

An environmental control box for serial crystallography enables multi-dimensional experiments

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Abstract

We present a new environmental enclosure for fixed-target, serial crystallography enabling full control of both the temperature and humidity. While maintaining the relative humidity to within a percent, this enclosure provides access to X-ray diffraction experiments in a wide temperature range from below 10 °C to above 80 °C. Coupled with the LAMA method, time-resolved serial crystallography experiments can now be carried out at truly physiological temperatures, providing fundamentally new insight into protein function. Using the hyperthermophile enzyme xylose isomerase, we demonstrate changes in the electron density as a function of increasing temperature and time. This method provides the necessary tools to successfully carry out multi-dimensional serial crystallography.

Keywords:

serial-crystallography, SSX, multi-temperature crystallography, time-resolved crystallography, multi-dimensional crystallography

Introduction

Life has evolved to occupy a wide range of different environmental niches, ranging from arctic deserts to the hot ponds at Yellowstone¹. Biochemical, bioinformatics and structural studies over many years have provided insight into how protein folds and function are maintained across, for example, such extreme ranges of temperature^{2,3}. Although enzymes operate over a wide range of temperatures, the vast majority of protein structural information, from X-ray crystallography and cryo-EM, has been acquired at cryogenic temperatures, using flash cooling to trap their structures close to the native, functional state. The main reason for data-collection at cryo-temperatures is to reduce the rate of radiation damage⁴. However, flash-cooling can lead to structural artefacts that are absent at ambient temperatures and that this may affect biological interpretation of the data⁵⁻⁸. In addition, as proteins are not single unique structures, but rather an ensemble of states that fluctuate around an energetic ground-state⁹, the conformational dynamics of enzymes are directly correlated with their catalytic rate¹⁰, and these dynamics are intimately linked to temperature.

This mismatch between standard data collection temperatures and the optimal temperature for function thus becomes an acute problem when carrying out time-resolved structural studies that seek to provide detailed insight into catalytic mechanisms and allosteric regulation. Although, with the advent of serial crystallographic methods at synchrotrons and XFELs (SSX, SFX), time-resolved structural studies are under going a resurgence, the majority of such experiments are still carried out at ambient temperatures (4°C - 25°C), regardless of the actual temperature optimum of the protein under study^{11,12}.

A notable exception is the pioneering work on photoactive yellow (PYP) protein, where time-resolved data were collected from -40 °C – 70°C, so called 5D-crystallography^{13,14}. Unfortunately, the considerable technological difficulties of this seemingly simple experiment have so far prevented routine structure determination as a function of physiologically relevant temperatures.

To address this challenge we have developed an environmental control system for serial crystallography that enables the precise control of humidity and temperature over a large temperature window from < 10°C to > 80°C. This temperature window is large enough to encompass physiologically relevant temperatures of hyperthermophile enzymes. Via full compatibility with our previously described tools for reaction initiation by rapid mixing (LAMA)¹⁵, this system permits multi-temperature, time-resolved serial crystallography experiments (5D-SSX), provided the crystals are stable at the temperatures of interest. These experiments enable a unique view into a hitherto inaccessible realm of protein function. To demonstrate the power of this approach we have used the hyperthermophile enzyme xylose isomerase (XI), which has an activity optimum at a temperature of ~80 °C¹⁶.

Results

Environmental control box design

A hallmark of protein crystals is their large solvent-content, which is typically in the range between 40% and 70% of the crystal volume. An advantage of this property is that proteins generally retain their function even in the crystalline state^{17,18}. A commonly known disadvantage is, however, their sensitivity to changes in environmental humidity, which in addition to the higher rates of radiation damage associated with higher temperatures makes routine data collection at even ambient temperatures a difficult task. Starting with traditional wax-enclosed glass capillaries, several solutions have been developed to maintain crystal hydration for single crystals^{19,20}. However, to maintain a controlled humidity environment for our *hit-and-return* (HARE)-chip setup including the liquid-application-method for time-resolved crystallography (LAMA)^{21,22}, required the development of a solution that could accommodate this larger experimental setup. To this end we have constructed a modular, portable environmental control box that encloses our previously described chip-setup including the LAMA droplet injector nozzle on a footprint of 118 mm x 283 mm (**Fig. 1, 2**)¹⁵.

Humidity control is achieved by flowing dry air either directly into the box, or first passing through a water bath (20-95 °C), with the proportion of gas through each channel controlled by a toggling (ball) valve in a (PID-) feedback loop. The water-bath is connected to the environmental control box via a silicone hose, to which a heating belt is attached preventing (re)condensation. Humidity control is achieved by flowing dry air either directly into the box, or first passing through a water bath (20-95 °C), with the proportion of gas through each channel controlled by a toggling ball valve in a PID-feedback loop. The water-bath is connected to the environmental control box via a silicone hose, to which a heating belt is attached preventing condensation. The target value can be achieved with high accuracy, enabling humidity control in the single point scale, while on the other hand as well as controlled crystal dehydration if that is required.

In addition to humidity, temperature within the enclosure can be set anywhere within the range ~7°C to above 80°C. An aluminum air-stream reflector directs the stream of humid air around the chip. In order to enable effective control over this wide temperature range, the box uses two modular temperature control units (**Fig. 2**). Module 1 covers a temperature range from ~7°C to ~55°C, while module 2 covers a temperature range from ~50°C to >80°C. Rapid exchange of the modules is possible without tools, enabling switching between different temperature regimes within a few minutes. Module 1 is a water-cooled Peltier-element that enables active cooling or heating of the interior of the box. To ensure the temperature is equilibrated across the box the module is equipped with fans. Cooling water and circuit control are fed in through the top side of the module. The heating element in module 2 is a power resistor network that disseminates sufficient heat to increase the interior temperature of the box to >80°C, while the relative humidity can be sustained at over 95 %. Temperature control is achieved by

a PID controller that sets the current through the Peltier elements and resistor network, respectively, to maintaining the target temperature.

The base of the box is made from durable polyether ether ketone (PEEK). To drain condensation water, several cotton wicks are fixed around the bottom corners and connected to an active pumping system that quickly drains excess liquid from the box. The sides and the lid of the box are made of 6 mm thick acrylic glass, while the rear panel (X-ray side) is made of acrylic glass and polyoxymethylene (POM). As an X-ray entrance window, an 8 mm opening in the POM is covered with two layers of COC foil, about 1 mm apart. To enable easy replacement of the X-ray entrance window, the COC-foil is placed on a magnetically mounted ring, that tightly seals the inside of the box.

While the right panel is solid, the left panel has a feedthrough for the humidity tube, and an access hatch through which HARE chips can be loaded onto the sample translation stage. The front panel (detector side) is an aluminum frame with a 190 mm opening. This is sealed with two X-ray transparent Mylar foils (6 μm), as an exit window for the diffracted beam. To reduce condensation on this window, the space between the Mylar-foils is continuously flushed with warm air. The lid of the box contains feedthroughs for the humidity- and temperature-sensors, as well as for the electropneumatic retractable, infrared (IR) backlight, the LAMA-nozzle lever, and an access port for heating module exchange.

The translation stage system implemented on T-REXX P14 EH2 is not humidity resistant and therefore has to be kept on the outside of the box. Hence, the translation stages are connected to the box via a flexible bellow, custom cut from two layers of commercial plastic wrap. On the inside, the bellow is sealed around the arm between the translation stages and the kinematic mount for the chip holder^{23,24}. To reduce the heat capacity of the chip holder, and thus maintain faster temperature equilibration the previously described aluminum chip holder design was fabricated from PEEK, providing the same functionality at a lighter weight²⁴.

The LAMA nozzle is attached via a kinematic mount to a retractable lever, that enables retraction of the nozzle from its injection position during chip exchange. To avoid the unnecessary opening of the box, which might lead to temperature and humidity fluctuations, the nozzle retracts into a parking position under the lid of the box. Fine LAMA nozzle alignment in the injector position is achieved via motorized translation stages (Thorlabs).

The whole box system is mounted on rails, residing on a stainless-steel baseplate that can be moved between “data collection” or “beam location” position, where the latter allows to use the X-ray scintillator built into the beam-shaping device (BSD; Arinax, Moirans France).

Operation of the serial crystallography environmental control enclosure is achieved via an external control unit where temperature and humidity parameters are electronically set (**Fig. 3**). Parameters needed to achieve certain conditions were calibrated and are listed in **Table 1**.

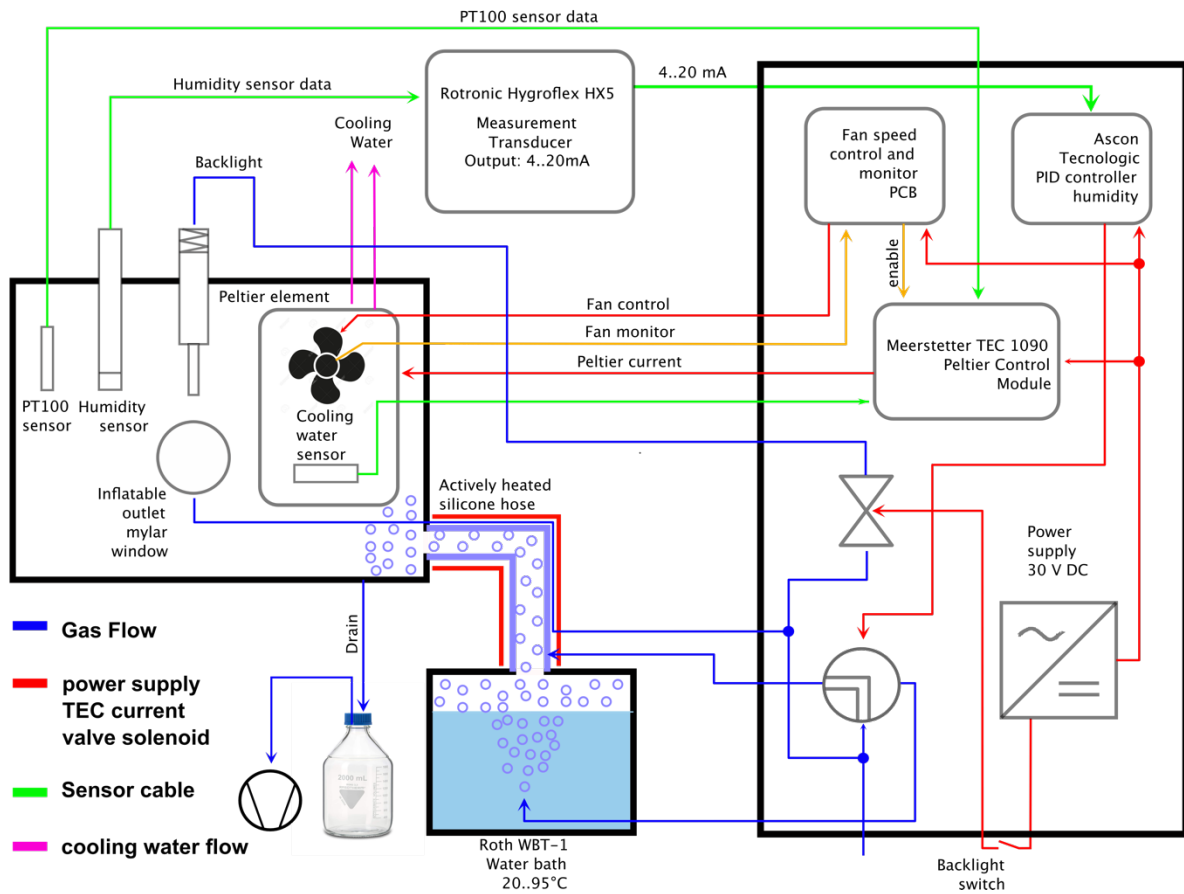


Figure 1: Schematic representation of the box-setup.

Displayed a schematic representation of the box-setup: gas flow is indicated by blue lines, power supply is red, while sensor cables are depicted in green, cooling water is displayed in pink..

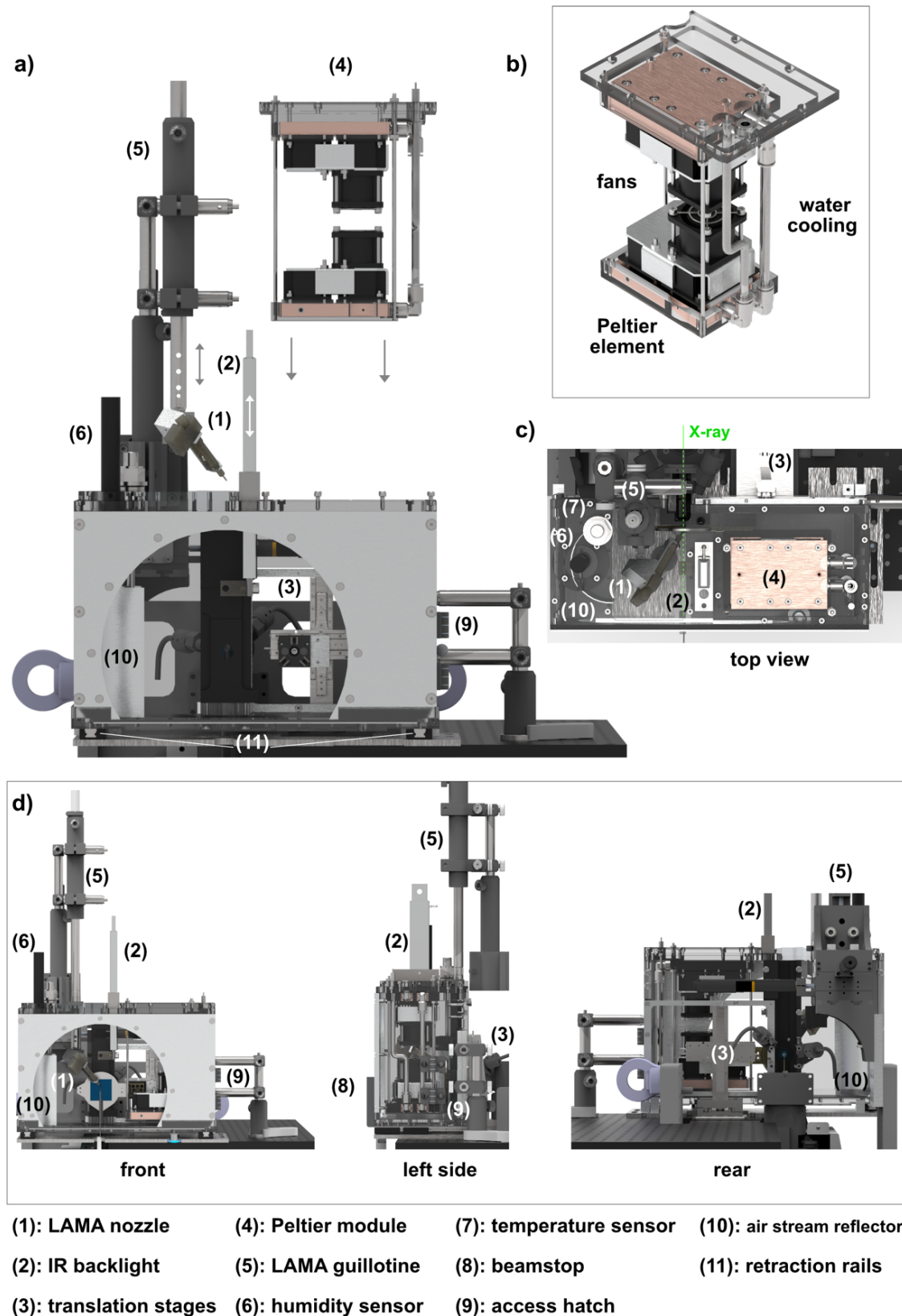


Figure 2: The 'Serial Crystallography Environmental Control Enclosure'. a) overview of the environmental control box, with retracted LAMA nozzle and Peltier module. b) closeup of the Peltier module, c) top-view providing an overview of the arrangement inside the box, d) front-, side- and rear-view of the box. Note: for clarity not all technical elements of the box (e.g. tubing, electric connections etc.) or the beamline are shown or described in detail. Elements mentioned in the text are numbered and shown in the figure legend.

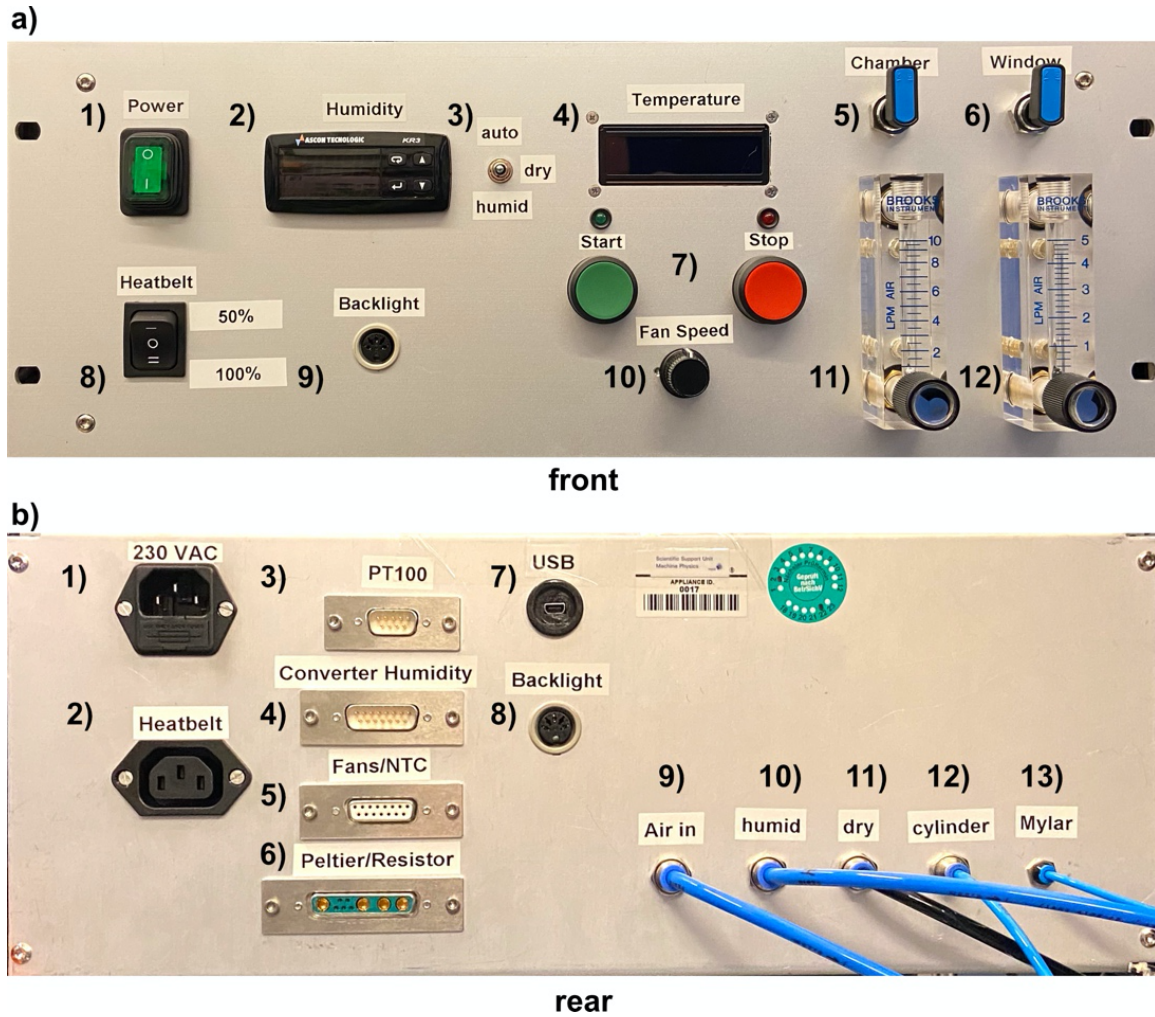


Figure 3: Control unit of the serial crystallography environmental control enclosure. A) front panel: 1) main power switch; 2) humidity control / display; 3) air flow destination switch; 4) temperature display; 5) airflow on/off switch; 6) inflatable front-window flow on/off switch; 7) Temperature control start/stop button; 8) heat-belt switch; 9) socket for IR backlight switch; 10) fan-speed regulator; 11) air-flow valve; 12) front-window flow valve. b) rear panel: 1) main power inlet; 2) heat belt power outlet; 3) PT100 temperature sensor; 4) humidity sensor; 5) fan connection; 6) Peltier/resistor connection; 7) USB connection to PC; 8) backlight outlet; 9) pressurized air in; 10) air flow to water-bath; 11) dry air outlet; 12) backlight cylinder air outlet; 13) inflatable front-window air outlet

Table 1: environmental control parameter settings

<i>Module 1</i>						
desired temp °C	flow rate L min ⁻¹	desired humidity %	actual humidity %	water bath temp °C	heat belt power %	measured current A
7.5	5	95	95	75	0	-12.3
15	5	95	95	75	50	-2.1
25	5	95	95	75	100	1.7
35	4	95	95	85	100	3.5
45	4	95	95	95	100	4.8
55	4	95	95	95	100	6

<i>Module 2</i>						
desired temp °C	flow rate L min ⁻¹	desired humidity %	actual humidity %	water bath temp °C	heat belt power %	current A
50	4	95	95	95	100	2.4
55	4	95	95	95	100	2.9
60	4	95	95	95	100	2.6
65	4	95	95	95	100	2.9
70	4	95	95	95	100	3.1
75	4	95	95	95	100	4.3
80	4	95	95	95	100	3.4

Characterization of the environmental control

To characterize how the environmental parameters can be controlled within the box, we recorded temperature and humidity changes as a function of time in 30 sec intervals. The temperature was modified as a step function, with the humidity target set to 95%. While the humidity rapidly equilibrates throughout the box, temperature gradients closer to the walls needs to be avoided. In addition to the overall temperature inside the box, we also monitored the temperature at two additional locations, directly on the surface of the backside of the HARE-chip and directly above the humidity sensor (**Fig. 4**). To assess how effectively different temperatures in the box can be achieved and maintained, we characterized the temperature increase from 7.5 °C to 80 °C. The data show that for both temperature control modules the humidity values quickly reach the target values. Over a temperature window of almost 50 °C the humidity remains stable within 2.5% of the target value. Analysis of the deviation of the chip temperature from the box temperature shows that the chip temperature follows the box temperature with a median difference of 0.7° C, over a temperature window of almost 50°C. Temperature and humidity typically equilibrate across the box and the chip 10-15 minutes.

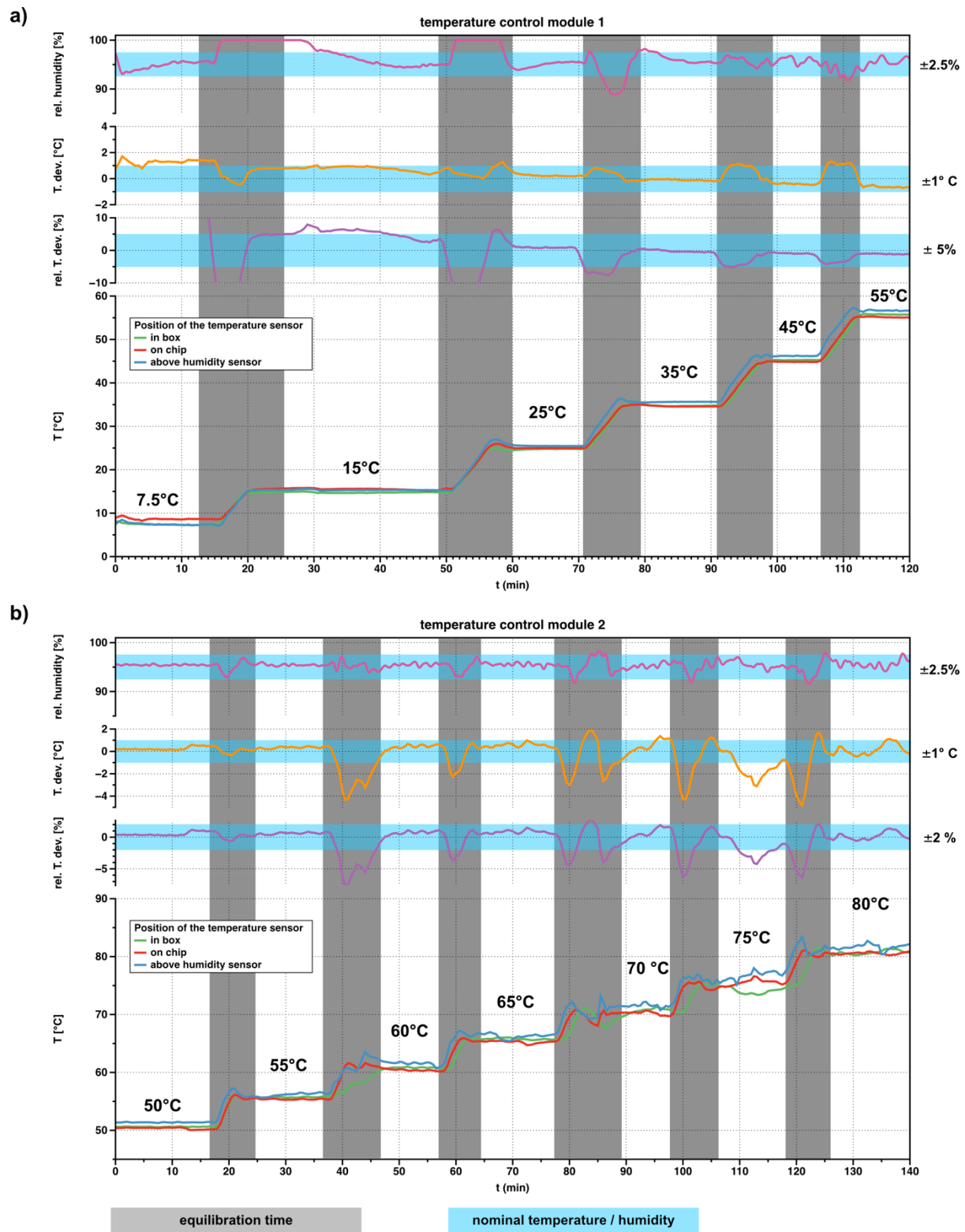


Figure 4: Characterization of the environmental control. a) module 1: 7.5 °C–55°C; b) module 2: 50–80°C; The temperature was successively increased from 7.5 to 55°C and 50 to 80°C respectively. The temperature was controlled at three independent positions inside of the chamber: green in the box, blue directly above the humidity sensor, red directly on the chip. The target humidity was set to 95%. The grey bars indicate the equilibration time, blue bars indicate the target window.

We also examined how well the environmental target values can be maintained during X-ray data-collection. To this end we collected X-ray diffraction data at 20 °C, 40 °C, 55 °C and 80 °C, for ~120 minutes and recorded temperature and humidity values in 30 sec intervals during this period (**Table 2**). Remarkably, during the data-collection the target humidity could be maintained within ~1%, while the temperature remained stable within ~0.5 °C. Clear deviations from this behavior are only observed during chip-exchange, when the hatch of the box is opened and during the subsequent reequilibration time while the environment restabilises. Reequilibration of the environmental parameters could be achieved within ~10-15 minutes, depending on the manual chip-exchange times.

In conclusion, these data show that after an equilibration time of ~10-15 minutes the environment in the control box has reached its target value, and that this can be maintained throughout extended periods of time, well beyond the typical data collection time of a chip (30 minutes). This enables collection of serial X-ray diffraction data at a variety of different temperatures with high accuracy and precision.

Table 2: Environmental parameters during X-ray diffraction data collection.

**values are derived from a long-term, (10 h) data-collection*

<i>nominal temperature (°C)</i>	<i>nominal rel. humidity (%)</i>	<i>measure rel. humidity (%)</i>	<i>rmsd (%)</i>	<i>measured temperature (°C)</i>	<i>rmsd (°C)</i>
20	95	95.4	0.5	19.8	0.2
40	95	95.5	0.9	40.3	0.3
55	95	95.5	0.7	55.3	0.7
80	95	95.2	0.7	80.1	0.3
20*	95*	95.2*	0.7*	19.8*	0.2*

Humidity dependent unit-cell modulation

Adjusting the relative humidity either prior to or during data-collection can improve several aspects of data-quality (resolution, mosaicity and anisotropy)²⁵⁻³³. To estimate the effect of the environmental humidity on diffraction data quality we monitored the unit-cell size of XI as a function of decreasing humidity. We started data-collection at a relative humidity of 95% and reduced the humidity in discrete steps of 5% per compartment row on the chip (**Fig. 5**). With decreasing humidity the crystals do not simply cease to diffract but undergo a change in unit-cell size. While at or above a relative humidity of 95% most diffraction patterns could be indexed with a large unit-cell ($a = 94.2 \text{ \AA}$, $b = 99.3 \text{ \AA}$, $c = 103.1 \text{ \AA}$; $\alpha, \beta, \gamma = 90.0^\circ$) the proportion rapidly changed to a smaller unit-cell ($a = 94.6 \text{ \AA}$, $b = 99.4 \text{ \AA}$, $c = 87.5 \text{ \AA}$; $\alpha, \beta, \gamma = 90.0^\circ$) as the relative humidity dropped from 90% to 75%. If the humidity is reduced even further the micro-crystals cease to diffract, presumably due to complete dehydration. This emphasizes the sensitivity of protein-microcrystals to environmental humidity, which must be precisely controlled to maintain their diffraction properties. On the other hand, this also opens the opportunity for crystals with large unit-cells to be specifically dehydrated to modulate their diffraction properties.

The response of protein crystals hydration to their environment has long been known³⁴ and chemical dehydration devices^{35–38} as well as dedicated dehumidification devices have successfully been used for this purpose on single, loop-mounted crystals^{27–29}. With this environmental control box these post-crystallization optimization protocols are now open to serial-crystallography.

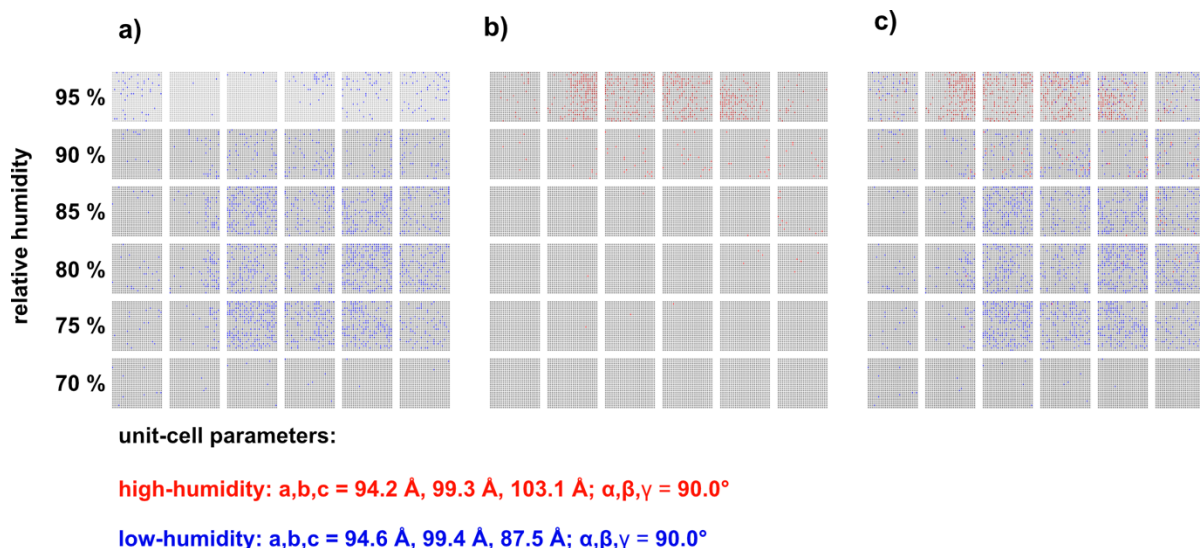


Figure 5: Humidity dependent unit-cell modulation displayed in a hit map. The HARE-chips consist of 6x6 compartments, divided into 24x24 features. Each feature is indicated by a small grey square; a blue-square indicates a diffraction pattern in the low humidity unit cell, a red square a diffraction pattern in the larger high-humidity unit-cell. Humidity has been reduced by 5% for each row of compartments. **a)** low-humidity diffraction patterns **b)** high-humidity diffraction patterns **c)** overlay of low- and high-humidity diffraction pattern hits.

Multi-temperature structural analysis of xylose isomerase

Exploiting the potential of this environment control box in a first *proof-of-principle* experiment, we have determined the structure of the hyperthermophile enzyme XI at six different temperatures: at 10° C, 20° C, 30° C, 40° C 50° C and 60° C, respectively. In general, all datasets could be processed to a resolution of 2 Å or better and display reasonable data-statistics (**Table 3**). In particular, the diffraction data do not indicate any obvious influence of the temperature on the diffraction properties.

All refined structures superimpose with a C-alpha r.m.s.d. of ~0.1 Å to the 20° C structure. With increasing temperature, the overall B-factors show a monotonically-increasing trend. However, while the B-factors of core residues including the active site remain moderate and relatively constant, the solvent exposed interface and the substrate entry cleft show increased B-factors with increasing temperature (**Table 3; Fig. 6**).

The high degree of conformational homogeneity is also reflected in the active site (**Fig. 7**). Here all residues superimpose almost perfectly, with a minimal amount of heterogeneity visible in Lys289, which adopts two alternate conformations and shows slightly increased B-factors as the temperature is increased. In addition to the amino acid side chains, the metal sites M1 and M2 remain in identical positions. The main differences that are observed lie in the position of the water molecules, in particular

those further away from the metal sites. These features emphasize the evolutionary adaptation of xylose isomerase to withstand higher temperatures and reflect the general structural rigidity of the enzyme.

Table 3: Data collection and refinement statistics of the apo data. Values in the highest resolution shell are shown in parenthesis.

Temperature (°C)	10	20	30	40	50	60
relative humidity (%)	98%	95%	98%	98%	98%	98%
Ligand	<i>apo</i>					
Space group	I222					
a, b, c (Å)	94.2, 103.1, 99.3					
α, β, γ (°)	90.0, 90.0, 90.0					
number of indexed images	7346	11115	8019	17004	5128	10007
Resolution (Å)	71.5 - 1.9 (1.97 - 1.90)	71.4 - 1.8 (1.86-1.80)	71.4 - 1.9 (1.97-1.90)	71.4 - 1.8 (1.86 - 1.80)	71.4 - 2.0 (2.07-2.00)	71.4 - 1.9 (1.97-1.90)
CC* (%)	93.09 (77.93)	96.3 (73.06)	94.43 (79.31)	97.90 (70.35)	91.38 (70.44)	96.70 (70.63)
R _{split} (%)	43.79 (89.24)	33.85 (113.4)	40.54 (89.36)	28.66 (129.23)	50.82 (111.71)	32.79 (124.08)
<1/ σ (I)> (SNR)	2.06 (1.11)	2.51 (0.90)	2.18 (1.11)	2.74 (0.82)	1.83 (0.90)	2.42 (0.81)
Multiplicity	69.5 (47.7)	424.9 (289.9)	154.0 (105.7)	1169.6 (801.8)	88.8 (61.1)	69.9 (47.8)
Completeness (%)	100.00 (100.00)					
Resolution range for refinement (Å)	71.49 - 1.90	71.49 - 1.80	71.49 - 1.90	71.49 - 1.80	71.49 - 2.00	71.49 - 1.90
No. Reflections	38388	45053	38388	41536	32995	38388
R _{work} (%)	20.54	17.67	18.93	16.91	21.04	18.32
R _{free} (%)	25.49	21.08	23.21	20.67	25.43	23.03
Mean B-factor (Å ²)						
overall	23.76	24.82	24.44	28.44	28.17	34.57
protein	22.42	23.55	23.33	27.53	27.18	33.31
water	36.37	36.37	35.54	38.87	37.46	48.31
other	19.82	20.5	18.1	20.53	21.3	21.26
Rms Deviations						
Bond Length (Å)	0.006	0.009	0.01	0.008	0.003	0.012
Bond Angles (°)	0.848	1.012	1.061	0.927	0.545	1.154
Ramachandran						
Favored (%)	97.14	97.14	97.4	97.14	96.87	96.59
Allowed (%)	2.6	2.6	2.34	2.6	2.87	3.15
Outliers (%)	0.26	0.26	0.26	0.26	0.26	0.26

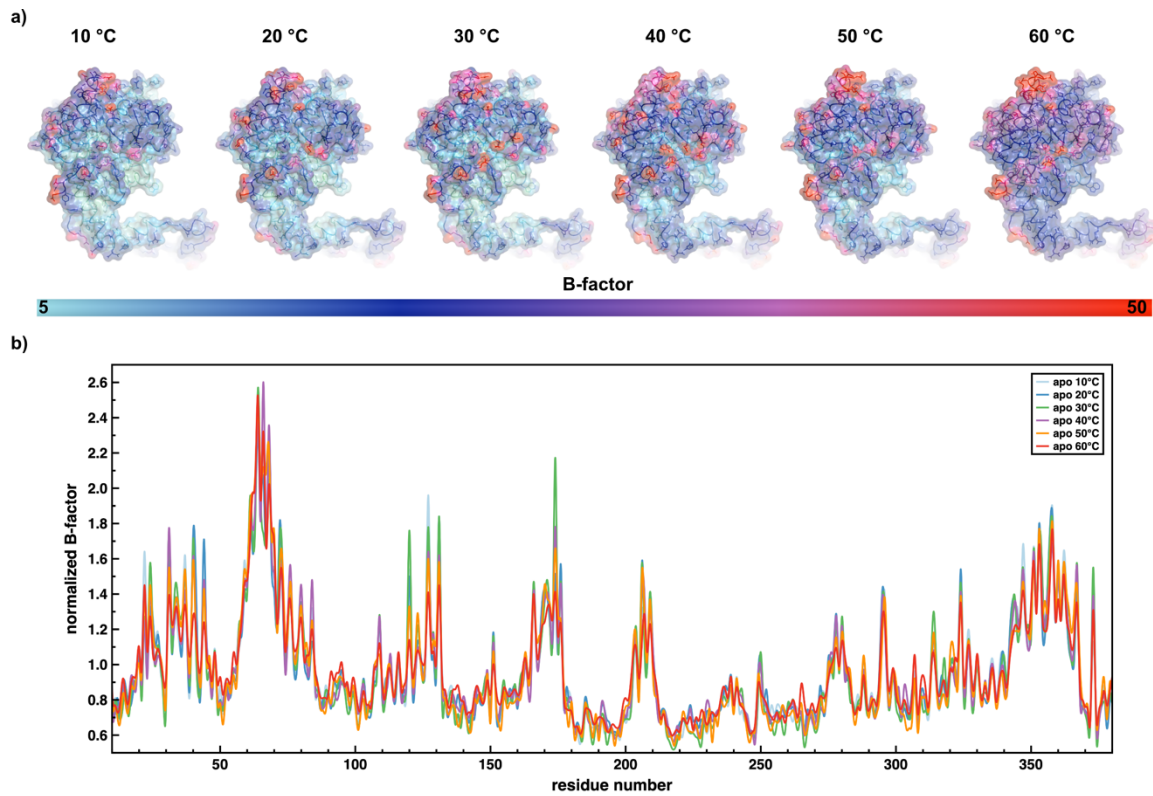


Figure 6: B-factor distribution as a function of temperature. a) A semi-transparent surface representation of the XI monomers, collected at 6 different temperatures. Coloring is according to the B-factor spectrum from 5 \AA^2 to 50 \AA^2 shown at the bottom. b) A plot of the normalized B-factors per residue for each structure. Ten residues on the N-terminus and eight residues at the C-terminus have been omitted for better visualization. Clearly no part of the apo structure is particularly sensitive to an increase in temperature.

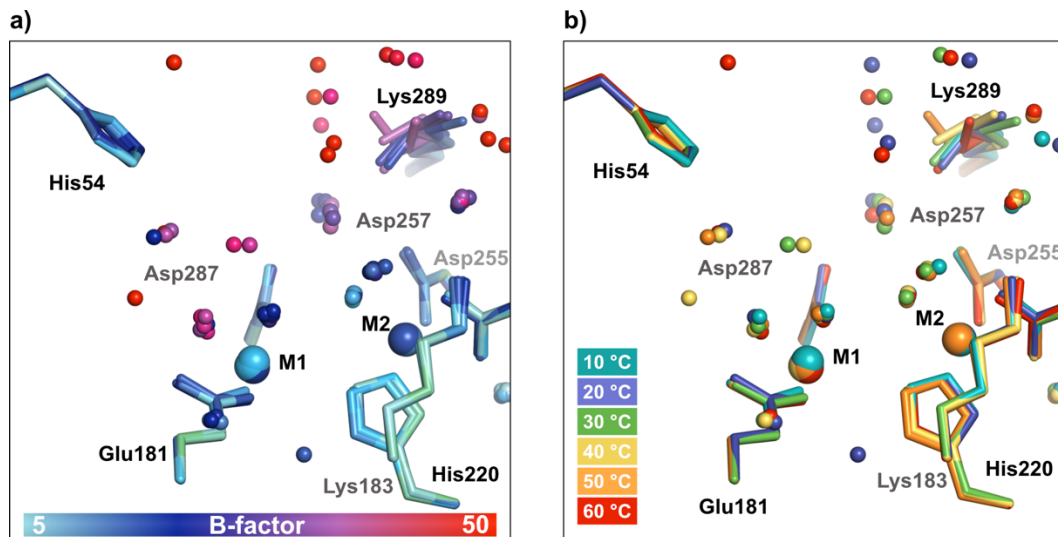


Figure 7: Active site of apo XI at temperatures from 10 °C to 60 °C. a) The active site of XI colored according to B-factor as indicated on the scale-bar. b) The active site residues of XI colored according to temperature, as indicated in the legend.

5D-SSX reveals temperature dependent electron density changes

To assess the effect of temperature on the catalytic reaction of XI, we triggered the reaction by adding the substrate glucose (alpha-D-glucose) using LAMA³⁹ and monitored its turnover at a constant time-delay of 60 seconds after reaction initiation at 20 °C, 40 °C, 45 °C and 50°C. Like the apo-data, the 5D datasets show good X-ray diffraction data-statistics (**Table 4**). A minimally lower resolution can be observed for the 50 °C dataset, but this is likely due to the lower number of images in this dataset rather than the increased temperature.

Table 4: Data collection and refinement statistics of the 60 sec data. Values in the highest resolution shell are shown in parenthesis.

Time-Delay (s)	60	60	60	60
Temperature (°C)	20	40	45	50
relative humidity (%)	99.0%	98.5%	95.0%	98.5%
Ligand		glucose		
Space group		I222		
a, b, c (Å)		94.20, 103.05, 99.25		
α, β, γ (°)		90.0, 90.0, 90.0		
number of images	9346	12349	9389	5224
Resolution (Å)	71.43-1.7 (1.76-1.70)	71.43-1.7 (1.76-1.70)	71.43-1.7 (1.86-1.80)	71.43-1.90 (1.97-1.90)
CC* (%)	94.63 (63.38)	95.45 (68.81)	95.39 (68.99)	92.10 (73.08)
R _{split} (%)	38.94 (141.80)	34.99 (126.20)	36.90 (117.50)	45.59 (102.08)
<I/σ(I)> (SNR)	2.27 (0.69)	2.65 (0.80)	2.36 (0.86)	2.11 (1.01)
Multiplicity	156.6 (93.8)	109.1 (65.6)	125.2 (85.4)	94.7 (64.9)
Completeness (%)		100.00 (100.00)		
Resolution range for refinement (Å)	71.49 - 1.70	71.49 - 1.70	71.49 - 1.80	71.49 - 1.90
No. Reflections	32995	53373	45053	38388
R _{work} (%)	18.38	18.05	17.68	19.48
R _{free} (%)	21.91	21.84	21.37	24.26
Mean B-factor (Å ²)				
overall	20.03	25.21	25.69	24.73
protein	19.12	23.87	24.82	23.81
water	29.5	38.13	35.6	34.55
other	20.84	25.7	23.56	21.14
Rms Deviations				
Bond Length (Å)	0.002	0.008	0.011	0.007
Bond Angles (°)	0.481	0.953	1.102	0.851
Ramachandran				
Favored (%)	97.14	96.35	96.61	96.88
Allowed (%)	2.6	3.39	3.12	2.86
Outliers (%)	0.26	0.26	0.26	0.26

Clearly, the 20 °C structure is in complex with a closed-ring glucose molecule, indicating that a ring opening reaction had not yet begun. The respective substrate complex structure at 20°C can be superimposed with an r.m.s.d. of 0.1 Å to previously reported substrate complexes at room temperature (PDB-ID: 3KCL)⁴⁰ (**Fig. 8**).

From temperatures above 40° C the electron density in the active site clearly changes from that of the lower temperature data and can no longer be explained by a closed ring substrate molecule alone. Thus this electron density can be interpreted best as a mixture between several different states comprising the closed ring glucose and an open-ring intermediate, and presumably also the apo state.

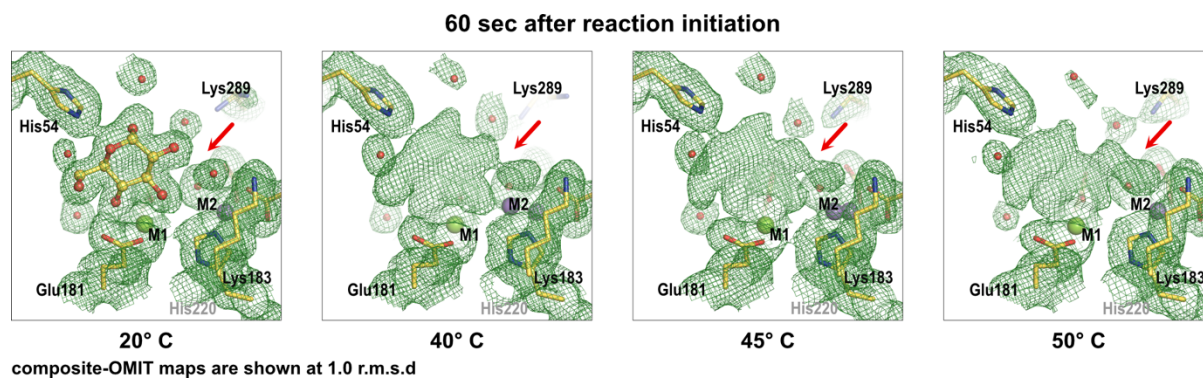


Figure 5: 5D-SSX reveals temperature dependent electron density changes. *Xylose isomerase active site, 60 sec after reaction initiation at various temperatures. Composite-OMIT maps are shown around the active site at an r.m.s.d. of 1.0. Clearly, with increasing temperature the electron density changes (red arrow) and can no longer be interpreted by a single state alone, and multiple, overlapping states have to be modelled.*

Discussion

Environmental control has consistently been a key aspect of macromolecular crystallography, as fragile protein crystals are sensitive to dehydration and physical stress. Open- and closed boundary environmental control solutions have been developed since the advent of structural biology. In the simplest instances, closed boundary devices include glass-capillaries, which retain protein crystals and typically a drop of mother liquor to sustain a humid atmosphere during data collection²⁰. More modern variations of this classic solution are the many fixed-target serial crystallography environments, which protect protein microcrystals against evaporation by some form of X-ray transparent window material⁴¹. For single crystal experiments a variety of solutions, sometimes for advanced parameter control, such as humidity, temperature, and electric fields have also been developed over the years^{42–44}. With the onset of cryo-crystallography, larger boxes were developed, which sometimes enclosed the stream of cryogenic gas in a dry atmosphere to prevent ice-formation during data-collection⁴⁵. However, for single-crystal experiments the majority of environmental control solutions fall into the open-boundary category, such as placing the crystal in an vapor stream with controlled humidity^{27,43,46–49}. Historically, the water content of protein crystals was controlled by post-crystallization treatments via chemical dehydration prior to crystal freezing^{34,36}. However, controlling the humidity around the mounted crystals enables the convenient identification of the optimal conditions for a particular sample during an X-ray diffraction experiment^{27,43,46–49}. Similar to humidity control, many beamlines are also equipped with a temperature control solution via a gas-stream that is directed at the sample, often encompassing wide temperature windows. Interestingly, however, a combined temperature-humidity

control is rarely found. Such a situation mandates the use of e.g., glass capillaries to maintain the crystals in a humid environment, which complicates time-resolved applications that are based on *in-situ* mixing. Historically, flow-cells were developed for time-resolved applications that allowed for *in-situ* mixing experiments with single-crystals and for trapping reaction intermediates on comparably slow time-scales⁵⁰⁻⁵². However, to the best of our knowledge these were not applied to multi-temperature experiments. That is, so far the only other multi-temperature time-resolved crystallographic experiment was limited to PYP, a well-studied reversible system that is triggered via photo-excitation^{13,14}.

Serial crystallography experiments can also be conducted in controlled, closed-boundary environments as demonstrated by the *drop-on-demand* device, which permits humidity control and for fully anaerobic experiments via the exchange of the surrounding atmosphere⁵³. However, with current sample delivery devices no temperature modulation has been achieved for serial crystallography experiments. In this regard, our environmental control box charts a new area, enabling enzymatic reaction characterization that comprises the full spectrum of physiological temperatures from psychrophilic to (hyper)-thermophile organisms. For the many irreversible mesophilic systems, we can now modulate enzyme kinetics through temperature variation, thereby accelerating or decelerating enzyme turnover and allowing for a more in-depth characterization of key enzymatic mechanisms. These can now be addressed at true physiological temperatures, enabling simultaneous correlation of mechanistic and thermodynamic aspects of protein function. Our solution combines environmental control with the advantages of serial crystallography in limiting radiation damage, as well as versatile delay-times provided by the HARE approach and the simplicity of reaction initiation of the LAMA-method^{15,22}. The observed glucose structure demonstrated in this study corroborates previous observations. The clear changes in electron density demonstrates a progression towards a 5-membered, closed-ring fructose product complex^{40,54,55}. However, the multiple overlapping states make an unambiguous interpretation of the electron density difficult, which mandates the use of more complex modelling and analysis protocols in future experiments^{56,57}. However, by increasing the temperature, different catalytic states, that is closed and likely open-chain intermediates can be captured at the same delay time, indicating an increase in the catalytic activity of xylose isomerase. Ergo, the modulation of the environmental temperature enables direct modulation of enzyme kinetics during a time-resolved X-ray diffraction experiment. Conducting similar experiments on other systems will allow us to understand how proteins exchange energy with their environment and how this is related to conformational dynamics and turnover. As described in the seminal paper by Schmidt et al., these multi-dimensional experiments will also permit determining experimental free-energy landscapes of proteins in action and thereby the elucidation of unique catalytic pathways^{13,58}. These experiments should fertilize the cross pollination different aspects of biophysical chemistry.

Material and Methods

Sample preparation

Crystals of *Streptomyces rubiginosus* Xylose Isomerase (XI) were obtained from Hampton Research (HR7-102) and stored at room temperature for several years. For this experiment the crystals were prepared as described previously¹⁵: commercial crystals were re-dissolved in water and concentrated to 80 mg/ml in xylose-isomerase buffer (10 mM Hepes/NaOH pH 7.5). This protein solution was re-diluted and concentrated again for 9 consecutive times to remove any remaining Tris/HCl from the storage buffer. Subsequently microcrystals were obtained by vacuum induced crystallization, as described previously by Martin et al.⁵⁹, in XI crystallization buffer (35% (w/v) PEG3350, 200 mM lithium sulfate and 10 mM Hepes/NaOH, pH 7.5). For a typical chip sufficient microcrystal were prepared from 25 μ L protein solution (80 mg/ml for xylose isomerase) combined with 25 μ L crystallization buffer. For droplet injection 1M glucose solution was prepared in ddH₂O prior to data collection and stored at room temperature.

Data processing and structure determination

Diffraction data were processed using the CrystFEL package⁶⁰. Structures were solved by molecular replacement using PHASER with our previously determined XI structure as a search model with one molecule in the asymmetric unit (PDB-ID: 6RNF)⁶¹. Structures were refined by iterative cycles in *phenix.refine* and manual model building of additional and disordered residues in COOT0.8⁶²⁻⁶⁴. POLDER-OMIT maps were generated using the PHENIX package⁶⁵. Molecular images were generated in PyMOL⁶⁶.

Experimental setup and data collection

Diffraction data were collected at endstation P14-2 (T-REXX) at EMBL Hamburg on an Eiger 4M detector (Dectris, Baden-Daettwil, Switzerland) and a micro-focussed beam of 7 x 10 μ m. Time delays for time-resolve data collection were generated via the HARE method, reaction initiation was achieved via in-situ droplet injection via the LAMA method as previously described in detail^{15,22}. Data-collection was conducted as previously described⁶⁷ within the environmental control box mounted on the T-REXX endstation. Briefly: chip-mounted microcrystals mounted in a HARE-chip solid target containing ~20,000 wells were moved through the X-ray beam using a 3-axis piezo translation stage (SmarAct, Oldenburg, Germany)²¹). As a broad guideline approximately 5000 still diffraction patterns were recorded per time-point as previously determined⁶⁸, the droplet deposition into each well-defined the start of the reaction.

Data interpretation

Normalized B-factors were determined via the structure comparison tool of the *phenix* software suite⁷².

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Author Contributions:

P.M. and E.C.S. designed the experiments; E.C.S. and P.M. performed the experiments with support from D.v.S. and F.T.; P.M. prepared the protein crystals; E.C.S., P.M., J.P.L. and F.T. designed and built the environmental control box; D.v.S, E.C.S. and P.M. processed and analyzed the diffraction data; P.M. and E.C.S. wrote the manuscript; All authors discussed and corrected the manuscript.

Competing Financial Interests Statement:

The authors declare no competing financial interests.

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