Best practices for time-resolved serial synchrotron crystallography

Eike C. Schulz,a* Briony A. Yorke,b Arwen R. Pearsona,c and Pedram Mehrabi*a

a Institute for Nanostructure and Solid State Physics, Universität Hamburg, HARBOR, Luruper Chaussee 149, 22761 Hamburg, Germany, b School of Chemistry and Bioscience, University of Bradford, Bradford BD7 1DP, United Kingdom, and c Hamburg Centre for Ultrafast Imaging, Universität Hamburg, HARBOR, Luruper Chaussee 149, 22761 Hamburg, Germany. *Correspondence e-mail: eike.schulz@uni-hamburg.de, pedram.mehrabi@uni-hamburg.de

With recent developments in X-ray sources, instrumentation and data-analysis tools, time-resolved crystallographic experiments, which were originally the preserve of a few expert groups, are becoming simpler and can be carried out at more radiation sources, and are thus increasingly accessible to a growing user base. However, these experiments are just that: discrete experiments, not just ‘data collections’. As such, careful planning and consideration of potential pitfalls is required to enable a successful experiment. Here, some of the key factors that should be considered during the planning and execution of a time-resolved structural study are outlined, with a particular focus on synchrotron-based experiments.

1. Introduction

Macromolecular X-ray crystallography has become near-ubiquitous at synchrotrons, which often house multiple heavily automated beamlines dedicated to macromolecular crystallography, permitting routine in-house and even remote data collection without extensive user training. This success has been possible as a result of the increasing levels of automation in the field and the development of standardized workflows. These have resulted in an explosion in the number of protein structures deposited annually into the PDB and have enabled the emergence of related fields such as structural genomics and structure-based drug discovery. Very recently, this wealth of protein structure information has allowed machine-learning approaches to essentially solve the challenge of protein structure prediction from primary-sequence data (Tunyasuvunakool et al., 2021; Baek et al., 2021). With the recent release of the AlphaFold protein structure database (https://alphafold.ebi.ac.uk), hundreds of thousands of (predicted) protein structure models have been made available covering the majority of published protein sequences (Tunyasuvunakool et al., 2021), and in time this is likely to increase to millions. This sudden abundance of structural information has highlighted that the main incentive for macromolecular structure determination is not the structure per se, but rather the increased understanding of the underlying biological function that the structure provides. We thus expect that mechanistic studies, and in particular those that focus on macromolecular dynamics, will see a resurgence in the coming years.

Combining conventional structural biology approaches with other biochemical or biophysical experiments has allowed the...
unravelling of biochemical mechanisms of proteins, for example via mutation studies, substrate analogues or freeze-trapping techniques (Moffat, 2001). However, while these classical X-ray crystallographic approaches have been extremely powerful in obtaining insight into equilibrium-state structures, structures obtained from mutants, or with substrate analogues or inhibitors, have the potential to reflect artefacts that only exist in the mutant or particular ligand complex, particularly when trying to probe a reaction mechanism by determining the structure of reaction intermediate-like states. These can go unnoticed when follow-up studies with other techniques or on wild-type proteins are not carried out (Moffat, 2001). In addition, while the stabilization of certain reaction intermediate states can be achieved with substantial modification of the target protein, many intermediate states are either impractical or simply impossible to obtain by static and trapping methods.

For a comprehensive insight into macromolecular function it is imperative to combine structural information with insight into the time-dependent changes, in other words the dynamics, of the protein of interest. This information can not only reveal metastable intermediates that are difficult to trap, but can also reveal hitherto invisible features of protein function, providing novel insights into catalysis, allostery, half-of-the-sites reactivity, oxidation states, side-chain motions, isomerization reactions and molecular breathing, as well as into molecular binding events (Weinert et al., 2019; Nango et al., 2016; Nogly et al., 2018; Nass et al., 2020; Arnlund et al., 2014; Coquelle et al., 2018; Barends et al., 2015; Kupitz et al., 2014; Tenboer et al., 2014; Stagno et al., 2017; Pande et al., 2016; Mehrabi, Schulz, Dsouza et al., 2019; Kern et al., 2013).

With the introduction of time-resolved crystallography nearly 30 years ago, pioneers in the field harnessed polychromatic X-rays at synchrotron sources to obtain Laue diffraction patterns, ultimately pushing the time resolutions for time-resolved protein crystallography into the picosecond regime (Ren & Moffat, 1994; Ren et al., 1999). However, it was not until the advent of X-ray free-electron lasers (XFELs) and the widespread adoption of serial crystallographic techniques that a wider interest in time-resolved macromolecular crystallography emerged (Levantino et al., 2015; Pearson & Mehrabi, 2020; Orville, 2020).

Soon after its exploitation at XFELs, serial crystallography was rapidly translated to microfocus beamlines at synchrotrons and serial synchrotron crystallography (SSX) was born (Boutet et al., 2012; Chapman et al., 2011; Redecke et al., 2013; Barends et al., 2014; Arnlund et al., 2014; Gati et al., 2014; Roedig et al., 2016; Botha et al., 2015; Stellato et al., 2014). Serial crystallography has a number of advantages for time-resolved diffraction data collection; for example, it avoids the need for reversible systems (Chapman, 2019; Weinert et al., 2017; Schulz et al., 2018; Mehrabi, Schulz, Agthe et al., 2019; Mehrabi, Schulz, Dsouza et al., 2019; Nango et al., 2019). Despite this increasing interest in time-resolved techniques in the past decade, the number of groups working on developing new methodologies remains small and the newly developed tools and techniques are currently mainly only accessible to experts in the field. In addition, in contrast to collecting static data from single crystals, time-resolved crystallography experiments require more careful planning and preparation to obtain successful results. As time-resolved crystallography methods continue to further evolve they will of course undergo simplification and automation and thus become more accessible to a larger user base. However, as this is not yet the case, in this article we aim to provide new users, who are already familiar with classical crystallographic approaches, with a systematic and didactic approach to preparing for and carrying out a time-resolved serial synchrotron crystallographic (TR-SSX) experiment (Fig. 1). Similar considerations are also required for SFX experiments, although these are not the focus of this review. We also highlight some of the common problems that we have encountered so far and how we have adjusted our approach to address these.

2. Preliminary considerations

Ultimately, the design of any time-resolved experiment depends on the scientific question that is being asked, the experimental data that already exist and whether the required experiment is, in principle, feasible.

2.1. The scientific question

We include an explicit consideration of the scientific question being asked as, while it is always tempting to ‘do what you know’, in some cases the question being posed can be answered much more easily with a noncrystallographic approach. Classical biochemical methods such as kinetics and spectroscopy, or alternate biophysical methods such as magnetic resonance and/or fluorescence techniques, can provide a considerable degree of insight into mechanisms of ligand binding, reaction turnover and protein conformational change and can often be performed in-house. SAXS and single-particle electron cryo-microscopy also provide structural data and are particularly well suited to the study of systems where large structural changes are expected. Finally, the tried and tested structural enzymology approaches of cryo-trapping and mechanistic trapping are extremely powerful and it is well worth testing whether such an approach can already answer the question at hand. The adage ‘choose your weapon wisely’ should be taken to heart here.

2.2. Feasibility

In spite of many recent improvements, serial crystallography experiments are still very sample-demanding, and this is even more the case for time-resolved studies where several structures are to be obtained, each of which requires at least ~5000 diffraction patterns (Moreno-Chicano et al., 2019; Weinert et al., 2017; Mehrabi et al., 2021; Gori et al., 2021). Thus, a critical question is whether a sufficient amount of protein is available. The protein should also crystalize readily in order to yield a sufficient supply of reproducible crystals. As in any crystallographic project, the crystals must also diffract
What do I need to do (be)for(e) a TR-SSX experiment?

Preliminary considerations
What is the scientific question?
Is sufficient protein available? Can the question be solved by TR-SSX?
Has the structure been solved? Can crystals be obtained (in large amounts)?
Will crystal packing allow the reaction to proceed?

Room-temperature data
- unit cell/size group?
- diffraction limit?
- pathologies (e.g. twinning)
- radiation damage?
- differences from cryo-data?

Conventional methods
- (bio)chemical trapping data?
- (inactive mutants, substrate analogues, product complexes)
- cryo-trapping data?

(Optimize) microcrystallization
Crystal size and homogeneity

Choose sample environment and characterize microcrystals
fixed target – tape drive – liquid jet – something else
1. tailor crystal size to sample environment
2. apo/dark-state SSX data set (diffraction limit?; differences from cryo-data?)
3. batch-soak SSX data set (diffusion possible?; do crystals survive reaction?; does the reaction happen?)

Choose method of reaction initiation
What timescales should be resolved?
SSX versus SFX?

Optical excitation (<μs)
- native photo-activity?
- photocages available?
- penetration depth?
- quantum yield and fraction excited?
- excitation wavelength?

Diffusion (>ms)
- Soaking feasible? (ligand and solvent channel size, diffusion coefficient)
- compatibility of buffer conditions (viscosity)?
- require ligand concentrations?

Alternatives
- use metal cofactors as triggers
- physical triggers (e.g. temp. jump; X-ray generated electrons)
- chemical triggers (e.g. pH jump)

TR-SSX data-collection strategy
1. acquire a long delay time point (e.g. 1 s) (can the reaction be triggered?)
2. acquire a short delay time point (e.g. 10 ms) (can the reaction be resolved via SSX?)
3. narrow down the time frame of interest (e.g. 30 ms, 100 ms, 300 ms …)
4. fill in the gaps with shorter time delays to trap stable and metastable intermediates

Data analysis
1. integration, scaling, merging
2. data-quality assessment
3. MR and refinement
4. data sufficient to provide reliable electron density that answers the scientific question?

Complementary methods
- kinetics from spectroscopy

no
yes

Figure 1
A workflow scheme for time-resolved serial synchrotron crystallography experiments.
2.3. Prior experimental knowledge

The question of how many preliminary data should be obtained before embarking on a time-resolved experiment is an important one that requires balancing the effort and time needed to generate the preliminary data with how much the data will help to streamline and optimize the time-resolved experiment. In our experience, however, this is always effort well spent.

2.3.1. The crystals. A good understanding of the crystal system will greatly enhance the likelihood of success of a TR-SSX experiment. For example, information on differences in the behaviour of the crystals at room temperature and 100 K, such as unit-cell changes or conformational differences, is important for optimal data processing and subsequent electron-density interpretation. A well-recorded, high-quality single-crystal data set is also invaluable here. Any crystal pathologies, such as twinning, are also best identified in single crystals before any serial data acquisition. If crystal pathologies are present, such as perfect twins or unresolvable indexing ambiguities, then new crystal conditions are needed before further progress can be made.

2.3.2. Soaks and trapped states. Mechanistically and cryo-trapped structures almost always provide valuable insight into (meta)stable intermediates that will aid in the interpretation of the data derived from the TR-SSX experiments. Such structures can be as simple as a substrate, product or inhibitor soak, or can be from mutant variants of the protein of interest. Differences between these trapped states and similar intermediates observed in the time-resolved data can provide important information about the reaction mechanism.

2.3.3. Reaction kinetics. Last, but very much not least, some insight into the reaction kinetics, ideally in the crystals themselves or under the crystallization buffer conditions, should be available. A convenient source to quickly obtain information on protein kinetics is the BRENDA database (https://www.brenda-enzymes.org; Chang et al., 2021). This information helps both with decisions on how to best initiate the reaction and the choice of time points to be measured.

3. Sample preparation

3.1. Microcrystal properties: size and homogeneity

Different sample-delivery and reaction-initiation methods have different crystal size requirements, and thus the crystal size has to be tailored to the planned experiment. While fixed-target and tape-drive approaches are relatively flexible with respect to crystal size, liquid-jet methods usually require smaller crystals (<10 μm; Mehrabi et al., 2020; Beyerlein et al., 2017; Martiel et al., 2019). The crystal size also directly affects the penetration depth of optical triggers and the diffusion time in mixing approaches (Mehrabi, Schulz, Agthe et al., 2019; Schmidt, 2013; Grünbein et al., 2020). It is also important that the microcrystals are as homogenous as possible. This is important in ensuring that the reaction initiation is similar for all crystals and also improves the downstream processing of the diffraction data.

3.2. Obtaining microcrystals

The crystallization conditions for previously obtained single crystals are an excellent starting point for obtaining microcrystals. Usually, these crystallization conditions can be tweaked to increase nucleation to yield more, smaller crystals. Some of the common methods for obtaining microcrystals will briefly be discussed here (Fig. 2).

3.2.1. Mechanical disruption. The simplest and quickest method to obtain a slurry of microcrystals and even nanocrystals is to apply mechanical force. This is typically performed using large single crystals obtained by standard hanging-drop or sitting-drop methods, isolating these crystals in sufficient number, adding glass beads and vortexing. The glass beads crush the large crystals and, depending on how long the sample is vortexed for, result in different size distributions of crystals. This method has been used to obtain nano-sized or near-nano-sized crystals for use in gas dynamic virtual focusing nozzle (GDVN) liquid jets at XFELs or for serial electron diffraction (Bücker et al., 2020; Cheng, 2020). The primary drawbacks of this method are that the shearing and crushing forces applied to the crystals can result in mechanical disruption of the crystal lattice, reducing the diffracting power, and that it usually results in a quite heterogeneous crystal size distribution that can require large crystal fragments to be removed by filtration.

3.2.2. Mass vapour-diffusion crystallization. Modifying the starting vapour-diffusion conditions, usually by increasing the precipitant or protein concentration, can result in reasonably sized microcrystals using standard vapour-diffusion methods. Subsequently, crystals from hundreds of hanging or sitting drops must be pooled together for use at the beamline. While this approach does not require any specialist hardware beyond that found in a normal crystallography laboratory, it is extremely tedious, and the harvesting and pooling of drops can be detrimental to crystals which are sensitive to handling.
3.2.3. Batch crystallization. Batch methods, in which large volumes (usually >50 μl) of both mother liquor and protein solution are mixed in a vial, are ideal for obtaining large numbers of homogenously sized microcrystals. Recently, a number of systematic approaches for obtaining microcrystals via batch-crystallization methods using different precipitation agents have been described (Beale et al., 2019; Stohrer et al., 2021). An extension to batch crystallization takes advantage of rapid evaporation from a vacuum environment (Martin & Zilm, 2003). This method was initially used to obtain nanocrystals for use in solid-state NMR and has two major advantages: (i) placing the mixed solution under vacuum results in very rapid nucleation, and thus microcrystals can usually be obtained within 10–20 min, for example immediately prior to data collection, and (ii) because nucleation occurs in a very short time frame, the resulting crystals generally have a homogeneous size distribution.

3.2.4. On-target crystallization. If fixed targets are used for sample delivery, then as an alternative to batch crystallization, crystals can be grown directly on the fixed target. Here, the protein and mother liquor are mixed and applied directly onto the chip surface. If the sample is viscous it can be forced into the features of the chip by drawing a razor blade across the chip surface. Crystals are then grown using a vapour-diffusion process very similar to the hanging-drop method (Lieske et al., 2019; Norton-Baker et al., 2021). In addition to allowing direct transfer of the starting vapour-diffusion conditions, this approach completely avoids crystal handling, produces a very homogeneous crystal size distribution and has the lowest sample consumption of all of the methods described here. Single data sets can be obtained from only micrograms of protein (Norton-Baker et al., 2021).

3.2.5. Seeding. If obtaining either the desired crystal size or a homogenous distribution is difficult, microcrystal seeding in batch is a powerful approach. By using relatively high concentrations of seeds, high yields of microcrystal crystals of the desired size and low size heterogeneity can be obtained.

3.3. Initial crystal characterization

3.3.1. Diffraction quality. To determine whether microcrystals are suitable for a TR-SSX experiment, minimally a static data set should be acquired. This can be performed simply using grid or mesh scans on a spine loop both at cryo and room temperatures to determine the resolution limits and general diffraction properties, ideally using the same beam parameters as will be used for the time-resolved experiment. The focus of this characterization should of course be on the room-temperature data. If possible, it is also advisable to test the full crystal-handling workflow planned for the time-resolved experiment, for example chip-loading conditions, liquid-flow parameters in microfluidic devices or jets etc., in order to avoid any surprises during the actual experiment.

Figure 2
Overview of commonly used microcrystallization techniques and SSX sample environments. Top: commonly applied microcrystallization approaches include (a) hanging-drop or sitting-drop crystallization, (b) batch crystallization, (c) vacuum-induced (batch) microcrystallization, (d) in-chip microcrystallization based on vapour-diffusion protocols and (e) LCP-based microcrystallization. Bottom: commonly used sample environments for serial synchrotron crystallography are (f) fixed-target-based approaches, (g) liquid or viscous jets, (h) hybrid methods such as tape-drive sample delivery and (i) microfluidic systems.
Obtaining suitable microcrystals for time-resolved studies often requires iterative optimization of both the crystal-growth conditions and the crystal handling (including sample delivery). An important parameter that must not be neglected is the humidity, as slight changes in hydration both improve and obliterate diffraction (Bowler et al., 2006; Sanchez-Weatherby et al., 2009; Russo Krauss et al., 2012).

3.3.2. Turnover. An important final control before the time-resolved experiment itself is a simple batch soak (for long enough for the reaction to complete) of the microcrystals with the substrate/ligand of interest or, for photocaging experiments, illumination, which allows the quick assessment of a number of important parameters. Do the microcrystals survive exposure to the ligand or optical trigger? Are diffraction properties affected? Does the reaction actually take place in the substrate/ligand of interest or, for photocaging experiments, activation? And is the microcrystalline material amenable to either mixing/diffusion or photo-activation?

4. Sample delivery

Serial crystallography approaches at both XFELs and synchrotrons generally encompass four types of sample-delivery methods, which include liquid-injection, microfluidics, fixed-target and hybrid approaches; these have recently been reviewed in detail (Martiel et al., 2019; Grünbein & Nass Kovacs, 2019). We will therefore limit the discussion here to sample environments which have successfully been used for time-resolved data collection at synchrotrons (Fig. 2). Assuming that the crystals are all well diffracting, isomorphous and randomly oriented with respect to the X-ray beam, the hit rate is the major consideration when choosing a sample-delivery method.

4.1. Liquid and viscous jets

The first serial sample-delivery devices employed at XFELs were liquid-injection devices using a GDVN. Due to their speed, GDVN injectors are well suited to XFEL experiments but are less ideal for synchrotron measurements, especially on monochromatic macromolecular crystallography (MX) beamlines. Similar devices have, however, been developed to extrude viscous media such as LCP; these deliver sample more slowly and are ideal for the longer exposure times required at synchrotron sources (Martiel et al., 2019). These can be combined with photoactivation for time-resolved experiments (Weinert et al., 2019).

4.2. Fixed targets

Fixed-target approaches generally come in one of two flavours, either using spine mounts with mesh loops or one of several different solid-support solutions (Martiel et al., 2019).

4.2.1. Mesh loops. The simplest fixed-target approach involves loading a mesh loop with a crystal slurry and either performing a simple grid scan, a helical scan or a MeshAndCollect-type data collection (Zander et al., 2015). While quick and simple, this approach generally requires cryo-conditions and is not amenable to either mixing/diffusion or photo-activation.

4.2.2. Solid supports. A number of solid-support designs have been reported that can be raster-scanned through the X-ray beam either using a modified goniometer (for example Roadrunner) or using (piezo-driven) three-axis stages (for example HARE/Oxford photochips) (Mehrabi et al., 2020; Oghbaey et al., 2016; Zarrine-Afsar et al., 2012; Sherrell et al., 2015; Roedig et al., 2017). The advantage of solid supports are a lower sample consumption relative to liquid-jet approaches (Norton-Baker et al., 2021; Mehrabi, Schulz, Agthe et al., 2019; Lieske et al., 2019) and a relatively high hit rate, which can be up to 100% if crystal locations can be mapped prior to data collection (Oghbaey et al., 2016). Additionally, many of the fixed-target solutions can accommodate a wide range of crystal sizes and morphologies without modification of the support. This allows easy screening of different samples or different crystals of the same sample within a single experiment.

Currently, the HARE/Oxford photochips are the only fixed-target chips that are routinely used for time-resolved measurements at both XFELs and synchrotrons (Schulz et al., 2018; Mehrabi, Schulz, Agthe et al., 2019; Mehrabi, Schulz, Dsouza et al., 2019). Both the HARE and Oxford photochips are amenable to experiments involving photoactivation or mixing/diffusion using the LAMA approach (Mehrabi, Schulz, Agthe et al., 2019). Because the features on these chips are well ordered and each position is known, crystals can be imaged more than once, allowing unique data-acquisition schemes such as the Hit-And-Return (HARE) approach (Schulz et al., 2018). This method allows data acquisition with a wide range of delay times from a few milliseconds to many seconds, while keeping the total data-collection times per chip low. A combination of the HARE and LAMA approaches is currently the primary method of choice at the T-REXX endstation (Beamline P14.EH2) at PETRA III, which is the world’s first dedicated endstation for time-resolved serial synchrotron crystallography.

4.2.3. Microfluidic devices. Microfluidic devices, such as 3D-MiXD, employ simple 3D-printed designs with multiple channels controlled by a series of syringe pumps where crystals and substrate can be mixed before flowing into the X-ray beam path for imaging (Monteiro et al., 2020). They are simple to set up, run reliably for many hours and are compatible with both photoactivation and rapid mixing, and their sample consumption is \(\sim 50\) µl h\(^{-1}\) (Monteiro et al., 2020). While some shortcomings do exist with microfluidic devices, such as clogging due to bubble formation or fouling of the Kapton windows, these can largely be avoided with the aid of a fast shutter.

4.2.4. Hybrid approaches. Hybrid approaches, such as the mix-and-diffuse or drop-on-tape approaches, employ a moving thin Kapton film onto which droplets containing crystals are deposited before being carried through the X-ray path (Roessler et al., 2013; Butryn et al., 2021; Beyerlein et al., 2017; Fuller et al., 2017). The drop-on-tape method uses a combination of microfluidics to mix both crystals and substrate...
5. Reaction initiation

All time-resolved crystallography experiments use the pump-probe approach in which a reaction is initiated (pumped) in a crystal, which is then probed, after a defined time delay, by X-rays to yield a diffraction pattern. A key aspect of the experimental design is therefore how the reaction will be initiated. Important here is that a sufficient fraction of molecules in the crystal are activated as synchronously as possible to ensure that the resulting time-dependent electron-density maps can be interpreted and that the temporal resolution is appropriate to the timescale being probed. We note here that many enzymes and biomolecular processes involving conformational changes are in fact relatively slow, with median turnover rates of the order of $k_{\text{cat}} = 13.7 \text{ s}^{-1}$ (Bar-Even et al., 2011). These timescales are ideally suited for study by time-resolved SSX.

5.1. Accessible timescales are a function of both the X-ray source and the reaction-initiation method

5.1.1. X-ray source. The minimal time needed to record a usable diffraction pattern from a macromolecular microcrystal is a function of the brightness of the X-ray beam. Third- and fourth-generation monochromatic microfocus synchrotron MX beamlines can reach a time resolution of at least 100 $\mu$s, and with multiplexing methods such as HATRX, can, in principle, access nanosecond timescales, although this has yet to be experimentally demonstrated (Yorke et al., 2014). Laue crystallography beamlines can achieve a time resolution of 100 ps. Sub-100 ps experiments are the preserve of XFELs, which can even address events on the order of tens of femtoseconds (Chapman, 2019). For time-resolved structural biology the availability of suitable X-ray sources is no longer a limiting factor.

5.1.2. Reaction-initiation method. The time required for reaction initiation is the true limitation on the timescales that can be probed by time-resolved crystallography. For systems where the reaction can be triggered by photoisomerization or direct photocleavage, femtosecond timescales are accessible. However, when a series of dark reactions must occur after the initial photoexcitation before the system becomes fully active, or where initiation is by rapid mixing, the accessible timescales become much longer.

5.2. Photocages

5.2.1. Photocages. As only a fraction of proteins are naturally photosensitive (<0.5%; Monteiro et al., 2021), caged compounds are an attractive option for reaction initiation in synchrotron-based time-resolved experiments, where the timescales of interest are usually >100 $\mu$s. Either a ligand or the protein can be made photosensitive by the addition of a photolabile protecting group, which can then be released with a laser pulse. The photocaged moiety should be inert and, if a small molecule has been caged, can be co-crystallized or soaked into the crystal prior to X-ray measurements. Selection of the optimal photocaging group for a particular experiment requires the consideration of a number of parameters, and these are discussed extensively in the recent review by Monteiro et al. (2021). A key consideration is the decaying rate, i.e. how quickly the uncaged, active state of the system is achieved. Obviously, this must be faster than the reaction steps to be studied in the time-resolved experiment. It is not always appreciated how slow decaging can be. While pH jumps using photocoids can occur on sub-microsecond timescales, many cages release in the millisecond time regime. The concentration of the caged compound should also be at least on parity with the concentration of the protein in the crystal, and ideally higher. For experiments using photoactivation, care must be taken when dealing with lasers of high power densities to avoid artefacts which can arise due to multiphoton effects; this is discussed in detail by Grünbein et al. (2020).

5.2.2. T-jumps. Excitation of the water within a crystal with an IR pulse can result in a temperature jump of several kelvin within a few nanoseconds as the energy is rapidly transferred from the water to the proteins. This can be sufficient to trigger reaction turnover, but has so far seen most application in SAXS/WAXS experiments rather than in TR-SSX (Levantino et al., 2015).

5.3. Diffusion or rapid mixing

While the diffusion times of small-molecule ligands into macroscopic crystals can be of the order of several seconds, the situation is quite different in microcrystals. Due to the surface-to-volume ratio, the diffusion time accelerates by two orders of magnitude when the crystal volume is reduced by only one order of magnitude; this relationship gives rise to low-millisecond diffusion times for microcrystals with dimensions of a few micrometres (Schmidt, 2013; Mehrabi, Schulz, Agthe et al., 2019; Butryn et al., 2021; Beyerlein et al., 2017; Schmidt, 2020). This means that mixing/diffusion-based methods not only become viable but are an excellent and simple method for reaction initiation in time-resolved SSX experiments.

In general, mixing methods require a simpler hardware setup compared with photoexcitation approaches and circumvent many of the difficulties associated with, for example, multiphoton effects. Nevertheless, some thought is still required when using mixing to initiate the reaction. As for photoactivation, the turnover rate of the target protein should be lower than the time required for the ligand to diffuse to the centre of the crystal. This critical depth parameter is a function of the kinetic properties of the enzyme and the diffusion coefficient of the ligand (Makinen & Fink, 1977). The diffusion coefficient can vary greatly depending on the size and hydrophobicity of the ligand, the viscosity of the crystallization
mother liquor and the size of the solvent channels within the crystal. As with caged compounds, the concentration of the ligand should be at least that of the protein in the crystal. In practice it is advisable to (greatly) exceed this concentration and work with the maximum concentration possible. This may require some screening of crystallization mother liquors to identify conditions which maintain the crystal while also allowing ligand solubilization at high concentrations.

Many proteins can be activated via secondary triggers, such as cations, which are required for protein function (Richter, 2013). This can be a convenient and elegant workaround to achieve rapid and homogeneous activation as diffusion of ions is quick and crystallizing a protein–ligand complex may prove to be easier than the protein alone.

6. Choice of time points

6.1. Crystals versus solution

The crystal environment can alter the kinetics of the system compared with those measured in dilute solution, and in extreme cases can result in deviations into dead-end or non-physiological pathways (Konold et al., 2020; Makinen & Fink, 1977). A simple test here is to remeasure the solution-phase kinetics but using a buffer system as close to the crystallization conditions as possible. Alternatively, microspectrophotometry can be used to directly follow the reaction progress in the microcrystalline slurry and precisely define the timescales of interest (Pearson et al., 2004; Wilmot et al., 2002; Pearson & Mozzarelli, 2011).

6.2. Shoot it and see?

If no data on the kinetics in the crystalline state are available, the simplest data-collection strategy, albeit high-risk, is to use the existing catalytic timescales obtained from solution measurements and observe whether intermediates obtained from the TR-SSX measurement agree. However, this can be risky, especially when beamtime is limited. We therefore suggest instead initially measuring one long time delay (for example >1 s) to determine whether the reaction has run to completion in the crystal. Next, one short delay (for example 10–20 ms) should be recorded to test whether the reaction is slow enough to be resolved by TR-SSX, i.e. whether the end state has already been reached or not. This approach makes good use of beamtime and allows mapping of the temporal boundary conditions.

Following this, a simple data-collection strategy can be employed whereby series of initially short time delays are measured (10, 50, 100 ms . . . ) with subsequent time delays becoming longer, representing a pseudo-logarithmic timescale that attempts to capture the entirety of the catalytic cycle. For mixing/diffusion-based approaches the initial time point that can be collected can be estimated based on knowledge of the crystal size and the expected or measured diffusion times. Following analysis of the data obtained, further measurements can fill in the gaps around key events.

6.3. Data-collection modes

6.3.1. Classical pump–probe. Pump–probe experiments display the simplest data-collection strategy: an individual crystal is ‘pumped’ via some trigger, that is the reaction is initiated, and after a desired delay time has been achieved the crystal is ‘probed’ via the X-ray pulse. This is the standard scheme for acquiring time points with a short time delay (<100 ms; Fig. 3b, i).

6.3.2. Burst pump–probe. In a burst experiment, taking advantage of high-repetition-rate detectors, each crystal is pumped and a sequential series of diffraction images are acquired with a \( \Delta t \) equaling the repetition rate of the detector before moving to the next crystal. The images are then stacked according to their respective time points (Ebrahim et al., 2019; de la Mora et al., 2020; Figs. 3a and 3b, iii).

6.3.3. HARE. The Hit-And-REturn (HARE) approach allows the collection of long time delays without the large increase in total data-collection time for a chip that would otherwise result from a standard pump–probe measurement with serial data collection (Schulz et al., 2018). Unlike a standard pump–probe experiment, in a HARE experiment a series of crystals are first pumped and subsequently probed in the same order. This means that the HARE method enables collection of time delays that can span up to several minutes, while the total data-collection time for a chip remains the same (Fig. 3b, ii).

7. Data-quality assessment

Obtaining high-quality data should be the goal when planning the TR-SSX experiment, and rapid feedback on the progression of the experiment in terms of hit rates, average resolution and ideally electron-density maps is invaluable in order to help to determine when sufficient data have been collected for each time point (Basu et al., 2019; Bieilsten-Edmands et al., 2020; Ke et al., 2018; Barty et al., 2014; White et al., 2016). Good-quality data are especially important for time-resolved studies, where dynamic and low-population states can complicate the interpretation of electron-density maps. Time-resolved XFEL-based experiments, which generally involve femtosecond to nanosecond timescales, require a more rigorous treatment of the experimental setup, especially in terms of timing tools and in terms of processing and analyzing XFEL data, where careful validation is required to avoid overestimating the small structural changes occurring on ultrafast timescales. The excellent recent review by Gorel and coworkers extensively discusses the best practices for XFEL-based experiments, many of which also hold true for synchrotron-based experiments (Gorel et al., 2021).

7.1. How many data are enough?

Arguably the first consideration when assessing the quality of any data set should be completeness; this single parameter is generally applicable to any crystallographic experiment and provides context for all other data-quality parameters. If the completeness of the data set is below an approximate cutoff of
95% then the meaning that can be derived from the scaling statistics is limited (Mehrabi et al., 2021). The pertinent question when collecting data may then appear to be ‘how many diffraction images do I need to collect to obtain a complete data set?’. However, particularly in a time-resolved experiment, completeness alone is not enough. The data should also be of high redundancy in order to ensure stable scaling and reliable error estimation.

The amount of data needed is ultimately dependent on both the sample and the experimental design. Although the minimum reported value for a good-quality data set is ~5000 indexable diffraction patterns, this number certainly increases the less populated the state one wants to observe is (Weinert et al., 2017; Moreno-Chicano et al., 2019; Mehrabi et al., 2021). Therefore, in order to determine when an experiment is complete, it is beneficial to monitor the hit rate (i.e. the ratio of indexable diffraction patterns to the number of X-ray pulses), scaling statistics (such as $CC_{1/2}$ or $R_{\text{split}}$), completeness, redundancy and resolution.

Prior knowledge of the space group and unit cell of the crystal are also essential when carrying out time-resolved experiments to help to avoid indexing ambiguities and support automatic data processing. In principle, higher symmetry point groups require fewer images for a complete data set (Dauter, 1999). However, the need for high redundancy and reliable error estimate remain and so in practice there is no real reduction in the number of diffraction patterns that are needed. On-the-fly data-processing methods utilizing the computational parallelization available at XFEL and synchrotron sources to allow the progress of the experiment to be monitored in real time are key to ensuring that data-collection strategies optimize data quality and experimental efficiency (Basu et al., 2019; Beilsten-Edmands et al., 2020; Ke et al., 2018; Barty et al., 2014; White et al., 2016). The hit rate of the experiment is a useful metric to give an indication of how many runs of the experiment will be required, but measuring this single metric does not indicate the completeness of a data set, only the efficiency of the delivery method.

7.2. Preferred orientation

In both fixed-target and flow-based sample-delivery approaches the crystal morphology can cause crystals to adopt a preferred orientation with respect to the X-ray beam. If this occurs it can be problematic, as serial crystallographic delivery systems are generally not designed to allow reorientation of the sample. Real-time feedback about the completeness and multiplicity of a data set can provide an early indication of this issue (Mariani et al., 2016), and methods have been developed to encourage random orientation in jets and fixed-target chips (Schlichting, 2015; Murray et al., 2015). Utilizing microfluidic and acoustic drop injection techniques can be useful in such cases (Zhao et al., 2020; Davy et al., 2019).

7.3. Isomorphism

The requirement that every crystal in a particular sample is isomorphous should not be taken lightly, as the presence of non-isomorphism can obscure the presence of weakly bound ligands and states with low populations. Separating data by grouping different isomorphs according to the unit-cell dimensions, cross-correlation statistics or by utilizing genetic algorithms can help to address this issue (Foadi et al., 2013; Giordano et al., 2012; Santoni et al., 2017; Zander et al., 2016; Ginn, 2020).

Figure 3

(a) Schematic representation of burst data collection. (b) Visual representation of various data-collection schemes on the HARE chip from standard pump–probe to the Hit-And-Return (HARE) approach to burst data collection.
It should also be noted that both laser- and diffusion-based initiation can have a profound effect on the isomorphism of the sample. Changes in unit-cell size and a gradual decline in diffraction quality may be observed, especially at longer time delays, due to a progressive increase in the disorder of the crystalline lattice and an increase in the superposition of intermediate states. Large amounts of energy are deposited into the crystal during laser initiation, and changes in temperature and pH can impact not only the diffraction quality but also the kinetics of the reaction that is being followed (Grünein et al., 2020).

7.4. Resolution

Resolution should be considered in terms of the minimum resolution necessary to reliably identify the important structural changes associated with the reaction of interest. A large protein conformational shift may be apparent at ~3 Å resolution; however, a higher resolution is needed if conclusions are to be drawn about bond formation/breakage, backbone flips, rearrangement of solvent molecules, oxidation state and isomerization, where small changes to specific interatomic distances are integral to the structural change being monitored.

7.5. Scaling statistics

In a single-crystal experiment $R_{\text{meas}} (|I|/\sigma(I))$ and completeness are the most commonly reported statistics, and a number of rules of thumb relating to ‘acceptable’ values of $R_{\text{meas}}$ and $|I|/\sigma(I)$ have emerged (Karplus & Diederichs, 2012) that are not necessarily directly applicable to serial data. $R_{\text{meas}}$ measures the spread of recorded intensities for a particular reflection around the average intensity, with a correction to account for redundancy. Since the strength of diffraction data decreases with resolution, a resolution cutoff point is often defined where $R_{\text{meas}}$ reaches a value of approximately 0.6 and $|I|/\sigma(I)$ is approximately equal to 2. The rationale behind these cutoff values was to ensure that weaker, noisy data are not included in structure refinement. However, with the almost ubiquitous use of photon-counting detectors the justification for excluding weak data [as defined by $|I|/\sigma(I)$] has become almost obsolete (Evans & Murshudov, 2013). In addition, $|I|/\sigma(I)$ as an indication of the signal-to-noise ratio (SNR) is not always reliable due to the various ways in which $\sigma(I)$ may be calculated.

7.5.1. $R_{\text{merge}}$ and $R_{\text{split}}$. $R_{\text{merge}}$ is particularly problematic as a data-quality indicator for serial data, since serial data reduction relies on sampling large distributions of intensities. In serial data the spread of intensities is large by design and so this metric provides no useful indication of data quality (White et al., 2012). In the case of serial data $R_{\text{merge}}$ is better replaced by $R_{\text{split}}$, where the data are split into two equal (and random) groups and the agreement between the intensities in each list is compared and corrected for multiplicity.

7.5.2. $CC_{1/2}$. The half-data correlation coefficient $CC_{1/2}$ is widely reported for serial data as it provides a useful metric for determining a suitable resolution cutoff and for identifying rogue images (Assmann et al., 2016; Karplus & Diederichs, 2015). Here the data are split into two randomly selected groups, or upwards of ten resolution shells, and the Pearson correlation coefficient is calculated between the average intensities in each subset. This method has the advantage of giving a clear indication regarding the precision of a data set with values ranging from 1 (perfectly correlated, precise) to 0 (no correlation, imprecise) (Assmann et al., 2016).

In summary, it is unwise to rely on a single scaling statistic to guide data-collection strategies. $R_{\text{split}}$, $|I|/\sigma(I)$ and $CC_{1/2}$ each provide useful insight into the overall quality of the model and should all be taken into account (Wang, Brudvig et al., 2017).

7.6. Electron density

Although scaling statistics provide an indication of the overall quality of the data sets for each time point, they do not provide information about the structure or the actual reaction being investigated. Only by inspection of the electron density of each individual data set is it possible to confirm the presence of a ligand or intermediate or the successful initiation of a reaction in the crystal. Although the use of complementary spectroscopic methods can aid with monitoring these changes in real time, a spectroscopic signal is no guarantee that the structural changes will be discernible in the electron density.

8. Radiation damage

8.1. Minimum exposure time

For a TR-SSX experiment the minimum exposure time to allow the required resolution to be achieved, whilst also ensuring that the data can be indexed and integrated reliably, should be determined before the time-resolved experiment begins. This is important to minimize the effects of radiation damage on the data.

8.2. Dose limits for room-temperature SSX

The radiation tolerance and the available dose budget for SSX experiments at room temperature has recently been addressed by de la Mora and coworkers, who systematically compared serial data collection from lysozyme crystals at cryo and room temperatures. They established a global damage dose limit of 15.3 MGy at 100 K and 0.57 MGy at room temperature (de la Mora et al., 2020). However, specific radiation damage, which could obscure or even initiate catalytic processes, is a greater concern in time-resolved studies, and they suggest keeping the cumulative dose per crystal below 80 kGy. This limit may be even lower if the system being investigated contains metal centres or redox-active cofactors that can be damaged even at very low doses. These dose limits can be directly translated into the number of exposures that a crystal can tolerate in an SSX burst experiment and thus put a boundary on the maximal achievable number of images per crystal.
8.3. Monitoring radiation damage

In the cases of proteins containing metal centres, chromophores and photocaged ligands site-specific damage is an extremely important consideration, although there is evidence that global and specific radiation damage are ‘re-coupled’ at room temperature and that specific damage may therefore be less of a concern than initially feared (Gotthard et al., 2019). Real-time monitoring of the sample by complementary methods such as UV–Vis and Raman spectroscopy can help to identify X-ray-induced changes to the structure that may not be linked to the reaction of interest.

In experiments using photocages where a ‘dark’ image is recorded from a crystal before photoactivation, spectroscopic measurements are also important to confirm that decaging only occurs upon laser illumination. There is some evidence indicating that photocleavage of caged ligands may be initiated by X-rays as well as UV light (Fig. 4; Specht et al., 2001). However, there are also cases where the photocaged group is clearly unaffected by X-ray exposure, suggesting that the susceptibility of the photocage to X-rays may be modulated by the buffer conditions, and by the local environment in cases where the photocaged compound is specifically bound within the crystal (Kesgin–Schaefer et al., 2019).

9. Assessing low-populated states

Electron-density maps derived from time-resolved structural data contain a superposition of unreacted and reacted molecules, especially at longer time points where the system has not yet reached a steady state but the individual molecules sample a distribution of states along the reaction coordinate. Careful experimental design that maximizes the fraction of molecules reacting can help, but the data will always contain a mixture of states. This means that the difference electron density can be very challenging to interpret and may not provide a clear picture of the dynamics and intermediate states involved in a particular reaction. This is the major challenge of time-resolved crystallographic data analysis: how to successfully identify, interpret and model low-population states or deconvolute a mixture of states?

9.1. Singular value decomposition

Singular value decomposition (SVD) is one of the most established methods for deconvoluting mixtures of states and was first used in Laue time-resolved crystallographic experiments (Ren et al., 2013; Schmidt et al., 2003). SVD deconvolutes the data into time-independent and time-dependent vectors, ultimately allowing weighted electron-density maps to be calculated that reflect the time-independent states contributing to the measured data. Combining SVD with data collection at a range of different temperatures and/or pH values, so-called five-dimensional crystallography, can be used to quantitatively analyse the reaction kinetics within a crystal and guide the optimization of an experiment to access low-populated states by altering the reaction kinetics and manipulating the lifetimes of short-lived intermediates (Schmidt et al., 2013).

9.2. Pan-data-set analysis

Pan-data-set density analysis (PANDDA) was originally developed to identify low-occupancy ligands in fragment-screening data sets (Pearce et al., 2017). It exploits the large number of data sets typically collected during a fragment-screening campaign to make a statistical analysis of the electron-density distributions in order to identify data sets with unique features that could be low-occupancy states and ligands. Time-resolved data sets can be analysed in the same way.
way in order to identify time-dependent changes in electron density.

9.3. Clustering

As discussed in Section 7.3, averaging of SSX data recorded from non-isomorphous crystals results in the loss of clear electron-density features for low-populated states. In this case, clustering of the data prior to scaling and merging can greatly improve the downstream analysis using tools such as PANDDA.

Cluster4x uses the correlation between the differences in Cα positions (or reflection amplitudes) to improve clustering prior to PANDDA analysis (Ginn, 2020), and we have found this to be particularly effective for the identification of weakly bound ligands in protein structures. When combined with SVD, it also aids in selecting the correct number of vectors and therefore intermediate states for time-resolved data analysis.

9.4. Modelling heterogeneity

Even at high resolutions, the modelling of conformational heterogeneity, or even only the modelling of multiple conformations, is often prone to subjectivity. It can therefore be beneficial to utilize large-scale computational analysis and parallelized refinement to sample large numbers of different conformation combinations in order to determine those which best describe the data. 

Qfit is a program that uses mixed integer quadratic programming (MIQP) to successfully model hidden conformations (Keedy et al., 2015; Fraser et al., 2011; Lang et al., 2014; Riley et al., 2021).

10. Complementary experiments

Although time-resolved crystallography can provide a huge amount of information, other biophysical and biochemical techniques are also required to obtain a full understanding of the mechanism of interest.

10.1. Neutron crystallography

Neutron crystallography (NMX) is a unique complement to TR-SSX experiments as it is able to experimentally determine protonation patterns. A neutron crystal structure of one or more stable or cryo-trapped intermediates can provide valuable insights, especially where the resolution of the data from TR-SSX is insufficient to reliably infer proton positions (Kono & Tamada, 2021).

10.2. NMR spectroscopy

Nuclear magnetic resonance (NMR) spectroscopy provides access to the thermodynamics and kinetics of protein−ligand interactions and enables a detailed picture of the equilibrium-state conditions. NMR relaxation methods can also provide insight into the populations of key states with relative populations down to ~1% (Vallurupalli et al., 2012). In addition, it can yield exchange rates and provide a wealth of other dynamic information on allosteric and conformational selection pathways (Kim et al., 2017). Moreover, NMR and electron paramagnetic resonance (EPR) are perfectly suited to provide information on intrinsically disordered regions of the protein that are otherwise completely invisible using structural techniques (Drescher, 2012; Gibbs et al., 2017).

10.3. Molecular simulations

Computational approaches such as molecular dynamics (MD) and hybrid quantum-mechanical/molecular-mechanics (QM/MM) simulations can be immensely helpful in filling in the gaps for otherwise inaccessible timescales from femtoseconds to microseconds and, with more rigorous approaches, providing energy landscapes for key chemical events along the reaction coordinate. Enhanced methods have been developed to calculate the rates of transitions between states and to include kinetic rate constants as constraints in MD simulations (Brotzakis et al., 2021; Wang, Martins et al., 2017). An approach that appears particularly interesting to TR-SSX experiments is to guide such simulations by experimental data (Bottaro et al., 2018).

10.4. Electronic and vibrational spectroscopy

Spectroscopic techniques, such as UV-Vis, Raman, IR etc., can provide key insights into the electronic states of metal cofactors and chromophores, radiation-damage processes and reaction kinetics and are perfectly suited to pinpointing the timings of state transitions (Balakrishnan et al., 2004). Multi-dimensional spectroscopic methods can also provide additional insights into how local changes are coupled to larger-scale dynamics and are extremely complementary to time-resolved structural data.

11. Summary and conclusions

Time-resolved structural studies can extend our understanding of protein function by making not just structure but also function visible. With recent developments in X-ray sources, instrumentation and data-analysis tools, time-resolved experiments, which were originally the preserve of a few expert groups, are becoming simpler, can be carried out at more radiation sources and are thus increasingly accessible to a growing user base. However, these experiments are just that: discrete experiments, not just ‘data collections’. As such, careful planning and consideration of potential pitfalls is required to enable a successful experiment.

Acknowledgements

Open access funding enabled and organized by Projekt DEAL.

Funding information

PM acknowledges support from the Deutsche Forschungsgemeinschaft (DFG) via grant No. 451079909 and from a Joachim Herz Stiftung add-on fellowship. ES acknowledges support from the Deutsche Forschungsgemeinschaft (DFG) via grant No. 458246365. ES and ARP acknowledge support
from the Joachim Herz Stiftung via the Biomedical Physics of Infection Consortium. BAY is supported by Academy of Medical Sciences Springboard via Grant No. SBF0061044. ARP acknowledges support from the Cluster of Excellence ‘The Hamburg Centre for Ultrafast Imaging – Structure, Dynamics and Control of Matter at the Atomic Scale’ of the Deutsche Forschungsgemeinschaft (DFG), EXC1074, and the Cluster of Excellence ‘CUI: Advanced Imaging of Matter’ of the Deutsche Forschungsgemeinschaft (DFG), EXC1074, and the project ID 390715994. The development of the T-REXX TR-SSX endstation is supported by the BMFB (‘Verbundforschung’, 0SK16GU1 and 0SK19GU1).

References

Eike C. Schulz et al. • Time-resolved serial synchrotron crystallography