



HAL
open science

Comparative study of the effects of high hydrostatic pressure per se and high argon pressure on urate oxidase ligand stabilization

Thierry Prangé, Philippe Carpentier, Anne-Claire Dhaussy, Peter van der Linden, Eric Girard, Nathalie Colloc'h

► To cite this version:

Thierry Prangé, Philippe Carpentier, Anne-Claire Dhaussy, Peter van der Linden, Eric Girard, et al.. Comparative study of the effects of high hydrostatic pressure per se and high argon pressure on urate oxidase ligand stabilization. *Acta crystallographica Section D: Structural biology* [1993-..], International Union of Crystallography, 2022, 78 (2), pp.162-173. 10.1107/S2059798321012134 . hal-03542387

HAL Id: hal-03542387

<https://hal-normandie-univ.archives-ouvertes.fr/hal-03542387>

Submitted on 25 Jan 2022

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Comparison study of the effects of high hydrostatic pressure *per se* and high argon pressure on urate oxidase-ligand stabilization

Authors

Thierry Prangé^{a*}, Philippe Carpentier^{bc}, Anne-Claire Dhaussy^d, Peter van der Linden^e, Eric Girard^f and Nathalie Colloc'h^{g*}

^aCiTCoM UMR 8038, CNRS, Université de Paris, Faculté de Pharmacie, 4, Av de l'Observatoire, Paris, 75006, France

^bUniversité Grenoble Alpes CEA CNRS, IRIG-LCBM UMR 5249, Grenoble, 38000, France

^cESRF, the European Synchrotron, Grenoble, 38000, France

^dNormandie Univ., Ensicaen, CNRS, CRISTMAT UMR 6508, Caen, France

^eESRF, PSCM (Partnership for Soft Condensed Matter), Grenoble, 38000, France

^fUniv. Grenoble-Alpes CEA CNRS, IBS, Grenoble, 38000, France

^gISTCT UMR 6030, CNRS Université de Caen-Normandie CEA, centre Cyceron, bd Becquerel, Caen, 14074, France

Correspondence email: thierry.prange@parisdescartes.fr; colloch@cyceron.fr

Synopsis Urate oxidase structures under high hydrostatic pressure *per se* and high argon pressure have been analysed, deciphering the basis of the protein-ligand stabilisation by pressure and the formation of a transient intermediate dimeric sub-state during the pressure-induced dissociation.

Abstract The stability of the tetrameric enzyme urate oxidase in complex with an excess of 8-azaxanthine was investigated either under high hydrostatic pressure *per se* or under a high pressure of argon. The active site is located at the interface of two subunits, and the catalytic activity is then directly related to the tetramer integrity. This study demonstrates that applying pressure to a protein-ligand complex drives the thermodynamic equilibrium towards the ligand saturation of the complex, revealing a new binding site. A transient dimeric intermediate occurring during the pressure-induced dissociation process is characterized under argon pressure and excited sub-states of the enzyme occurring during the catalytic cycle can be trapped by pressure. The comparison of the different structures under pressure infers an allosteric role of the internal hydrophobic cavity where argon is bound, since this cavity gives the necessary flexibility for the active site to function.

Keywords: homotetramer, dissociation, enzymatic mechanism, argon, high pressure, HPMX

1. Introduction

Pressure is a reversible thermodynamic variable, which increases the energy of biochemical systems and hence allows access to their low-lying excited states (Akasaka, 2006; Akasaka *et al.*, 2013; Espada *et al.*, 2016; Frauenfelder *et al.*, 1990, 2007; Winter, 2019; Frauenfelder *et al.*, 2009). Studies of proteins under pressure also give access to their dynamic behavior in excited states, and to properties of their transient conformational states. Many studies using NMR or crystallography under high hydrostatic pressure have demonstrated that these high energy conformers are functionally relevant, designed either as functional or kinetic intermediates (Louis & Roche, 2016; Nielsen *et al.*, 2013; Williamson & Kitahara, 2019; Kalbitzer *et al.*, 2013; Fourme, Girard *et al.*, 2012; Collins *et al.*, 2011; Dhaussy & Girard, 2015; Fourme *et al.*, 2009). These conformational states correspond to local minima in the energy landscape of a given protein along its reaction pathways.

Internal protein cavities present in the native state play an important role in pressure-induced structural modifications. Upon increasing pressure, above a critical value, cavities either collapse or become hydrated. Both processes drive pressure-induced denaturation, and trigger the unfolding process (Collins *et al.*, 2005; Nagae *et al.*, 2012; Xue *et al.*, 2019). However, proteins partial unfolding behavior is already observed at lower pressure (Fuglestad *et al.*, 2017; Nagae *et al.*, 2012; Silva & Weber, 1993; Xue *et al.*, 2019). More specifically, when high pressure is applied to multimeric structures, denaturation usually begins by the dissociation of subunits, leading to lower order oligomeric states (Winter, 2019). This process is likely due to the fact that the association of subunits is stabilized by polar and ionic bonds that are the first to be disrupted by pressure.

Noble gases such as xenon, krypton and more recently argon are known to map hydrophobic tunnels and cavities within proteins (Prangé *et al.*, 1998), deciphering pathways for gaseous ligands like molecular oxygen, carbon monoxide or hydrogen (Colloc'h *et al.*, 2017; Knapp *et al.*, 2009; Kalms *et al.*, 2016; Engilberge *et al.*, 2020; Luna *et al.*, 2012, 2008; Tilton *et al.*, 1984; Whittington *et al.*, 2001; Hahn *et al.*, 2021; Winter *et al.*, 2011; Montet *et al.*, 1997).

In the present study, we investigate the pressure-induced structural effects on the tetrameric enzyme urate oxidase (uricase; EC 1.7.3.3; UOX), with a competitive inhibitor (8-azaxanthine) bound at the interface between subunits. The UOX/8-azaxanthine complex appears to be a suitable model system to study the effect of pressure on the stability of an enzyme-ligand complex.

UOX is involved in the purine degradation pathway and is active as a homotetramer. Without any cofactor, it catalyzes the oxidation of uric acid to 5-hydroxy-isourate in the presence of molecular oxygen, and generates hydrogen peroxide as by-product (Scheme 1). UOX from *Aspergillus flavus* in complex with the competitive inhibitor 8-azaxanthine (8-aza) crystallizes in the orthorhombic space

group I222 with one monomer per asymmetric unit (Colloc'h *et al.*, 1997). The tetramer is composed of two face-to-face 16-stranded β -barrel dimers, delimiting a central tunnel of 50 Å long and 12 Å in diameter. The dimer composed of two face-to-face monomers is termed A-B, while the dimer that forms the 16-stranded β -barrel is termed A-C (Fig. 1, upper part). The active site is located at the interface between monomers A and C, where the ligand (here the 8-aza molecule) is hydrogen-bonded to the side chains of the two tweezer amino acids Arg 176 and Gln 228, and to the main chain nitrogen of the Thr 57* (* strands for the symmetric monomer), and forms a π - π stacking interaction with Phe 159 (Colloc'h *et al.*, 1997). In the peroxo hole, on the opposite side of the phenylalanine, the water molecule responsible for the catalytic hydration step is hydrogen-bonded between side chains of Asn 254 and Thr 57*, acting as peroxo hole tweezers. A well conserved H-bond network connects the ligand to the catalytic water molecule, through two water molecules, the His 256 side chain, and the catalytic diad Lys 10* and Thr 57* side chains. This network was proposed to act as a proton shuttle during the different steps of the reaction mechanism (Gabison *et al.*, 2008, 2010, 2011; Wei *et al.*, 2017; Bui *et al.*, 2014; McGregor *et al.*, 2021).

The functional relevance of a large buried hydrophobic cavity close to the active site in each monomer was probed using both crystallography under moderate pressure of noble gas (less than 5 MPa) and high pressure (Colloc'h & Prangé, 2014; Girard *et al.*, 2010). This cavity was proposed to have an allosteric role acting as a connecting vessel to provide the flexibility necessary to the neighboring active site for function.

Above a critical hydrostatic pressure, which depends on the ligand presence and concentration, the tetramer in complex with 8-aza slowly begins to unfold (Girard *et al.*, 2010). In order to determine the different factors involved in UOX pressure-induced destabilization, we compare the direct effects of hydrostatic pressure *per se* with the effects of argon-mediated pressure on the structure of a complex UOX/8-aza produced in excess of ligand. In the following, the two types of pressure, either hydrostatic pressure *per se* and high argon pressure, will be referred to as HHP and HArP respectively, and each (P, T) state of UOX crystals will be designated by the acronym "UOX-type-pressure in MPa-temperature in K or room temperature (RT)".

This comparative study aims at investigating the first glimpse of the pressure-induced UOX tetramer dissociation, to identify a potential intermediate sub-state, and to confirm the allosteric role of the cavity in the enlargement of the active site necessary for efficient enzymatic activity.

2. Materials and methods

2.1. Crystallization

Purified urate oxidase from *Aspergillus flavus* expressed in *S. cerevisiae* was a gift from the Sanofi Company (Montpellier, France). 8-azaxanthine was purchased from Sigma-Aldrich (Lyon, France). Crystals were routinely prepared at room temperature (291K) by mixing 20 μ L of enzyme (20 mg/mL at pH 7.5-8.0) with 20 μ L of a 8% PEG 4000 solution at the same pH (buffer = Tris/acetate 50 mM), in the presence of a large excess of 8-azaxanthine (10 mg). Large diamond-shape colorless crystals usually grow in about two days. They were stabilized by increasing the PEG concentration to 10 % in the final solution. They usually grow with approximate size 0.4 x 0.6 mm, and were stable at 291 K over several weeks.

2.2. Pressurization cells

High hydrostatic pressure macromolecular crystallography (HPMX) were performed using a Diamond Anvil Cell (DAC) described previously (Fourme *et al.*, 2001; Fourme, Girard *et al.*, 2012; Girard *et al.*, 2007). We used two large aperture DACs designed by Roger Fourme and colleagues, and manufactured by the BETSA Company (Nangis, France). The DACs have an Inconel gasket elastic enough to allow a reversible decrease of pressure, a drilled hole of 400 μ m in diameter and 200 μ m depth and an aperture of 90° allowing a complete data set to be recorded on crystals belonging to highly symmetric space groups in a single rotation. The pressure within the DAC compression chamber was monitored through the pressure-dependent fluorescence of a ruby chip used as an internal probe. We used the DACs to compress UOX crystals at 210 and 310 MPa, for which the pressure transmitter is solely the mother liquor.

At low argon pressure, 3.5 and 6.5 MPa, we used a quartz capillary pressure cell, in which the pressure is transmitted by the static pressurized argon gas (Schiltz *et al.*, 1994). At moderate argon pressure, 12 and 22 MPa, we used the “soak-and-freeze” method as described in (Lafumat *et al.*, 2016). At high argon pressure (60 MPa, 100 MPa, 150 MPa and 200 MPa), we used the same approach but with a two-stage compressor apparatus to reach higher pressure (van der Linden *et al.*, 2014). For these two latter techniques, UOX crystals are compressed by static gaseous argon prior to being frozen in the liquid phase of argon at 77 K while they are still under pressure. They were thereafter depressurized at cryogenic temperature.

2.3. Data recording and processing

Three high hydrostatic pressure (HHP) data collections with a DAC were recorded at room temperature on UOX in the presence of an excess of 8-aza (UOX-HHP-0.1-RT as a reference, UOX-HHP-210-RT and UOX-HHP-310-RT) at the CRISTAL beamline (SOLEIL synchrotron, Gif sur Yvette, France), with a Rayonix SX-165 detector at an incident X-ray wavelength $\lambda = 0.58183$ Å (21 keV), a beam size of 50 μ m x 50 μ m and a crystal-to-detector distance of 124.5 mm. Data collection at short wavelengths

reduces the absorption of the diamond windows and increases the accessible reciprocal space allowing higher resolution to be reached (Fourme, Honkimäki *et al.*, 2012).

Particular attention was paid to minimize radiation damage for data collections at room temperature, in order to avoid misinterpretations due to potential unwanted X-ray induced structural changes. Figure S1 illustrates the typical strategy used to collect data on a large crystal which fitted the DAC cavity well. The DAC is placed on a support and is vertically rotated with an available rotation range of -45° to 45° thanks to the large geometrical opening of our pneumatic DAC (Fourme *et al.*, 2001; Fourme, Girard *et al.*, 2012). Prior to data collection, the cavity is centered with X-rays using a highly attenuated beam (dashed square in Fig S1). A typical data collection is thus performed with $50\ \mu\text{m} \times 50\ \mu\text{m}$ beam size on several zones (blue squares) using to a 10° rotation at each position.

Hence, large crystals of UOX ($300 \times 300 \times 180\ \mu\text{m}^3$) that fit optimally into the DAC cavity were selected to have maximum of diffracting material and power. With such large crystals compared to the small beam, full data collection on a single crystal at a given pressure were performed by merging 3 to 5 partial datasets of 20 to 30° rotation ranges collected on well separated zones with at least $50\ \mu\text{m}$ of translation between each of them. In the present study, a single crystal was exploited for data measurements at ambient pressure by collecting 20° data from each of 5 zones, and a second crystal was exploited for data measurements at high pressure by collecting 30° data on each of 3 zones situated on one side of the crystal, at 210 MPa and 30° each from 3 zones on the other side of the sample at 310 MPa. This corresponds to a translation of more than $100\ \mu\text{m}$ between the two sample parts used for data collection. Moreover, owing to the diffracting power of the bulky crystals and to the orthorhombic I222 space group, the complete data set was collected with an exposure time of 4 s, an oscillation range of 1° per frame over a full rotation of 90° , and an X-ray beam flux of 1.9×10^{10} photon/s (26 fold attenuated) at 21keV.

The two data collections under argon pressure in quartz capillaries were performed at room temperature on UOX in the presence of an excess of 8-aza (UOX-HArP-3.5-RT and UOX-HArP-6.5-RT) with a MAR CCD Mosaic detector at the MAX II synchrotron, beamline I711 (Lund, Sweden), with 1° rotation per frame and a full rotation of 120° . The six other data collections under argon pressure (12, 22, 60, 100, 150 and 200 MPa) were performed at 100 K on UOX in the presence of an excess of 8-aza (UOX-HArP-12-T100, UOX-HArP-22-T100, UOX-HArP-60-T100, UOX-HArP-100-T100, UOX-HArP-150-T100, and UOX-HArP-200-T100) using the flash-cooled crystals recovered from the pressurization devices, with a Pilatus 6M detector at the ESRF beamline ID30B (Grenoble, France)(McCarthy *et al.*, 2018), with 0.1° rotation per frame and the same range of 120° . To discriminate argon atoms from water molecules or other ions, the six data sets were collected at a wavelength of $1.7\ \text{Å}$, to locate argon atoms using their anomalous signal at low energy (Fig. S2).

All data sets were processed using the XDS software (Kabsch, 2010). The resolution limits were determined using various criteria in the highest bin, including a $I/\sigma(I)$ ratio around 2, and a R_{meas} lower

than 80 %. The integrated intensities were scaled and merged with SCALA and TRUNCATE, and the structures were solved by molecular replacement with MOLREP, all from the CCP4 suite of programs (Winn *et al.*, 2011). Molecular replacement was performed using the pdb *2iba* structure as the starting model for the data in the I222 space group and a reconstructed dimer of *2iba* for the data in the P2₁2₁2 space group. Argon sites and their occupancy were initially estimated from their peaks in anomalous $\Delta f'' \pm$ maps, using the neighboring sulfur atoms (from Met and Cys residues) as calibration (Fig. S2). All data were refined with REFMAC5 (Murshudov *et al.*, 1997) and the resulting models analyzed and rebuilt with the graphic tool Coot (Emsley *et al.*, 2010). All water molecules in the final models were kept only if they were located at correct H-bond distances and if their unconstrained *B*-factors remained lower than 60 Å².

The data processing and refinement statistics are given in the Tables 1 and 2, respectively. The final models and associated data sets were deposited in the Protein Data Bank (the pdb files IDs are reported at the end of Table 2).

Calculations of cavity volumes were performed using VOIDOO (Kleywegt & Jones, 1994) with a grid spacing of 0.6 Å and a probe radius of 1.1 Å, using 20 randomly generated orientations of the protein model. Calculations of thermodynamic parameters were performed using the PISA program (Krissinel, 2015) from the CCP4 package. Figures were prepared using PYMOL (Version 2.4 Schrodinger, LLC).

3. Results

3.1. UOX structures under high hydrostatic pressure *per se* (HHP)

Although recorded at different synchrotrons, the consistency and reproducibility of all the experiments under pressure are illustrated by the similar linear dependencies of the unit cell volume of UOX with pressure (Fig. S3).

In the present study, three structures of UOX in the presence of an excess of 8-aza were refined from data sets collected at the CRISTAL beamline: a reference data set at ambient pressure (UOX-HHP-0.1-RT), and two high pressure data sets at 210 and 310 MPa (UOX-HHP-210-RT and UOX-HHP-310-RT respectively). The pressure degrades the quality of the crystal, with a resolution decrease from 1.9 Å to 2.15 Å and 2.4 Å for the UOX-HHP-0.1-RT, UOX-HHP-210-RT and UOX-HHP-310-RT structures respectively, while the average thermal *B*-factors increase from 21 Å² to 29 Å² and 49 Å². Consistently, the number of modelled water molecules decreases from 261 to 179 and 115. We observe a clear destabilization of UOX at 310 MPa, which is close to the pressure of denaturation since at 340 MPa UOX crystals weakly diffract and dissolve at 400 MPa.

The estimated dose per irradiation zone was calculated using the program RADDPOSE-3D (Bury *et al.*, 2018), taking the experimental parameters described above into account. The average dose on each exposed region was evaluated to be about 11 kGy (Average Dose Exposed region) and the maximum

dose to be 95 kGy. Hence, the present data collections were recorded with absorbed doses one order of magnitude below the established values that should not be exceeded to solve structures at room temperature (100 – 500 kGy) (Leal *et al.*, 2013; de la Mora *et al.*, 2020). A recent study experimentally demonstrated that, when structures are solved at room temperature with low dose, minimizing global radiation damage, the specific radiation damage that would modify the structures are of much less concern (Gotthard *et al.*, 2019). This ensures that the differences between the three UOX-HHP-RT structures, which are discussed below, are unlikely to be related to radiation damage issues. This strategy was reinforced by a recent study experimentally showing the benefits of using high X-ray energies for macromolecular crystallography (Storm *et al.*, 2021).

The structural differences between ambient and high-pressure structures remain small. The UOX-HHP-0.1-RT structure displays an r.m.s deviation on the C α of 0.18 Å with UOX-HHP-210-RT and 0.19 Å with UOX-HHP-310-RT (Fig. S4). The main difference between these structures is, as expected, the diminution of the volume of the internal hydrophobic cavity, of 38 % between UOX-HHP-210-RT and UOX-HHP-0.1-RT. This decrease is consistent with that of 16 % at 150 MPa compared to ambient pressure described in previous work (Girard *et al.*, 2010). At 310 MPa, there is no further contraction, but a slight expansion of 8 % compared to UOX-HHP-210-RT. However, the calculation of the cavity volume might be imprecise for a structure at such a moderate resolution. No electron density for any water molecules that could have penetrated the cavity was evidenced whatever the applied pressure.

The main chain overall thermal *B*-factors increased on average by 9 Å² between the two UOX-HHP-0.1-RT and UOX-HHP-210-RT structures. Interestingly, amongst the 47 least pressure-sensitive residues (*i.e.* with *B*-factors increasing by less than 5 Å²), 13 are located along the central tunnel and 19 belong to the A-C interface forming the β -barrel, among which 10 are specifically located in the active site. In contrast, 23 amongst the 44 most pressure-sensitive residues (*i.e.* with *B*-factors increasing by more than 13 Å²) belong to the face-to-face A-B interface (Fig. 1, lower part). There is a large increase of average *B*-factor (19 Å²) between the two UOX-HHP-210-RT and UOX-HHP-310-RT structures, revealing that the tetramer becomes destabilized by pressure. Here also, half of the most sensitive residues (*i.e.* with *B*-factors increasing by more than 24 Å²) belong to the A-B interface (Fig. S5). This behavior strongly suggests that the pressure-induced destabilization appears precisely at this face-to-face interface, which therefore represents the weak point of the quaternary structure leading to the disruption of the tetramer.

The active site, located at the β -barrel A-C interface, is clearly enlarged by pressure. This expansion can be quantified by the distance between the residues Asn 254 and Thr 57* (the clamp holding the catalytic water termed W1) that increases from 5.35 Å (UOX-HHP-0.1-RT) to 5.63 Å (UOX-HHP-210-RT) and to 5.88 Å (UOX-HHP-310-RT). Notably, there is a backward displacement of Asn 254 side chain that disrupts the H-bond with W1 (distance of 2.8, 3.2 Å and 3.4 Å for UOX-HHP-0.1-RT, UOX-HHP-210-RT and UOX-HHP-310-RT respectively), as shown in Fig. 2.

On the other hand, the positions of 8-aza and the catalytic water W1 above the inhibitor are observed to be very stable. Similarly, the conserved network of H-bonds linking 8-aza to the catalytic water molecule W1, through two conserved water molecules (8-aza→W2→W3→ His256→Lys 10*→ Thr 57*→W1), remains almost unmodified, except the W2-W3 distance that expands from 3 Å (UOX-HHP-0.1-RT) to 3.3 Å (UOX-HHP-210-RT and UOX-HHP-310-RT).

Interestingly, at high pressure, a second 8-aza molecule is observed at the entrance of the channel leading to the active site, while this latter is still occupied by the first ligand. The occupancies of this second ligand are estimated to be 50 % and 40 % in the refined structures UOX-HHP-210-RT and UOX-HHP-310-RT respectively. This second inhibitor is undetectable at 0.1 MPa. It is located between the two subunits of the β -barrel and stabilized through a π -stacking interaction with Phe 258, and also loosely stabilized by the formation of H-bonds with Asp 58* and Lys 61* side chains. A water molecule located above the ligand is H-bonded to the Asp 58* side chain, an environment that closely mimics the active site. In addition, a network of hydrogen-bonds links the two inhibitors, from the N9 atom of the first 8-aza molecule to the O6 atom of the second one through two water molecules, W2 and a new one termed here W4 (Fig. 3).

3.2. UOX structures under argon pressure (HArP)

Previous studies of UOX structures under pressurized noble gas have shown that the main gas binding site is the large internal hydrophobic cavity close to the active site. At higher pressure, Xe and Kr also bind at the bottom of a surface pocket within each subunit with a low occupancy (Colloc'h *et al.*, 2011; Marassio *et al.*, 2011). In the present study, 8 structures of UOX in presence of an excess of 8-aza under Ar pressure were refined (UOX-HArP-3.5-RT, UOX-HArP-6.5-RT, UOX-HArP-12-T100, UOX-HArP-22-T100, UOX-HArP-60-T100, UOX-HArP-100-T100, UOX-HArP-150-T100, and UOX-HArP-200-T100) showing that argon binds to the internal cavity with occupancies increasing with the applied pressure from 20 % at 3.5 MPa to nearly saturation at 150 MPa (Fig. S6). Because of its high critical pressure, argon becomes visible in the minor binding site only at 12 MPa with an occupancy of 25 % that increases to 50 % at 100 MPa and 80 % at 150 MPa. No other binding sites are evidenced in the anomalous maps, demonstrating that noble gases have mainly two binding sites in the monomer of UOX (Fig. S2 and S7).

Interestingly, a high pressure of argon of 60 MPa and above induces a loss of two of the 222-fold symmetries in the α_4 homotetramer, leading to a less symmetric $(\alpha\beta)_2$ homotetramer. Consequently, the crystal space group changes from I222 to P2₁2₁2 (Fig. 4, upper part). This HArP-induced disruption of the symmetry at the A-B interface is interpreted as the very first step of pressure-induced dissociation of the tetrameric organization into a transient dimeric state constituted by the β -barrel A-C. The structural differences related to the $\alpha_4 \rightarrow (\alpha\beta)_2$ transition within the tetramer are quantified by the least-

squares deviations between main chain atoms of the A subunit with those of the B subunit within the AB dimer (Fig. 4, lower part). Surprisingly at a local level, the r.m.s. displacements appear rather low, while noticeable shifts are observed in five short regions, with a prominence for residues 123-125 in the A-B interface. In the I222 habit of the α_4 tetramer, the 123-125 residues are in close contact with their symmetric mates. Hence, this segment needs to be refined as a two-orientation disordered loop to avoid unwanted steric clashes. This disorder is likely due to the averaging of the 222 local symmetry of the α_4 tetramer over the 222 crystallographic symmetry. In the $(\alpha\beta)_2$ tetramer, the two-fold symmetry related to the segment 123-125 is no longer affected by a steric clash. This disorder is thus fully resolved and the segment 123-125 can be perfectly traced in the two independent chains.

Regarding the other striking regions, the two loops 50-52 and 264-267 are involved in the crystalline packing interface and the loop 196-202 is involved in the A-B interface. These three loops can be considered as the mobile parts of the monomer that further make the tetramer inner 222 symmetries to fit the crystallographic symmetries as well as possible. The residue Glu 22 is embedded in the loop 21-26 which is always very disordered.

In the eight HArP structures, the active site remains unmodified whatever the argon pressure. The distance between the peroxo hole tweezers also remains unchanged (distance of 5.4 Å between Asn 254 and Thr 57* side chains), the H-bond between the catalytic water W1 and Asn 254 is unaffected, and the H-bond between W2 and W3 is conserved, contrary to what is observed in the HHP structures. However, and similarly to the HHP structures, a second 8-aza molecule becomes visible in the electron density map at 150 MPa and 200 MPa, bound at the entrance of the channel leading to the active site. The occupancy of this second ligand is estimated to be about 30 % in UOX-HArP-150-T100 and about 60 % in UOX-HArP-200-T100. The same H-bond network links the two 8-aza molecules through W2 and W4 (Fig. 3) similarly to what is observed in the HHP structures.

4. Discussion

4.1. Argon labelling

Usually, argon labelling, like other noble gases, induces only marginal structural changes since it maintains a good isomorphism between labelled and non-labelled structures (Schiltz *et al.*, 2003; Vitali *et al.*, 1991). This results in very low r.m.s. deviations when comparing their C α traces. The noble gas occupancies in binding sites of a given protein depend on both the atom physicochemical properties and the applied pressure. It can usually be modelled by the Hill-Langmuir isotherm saturation curve (Marassio *et al.*, 2011) that can be written as:

$$f = \frac{f_{\max} \left(\frac{P}{P_c}\right)}{1 + \left(\frac{P}{P_c}\right)^a}$$

where P_c is the critical pressure at which the occupancy f reaches half of its saturation ($f_{max}/2$) and α is the curve steepness coefficient (or sigmoidality that reflects a cooperative binding). For a given site, the value of P_c depends essentially on both the Henry's coefficient of the dissolved gas in the protein solution and on the gas polarizability (*i.e.* the protein/noble-gas interaction strength). P_c is thus ranked as: $P_c^{Xe} \ll P_c^{Kr} \ll P_c^{Ar}$. Experimentally, the critical pressure to populate a given protein binding site is observed to be rather low for xenon ($\sim 0.5 - 3$ MPa), intermediate for krypton ($\sim 1 - 20$ MPa), and high for argon ($\sim 50 - 200$ MPa). In a given binding site, the noble gas occupancies in UOX are, as expected, in the order of their polarizabilities (Fig. S6)(Quillin *et al.*, 2000; Colloc'h *et al.*, 2011).

Contrary to what happens in many cases where the number of binding sites increases with pressure (Colloc'h *et al.*, 2017; Kalms *et al.*, 2016; Engilberge *et al.*, 2020; Prangé *et al.*, 1998), we only observed two noble gas binding sites in UOX whatever the pressure. The main noble gas binding site highlights a putative dioxygen storage site, since the hydrophobic cavity where it binds was proposed to serve as a transient reservoir for dioxygen on its way to the active site (Colloc'h & Prangé, 2014). The minor binding site is a surface pocket largely deepened in the presence of the gas and located in the vicinity of residues lining the central tunnel. Interestingly, this pocket is also deepened in UOX-HHP-210-RT structure (but not in UOX-HHP-310-RT), with however no exogenous molecule bound into it, revealing the intrinsic dynamics of this pocket.

4.2. HARp reveals the first step of tetramer dissociation

The most important feature observed in UOX HARp structures is a transition from the body-centered space group of the crystals to a less-ordered primitive space group. The transition $I222 \rightarrow P2_12_12$ is triggered by argon at a pressure between 22 MPa and 60 MPa. This corresponds to a loss of a symmetry between the four equivalent monomers, according to the $(\alpha)_4 \rightarrow (\alpha\beta)_2$ scheme.

This transition is not observed in the HHP structures when the pressure transmitter is solely the mother liquor. However, we can speculate that the cluster of pressure-sensitive residues along the A-B interface could be indicative of the future loss of the two-fold symmetry between these two monomers. Subdomains that are the most pressure sensitive usually indicate the presence of a mispacked state. In our case, this would correspond to the region encompassing the Asp 123 residue.

A thermodynamic analysis of the UOX monomer by PISA shows that the tetramer is, as expected, the most stable assembly ($\Delta G = -164$ kcal/mol), and that both dimers are equally stable with a $\Delta G = -50$ kcal/mol. Interestingly, their free energy of assembly dissociation is lower for the face-to-face A-B dimer ($\Delta G_{diss} = 30$ kcal/mol) than for the β -barrel A-C dimer ($\Delta G_{diss} = 42$ kcal/mol), showing that the face-to-face dimer is thermodynamically less stable than the β -barrel dimer. In line with these calculations, the structures under very high argon pressure (60 MPa and above) confirm that the face-to-face dimer A-B seems to be less stable than the β -barrel dimer A-C, since pressure induces the

disruption of the tetramer symmetry precisely at the A-B interface. We can then suggest, based on the loss of symmetry induced by a high pressure of argon, that pressure would begin to dissociate the tetrameric UOX *via* the formation of an intermediate A-C sub-state constituted by the most stable 16-stranded β -barrel.

In contrast, it has been shown recently that controlled dehydration of UOX crystals from *bacillus sp. TB-90*, using the humid air and glue-coating method, induces a phase transition from a $P2_12_12$ space group to a $I222$ space group (Hibi *et al.*, 2014, 2016; Hibi & Itoh, 2021). Dehydration has an effect opposite to pressure, inducing a modification from a homotetramer with the four chains in the asymmetric unit to a more ordered tetramer with the 16-stranded β -barrel dimer in the asymmetric unit.

4.3. Comparison between HHP and HArP effects at 200 MPa

The difference between hydrostatic pressure *per se* and high pressure induced by noble gas has been mainly studied in the context of high pressure nervous syndrome studies, to discriminate between the pure pressure effect and the pressure effect of breathing helium by divers. Theoretical studies on the NMDA receptor have shown that a pressure of 2.5 MPa helium induces more deformation than a 2.5 MPa pressure *per se*, mainly through protein-lipid interactions (Bliznyuk *et al.*, 2019). Although our globular model cannot be compared to a membrane-embedded protein, it is interesting to note that argon pressure induces a disruption of the symmetry not observed at a similar high pressure *per se*.

The UOX-HArP-200-T100 structure is closer to the UOX-HHP-210 RT structure than to the UOX-HHP-310 RT structure (Fig. S4), suggesting that a part of the Ar pressure-induced structural modifications at 200 MPa may arise from the direct effect of pressure (*per se*), even if it is difficult to take into account the effect due to the difference in data collection temperatures (100 K and RT).

4.4. Tetramer stabilization by the inhibitor bound at interface

A protein interacts with a ligand through a spatial reorganization, implying that the complexation process is associated with a volume change. Thus, applying pressure, which is thermodynamically conjugated to volume, modifies *de facto* the protein/ligand interaction by adjusting voids at the interface between the protein and its ligand. Pressure is a way to induce a denser packing, with a smaller volume at the interface, leading to a more stable complex. Moreover, pressure by exploring the conformational landscape could favor a sub-state corresponding to a local minima which in turn favors ligand binding (Levin *et al.*, 2019; Luong *et al.*, 2015; Winter, 2019).

We have previously shown by fluorescence spectroscopy experiments that UOX displays an enhanced pressure stability upon inhibitor binding, in a concentration-dependent manner (Girard *et al.*, 2010). Without inhibitor, the pressure-induced dissociation of the tetramer (denaturation) was observed around 175-200 MPa. This limit is shifted toward higher pressure depending on the concentration of the

inhibitor, around 220–250 MPa with an equimolar concentration and around 350 MPa with an excess of 8-aza. The previously determined crystallographic structure of UOX under a hydrostatic pressure of 150 MPa (pdb *3f2m*) was obtained in the presence of an equimolar concentration of 8-aza. With such a ratio, the diffraction was lost above 200 MPa. The stabilizing effect of 8-aza in excess is here confirmed, since diffraction was now lost at about 400 MPa. This result highlights that the ligand bound at the interface strengthens the dimer and further protects the tetrameric organization against pressure effects. Moreover, both HHP and HArP show that a second inhibitor molecule is stabilized in the channel connecting the solvent to the active site. The Lys 61* residue, which is H-bonded to this second 8-aza, is amongst the least pressure-sensitive residues, confirming that pressure consolidates this area.

In the previous study of UOX at 150 MPa in the presence of an equimolar concentration of 8-aza, we observed that three loops surrounding the channel leading to the active site were particularly pressure-sensitive with the largest *B*-factor increases (Girard *et al.*, 2010). In the presence of an excess of 8-aza, the *B*-factor increase of two of these loops (51 – 55 and 161 – 173) remains average, confirming that 8-aza in excess stabilizes the active site against the effect of pressure. On the other hand, the loop 260 – 275, which is part of the face-to-face A-B interface, still displays a high pressure sensitivity, in line with a destabilization of this interface. It is interesting to note that the two loops 51–55 and 260–275 are also those which differ between the two A and B monomers when the symmetry is disrupted.

4.5. Differences of temperature in data collections

HHP data collections within a DAC and HArP data collections in capillaries below 7 MPa were performed at room temperature, but at 100 K for HArP data collections at 12 MPa and above. The mosaicity of the crystals is indeed affected upon freezing (Table 1). The structural modifications and the intrinsic dynamics of flexible loops induced by pressure lead to a degradation of the order and resolution together with an associated increase of thermal *B*-factors. This is especially visible at 310 MPa, a pressure quite close to that which leads to a total loss of diffraction. On the other hand, the freezing of crystals under high pressure of Ar leads experimentally to immobilise transient sub-states in crystals, which thus display diffraction to higher resolution. Hence, the second 8-aza becomes clearly more visible in UOX-HArP-200-T100 structure than in both UOX-HHP-210-RT and UOX-HHP-310-RT MPa structures. However, the treatment under argon pressurization followed by a depressurization step then a freezing step is quite difficult to conduct, leading sometimes to a degradation of the quality of diffraction.

4.6. HHP reveals an excited sub-state of UOX catalytic cycle

The enzyme conformation in the UOX-HHP-210-RT and UOX-HHP-310-RT structures, that display a broader active site characterized by an elongated peroxo hole tweezers (Asn 254 – Thr 57*), can

correspond to a sub-state where the active site needs to accommodate a larger ligand. This occurs during the catalytic mechanism, when the non-planar 5-hydroxy-isourate product is bound before release. The larger distance between Asn 254 and Thr 57* also corresponds to a sub-state in which molecular oxygen is bound in the active site in place of the catalytic water (Colloc'h & Prangé, 2014).

In previous work, it was proposed through the dissection of the complete UOX catalytic reaction that a base-oxidative-system (BOS) plays a central role in the mechanism (Gabison *et al.*, 2011). It consists of the peroxo hole tweezers Thr 57*/Asn 254 acting together with the triad Lys 10*/His 256/Lys 61* which would deprotonate and then extract an electron from uric acid. The clear stabilization of Lys 61* upon pressurization suggests again that pressure stabilizes an active sub-state of UOX that occurs along the catalytic cycle.

Interestingly, the proton shuttle mechanism (Gabison *et al.*, 2010, 2011), which runs from 8-aza to W1 through W2, W3, His 256, Lys 10* and Thr 57* is weakened at 210 and 310 MPa. Indeed, at these pressures, the H-bond between W2 and W3 becomes elongated, since W2 is engaged in an H-bond with W4 that interlinks the two inhibitors. The HHP structures thus suggest the existence of a mechanism with two sub-states, a sub-state that processes the first ligand through the proton pathway exchange (the proton shuttle) and a sub-state where an H-bond system stabilizes the entrance of a second ligand.

Similarly, Asn 254, which is displaced backward by pressure, has been shown to explore different conformations. It has been proposed that Asn 254 dynamics could modulate W1 stability in the peroxo hole and favor its displacement out of the peroxo hole (McGregor *et al.*, 2021). It is worth noting that this Asn residue is located between the active site and the hydrophobic cavity proposed to be a transient reservoir for di-oxygen close to the peroxo hole (Colloc'h & Prangé, 2014). The HHP structures of UOX could then represent excited substates where Asn 254 moves backward to allow the di-oxygen to migrate from the hydrophobic cavity to the peroxo-hole.

We have suggested that the swelling of the active site can occur through the contraction of the neighboring hydrophobic cavity that has been proposed to act as a connecting vessel to give flexibility to the active site (Colloc'h & Prangé, 2014; Girard *et al.*, 2010). Its role is confirmed here in both the HHP and HARp structures. In the UOX-HHP-210-RT structure, the active site is enlarged, while the hydrophobic cavity volume is reduced. Conversely, in the UOX-HARp-200-T100 structure, the binding of argon in the cavity prevents its contraction and thus induces no enlargement of the active site. This confirms that argon restrains the intrinsic flexibility of the cavity. Consequently, the proton shuttle is not weakened when Ar is bound in the internal cavity, as was the case in HHP structures.

Accordingly, the argon atom bound in the hydrophobic cavity prevents the enlargement of the active site, thus stabilizing the UOX tetramer against pressure effects.

4.7. Pressure inhibits substrate affinity of the enzyme

Pressure has been shown to both enhance or inhibit enzymatic activities (Eisenmenger & Reyes-De-Corcuera, 2009; Akasaka *et al.*, 2008; Winter, 2019). For example, it can promote or hamper a

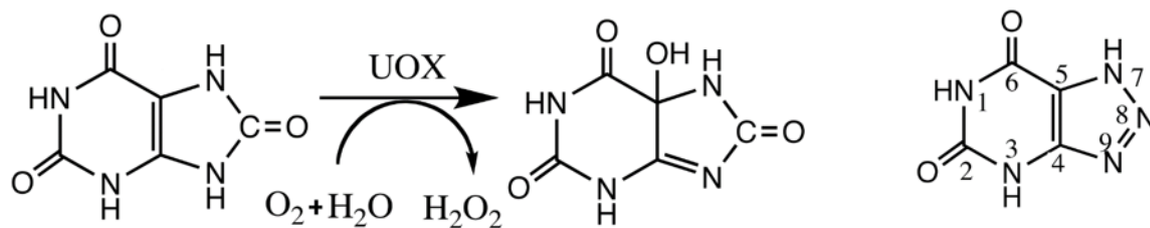
catalytically active state of an enzyme with a better/lower ligand binding affinity or improve/reduce reaction rates by increasing/decreasing conformational flexibility. In the present case, enzymatic assays under pressure have previously shown that pressure decreases the enzymatic activity of UOX with a loss of substrate affinity, while the catalytic efficiency was unmodified and the protein was still tetrameric (Girard *et al.*, 2010). Moreover, it was shown by electron spin resonance spectroscopy and quantum mechanical studies that residues from the face-to-face A-B interface participate in the stabilization of the active site (Gabison *et al.*, 2011). The loss of substrate affinity induced by pressure might thus be explained by both the contraction of the allosteric cavity leading to the swelling in the active site and the destabilization of the face-to-face A-B interface.

5. Conclusion

Both ways of applying pressure, either *per se* (HHP) or noble-gas mediated (HArP), provide complementary information about the UOX mechanism. The UOX-HArP structures show that upon pressure, UOX does not transit directly from a tetrameric association to a monomeric form, but *via* a transient dimeric intermediate state constituted of the 16-stranded β -barrel built by the A and C chains. Very high argon pressures (above 22 MPa) thus allow the characterization of this intermediate excited sub-state with a partial unfolding behavior.

This study also highlights that high pressure applied to a protein-ligand complex, whatever the transmission medium, shifts the thermodynamic equilibrium towards the completion of the complex formation, according to Le Chatelier's principle. Pressure leads to the saturation of this complex and populates new specific sites within the protein. The comparison between the structures under high hydrostatic pressure *per se*, in which the cavity is empty, and the structures under high argon pressure, in which argon is bound, demonstrates the role of the allosteric cavity that provides the functional flexibility to the active site, necessary for the catalytic process.

High pressure NMR experiments have widely been used to characterize oligomeric states upon unfolding and to analyze structural perturbations and unfolding transitions. Here we show that crystallography under pressure (HPMX) is a valuable complementary tool which gives additional understanding in analyzing the structural perturbations at atomic level, and is even able to catch the first steps of the oligomeric dissociation preceding denaturation in the case of multimeric proteins like urate oxidase.



Scheme 1 *Left*: The reaction catalysed by urate oxidase (UOX) from uric acid to 5-hydroxy-isourate.

Right: The analogue 8-azaxanthine in which the 8-carbonyl group is replaced by a nitrogen atom.

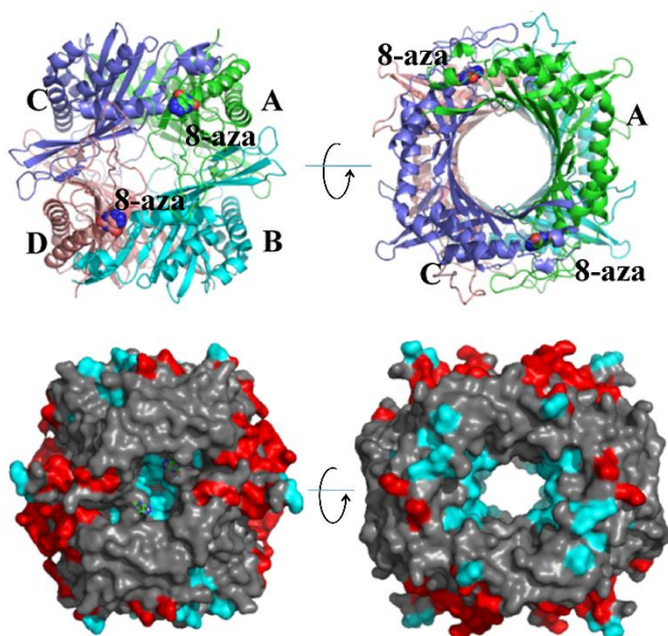


Figure 1 *Upper left*: The tetramer of urate oxidase in cartoon representation with the A chain in green, the B chain in cyan, the C chain in slate, and the D chain in salmon. The 8-azaxanthine molecules are shown in sphere representation. *Upper right*: Same representation, viewed along the central tunnel. *Lower left*: Molecular surface of UOX tetramer, colored in red where the B -factors increase more than 13 \AA^2 between the UOX-HHP-0.1-RT and UOX-HHP-210-RT structures, in cyan where the B -factors increase less than 5 \AA^2 and in grey otherwise. The 8-azaxanthine molecules are shown in ball-and-stick representation. *Lower right*: Same representation, viewed along the central tunnel.

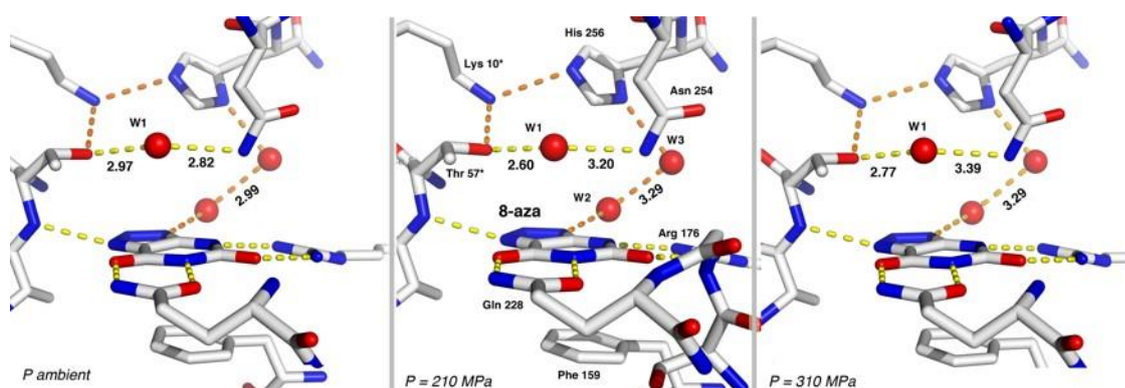


Figure 2 From left to right, the active site in the UOX-HPP-0.1-RT, UOX-HPP-210-RT and UOX-HPP-310-RT structures, showing the progressive elongation of the catalytic water W1 tweezers and of the hydrogen bond between W2 and W3.

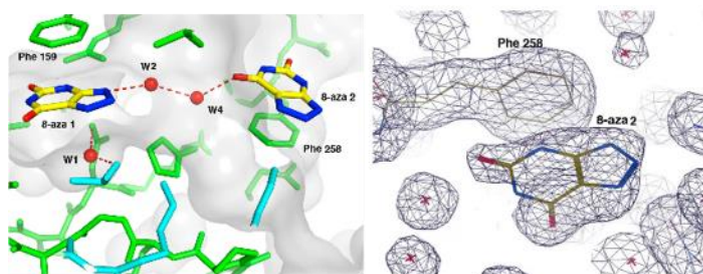


Figure 3 *Left*: In the UOX-HHP-210-RT, UOX-HHP-310-RT and UOX-HArP-200-T100 structures, the second ligand molecule (8-aza 2) is located within the channel ending at the active site occupied by 8-aza 1. It is connected to the first ligand through the two water molecules W2 and W4 and is stacked over Phe 258. *Right*: The electron density of the second ligand observed in the HARp-200-T100 structure (2Fo-Fc map, level at 1.5 sigmas above background).

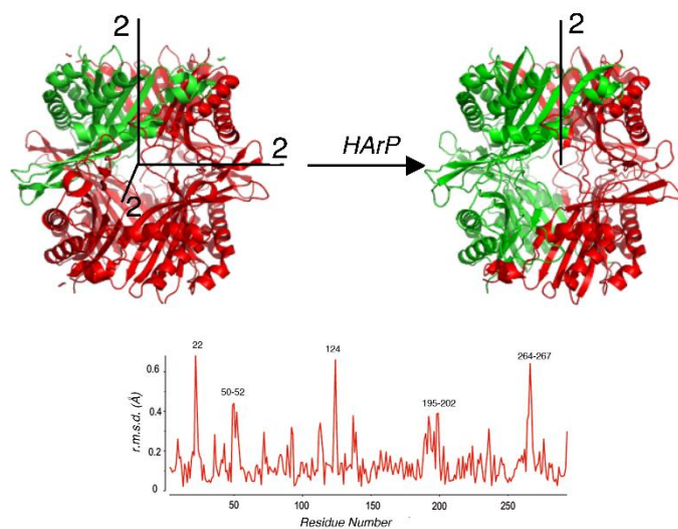


Figure 4 *Upper:* View of the HARp transition from an α_4 tetramer with three 2-fold axes (left, either the UOX-HPP-RT structures or the UOX-HArP structures up to 22 MPa) to a less-ordered ($\alpha\beta$)₂ tetramer with one two-fold axis (right, the UOX-HArP structures at 60 MPa and above). The asymmetric units are coloured in green and the symmetry-generated subunits in the tetramer are coloured in red. *Lower:* The r.m.s. deviations of the two α and β subunits of the UOX-HArP-200-T100 structure, relevant to the loss of the 2-fold symmetry between them.

Table 1 Data collection and processing statistics

Values for the outer shell are given in parentheses.

Acronym	UOX- HHP- 0.1- RT	UOX- HHP- 210- RT	UOX- HHP- 310- RT	UOX- HArP- 3.5- RT	UOX- HArP- 6.5- RT	UOX- HArP- 12- T100	UOX- HArP- 22- T100	UOX- HArP- 60- T100	UOX- HArP- 100- T100	UOX- HArP- 150- T100	UOX- HArP- 200- T100
Diffraction source	SOLEIL, CRISTAL beamline			MAXII, I711 beamline			ESRF, ID30 beamline				
Wavelength (Å)	0.5818	0.5818	0.5818	0.9775	0.9775	1.77	1.77	1.77	1.77	1.77	1.77
Pressure (MPa)	0.1	210	310	3.5	6.5	12	22	60	100	150	200
Temperature (K)	291	291	291	291	291	100	100	100	100	100	100
Space group	I222	I222	I222	I222	I222	I222	I222	P2 ₁ 2 ₁ 2			
<i>a</i> (Å)	80.34	79.53	79.13	80.23	80.32	80.89	79.79	78.23	78.43	77.40	78.55
<i>b</i> (Å)	96.16	95.63	95.40	96.11	96.12	95.50	95.15	95.27	95.18	95.34	96.68
<i>c</i> (Å)	105.51	104.75	104.67	105.39	105.41	105.23	104.28	104.5	104.4	104.66	105.58
Mosaicity (°)	0.07	0.15	0.09	0.09	0.11	0.36	0.28	0.20	0.19	0.22	0.34
Resolution range (Å)	29.77 - 1.90	29.59 - 2.15	29.51 - 2.40	4.78 - 1.60	20.00 - 1.60	40.50 - 2.36	70.00 - 1.69	47.60 - 1.86	47.60 - 1.64	47.70 - 2.19	71.32 - 1.72
Total No of reflections	107 997 (150493)	69 635 (10 163)	51 523 (7 586)	220 558 (21 323)	187 218 (18 022)	175 236 (15 472)	490 648 (7 193)	607 475 (1 545)	908 151 (19 326)	405 680 (11 521)	531 793 (11 199)
No of unique reflections	30 145 (4 400)	21 075 (3 071)	15 109 (2 216)	53 681 (5 277)	52 954 (5 301)	47 439 (4 120)	39 125 (2 310)	53 842 (499)	84 438 (4955)	39 208 (11 521)	73 215 (2 671)

Completeness (%)	93.3 (94.2)	96.0 (97.1)	96.0 (97.2)	99.5 (99.6)	98.0 (99.5)	97.3 (98.9)	87.8 (36.3)	81.5 (21.1)	88.6 (36.3)	97.2 (76.9)	93.8 (56.5)
Multiplicity	3.6 (3.5)	3.3 (3.3)	3.4 (3.4)	4.1 (4.1)	3.5 (3.3)	4.8 (4.4)	12.5 (3.1)	11.3 (2.8)	10.8 (3.9)	10.3 (4.4)	11.4 (5.3)
Mean I/ σ (I)	6.5 (2.2)	5.6 (1.8)	7.0 (1.9)	18.5 (2.8)	19.3 (2.2)	8.2 (1.4)	31.3 (2.5)	21.4 (3.1)	16.0 (1.9)	9.2 (1.2)	22.8 (2.0)
R_{meas}	15.8 (60.2)	21.2 (83.5)	16.5 (75.1)	4.0 (26.0)	6.3 (59.9)	9.9 (85.3)	6.3 (20.9)	6.6 (35.6)	8.5 (58.6)	16.2 (81.1)	12.7 (55.2)
$R_{r.i.m.}^\dagger$	0.158 (0.608)	0.208 (0.840)	0.165 (0.756)	0.040 (0.260)	0.063 (0.599)	0.099 (0.853)	0.063 (0.209)	0.064 (0.202)	0.085 (0.591)	0.170 (1.044)	0.062 (0.63)
CC $_{1/2}$	99.0 (80.5)	98.5 (69.8)	98.8 (69.8)			99.9 (44.4)	99.9 (97.9)	99.8 (96.9)	99.7 (80.0)	99.0 (77.3)	99.6 (80.8)
Overall B factor (\AA^2)	17.5	26.7	37.7	24.5	25.5	53.4	18.1	16.9	19.1	25.4	33.2

† Estimated $R_{r.i.m.} = R_{merge}[N/(N-1)]^{1/2}$, where N = data multiplicity.

Table 2 Structure solution and refinement statistics

Values for the outer shell are given in parentheses.

Acronym	UOX- HHP- 0.1-RT	UOX- HHP- 210-RT	UOX- HHP- 310-RT	UOX- HArP- 3.5-RT	UOX- HArP- 6.5-RT	UOX- HArP- 12-T100	UOX- HArP- 22-T100	UOX- HArP- 60-T100	UOX- HArP- 100-T100	UOX- HArP- 150-T100	UOX- HArP- 200-T100
Resolution range (Å)	20.0 - 1.90	20.0 - 2.15	20.0 - 2.40	14.78 - 1.60	14.42 - 1.60	40.48 - 2.36	70.0 - 1.69	47.70 - 1.86	47.65 - 1.64	47.70 - 2.19	46.38 - 1.80
No of reflections, working set	25 754 (1 746)	17 795 (1 122)	13 313 (869)	50 892 (3 677)	50 192 (3 714)	14 823 (72)	37 292 (668)	51 064 (659)	80 214 (1 305)	37 198 (2 053)	68 695 (3 402)
No of reflections, test set	1390 (83)	944 (57)	693 (50)	2 729 (187)	2 707 (173)	739 (8)	1 831 (34)	2 726 (34)	4 143 (56)	1 959 (124)	3 583 (170)
Final Rcryst	0.174 (0.269)	0.189 (0.309)	0.188 (0.349)	0.151 (0.206)	0.156 (0.300)	0.198 (0.863)	0.154 (0.646)	0.165 (0.227)	0.177 (0.297)	0.174 (0.317)	0.212 (0.386)
Final Rfree	0.222 (0.316)	0.248 (0.340)	0.270 (0.366)	0.178 (0.245)	0.183 (0.305)	0.264 (1.154)	0.196 (0.673)	0.208 (0.225)	0.205 (0.361)	0.226 (0.333)	0.243 (0.391)
No of non-H atoms											
Proteins	2362	2362	2362	2362	2362	2362	2362	4724	4824	4829	4724
Ion	1	1	1	1	1	1	6	2	2	2	2
Ligand	11	22	22	11	11	20	16	17	22	44	44
Water	261	179	115	229	221	203	435	531	652	580	473
Argon	-	-	-	1	1	2	2	4	4	4	4
Total	2 635	2 564	2 500	2 604	2 596	2 588	2 821	5 277	5 504	5 459	5 247
r.m.s. deviations											

Bonds (Å)	0.019	0.014	0.013	0.013	0.013	0.003	0.016	0.007	0.008	0.006	0.006
Angles (°)	1.808	1.66	1.461	1.873	1.917	0.999	2.512	1.31	1.33	1.29	1.324
Average B factors (Å ²)											
Proteins	19.91	29.41	49.03	26	26.4	54.6	16.6	17.9	19.9	30.0	34.2
Ion	22.26	31.35	58.77	37.5	40.1	65.2	24.2	19.4	21	35.2	42.3
Ligand	11.88	31	41.18	26.3	28.2	54.1	19.3	11.0	14.2	30.1	37.4
Water	34.67	35.04	44.6	39.2	38.7	53.5	28.5	27.3	29	39.7	37.3
Ramachandran diagram											
Most favored (%)	96.8	97.5	95.4	96.6	96.9	94.7	96.8	96.1	96.1	94.6	96.2
Allowed (%)	3.2	2.5	4.3	3.0	2.7	4.9	3.2	3.6	3.6	5.4	3.4
<i>PDB accession ID</i>	<i>7p0g</i>	<i>7p0c</i>	<i>7p0d</i>	<i>6i9x</i>	<i>6i9z</i>	<i>6ia1</i>	<i>6ia3</i>	<i>7puf</i>	<i>7pwn</i>	<i>7q09</i>	<i>6ia9</i>

Acknowledgements The authors acknowledge the SOLEIL (Gif-sur-Yvette, France), the European Synchrotron Radiation Facility (ESRF, Grenoble, France) and the MAX II synchrotrons (Lund, Sweden) for provision of synchrotron radiation facilities. They thank Pierre Fertey of the CRISTAL beamline (SOLEIL) for his technical help and assistance during data collection under high pressure *per se*, Yngve Cerenius and the staff members of I117 beamline (MAX II) for data collection in quartz capillaries under moderate argon pressure, and the staff members of ID30-B beamline (ESRF) for data collection under high argon pressure. The authors acknowledge access to the ESRF MX high-pressure laboratory for the preparation of the frozen UOX crystals under argon pressure. The use of the different large aperture DACs built by BETSA has been made possible through the IBS High Pressure platform. IBS acknowledges integration into the interdisciplinary Research Institute of Grenoble (IRIG, CEA). The authors also have special thoughts for Roger Fourme, deceased in 2012, who developed HPMX with such great enthusiasm.

References

- Akasaka, K. (2006). *Chem. Rev.* **106**, 1814–1835.
- Akasaka, K., Kitahara, R. & Kamatari, Y. O. (2013). *Arch. Biochem. Biophys.* **531**, 110–115.
- Akasaka, K., Nagahata, H., Maeno, A. & Sasaki, K. (2008). *Biophysics (Nagoya-Shi)*. **4**, 29–32.
- Bliznyuk, A., Grossman, Y. & Moskovitz, Y. (2019). *Sci Rep.* **9**, 10814.
- Bui, S., von Stetten, D., Jambrina, P. G., Prangé, T., Colloc'h, N., de Sanctis, D., Royant, A., Rosta, E. & Steiner, R. A. (2014). *Angew. Chem. Int. Ed. Engl.* **53**, 13710–13714.
- Bury, C. S., Brooks-Bartlett, J. C., Walsh, S. P. & Garman, E. F. (2018). *Protein Sci.* **27**, 217–228.
- Collins, M. D., Hummer, G., Quillin, M. L., Matthews, B. W. & Gruner, S. M. (2005). *Proc. Natl. Acad. Sci. U.S.A.* **102**, 16668–16671.
- Collins, M. D., Kim, C. U. & Gruner, S. M. (2011). *Annu Rev Biophys.* **40**, 81–98.
- Colloc'h, N., Carpentier, P., Montemiglio, L. C., Vallone, B. & Prangé, T. (2017). *Biophys. J.* **113**, 2199–2206.
- Colloc'h, N., el Hajji, M., Bachet, B., L'Hermite, G., Schiltz, M., Prangé, T., Castro, B. & Mornon, J. P. (1997). *Nat. Struct. Biol.* **4**, 947–952.

- Colloc'h, N., Marassio, G. & Prangé, T. (2011). *Current Trends in X-Ray Crystallography*, Vol. edited by A. Chandrasekaran, pp. 285–308. InTech.
- Colloc'h, N. & Prangé, T. (2014). *FEBS Lett.* **588**, 1715–1719.
- Dhaussy, A.-C. & Girard, E. (2015). *Subcell Biochem.* **72**, 215–235.
- Eisenmenger, M. J. & Reyes-De-Corcuera, J. I. (2009). *Enzyme and Microbial Technology.* **45**, 331–347.
- Emsley, P., Lohkamp, B., Scott, W. G. & Cowtan, K. (2010). *Acta Crystallogr D Biol Crystallogr.* **66**, 486–501.
- Engilberge, S., Wagner, T., Carpentier, P., Girard, E. & Shima, S. (2020). *Chem Commun (Camb).* **56**, 10863–10866.
- Espada, R., Sánchez, I. E. & Ferreiro, D. U. (2016). *Biophys. J.* **111**, 2339–2341.
- Fourme, R., Girard, E. & Akasaka, K. (2012). *Curr. Opin. Struct. Biol.* **22**, 636–642.
- Fourme, R., Girard, E., Kahn, R., Dhaussy, A.-C. & Ascione, I. (2009). *Annu Rev Biophys.* **38**, 153–171.
- Fourme, R., Honkimäki, V., Girard, E., Medjoubi, K., Dhaussy, A.-C. & Kahn, R. (2012). *J Appl Cryst.* **45**, 652–661.
- Fourme, R., Kahn, R., Mezouar, M., Girard, E., Hoerentrup, C., Prangé, T. & Ascione, I. (2001). *J Synchrotron Radiat.* **8**, 1149–1156.
- Frauenfelder, H., Alberding, N., Ansari, A., Braunstein, D., Cowen, B., Hong, M., Iben, I., Johnson, J., Luck, S., Marden, M., Mourant, J., Ormos, P., Reinisch, L., Scholl, R., Schulte, A., Shyamsunder, E., Sorensen, L., Steinbach, P., Xie, A., Young, R. & Yue, K. (1990). *J. Phys. Chem.* 1024–1037.
- Frauenfelder, H., Chen, G., Berendzen, J., Fenimore, P. W., Jansson, H., McMahon, B. H., Stroe, I. R., Swenson, J. & Young, R. D. (2009). *Proc. Natl. Acad. Sci. U.S.A.* **106**, 5129–5134.
- Frauenfelder, H., Fenimore, P. W. & Young, R. D. (2007). *IUBMB Life.* **59**, 506–512.
- Fuglestad, B., Stetz, M. A., Belnavis, Z. & Wand, A. J. (2017). *Biophysical Chemistry.* **231**, 39–44.
- Gabison, L., Chiadmi, M., El Hajji, M., Castro, B., Colloc'h, N. & Prangé, T. (2010). *Acta Crystallogr. D Biol. Crystallogr.* **66**, 714–724.
- Gabison, L., Chopard, C., Colloc'h, N., Peyrot, F., Castro, B., El Hajji, M., Altarsha, M., Monard, G., Chiadmi, M. & Prangé, T. (2011). *Proteins.* **79**, 1964–1976.
- Gabison, L., Prangé, T., Colloc'h, N., El Hajji, M., Castro, B. & Chiadmi, M. (2008). *BMC Struct. Biol.* **8**, 32.

- Girard, E., Dhaussy, A.-C., Couzinet, B., Chervin, J.-C., Mezouar, M., Kahn, R., Ascone, I. & Fourme, R. (2007). *J Appl Crystallogr.* **40**, 912–918.
- Girard, E., Marchal, S., Perez, J., Finet, S., Kahn, R., Fourme, R., Marassio, G., Dhaussy, A.-C., Prangé, T., Giffard, M., Dulin, F., Bonneté, F., Lange, R., Abraini, J. H., Mezouar, M. & Colloc'h, N. (2010). *Biophys. J.* **98**, 2365–2373.
- Gotthard, G., Aumonier, S., De Sanctis, D., Leonard, G., von Stetten, D. & Royant, A. (2019). *IUCrJ.* **6**, 665–680.
- Hahn, C. J., Lemaire, O. N., Kahnt, J., Engilberge, S., Wegener, G. & Wagner, T. (2021). *Science.* **373**, 118–121.
- Hibi, T., Hayashi, Y., Fukada, H., Itoh, T., Nago, T. & Nishiya, Y. (2014). *Biochemistry.* **53**, 3879–3888.
- Hibi, T. & Itoh, T. (2021). *J Biochem.* **169**, 15–23.
- Hibi, T., Kume, A., Kawamura, A., Itoh, T., Fukada, H. & Nishiya, Y. (2016). *Biochemistry.* **55**, 724–732.
- Kabsch, W. (2010). *Acta Crystallogr. D Biol. Crystallogr.* **66**, 125–132.
- Kalbitzer, H. R., Rosnizeck, I. C., Munte, C. E., Narayanan, S. P., Kropf, V. & Spoerner, M. (2013). *Angew. Chem. Int. Ed. Engl.* **52**, 14242–14246.
- Kalms, J., Schmidt, A., Frielingsdorf, S., van der Linden, P., von Stetten, D., Lenz, O., Carpentier, P. & Scheerer, P. (2016). *Angew. Chem. Int. Ed. Engl.* **55**, 5586–5590.
- Kleywegt, G. J. & Jones, T. A. (1994). *Acta Crystallogr. D Biol. Crystallogr.* **50**, 178–185.
- Knapp, J. E., Pahl, R., Cohen, J., Nichols, J. C., Schulten, K., Gibson, Q. H., Srajer, V. & Royer, W. E., Jr (2009). *Structure.* **17**, 1494–1504.
- Krissinel, E. (2015). *Nucleic Acids Res.* **43**, W314–319.
- Lafumat, B., Mueller-Dieckmann, C., Leonard, G., Colloc'h, N., Prangé, T., Giraud, T., Dobias, F., Royant, A., van der Linden, P. & Carpentier, P. (2016). *Journal of Applied Crystallography.* **49**, 1478–1487.
- Leal, R. M. F., Bourenkov, G., Russi, S. & Popov, A. N. (2013). *J Synchrotron Radiat.* **20**, 14–22.
- Levin, A., Cinar, S., Paulus, M., Nase, J., Winter, R. & Czeslik, C. (2019). *Biophys Chem.* **252**, 106194.
- van der Linden, P., Dobias, F., Vitoux, H., Kapp, U., Jacobs, J., Sweeney, S. M., Mueller-Dieckmann, C. & Carpentier, P. (2014). *Journal of Applied Crystallography.* **47**, 584–592.
- Louis, J. M. & Roche, J. (2016). *J Mol Biol.* **428**, 2780–2792.

- Luna, V. M., Chen, Y., Fee, J. A. & Stout, C. D. (2008). *Biochemistry*. **47**, 4657–4665.
- Luna, V. M., Fee, J. A., Deniz, A. A. & Stout, C. D. (2012). *Biochemistry*. **51**, 4669–4676.
- Luong, T. Q., Kapoor, S. & Winter, R. (2015). *Chemphyschem*. **16**, 3555–3571.
- Marassio, G., Prangé, T., David, H. N., Sopkova-de Oliveira Santos, J., Gabison, L., Delcroix, N., Abraini, J. H. & Colloc'h, N. (2011). *FASEB J*. **25**, 2266–2275.
- McCarthy, A. A., Barrett, R., Beteva, A., Caserotto, H., Dobias, F., Felisaz, F., Giraud, T., Guijarro, M., Janocha, R., Khadrouche, A., Lentini, M., Leonard, G. A., Lopez Marrero, M., Malbet-Monaco, S., McSweeney, S., Nurizzo, D., Papp, G., Rossi, C., Sinoir, J., Sorez, C., Surr, J., Svensson, O., Zander, U., Cipriani, F., Theveneau, P. & Mueller-Dieckmann, C. (2018). *J Synchrotron Radiat*. **25**, 1249–1260.
- McGregor, L., Földes, T., Bui, S., Moulin, M., Coquelle, N., Blakeley, M. P., Rosta, E. & Steiner, R. A. (2021). *IUCrJ*. **8**, 46–59.
- Montet, Y., Amara, P., Volbeda, A., Vernede, X., Hatchikian, E. C., Field, M. J., Frey, M. & Fontecilla-Camps, J. C. (1997). *Nat Struct Biol*. **4**, 523–526.
- de la Mora, E., Coquelle, N., Bury, C. S., Rosenthal, M., Holton, J. M., Carmichael, I., Garman, E. F., Burghammer, M., Colletier, J.-P. & Weik, M. (2020). *Proc Natl Acad Sci U S A*. **117**, 4142–4151.
- Murshudov, G. N., Vagin, A. A. & Dodson, E. J. (1997). *Acta Crystallogr D Biol Crystallogr*. **53**, 240–255.
- Nagae, T., Kawamura, T., Chavas, L. M. G., Niwa, K., Hasegawa, M., Kato, C. & Watanabe, N. (2012). *Acta Crystallogr. D Biol. Crystallogr*. **68**, 300–309.
- Nielsen, G., Jonker, H. R. A., Vajpai, N., Grzesiek, S. & Schwalbe, H. (2013). *Chembiochem*. **14**, 1799–1806.
- Prangé, T., Schiltz, M., Pernot, L., Colloc'h, N., Longhi, S., Bourguet, W. & Fourme, R. (1998). *Proteins*. **30**, 61–73.
- Quillin, M. L., Breyer, W. A., Griswold, I. J. & Matthews, B. W. (2000). *J. Mol. Biol*. **302**, 955–977.
- Schiltz, M., Fourme, R. & Prangé, T. (2003). *Methods Enzymol*. **374**, 83–119.
- Schiltz, M., Prangé, T. & Fourme, R. (1994). *J Appl Cryst*. **27**, 950–960.
- Silva, J. L. & Weber, G. (1993). *Annu Rev Phys Chem*. **44**, 89–113.
- Storm, S. L. S., Axford, D. & Owen, R. L. (2021). *IUCrJ*. **8**,.
- Tilton, R. F., Kuntz, I. D. & Petsko, G. A. (1984). *Biochemistry*. **23**, 2849–2857.
- Vitali, J., Robbins, A. H., Almo, S. C. & Tilton, R. F. (1991). *J Appl Cryst*. **24**, 931–935.

- Wei, D., Huang, X., Qiao, Y., Rao, J., Wang, L., Liao, F. & Zhan, C.-G. (2017). *ACS Catal.* **7**, 4623–4636.
- Whittington, D. A., Rosenzweig, A. C., Frederick, C. A. & Lippard, S. J. (2001). *Biochemistry*. **40**, 3476–3482.
- Williamson, M. P. & Kitahara, R. (2019). *Biochim Biophys Acta Proteins Proteom.* **1867**, 350–358.
- Winn, M. D., Ballard, C. C., Cowtan, K. D., Dodson, E. J., Emsley, P., Evans, P. R., Keegan, R. M., Krissinel, E. B., Leslie, A. G. W., McCoy, A., McNicholas, S. J., Murshudov, G. N., Pannu, N. S., Potterton, E. A., Powell, H. R., Read, R. J., Vagin, A. & Wilson, K. S. (2011). *Acta Crystallogr D Biol Crystallogr.* **67**, 235–242.
- Winter, M. B., Herzik, M. A., Kuriyan, J. & Marletta, M. A. (2011). *Proc. Natl. Acad. Sci. U.S.A.* **108**, E881–889.
- Winter, R. (2019). *Annu Rev Biophys.* **48**, 441–463.
- Xue, M., Wakamoto, T., Kejlberg, C., Yoshimura, Y., Nielsen, T. A., Risør, M. W., Sanggaard, K. W., Kitahara, R. & Mulder, F. A. A. (2019). *Proc Natl Acad Sci U S A.* **116**, 21031–21036.