic data collection

Room temperature diffraction data for microcrystal slurries were collected at the MFX (Macromolecular Femtosecond Crystallography) instrument of Linac Coherent Light Source (LCLS) (46, 47) (proposals LU50/P143 and LS34/P110) and at the BL2 beamline at SACLA (proposal 2017B8085) (table S1). The drop-on-tape (DOT) sample delivery method (23) was used combined with ADE at LCLS to obtain anaerobic IPNS:Fe(II):ACV complex and tr O₂-exposed structures (23). The ejected droplets (~3.5 to 4 nl; flow rate, 7 µl min⁻¹) are deposited onto the conveyor belt at room temperature in a helium atmosphere. To trigger the reaction in crystallo, our method exposes the microdroplets of the anaerobic crystal slurry to a 100% O₂ atmosphere as they pass through a 60-mm-long reaction chamber for a varied time regulated by the Kapton belt speed (table S1) (23). We collected tr-SFX datasets after exposing droplets to O₂ and then applying additional reaction times of 400, 500, 800, 1600, and 3000 ms. Shorter reaction times derive from faster belt speeds, which reduces the overall time for O2 diffusion into the microdroplets and hence reaction within the microcrystals. Longer reaction times derive from a slower belt speed and commensurately longer O2 droplet equilibration and reaction times. Detailed information about the Kapton tape speed (millimeter per second) and the O₂ incubation times (millisecond, including incubation time in the reaction chamber and additional travel time of microdroplets from the reaction chamber until they arrive in the x-ray interaction zone) are shown in table S1. The x-ray wavelength for experiments obtained under proposals LU50/P143 was kept at 1.30 Å (9.537 keV) and LS34/P110 at 1.31 Å (9.464 keV) with a data collection rate of 30 Hz, 4 mJ/ pulse, a pulse duration of ~35 fs and an x-ray beam size at the sample of ~3 µm in diameter. XRD data were collected using a Rayonix MX340-HS detector for the LU50/P143 beam time and on a Rayonix MX170-HS detector for LS34/P110 with 4×4 binning.

The IPNS:Fe:IPN product complex SFX data were collected using the viscous extruder and a grease-matrix carrier by first exposing anaerobic slurries of the IPNS:Fe(II):ACV complex (crystal density > 5 × 10⁷ ml⁻¹) to atmospheric O₂ for 30 min and then mixing the slurry with grease in the ratio 10 µl slurry:90 µl grease at SACLA (48). Mixing of grease with slurry was performed by using two 100-µl Hamilton syringes connected over an extruder-based connector. The obtained grease matrix containing randomly oriented protein microcrystals was applied to the sample reservoir (60 µl of reservoir and 2 mm in diameter) of the sample delivery system and mounted in the setup for data collection. A flow rate of 1 to 1.5 μ l min⁻¹ and a nozzle dimension of 100 μ m were used to obtain a stable flow for the protein grease matrix. Data were collected at room temperature with a collection rate of 30 Hz and an x-ray beam size at the sample of ~1.7 µm in diameter. The x-ray wavelength for the experiment was kept at 1.14 Å (11 keV) with 0.34 mJ/

pulse and 10-fs pulse length. XRD data were collected on a MPCCD (MultiPort Charge-Coupled Device) octal detector.

Data processing, model building, and refinement of SFX datasets

During the SFX experiments, data acquisition was tracked with the cctbx.xfel graphical user interface (49), which monitors for new data and submits processing jobs to the computing cluster. The jobs run the core program dials.stills_process to index and integrate the images while providing real-time feedback, as part of the larger cctbx. xfel and DIALS processing suite (50-55). A first estimate of the detector position (distance and beam center) was obtained from a powder diffraction pattern of silver(I) behenate (Alfa Aesar). After initial spot finding of strong reflections, followed by indexing of strong reflections and integration using dials.stills_process, a round of metrology refinement was done (56). A second round of indexing and integration was performed with the refined detector position. Initial merging was performed using PRIME (57) followed by molecular replacement with PHASER (58). This provided a reference model using a target unit cell of a = 41.9 Å, b = 75.7 Å, c = 102 Å, $\alpha = \beta = \gamma = 90^{\circ}$ (space group *P* 2₁2₁2₁). After correction of the integrated intensities for absorption by the Kapton conveyor belt, final data integration and merging were performed using cxi.merge (21). Justification of the resolution cutoff for the merged data was determined on the basis of multiplicity in the highest-resolution shell (>10-fold) and on $CC_{1/2}$ (monotonic decrease) (see tables at the end of the Supplementary Materials with the merging statistics justifying the resolution cutoff) (20, 23).

Structures were solved by isomorphous molecular replacement using the reported structural data file of IPNS [PDB: 1BLZ (26)] as a search model. All structures were iteratively fitted and refined using PHENIX (59) and Coot (60). Processing and refinement statistics for all anaerobically and O2-exposed IPNS structures are given in table S2.

XES analysis

The on-demand ADE/tape drive sample delivery setup enables the simultaneous collection of tr-XES and tr-SFX (23) from the same sample and x-ray pulse to study the Fe oxidation state. We used a wavelength-dispersive von Hamos spectrometer with four cylindrically bent (R = 250 mm) germanium (440) crystals orthogonal to the sample with the center of the crystals located at 75.41° with respect to the interaction point and each focusing the emitted Fe K $\alpha_{1,2}$ x-ray photons onto an ePix100 detector located below the sample x-ray interaction region. Calibration was performed using aqueous solutions of 10 mM Fe(III)(NO₃)₃ and Fe(II)Cl₂ (anaerobically presolutions of 10 mM Fe(III)(NO₃)₃ and Fe(II)Cl₂ (anaerobically prepared and measured) as references. Detector images were sorted by $\begin{bmatrix} 2 & 0 \\ 0 & 0 \end{bmatrix}$ the sample hit rate as described (25). Briefly, for anaerobic, 400-, 500-, 800-, and 1600-ms datasets, two threshold values were set as 3 and 2 for thresholds I and II, respectively. For the 3000-ms dataset, the value for threshold II, which compares the number of photons per unit area inside and outside the region of interest, was reduced to 1. A linear function interpolated across two points outside the region of interest was subtracted from the images for background correction. For spectral analysis, the spectra were area-normalized in the 6380- to 6415-eV range. In this range, the recorded energy was initially spread out by 0.146- to 0.159-eV intervals. To create equal intervals and apply smoothing, the spectra were first linearly splined to a 0.001-eV interval and then processed by a second-order

Savitzky-Golay filter with a 3003-point window size. This window size covers approximately the same range of points as a 19-point window applied to the original intervals. Difference spectra were calculated after smoothing. For calculation of FWHM, the spectral range for K α_1 was selected as 6396 to 6407 eV. Error bars for the calculated FWHM were obtained by a bootstrapping procedure as described (25).

Exposure of anaerobically obtained single IPNS:Fe:ACV crystals to O_2

IPNS:Fe:ACV crystals (needle morphology) were grown anaerobically to the appropriate size $[-5 \times 5 \times (100 \text{ to } 150) \,\mu\text{m}^3]$ (27); the anaerobic microcrystal slurry (10 µl) was transferred into a 500-µl tube. A second 500-µl tube was filled with the IPNS crystallization buffer [100 µl, 1.7 M Li₂SO₄, and 0.1 M tris (pH 8.5) supplemented with 20% glycerol (v/v)], and the solution was saturated with O₂. The anaerobic crystal slurry $(2 \mu l)$ was removed from the glovebox and immediately mixed with the O2-saturated crystallization solution $(8 \mu l)$. The samples were incubated at room temperature and at varying timepoints (30 s to 10 min). Single crystals were mounted on nylon loops and then cryo-cooled by rapid plunging into liquid N₂. Data for a total of ~100 cryo-cooled single crystals of IPNS:Fe:ACV exposed to O2 were collected at the MX beamlines of the Diamond Light Source, UK. From the IPNS:Fe:ACV:O2 datasets, one representative structure was selected for deposition (PDB: 6ZAP), which was refined to 1.36-Å resolution. In this complex, both confs. A and B of α 3 are observed and were refined in 50:50 occupancy.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/ content/full/7/34/eabh0250/DC1

REFERENCES AND NOTES

- H. R. Arnstein, D. Morris, The structure of a peptide, containing α-aminoadipic acid, cystine and valine, present in the mycelium of *Penicillium chrysogenum*. *Biochem. J.* 76, 357–361 (1960).
- N. I. Burzlaff, P. J. Rutledge, I. J. Clifton, C. M. H. Hensgens, M. Pickford, R. M. Adlington, P. L. Roach, J. E. Baldwin, The reaction cycle of isopenicillin N synthase observed by X-ray diffraction. *Nature* 401, 721–724 (1999).
- P. L. Roach, I. J. Clifton, V. Fülöp, K. Harlos, G. J. Barton, J. Hajdu, I. Andersson, C. J. Schofield, J. E. Baldwin, Crystal structure of isopenicillin N synthase is the first from a new structural family of enzymes. *Nature* **375**, 700–704 (1995).
- P. Rabe, J. Kamps, C. J. Schofield, C. T. Lohans, Roles of 2-oxoglutarate oxygenases and isopenicillin N synthase in β-lactam biosynthesis. *Nat. Prod. Rep.* 35, 735–756 (2018).
- C. Q. Herr, R. P. Hausinger, Amazing diversity in biochemical roles of Fe(II)/2-oxoglutarate oxygenases. Trends Biochem. Sci. 43, 517–532 (2018).
- 6. C. J. Schofield, R. P. Hausinger, 2-Oxoglutarate-Dependent Oxygenases, C. J. Schofield, R. P. Hausinger, Eds. (The Royal Society of Chemistry, 2015), pp. 1–487.
- R. Chowdhury, I. K. H. Leung, Y. M. Tian, M. I. Abboud, W. Ge, C. Domene, F. X. Cantrelle, I. Landrieu, A. P. Hardy, C. W. Pugh, P. J. Ratcliffe, T. D. W. Claridge, C. J. Schofield, Structural basis for oxygen degradation domain selectivity of the HIF prolyl hydroxylases. *Nat. Commun.* 7, 12673 (2016).
- C. J. Schofield, P. J. Ratcliffe, Oxygen sensing by HIF hydroxylases. *Nat. Rev. Mol. Cell Biol.* 5, 343–354 (2004).
- J. E. Baldwin, M. Bradley, Isopenicillin N synthase: Mechanistic studies. Chem. Rev. 90, 1079–1088 (1990).
- E. Tamanaha, B. Zhang, Y. Guo, W. C. Chang, E. W. Barr, G. Xing, J. St. Clair, S. Ye, F. Neese, J. M. Bollinger Jr., C. Krebs, Spectroscopic evidence for the two C-H-Cleaving Intermediates of *Aspergillus nidulans* Isopenicillin N Synthase. J. Am. Chem. Soc. 138, 8862–8874 (2016).
- C. D. Brown, M. L. Neidig, M. B. Neibergall, J. D. Lipscomb, E. I. Solomon, VTVH-MCD and DFT studies of thiolate bonding to [FeNO]₇/[FeO₂]₈ complexes of isopenicillin N synthase: Substrate determination of oxidase versus oxygenase activity in nonheme Fe enzymes. J. Am. Chem. Soc. **129**, 7427–7438 (2007).

- V. J. Chen, A. M. Orviller, M. R. Harpel, C. A. Frolik, K. K. Surerus, E. Münck, J. D. Lipscomb, Spectroscopic studies of isopenicillin N synthase. A mononuclear nonheme Fe²⁺ oxidase with metal coordination sites for small molecules and substrate. *J. Biol. Chem.* 264, 21677–21681 (1989).
- M. Lundberg, P. E. Siegbahn, K. Morokuma, The mechanism for isopenicillin N synthase from density-functional modeling highlights the similarities with other enzymes in the 2-His-1-carboxylate family. *Biochemistry* 47, 1031–1042 (2008).
- M. N. Blakely, M. A. Dedushko, P. Chaau Yan Poon, G. Villar-Acevedo, J. Kovacs, Formation of a reactive, alklyl thiolate-ligated FellI-superoxo intermediate derived from dioxygen. J. Am. Chem. Soc. 141, 1867–1870 (2019).
- M. Lundberg, K. Morokuma, Protein environment facilitates O₂ binding in non-heme iron enzyme. An insight from ONIOM calculations on isopenicillin N synthase (IPNS). J. Phys. Chem. B 111, 9380–9389 (2007).
- M. Lundberg, T. Kawatsu, T. Vreven, M. J. Frisch, K. Morokuma, Transition states in a protein environment—ONIOM QM:MM Modeling of Isopenicillin N Synthesis. J. Chem. Theory Comput. 5, 222–234 (2009).
- C. D. Brown-Marshall, A. R. Diebold, E. I. Solomon, Reaction coordinate of isopenicillin N synthase: Oxidase versus oxygenase activity. *Biochemistry* 49, 1176–1182 (2010).
- M. Unno, H. Chen, S. Kusama, S. Shaik, M. Ikeda-Saito, Structural characterization of the fleeting ferric peroxo species in myoglobin: Experiment and theory. *J. Am. Chem. Soc.* **129**, 13394–13395 (2007).
- K. Nass, A. Gorel, M. M. Abdullah, A. V. Martin, M. Kloos, A. Marinelli, A. Aquila,
 T. R. M. Barends, F. J. Decker, R. Bruce Doak, L. Foucar, E. Hartmann, M. Hilpert,
 M. S. Hunter, Z. Jurek, J. E. Koglin, A. Kozlov, A. A. Lutman, G. N. Kovacs, C. M. Roome,
 R. L. Shoeman, R. Santra, H. M. Quiney, B. Ziaja, S. Boutet, I. Schlichting, Structural
 dynamics in proteins induced by and probed with X-ray free-electron laser pulses.
 Nat. Commun. 11, 1814 (2020).
- J. Kern, R. Chatterjee, I. D. Young, F. D. Fuller, L. Lassalle, M. Ibrahim, S. Gul, T. Fransson, A. S. Brewster, R. Alonso-Mori, R. Hussein, M. Zhang, L. Douthit, C. de Lichtenberg, M. H. Cheah, D. Shevela, J. Wersig, I. Seuffert, D. Sokaras, E. Pastor, C. Weninger, T. Kroll, R. G. Sierra, P. Aller, A. Butryn, A. M. Orville, M. Liang, A. Batyuk, J. E. Koglin, S. Carbajo, S. Boutet, N. W. Moriarty, J. M. Holton, H. Dobbek, P. D. Adams, U. Bergmann, N. K. Sauter, A. Zouni, J. Messinger, J. Yano, V. K. Yachandra, Structures of the intermediates of Kok's photosynthetic water oxidation clock. *Nature* 563, 421–425 (2018).
- I. D. Young, M. Ibrahim, R. Chatterjee, S. Gul, F. D. Fuller, S. Koroidov, A. S. Brewster, R. Tran, R. Alonso-Mori, T. Kroll, T. Michels-Clark, H. Laksmono, R. G. Sierra, C. A. Stan, R. Hussein, M. Zhang, L. Douthit, M. Kubin, C. de Lichtenberg, L. Vo Pham, H. Nilsson, M. H. Cheah, D. Shevela, C. Saracini, M. A. Bean, I. Seuffert, D. Sokaras, T. C. Weng, E. Pastor, C. Weninger, T. Fransson, L. Lassalle, P. Bräuer, P. Aller, P. T. Docker, B. Andi, A. M. Orville, J. M. Glownia, S. Nelson, M. Sikorski, D. Zhu, M. S. Hunter, T. J. Lane, A. Aquila, J. E. Koglin, J. Robinson, M. Liang, S. Boutet, A. Y. Lyubimov, M. Uervirojnangkoorn, N. W. Moriarty, D. Liebschner, P. V. Afonine, D. G. Waterman, G. Evans, P. Wernet, H. Dobbek, W. I. Weis, A. T. Brunger, P. H. Zwart, P. D. Adams, A. Zouni, J. Messinger, U. Bergmann, N. K. Sauter, J. Kern, V. K. Yachandra, J. Yano, Structure of photosystem II and substrate binding at room temperature. *Nature* 540, 453–457 (2016).
- C. Gisriel, J. Coe, R. Letrun, O. M. Yefanov, C. Luna-Chavez, N. E. Stander, S. Lisova, V. Mariani, M. Kuhn, S. Aplin, T. D. Grant, K. Dörner, T. Sato, A. Echelmeier, J. Cruz Villarreal, M. S. Hunter, M. O. Wiedorn, J. Knoska, V. Mazalova, S. Roy-Chowdhury, J. H. Yang, A. Jones, R. Bean, J. Bielecki, Y. Kim, G. Mills, B. Weinhausen, J. D. Meza, N. al-Qudami, S. Bajt, G. Brehm, S. Botha, D. Boukhelef, S. Brockhauser, B. D. Bruce, M. A. Coleman, C. Danilevski, E. Discianno, Z. Dobson, H. Fangohr, J. M. Martin-Garcia, Y. Gevorkov, S. Hauf, A. Hosseinizadeh, F. Januschek, G. K. Ketawala, C. Kupitz, L. Maia, M. Manetti, M. Messerschmidt, T. Michelat, J. Mondal, A. Ourmazd, G. Previtali, I. Sarrou, S. Schön, P. Schwander, M. L. Shelby, A. Silenzi, J. Sztuk-Dambietz, J. Szuba, M. Turcato, T. A. White, K. Wrona, C. Xu, M. H. Abdellatif, J. D. Zook, J. C. H. Spence, H. N. Chapman, A. Barty, R. A. Kirian, M. Frank, A. Ros, M. Schmidt, R. Fromme, A. P. Mancuso, P. Fromme, N. A. Zatsepin, Membrane protein megahertz crystallography at the European XFEL. *Nat. Commun.* **10**, 5021 (2019).
- F. D. Fuller, S. Gul, R. Chatterjee, E. S. Burgie, I. D. Young, H. Lebrette, V. Srinivas, A. S. Brewster, T. Michels-Clark, J. A. Clinger, B. Andi, M. Ibrahim, E. Pastor, C. de Lichtenberg, R. Hussein, C. J. Pollock, M. Zhang, C. A. Stan, T. Kroll, T. Fransson, C. Weninger, M. Kubin, P. Aller, L. Lassalle, P. Bräuer, M. D. Miller, M. Amin, S. Koroidov, C. G. Roessler, M. Allaire, R. G. Sierra, P. T. Docker, J. M. Glownia, S. Nelson, J. E. Koglin, D. Zhu, M. Chollet, S. Song, H. Lemke, M. Liang, D. Sokaras, R. Alonso-Mori, A. Zouni, J. Messinger, U. Bergmann, A. K. Boal, J. M. Bollinger Jr., C. Krebs, M. Högbom, G. N. Phillips Jr., R. D. Vierstra, N. K. Sauter, A. M. Orville, J. Kern, V. K. Yachandra, J. Yano, Drop-on-demand sample delivery for studying biocatalysts in action at X-ray free-electron lasers. *Nat. Methods* 14, 443–449 (2017).
- J. Kern, R. Alonso-Mori, R. Tran, J. Hattne, R. J. Gildea, N. Echols, C. Glockner, J. Hellmich, H. Laksmono, R. G. Sierra, B. Lassalle-Kaiser, S. Koroidov, A. Lampe, G. Han, S. Gul, D. DiFiore, D. Milathianaki, A. R. Fry, A. Miahnahri, D. W. Schafer, M. Messerschmidt,

M. M. Seibert, J. E. Koglin, D. Sokaras, T. C. Weng, J. Sellberg, M. J. Latimer, R. W. Grosse-Kunstleve, P. H. Zwart, W. E. White, P. Glatzel, P. D. Adams, M. J. Bogan, G. J. Williams, S. Boutet, J. Messinger, A. Zouni, N. K. Sauter, V. K. Yachandra, U. Bergmann, J. Yano, Simultaneous femtosecond X-ray spectroscopy and diffraction of photosystem II at room temperature. *Science* **340**, 491–495 (2013).

- T. Fransson, R. Chatterjee, F. D. Fuller, S. Gul, C. Weninger, D. Sokaras, T. Kroll, R. Alonso-Mori, U. Bergmann, J. Kern, V. K. Yachandra, J. Yano, X-ray emission spectroscopy as an in situ diagnostic tool for X-ray crystallography of metalloproteins using an X-ray free-electron laser. *Biochemistry* 57, 4629–4637 (2018).
- P. L. Roach, I. J. Clifton, C. M. H. Hensgens, N. Shibata, C. J. Schoffeld, J. Hajdu, J. E. Baldwin, Structure of isopenicillin N synthase complexed with substrate and the mechanism of penicillin formation. *Nature* **387**, 827–830 (1997).
- P. Rabe, J. H. Beale, A. Butryn, P. Aller, A. Dirr, P. A. Lang, D. N. Axford, S. B. Carr,
 T. M. Leissing, M. A. McDonough, B. Davy, A. Ebrahim, J. Orlans, S. L. S. Storm, A. M. Orville,
 C. J. Schofield, R. L. Owen, Anaerobic fixed-target serial crystallography. *IUCrJ* 7, 901–912 (2020).
- P. L. Roach, I. J. Clifton, C. M. Hensgens, N. Shibata, A. J. Long, R. W. Strange, S. S. Hasnain, C. J. Schofield, J. E. Baldwin, J. Hajdu, Anaerobic crystallisation of an isopenicillin N synthase-Fe(II)-substrate complex demonstrated by X-ray studies. *Eur. J. Biochem.* 242, 736–740 (1996).
- H. Nogami, Y. Kato, Diffusion coefficient of oxygen in propylene glycol-water mixtures and glycerol-water mixtures. Yakugaku Zasshi 82, 120–126 (1962).
- G. Vankó, T. Neisius, G. Molnár, F. Renz, S. Kárpáti, A. Shukla, F. M. F. de Groot, Probing the 3D spin momentum with X-ray emission spectroscopy: The case of molecular-spin transitions. J. Phys. Chem. B 110, 11647–11653 (2006).
- V. Srinivas, R. Banerjee, H. Lebrette, J. C. Jones, O. Aurelius, I. S. Kim, C. C. Pham, S. Gul, K. D. Sutherlin, A. Bhowmick, J. John, E. Bozkurt, T. Fransson, P. Aller, A. Butryn, I. Bogacz, P. Simon, S. Keable, A. Britz, K. Tono, K. S. Kim, S. Y. Park, S. J. Lee, J. Park, R. Alonso-Mori, F. D. Fuller, A. Batyuk, A. S. Brewster, U. Bergmann, N. K. Sauter, A. M. Orville, V. K. Yachandra, J. Yano, J. D. Lipscomb, J. Kern, M. Högbom, High-resolution XFEL structure of the soluble methane monooxygenase hydroxylase complex with its regulatory component at ambient temperature in two oxidation states. J. Am. Chem. Soc. 142, 14249–14266 (2020).
- S. Lafuerza, A. Carlantuono, M. Retegan, P. Glatzel, Chemical sensitivity of Kβ and Kα X-ray emission from a systematic investigation of iron compounds. *Inorg. Chem.* 59, 12518–12535 (2020).
- L. A. McNeill, T. J. N. Brown, M. Sami, I. J. Clifton, N. I. Burzlaff, T. D. W. Claridge, R. M. Adlington, J. E. Baldwin, P. J. Rutledge, C. J. Schofield, Terminally truncated isopenicillin N synthase generates a dithioester product: Evidence for a thioaldehyde intermediate during catalysis and a new mode of reaction for non-heme iron oxidases. *Chemistry* 23, 12815–12824 (2017).
- A. M. Rydzik, J. Brem, S. S. van Berkel, I. Pfeffer, A. Makena, T. D. W. Claridge, C. J. Schofield, Monitoring conformational changes in the NDM-1 Metallo-β-lactamase by ¹⁹F NMR spectroscopy. *Angew. Chem. Int. Ed.* **53**, 3129–3133 (2014).
- D. Roelofs, M. J. T. N. Timmermans, P. Hensbergen, H. van Leeuwen, J. Koopman,
 A. Faddeeva, W. Suring, T. E. de Boer, J. Mariën, R. Boer, R. Bovenberg, N. M. van Straalen,
 A functional isopenicillin N synthase in an animal genome. *Mol. Biol. Evol.* 30, 541–548 (2013).
- J. P. Klinman, Dynamically achieved active site precision in enzyme catalysis. Acc. Chem. Res. 48, 449–456 (2015).
- J. E. Baldwin, J. M. Blackburn, J. D. Sutherland, M. C. Wright, High-level soluble expression of isopenicillin N synthase isozymes in *E. coli. Tetrahedron* 47, 5991–6002 (1991).
- I. Pettinati, P. Grzechnik, C. Ribeiro de Almeida, J. Brem, M. A. McDonough, S. Dhir, N. J. Proudfoot, C. J. Schofield, Biosynthesis of histone messenger RNA employs a specific 3' end endonuclease. *eLife* 7, (2018).
- M. R. Thomas, S. G. Boxer, 19F NMR of trifluoroacetyl-labeled cysteine mutants of myoglobin: Structural probes of nitric oxide bound to the H93G cavity mutant. *Biochemistry* 40, 8588–8596 (2001).
- R. K. Lesniak, A. M. Rydzik, J. J. A. G. Kamps, A. Kahn, T. D. W. Claridge, C. J. Schofield, ¹⁹F NMR studies on γ-butyrobetaine hydroxylase provide mechanistic insights and suggest a dual inhibition mode. *Chem. Commun.* 55, 14717–14720 (2019).
- E. van Groesen, C. T. Lohans, J. Brem, K. M. J. Aertker, T. D. W. Claridge, C. J. Schofield, ¹⁹F NMR monitoring of reversible protein post-translational modifications: Class D β-lactamase carbamylation and inhibition. *Chemistry* 25, 11837–11841 (2019).
- S. J. Walmsley, P. A. Rudnick, Y. Liang, Q. Dong, S. E. Stein, A. I. Nesvizhskii, Comprehensive analysis of protein digestion using six trypsins reveals the origin of trypsin as a significant source of variability in proteomics. *J. Proteome Res.* 12, 5666–5680 (2013).
- J. Gault, J. A. C. Donlan, I. Liko, J. T. S. Hopper, K. Gupta, N. G. Housden, W. B. Struwe, M. T. Marty, T. Mize, C. Bechara, Y. Zhu, B. Wu, C. Kleanthous, M. Belov, E. Damoc, A. Makarov, C. V. Robinson, High-resolution mass spectrometry of small molecules bound to membrane proteins. *Nat. Methods* **13**, 333–336 (2016).

- G. Winter, C. M. C. Lobley, S. M. Prince, Decision making in xia2. Acta Crystallogr D 69, 1260–1273 (2013).
- A. Dubus, M. Sami, T. J. N. Brown, C. J. Schofield, J. E. Baldwin, J. M. Frère, Studies of isopenicillin N synthase enzymatic properties using a continuous spectrophotometric assay. *FEBS Lett.* **485**, 142–146 (2000).
- P. Emma, R. Akre, J. Arthur, R. Bionta, C. Bostedt, J. Bozek, A. Brachmann, P. Bucksbaum, R. Coffee, F. J. Decker, Y. Ding, D. Dowell, S. Edstrom, A. Fisher, J. Frisch, S. Gilevich, J. Hastings, G. Hays, P. Hering, Z. Huang, R. Iverson, H. Loos, M. Messerschmidt, A. Miahnahri, S. Moeller, H. D. Nuhn, G. Pile, D. Ratner, J. Rzepiela, D. Schultz, T. Smith, P. Stefan, H. Tompkins, J. Turner, J. Welch, W. White, J. Wu, G. Yocky, J. Galayda, First lasing and operation of an Ångstrom-wavelength free-electron laser. *Nat. Photonics* 4, 641–647 (2010).
- S. Boutet, A. Cohen, S. Wakatsuki, The new macromolecular femtosecond crystallography (mfx) instrument at LCLS. *Synchrotron. Radiat. News* 29, 23–28 (2016).
- M. Sugahara, E. Mizohata, E. Nango, M. Suzuki, T. Tanaka, T. Masuda, R. Tanaka, T. Shimamura, Y. Tanaka, C. Suno, K. Ihara, D. Pan, K. Kakinouchi, S. Sugiyama, M. Murata, T. Inoue, K. Tono, C. Song, J. Park, T. Kameshima, T. Hatsui, Y. Joti, M. Yabashi, S. Iwata, Grease matrix as a versatile carrier of proteins for serial crystallography. *Nat. Methods* 12, 61–63 (2015).
- A. S. Brewster, I. D. Young, A. Lyubimov, A. Bhowmick, N. K. Sauter, Processing serial crystallographic data from XFELs or synchrotrons using the *cctbx.xfel* GUI. *Comput. Crystallogr. News.* **10**, 22–39 (2019).
- N. K. Sauter, J. Hattne, R. W. Grosse-Kunstleve, N. Echols, New Python-based methods for data processing. Acta Crystallogr. D Biol. Crystallogr. 69, 1274–1282 (2013).
- J. Hattne, N. Echols, R. Tran, J. Kern, R. J. Gildea, A. S. Brewster, R. Alonso-Mori, C. Glöckner, J. Hellmich, H. Laksmono, R. G. Sierra, B. Lassalle-Kaiser, A. Lampe, G. Han, S. Gul, D. DiFiore, D. Milathianaki, A. R. Fry, A. Miahnahri, W. E. White, D. W. Schafer, M. M. Seibert, J. E. Koglin, D. Sokaras, T. C. Weng, J. Sellberg, M. J. Latimer, P. Glatzel, P. H. Zwart, R. W. Grosse-Kunstleve, M. J. Bogan, M. Messerschmidt, G. J. Williams, S. Boutet, J. Messinger, A. Zouni, J. Yano, U. Bergmann, V. K. Yachandra, P. D. Adams, N. K. Sauter, Accurate macromolecular structures using minimal measurements from X-ray free-electron lasers. *Nat. Methods* **11**, 545–548 (2014).
- N. K. Sauter, XFEL diffraction: Developing processing methods to optimize data quality. J. Synchrotron Radiat. 22, 239–248 (2015).
- G. Winter, D. G. Waterman, J. M. Parkhurst, A. S. Brewster, R. J. Gildea, M. Gerstel, L. Fuentes-Montero, M. Vollmar, T. Michels-Clark, I. D. Young, N. K. Sauter, G. Evans, DIALS: Implementation and evaluation of a new integration package. *Acta Crystallogr. D Struct. Biol.* 74, 85–97 (2018).
- N. K. Sauter, J. Hattne, A. S. Brewster, N. Echols, P. H. Zwart, P. D. Adams, Improved crystal orientation and physical properties from single-shot XFEL stills. *Acta Crystallogr. D* 70, 3299–3309 (2014).
- D. G. Waterman, G. Winter, R. J. Gildea, J. M. Parkhurst, A. S. Brewster, N. K. Sauter, G. Evans, Diffraction-geometry refinement in the *DIALS* framework. *Acta Crystallogr. D Struct. Biol.* **72**, 558–575 (2016).
- A. S. Brewster, D. G. Waterman, J. M. Parkhurst, R. J. Gildea, I. D. Young, L. J. O'Riordan, J. Yano, G. Winter, G. Evans, N. K. Sauter, Improving signal strength in serial crystallography with *DIALS* geometry refinement. *Acta Crystallogr. D Struct. Biol.* 74, 877–894 (2018).
- M. Uervirojnangkoorn, O. B. Zeldin, A. Y. Lyubimov, J. Hattne, A. S. Brewster, N. K. Sauter, A. T. Brunger, W. I. Weis, Enabling X-ray free electron laser crystallography for challenging biological systems from a limited number of crystals. *eLife* 4, e05421 (2015).
- A. J. McCoy, R. W. Grosse-Kunstleve, P. D. Adams, M. D. Winn, L. C. Storoni, R. J. Read, Phaser crystallographic software. J. Appl. Cryst. 40, 658–674 (2007).
- P. D. Adams, R. W. Grosse-Kunstleve, L. W. Hung, T. R. loerger, A. J. McCoy, N. W. Moriarty, R. J. Read, J. C. Sacchettini, N. K. Sauter, T. C. Terwilliger, PHENIX: Building new software for automated crystallographic structure determination. *Acta Crystallogr. D Biol. Crystallogr.* 58, 1948–1954 (2002).
- P. Emsley, B. Lohkamp, W. G. Scott, K. Cowtan, Features and development of Coot. Acta Crystallogr. D Biol. Crystallogr. 66, 486–501 (2010).
- R. H. B. Smith, A. C. Dar, A. Schlessinger, PyVOL: A PyMOL plugin for visualization, comparison, and volume calculation of drug-binding sites. *bioRxiv* 2019, 816702 (2019).
- R. L. Owen, D. Axford, D. A. Sherrell, A. Kuo, O. P. Ernst, E. C. Schulz, R. J. D. Miller, H. M. Mueller-Werkmeister, Low-dose fixed-target serial synchrotron crystallography. *Acta Crystallogr. D Struct. Biol.* **73**, 373–378 (2017).
- D. Liebschner, P. V. Afonine, N. W. Moriarty, B. K. Poon, O. V. Sobolev, T. C. Terwilliger, P. D. Adams, Polder maps: Improving OMIT maps by excluding bulk solvent. *Acta Crystallogr. D Struct. Biol.* 73, 148–157 (2017).
- C. R. Randall, Y. Zang, A. E. True, L. Que Jr., J. M. Charnock, C. D. Garner, Y. Fujishima, C. J. Schofield, J. E. Baldwin, X-ray absorption studies of the ferrous active site of isopenicillin N synthase and related model complexes. *Biochemistry* **32**, 6664–6673 (1993).

- 65. A. M. Orville, V. J. Chen, A. Kriauciunas, M. R. Harpel, B. G. Fox, E. Munck, J. D. Lipscomb, Thiolate ligation of the active site iron(II) of isopenicillin N synthase derives from substrate rather than endogenous cysteine: Spectroscopic studies of site-specific Cys → Ser mutated enzymes. *Biochemistry* **31**, 4602–4612 (1992).
- S. F. Altschul, W. Gish, W. Miller, E. W. Myers, D. J. Lipman, Basic local alignment search tool. J. Mol. Biol. 215, 403–410 (1990).
- P. Gouet, X. Robert, E. Courcelle, ESPript/ENDscript: Extracting and rendering sequence and 3D information from atomic structures of proteins. *Nucleic Acids Res.* **31**, 3320–3323 (2003).
- K. Valegard, A. C. van Scheltinga, M. D. Lloyd, T. Hara, S. Ramaswamy, A. Perrakis, A. Thompson, H. J. Lee, J. E. Baldwin, C. J. Schofield, J. Hajdu, I. Andersson, Structure of a cephalosporin synthase. *Nature* **394**, 805–809 (1998).
- C. G. Yang, C. Yi, E. M. Duguid, C. T. Sullivan, X. Jian, P. A. Rice, C. He, Crystal structures of DNA/RNA repair enzymes AlkB and ABH2 bound to dsDNA. *Nature* 452, 961–965 (2008).
- R. Chowdhury, M. A. McDonough, J. Mecinović, C. Loenarz, E. Flashman, K. S. Hewitson, C. Domene, C. J. Schofield, Structural basis for binding of hypoxia-inducible factor to the oxygen-sensing prolyl hydroxylases. *Structure* **17**, 981–989 (2009).
- J. R. Horton, A. K. Upadhyay, H. H. Qi, X. Zhang, Y. Shi, X. Cheng, Enzymatic and structural insights for substrate specificity of a family of jumonji histone lysine demethylases. *Nat. Struct. Mol. Biol.* **17**, 38–43 (2010).
- H. Hashimoto, J. E. Pais, X. Zhang, L. Saleh, Z. Q. Fu, N. Dai, I. R. Corrêa, Y. Zheng, X. Cheng, Structure of a Naegleria Tet-like dioxygenase in complex with 5-methylcytosine DNA. *Nature* 506, 391–395 (2014).
- M. A. McDonough, K. L. Kavanagh, D. Butler, T. Searls, U. Oppermann, C. J. Schofield, Structure of human phytanoyl-CoA 2-hydroxylase identifies molecular mechanisms of Refsum disease. J. Biol. Chem. 280, 41101–41110 (2005).
- D. H. Juers, J. Ruffin, MAP_CHANNELS: A computation tool to aid in the visualization and characterization of solvent channels in macromolecular crystals. J. Appl. Cryst. 47, 2105–2108 (2014).

Acknowledgments: We thank the staff at LCLS and SACLA. We also thank the staff at Diamond Light Source UK for beam time at beamline 103, 104, 104-1, and 124 (proposals MX-23459 and MX-19458). The Oxford authors dedicate their work in this manuscript to the late professor J. E. Baldwin who was a massive enthusiast for reactions such as that catalyzed by IPNS which lead to "molecules from Mars." Funding: P.R. thanks the Deutsche Akademie für Naturforscher Leopoldina, Germany, for a postdoctoral fellowship. P.A., A.Bu., and A.M.O. were supported by the Strategic Award from the Wellcome Trust and the Biotechnology and Biological Sciences Research Council and a Wellcome Investigator Award (210734/Z/18/Z) and a Royal Society Wolfson Fellowship (RSWF\R2\182017) to A.M.O. C.J.S. thanks the Ineos Oxford Institute, the Wellcome Trust (106244/Z/14/Z), the Biotechnology and Biological Sciences Research Council and Cancer Research UK for supporting work on oxygenases. J.J.A.G.K. was funded by the EPSRC Synthesis for Biology and Medicine CDT(EP/L015838/1) and a Clarendon Scholarship. J.D.S.L. was funded by the BBSRC (BB/M011224/1) and St. John's College, Oxford. P.A.L. was funded by the National PhD Training Program in Antimicrobial Resistance Research by the Medical Research Foundation (MRF-145-0004-TPG-AVISO). This work was supported by NIH grants GM133081 (to K.D.S.), 1P41GM139687 (to R.A.-M.), GM117126 (to N.K.S.), GM55302 (to V.K.Y.), GM110501 (to J.Y.), and GM126289 (to J.F.K.). The DOT instrument used in this research was funded by Department of Energy (DOE), Office of Science, Office of Basic

Energy Sciences (BES), Division of Chemical Sciences, Geosciences, and Biosciences (to J.F.K., J.Y., and V.K.Y.). XFEL data were collected under proposals LS34, LU50, P143, and P110 at the LCLS, SLAC, Stanford, USA, and under proposal 2017B8085 at the Sub Ångstrom Compact Light Source, SACLA, Japan. The Ravonix detector used at LCLS was supported by the NIH grant S10 OD023453. Use of the LCLS, SLAC National Accelerator Laboratory, is supported by the U.S. DOE, Office of Science, BES, under contract no. DE-AC02-76SF00515. Data processing was performed in part at the National Energy Research Scientific Computing Center, a user facility supported by the DOE, Office of Science, under contract no. DE-AC02-05CH11231. Author contributions: A.M.O., C.J.S., and J.F.K. conceived, planned, and executed the experiments and analyzed the data. P.R. and J.J.A.G.K. prepared the microcrystalline slurries for SFX analysis. P.R. and J.J.A.G.K. prepared the substrates. P.R. prepared the wild-type enzyme with support from J.J.A.G.K. P.R., J.J.A.G.K., and J.D.S.L. prepared the mutant enzyme. I.P. was involved in cloning/optimization of pCold_IPNS. P.R. performed crystallographic data analysis with support from I.C. and M.A.M. P.R., J.J.A.G.K., K.D.S., P.A., C.C.P., T.M.L., P.A.L., A.Bu., S.G., F.D.F., I.-S.K., M.H.C., A.Bh., I.D.Y., L.O.'R., A.S.B., Y.J., S.O., K.T., A.Ba., M.S.H., R.A.-M., V.K.Y., J.Y., J.F.K., and A.M.O. participated in data collections at XFELs, P.R., P.A.L., J.F.K., A.M.O., and C.J.S. were involved in data visualization. P.R. was producing the video. P.R., J.J.A.G.K., P.A., A.Bu., P.A.L., and R.L.O. participated in serial data collections at Diamond Light Source. P.R. performed crvo-data collection at Diamond Light Source with support from J.B. K.D.S., P.A. A.Bu., A.Bh., I.D.Y., A.S.B., and N.K.S. participated in serial crystallography data processing and analysis. K.D.S., C.C.P., H.M., T.F., M.D., U.B., J.Y., J.F.K., and A.M.O. participated in x-ray emission spectroscopy data processing and analysis. C.C.P., S.G., F.D.F., I.-S.K., R.A.-M., V.K.Y., J.Y., and J.F.K. were operating the ADE tape drive. D.S., P.R., and J.J.A.G.K. performed the nondenaturing MS experiments supported by C.V.R. J.J.A.G.K. performed the NMR data collection, processing, and analysis with support of J.D.S.L. and T.D.W.C. P.R., J.F.K., A.M.O., and C.J.S. wrote the manuscript with edits and suggestions from all authors. Competing interests: The authors declare that they have no competing interests. Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. The atomic coordinates and structure factors are deposited in the PDB accession codes: 6ZAE (IPNS:Fe:ACV, SFX, anaerobic), 6ZAF (IPNS:Fe:ACV, SFX, 400-ms O2 exposure), 6ZAG (IPNS:Fe:ACV, SFX, 500-ms O2 exposure), 6ZAH (IPNS:Fe:ACV, SFX, 800-ms O2 exposure), 6ZAI (IPNS:Fe:ACV:O2, SFX, 1600-ms O2 exposure), 6ZAJ (IPNS:Fe:ACV:O2, SFX, 3000-ms O2 exposure, ACV conf. C), 6ZAL (IPNS:Fe:ACV:O2, SFX, 500-ms O2 exposure, no glycerol), 6ZAQ (IPNS:Fe:IPN, SFX), 6Y0P (IPNS:Fe:IPN, SSX), 6ZAO (IPNS:Fe:IPN, MX), 6ZAN (IPNS:Fe:ACV:NO, MX), 6ZAP (IPNS:Fe:ACV:O2, MX), 6ZAM (19F-labeled IPNS*:Fe:ACV, MX), and 6ZW8 (IPNS:Cd:ACV, MX).

Submitted 10 February 2021 Accepted 29 June 2021 Published 20 August 2021 10.1126/sciadv.abh0250

Citation: P. Rabe, J. J. A. G. Kamps, K. D. Sutherlin, J. D. S. Linyard, P. Aller, C. C. Pham, H. Makita, I. Clifton, M. A. McDonough, T. M. Leissing, D. Shutin, P. A. Lang, A. Butryn, J. Brem, S. Gul, F. D. Fuller, I.-S. Kim, M. H. Cheah, T. Fransson, A. Bhowmick, I. D. Young, L. O'Riordan, A. S. Brewster, I. Pettinati, M. Doyle, Y. Joti, S. Owada, K. Tono, A. Batyuk, M. S. Hunter, R. Alonso-Mori, U. Bergmann, R. L. Owen, N. K. Sauter, T. D. W. Claridge, C. V. Robinson, V. K. Yachandra, J. Yano, J. F. Kern, A. M. Orville, C. J. Schofield, X-ray free-electron laser studies reveal correlated motion during isopenicillin N synthase catalysis. *Sci. Adv. 7*, eabh0250 (2021).

ScienceAdvances

X-ray free-electron laser studies reveal correlated motion during isopenicillin N synthase catalysis

Patrick RabeJos J. A. G. KampsKyle D. SutherlinJames D. S. LinyardPierre AllerCindy C. PhamHiroki Makitalan CliftonMichael A. McDonoughThomas M. LeissingDenis ShutinPauline A. LangAgata ButrynJürgen BremSheraz GulFranklin D. FullerIn-Sik KimMun Hon CheahThomas FranssonAsmit BhowmickIris D. YoungLee O'RiordanAaron S. Brewsterllaria PettinatiMargaret DoyleYasumasa JotiShigeki OwadaKensuke TonoAlexander BatyukMark S. HunterRoberto Alonso-MoriUwe BergmannRobin L. OwenNicholas K. SauterTimothy D. W. ClaridgeCarol V. RobinsonVittal K. YachandraJunko YanoJan F. KernAllen M. OrvilleChristopher J. Schofield

Sci. Adv., 7 (34), eabh0250. • DOI: 10.1126/sciadv.abh0250

View the article online https://www.science.org/doi/10.1126/sciadv.abh0250 Permissions https://www.science.org/help/reprints-and-permissions

Use of this article is subject to the Terms of service

Science Advances (ISSN) is published by the American Association for the Advancement of Science. 1200 New York Avenue NW, Washington, DC 20005. The title *Science Advances* is a registered trademark of AAAS.

Copyright © 2021 The Authors, some rights reserved; exclusive licensee American Association for the Advancement of Science. No claim to original U.S. Government Works. Distributed under a Creative Commons Attribution License 4.0 (CC BY).