Abstract

X-ray crystallography at X-ray free-electron laser (XFEL) sources is a powerful method for studying macromolecules at biologically relevant temperatures. Moreover, when combined with complementary techniques like X-ray emission spectroscopy (XES), both global structures and chemical properties of metalloenzymes can be obtained concurrently, providing new insights into the interplay between the protein structure/dynamics and chemistry at an active site. Implementing such a multimodal approach can be compromised by conflicting requirements to optimize each individual method. In particular, the method used for sample delivery greatly impacts the data quality. We present here a new, robust way of delivering controlled sample amounts on demand using acoustic droplet ejection coupled with a conveyor belt drive that is optimized for crystallography and spectroscopy measurements of photochemical and chemical reactions over a wide range of time scales. Studies with photosystem II, the phytochrome photoreceptor, and ribonucleotide reductase R2 illustrate the power and versatility of this method.

INTRODUCTION

X-ray scattering/diffraction and spectroscopy are valuable tools for studying biomolecular mechanisms because they yield direct and element-specific molecular geometry and electronic state information. X-ray signals generated by excitation with <50 femtosecond pulses from an X-ray Free Electron Laser (XFEL) are produced before radiation-induced changes occur (picosecond time-scale)\(^1\)–\(^4\). Consequently, XFELs provide new opportunities to study biomolecules at ambient temperature and probe enzymatic reaction intermediates prepared \textit{in situ}. To understand enzymatic mechanisms in detail, both electronic and...
molecular structural information are needed for reaction intermediates and in practice this requires that complementary time-resolved methods be used to probe the reaction. Time-resolved studies of biomolecules, however, are challenging at an XFEL because the focused XFEL pulses destroy the sample after each shot. This requires that the sample be continually replenished in a consistent state, with time-delays from reaction initiation to the X-ray probe in the picoseconds to seconds time scale\textsuperscript{5,6}.

Recently, we reported an approach for simultaneously collecting X-ray crystal diffraction (XRD) data and X-ray emission spectroscopy (XES) data, with the goal of investigating catalysis in metalloenzymes\textsuperscript{4}. XES directly probes the redox and spin state of the active site metal(s) and ensures that the atomic structure revealed by XRD is in the correct catalytic state. However, optimal sample delivery conditions (crystal size, sample path lengths for spectroscopy, overall sample consumption, etc.) for collecting XRD and XES datasets are not the same. Several continuous-flow sample delivery strategies for XFEL experiments have been established (Fig. 1a)\textsuperscript{7–10}. Liquid jet systems like the gas dynamic virtual nozzle (GDVN)\textsuperscript{7} and the microfluidic electrokinetic sample holder (MESH)\textsuperscript{8} use small crystals to obtain the flow rates reported, as a consequence of the narrow feed capillaries they use, whereas lipidic cubic phase (LCP) or grease matrix injectors\textsuperscript{9,10} use highly viscous media to achieve low flow rates with medium size capillaries. Ejection from an ADE has a high upper bound for the crystal size, while keeping sample consumption at low levels, orders of magnitude lower than Rayleigh jets of the same pathlength. Fixed-target systems have also been successfully used at XFEL facilities\textsuperscript{11–14}, but such systems have not yet been demonstrated for multi-modal data collection.

Acoustic Droplet Ejection (ADE) offers a sweet spot in sample consumption rate and available pathlength to accommodate both XES and XRD simultaneously for biomolecules\textsuperscript{15,16}. To extend the ADE method to probe longer time delays (milliseconds to a few seconds) after the initiation of reactions and/or more complex reaction initiation schemes, we describe here a novel conveyor belt system. The conveyor belt continuously transports droplets containing crystal slurry or protein solution through the reaction region, consisting of an array of laser excitations for photochemical reactions, or gas-activation area for chemically-triggered reactions (Fig. 1b,c and Supplemental Fig. 1). The sample delivery parameters employed here were optimized for crystals of diameter 20–100 μm, solvent pathlengths of 200–300 μm, and sample consumption of 0.8–6 nL/shot with sample replacement rates of up to 120 Hz.

To demonstrate that this acoustic droplet ejection-drop on tape (ADE-DOT) method is versatile for capturing reaction intermediates, we show here two types of reactions triggered and observed \textit{in situ} by Kβ\textsubscript{1,3} or Kα. XES using solution samples; (i) the water oxidation reaction of photosystem II (PS II), which requires multiple timed laser excitations, and (ii) an O\textsubscript{2}-activation reaction at the Mn\textsuperscript{II}/Fe\textsuperscript{II} cluster of the class Ic ribonucleotide reductase R2 (RNR) from \textit{Chlamydia trachomatis} (Ct), which yields the active Mn\textsuperscript{IV}/Fe\textsuperscript{III} cluster via a Mn\textsuperscript{IV}/Fe\textsuperscript{IV} intermediate\textsuperscript{17–19}. The application of DOT to crystallography is further demonstrated by (iii) atomic-resolution XRD data of a bilin containing phytochrome photoreceptor and (iv) simultaneous XRD and XES of the Mn/Fe RNR.
RESULTS
The Drop-on-Tape setup
The conveyor belt is the core component of our DOT sample delivery technique and is used to build a queue of droplets (Fig. 1b, top right) that arrive at the XFEL beam in a desired state of preparation. Synchronized with the XFEL, droplets are deposited from an open 2.5 mm diameter reservoir onto a polyimide belt using an ADE. ADE can eject large crystals without complications such as orifice clogging, which commonly plagues delivery systems using narrow (<100 micron) capillaries. A wide (~250 micron) capillary supply line maintains constant reservoir levels (Fig. 1b top left), allowing ejection conditions to remain stable and maintenance-free for hours. A focused XFEL beam (<25 \( \mu \)m) that passes parallel to the belt surface probes the droplet atop the belt. After X-rays probe the sample, the belt is cleaned before it loops back to the start.

DOT is well-suited to probe a wide range of photochemical reactions in enzymes and inorganic systems. Excitation periods ranging from seconds to hundreds of milliseconds are accessible by positioning an evenly spaced array of laser excitation points along the belt (Fig. 2a). Shorter delays (femtoseconds to hundreds of milliseconds) are accessed by a movable excitation point near the X-ray interaction region (Fig. 1b). The length of the time delay is the ratio of the distance \( D \) between the points of reaction initiation and X-ray interaction to the tangential velocity of the belt \( v \). The precision of the belt velocity required to achieve a desired time delay is determined by the ratio of the droplet diameter to the delay, where longer delay times and smaller droplets require higher precision. Delays of at least 3 seconds were readily achieved with 270 \( \mu \)m droplets (see Fig. 2a). In liquid jet systems, by contrast, excitation pulses for delay times longer than 10 ms need to be coupled into the capillary that supplies the jet, as was implemented for previous PS II studies. The current method, with discrete excitation in a droplet-based system, is more flexible for multi-step excitation experiments and excludes population mixing resulting from laminar flow in a capillary.

Photoactivation of photosystem II
Fig. 2a shows the DOT setup used to study PS II catalytic intermediates in solution sample. The setup was also used for recently reported studies of PS II crystal samples. The oxidation of water by PS II requires multiple laser flashes to complete the four-electron redox chemistry that is catalyzed by the Mn4CaO5 cluster. This complex reaction progresses through four intermediates, called S-states (Fig. 2b). Long delay times between flashes (0.1–1.0 seconds) are required to ensure nearly complete production of each successive S-state due to a slow rate-limiting quinone movement. To accommodate this intricate reaction mechanism, we implemented a feedback control system of the belt speed and deposition delay, which affected the flashing period and droplet phase, respectively. This was provided by a pair of optical gates that reported the arrival time of the droplet stream in the excitation grid (Fig. 2a and Supplementary Fig. 1).

\( K_{\beta 1,3} \) emission (metal 3p to 1s transition) is sensitive to oxidation and spin state and serves as an indicator of the S-state advancement in PS II. Fig. 2c shows the Mn \( K_{\beta 1,3} \)
XES difference spectra of PS II in solution collected with DOT. 270 μm diameter droplets (~5 nL) were injected onto the conveyor belt at room temperature in a helium atmosphere (Fig. 1b). The emission signal was collected with a 16 crystal wavelength-dispersive von Hamos spectrometer\textsuperscript{29,30}, which collects the full Kβ\textsubscript{1,3} energy range\textsuperscript{31} without scanning. These data are in good agreement with those collected at a synchrotron source using the freeze quench method for preparing PS II solution in different S-states\textsuperscript{26} (Fig. 2c). In the transition from the dark S\textsubscript{1} state (0F, F: flash) to the most oxidized 2F (S\textsubscript{3}-rich) state, a negative difference on the high-energy side of the peak is seen, indicating oxidation. On the other hand, a clear positive difference relative to 2F at the high energy side is seen upon illumination to 3F (S\textsubscript{0}-rich), indicating transition of the major fraction of PS II from S\textsubscript{3} to the most reduced S\textsubscript{0} state. The purity of the states prepared, witnessed by the difference amplitude, on the belt is comparable with the best case S-state population acquired by freeze quench methods, while operating at room temperature and in high-throughput mode.

**Chemical activation of ribonucleotide reductase by gas exposure**

DOT can also be used to study chemically activated reactions. To demonstrate this, we examined O\textsubscript{2} activation of Ct RNR by replacing the aforementioned laser excitation grid with an incubation chamber that exposes the drops to oxygen gas as they travel to the X-ray beam (Fig 3a,b). We built a chamber that exposes a 60 mm portion of the conveyor belt to a 4.7 L/min O\textsubscript{2} flow. Leakage of O\textsubscript{2} to the surrounding Helium environment was mitigated by using a flow-through exhaust design, guard baffles, and differential pumping. Tape speed through the chamber determines the gas exposure time, and we explored a range of exposures from 0.5 to 8 seconds.

Fig. 3c shows the Fe Kα\textsubscript{1} and Kα\textsubscript{2} XES difference spectra of Fe/Mn RNR solution in which we monitored the advancement of the Fe site from its reduced form (Fe\textsuperscript{II}) to the oxidized form by varying O\textsubscript{2} exposure time. Both Kα\textsubscript{1} and Kα\textsubscript{2} exhibit spectral changes as a function of O\textsubscript{2} exposure time. The full width half max (FWHM) of the Kα\textsubscript{1} line, which serves as a proxy for oxidation state\textsuperscript{32,33}, shows the reaction progressing towards peak oxidation at 2 s and then reduction again at 8 s exposure, which roughly matches expected changes for the formation of Mn\textsuperscript{IV}/Fe\textsuperscript{IV} intermediate and subsequent decay to Mn\textsuperscript{IV}/Fe\textsuperscript{III} (Fig. 3b)\textsuperscript{17–19}. The difference spectra with respect to the 8 s O\textsubscript{2} exposure show that the Fe Kα spectra contain complex lineshape changes, with both shifts and width changes throughout the reaction cycle. We note that no significant signs of droplet dehydration were detected for these studies of RNR solution. In samples containing a significant amount of Polyethylene Glycol (PEG), like many protein crystal buffers, PEG diffraction appears if the belt droplets are exposed to Helium for too long. The lower bound belt speed is thus limited by sample dehydration and needs to be evaluated on a per-sample basis.

For X-ray spectroscopy studies as described above, the ability to control pathlengths (e.g. 1/e is 520 μm at 6500 eV for water) is important for efficient data collection, by balancing the sample consumption rate and the data acquisition time. The acoustic pulse in the ADE system can be tuned to provide a range of droplet sizes (50–400 μm diameter), with larger droplets simply requiring a lower central frequency and/or higher driving force. The ability to access larger droplets is particularly beneficial for collecting signals from weaker...
transitions like $K\beta_{2,5}/K\beta''$ that can uniquely probe changes in the metal-ligand environment while its intensity is $\sim 60$ times lower than that of $K\beta_{1,3}$

Femtosecond crystallography using the DOT setup

ADE’s ability to adjust droplet size also allows a wide range of crystal sizes to be analyzed, while the discrete nature of delivery keeps sample consumption at moderate levels. Crystals $>20$ μm in size are of general interest for Serial Femtosecond Crystallography (SFX)$^{2-4}$, as their larger crystal volume, in comparison to $<5$ μm crystals used in many liquid jet systems, can translate to stronger diffraction signals and potentially higher resolution. Also, diffraction quality may be more easily optimized at synchrotron beamlines prior to XFEL experiments for crystals in this size range. For the XRD results presented in this study, we used 5.5 nL droplets ($\sim 270–290$ μm hemisphere) with crystal sizes ranging from 20 to 100 μm. With this size a 100% droplet hit rate was achieved with favorable signal collection rates and reasonable sample consumption (3.3 μL/min at 10 Hz). XRD solvent background was present, but manageable with a beam focus of $\sim 5$ μm. Thus, the solvent pathlength that can be achieved with the DOT (100 to 300 μm) is practical, vide infra, for the study of crystals surveyed here. However, practical considerations of what droplet size to select for which crystal size must be made for balancing the diffractivity of crystals and the solvent background; there is a pragmatic trade-off between factors favoring smaller droplets (solvent background, sample consumption) and those favoring larger droplets (instrument precision and distance of the X-ray beam from the belt). A consideration for XRD, which is unique to this setup, is polymer scatter and absorption from the polyimide conveyor belt. Polymer scatter was typically lower in amplitude than solvent scatter, and absorption was successfully treated in post-analysis (see Supplementary Fig. 2).

To explore the applicability of DOT for different protein crystals, we examined four samples: two different phytochrome constructs (50 μm plates or 100 μm needles), RNR (20–30 μm, bipyramidal), and PS II (20–50 μm, bipyramidal) (Supplementary Fig. 3, Supplementary Table 1). To collect diffraction from crystals of various sizes without saturating the detector, we used a high dynamic range Rayonix MX 170 HS detector, operated at its maximum data rate of 10 Hz in the 2×2 binning mode. For all crystallography data presented here, an incident X-ray energy of 9.5–9.8 keV was used. High-resolution room temperature structures were obtained in all cases (Supplementary Table 2)$^{21}$. Data collection time varied depending on the crystal number density, but complete data sets could be collected in $\sim 40–80$ min with between 10 and 50% of all shots giving indexed diffraction patterns (Supplementary Table 1 and Supplementary Fig. 4).

Room temperature structures of phytochromes

XRD data were collected for the chromophore-binding region of Deinococcus radiodurans bacterial phytochrome (BphP) containing only the PAS (Per/Arnt/Sim) and GAF (cGMP phosphodiesterase/adenyl cyclase/FhlA) domains (residues 1–321) or the entire photosensory module (PSM, residues 1–501) in their dark-adapted (Pr) states (Supplementary Table 2). These data yielded novel room temperature structures of DrBphP dimers at 1.65 and 3.3 Å resolution, respectively (Fig. 4a,b). The chromophore binding pocket of the PAS-GAF bili-protein was highly congruent with the best resolved structure of

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the DrBphP-PSM to date\textsuperscript{34}, but diverged from an alternate binding pocket orientation seen for recently reported monomeric crystal assemblies of DrBphP\textsuperscript{35,36}, and the photochemically compromised D207A mutant (Supplementary Fig. 5)\textsuperscript{34}. Whether both bilin configurations represent authentic structural microstates of the Pr photostate in BphP dimers is not yet known. Additionally, we detected a significant shift of the PHY domain within the crystal lattice of the PSM dimer as compared to models generated with diffraction data collected at 100 K, suggesting that cryopreservation and/or cryogenic temperatures perturb the structure and/or dynamics of the GAF and PHY domains (Fig. 4b).

Phytochromes undergo a cascade of conformational changes upon photoexcitation of the bilin chromophore, with dynamics that span from picoseconds to milliseconds\textsuperscript{37–39}. However, the electronic state of the bilin chromophore is exceedingly sensitive to radiation-induced changes from synchrotron X-ray sources\textsuperscript{40}, culminating in rupture of the thioether linkage between the Cys24 and the bilin at high dose. These complications were eliminated in our XFEL data (Fig. 4c,d), with both structures clearly showing continuous electron density along the thioether-bilin linkage, visible already in the initial F\textsubscript{o}-F\textsubscript{c} omit maps (Supplementary Fig. 6). The ability to collect RT structures of phytochrome that are free of radiation induced artifacts thus opens the possibility of elucidating bona fide structures of phytochrome photoconversion intermediates.

**Combined X-ray emission and crystallography of ribonucleotide reductase**

We also collected a high-resolution room temperature structure of *S. erythraea* Mn/Fe RNR (Fig. 4e). Ribonucleotide reductases are the only enzymes that catalyze the reduction of ribonucleoside-5′-diphosphates or triphosphates (NDP or NTP) to the 2′-deoxynucleotides (dNDP or dNTP). Class I RNRs generate an essential catalytic radical via a dinuclear metal site, involving redox state changes at the metal site\textsuperscript{41}. XES can be collected simultaneously with XRD for many metalloproteins serving as a sensitive in-situ verification of state preparation. To demonstrate this, we collected a Mn K\textsubscript{β\textsubscript{1,3}} and Fe K\textsubscript{α} spectrum of Mn/Fe RNR, simultaneous with XRD, from crystal slurries (Fig. 4e). Mn XES from RNR protein crystals was overlaid with a solution spectrum, using the same dispersive X-ray emission spectrometer system. The results confirmed that the XRD data collected, and thus the electron density obtained, represent the metal site in its as-prepared oxidized form and that the Mn ion was not photo-reduced to the (II) state before probing the scattering data. Model building and refinement is currently underway and will be reported elsewhere.

**DISCUSSION**

Through the XRD and XES results reported here, we demonstrate that the ADE-DOT sample delivery method provides a foundation for multimodal data collection in both solution and crystalline samples. The conveyor belt in DOT is flexible and extensible, permitting either photochemical or chemical activation of samples in situ, and thus enables it to study a wide variety of transient reaction intermediates. For photochemical reactions, in particular, the ability to precisely illuminate a discrete volume eliminates potential flow-induced state-mixing prior to probing the sample. Here we carried out the gas activation
studies of a solution sample, but the method can also be applied to crystalline samples; the influence of substrate transport speed on the observed kinetics would depend on crystal size.

Another extension of the current work that is envisioned is liquid mixing prior to probing the droplet. This would enable a much broader class of enzyme-substrate studies. For reactions with relatively slow kinetics (seconds timescale) where liquids can be mixed right before droplet ejection, the method as it is shown can already be applied. Faster reactions can be probed by incorporating a rapid mixing scheme directly either in droplet creation or shortly after deposition on to the belt.

**ONLINE METHODS**

**Samples**

(a) **PS II**—PS II dimer was extracted and purified from *T. elongatus* as reported previously. PS II crystals ranging in size from 10–60 μm were then prepared using an established seeding protocol. The crystals were transferred into a final buffer consisting of 0.1 M MES pH 6.5, 0.1 M ammonium chloride, 10% (w/v) ethylene glycol and 35% (w/v) polyethylene glycol (PEG) 5000, with ~0.5 mM chlorophyll concentration, before loading into the sample delivery syringe (Hamilton gastight syringe, 500 μl).

Dimeric PS II solution samples were extracted and purified as described previously. The final chlorophyll concentration was adjusted to ~7.0 mM (70 mg/mL protein) in a final buffer comprising 0.1 M PIPES pH 7.0, 5 mM calcium chloride, 30% (v/v) ethylene glycol and 0.02% (w/v) detergent β-dodecyl maltoside (βDM).

(b) **Phytochrome**—*D. radiodurans* BphP apoproteins bearing an N-terminal T7 tag (MASMTGQQMGRGS) and a C-terminal hexahistidine tag were expressed in *Escherichia coli* BL21(DE3) cells as the PSM (residues 1–501) or the PAS-GAF region (residues 1–321 and including the Y307S mutation using the pET21b plasmid (Novagen, Madison, WI). Cultures were grown at 16 °C in terrific broth and harvested by centrifugation 16 h after inducing apoprotein expression with isopropyl-D-1-thiogalactopyranoside. The cell pellets were frozen in liquid nitrogen and stored at −80 °C.

Frozen pellets were resuspended in lysis buffer (10% (w/v) glycerol, 20 mM HEPES-NaOH (pH 7.8), 500 mM NaCl, 0.05% Tween 20, 30 mM imidazole, 1 mM phenylmethanesulfonyl fluoride, and 1 mM 2-mercaptoethanol), sonicated, and clarified by centrifugation. *DsbPhP* apoproteins were enriched from the lysates by nickel-nitrilotriacetic acid (Qiagen, Valencia, CA) chromatography, using the lysis buffer for washing and lysis buffer with 300 mM imidazole for elution. The *DsbPhP*-containing eluates were incubated overnight with a molar excess of biliverdin (BV) (Frontier Scientific, Logan, UT), and the free BV was removed by filtration through a G-25 Sephadex column (GE Healthcare) equilibrated in the ion-exchange buffer (10% (w/v) glycerol, 20 mM HEPES-NaOH (pH 7.8), 20 mM NaCl, and 10 mM 2-mercaptoethanol). The chromoproteins were then purified by ion-exchange chromatography with a Q-Sepharose HP column (GE Healthcare) and employing a 20–1000 mM linear NaCl gradient for elution. The PAS-GAF chromoprotein was exchanged into 30 mM Trizma base-HCl (pH 8.0), and the PSM was exchanged into 10 mM HEPES-NaOH.
(pH 7.8) using a 50-mL G-25 Sephadex column (GE Healthcare), frozen as 30–50 μL droplets in liquid nitrogen, and stored at −80 °C.

Crystallization of DrBphP chromoproteins was conducted in batch mode. PAS-GAF chromoproteins at 40 mg/mL were mixed at a 1:1 volumetric ratio with 15–16% (w/v) PEG 3350, 19% (v/v) isopropanol, 5% (w/v) glycerol, and 100 mM tetrasodium citrate-citric acid (pH 5.6). 2 mL batches were then seeded with 10 μL of a finely crushed PAS-GAF crystal slurry. PSM crystals were produced in the same manner except that the crystallization solution contained 22–28% (w/v) PEG 3350, 2.5% (v/v) Tacsimate (pH 6.0), 2.5% (v/v) Tacsimate (pH 7.0), 2% (v/v) ethylene glycol, and 100 mM PIPES-NaOH (pH 6.5).

Crystalline samples were mobilized by scraping with a pipet tip and suspended by aspiration. The crystal slurry was then loaded into a Hamilton syringe for sample delivery.

**(c) RNR**—A crystallizable construct of the RNR R2 from *S. erythraea* was purified and prepared as described in ref.46. Crystals suitable for this XFEL experiment were obtained at 20 °C under aerobic conditions using the batch crystallization method, in a 1:1 protein solution to crystallization buffer ratio. The crystallization buffer (10% (w/v) PEG3350, 1% (v/v) Tacsimate pH 4.0) was slowly released at the bottom of a 20 mg/mL protein solution containing tube such that phase separation occurred between the upper protein solution and the lower crystallization buffer. The tube was incubated upright without mixing for crystal growth. A pellet of up to 30 μm sized square bipyramidal shaped crystals appeared overnight. This crystal pellet was then resuspended in the crystallization buffer supplemented with 10% glycerol and loaded into the sample delivery syringe used for data collection.

For solution XES measurements, a histidine-tagged construct of *CyrR*2 was expressed and purified as previously described19,47; the resulting apo-protein was anaerobically reconstituted with 0.75 equivalents of Fe(II) and Mn(II) per monomer.

Representative images of crystals of all four samples used for the data collection are shown in Supplemental Fig. 3.

**Cleaning and drying the belt for continuous use**—During operation of the DOT, residual sample was removed continuously from the PI belt. After passing the X-ray probe point, the belt was immersed in a water tank where 4 high pressure water jets were used to remove sample debris. To remove waste and maintain a constant level in the tank, water was drained using a peristaltic pump (Masterflex L/S digital pump, HV-77921-75, Cole-Parmer Instrument Company, LLC, USA). Water jets were produced through 250 μm inner diameter PEEK tubing (Part # 1531, IDEX Health & Science LLC, USA), and were generated by an HPLC pump (Prep 100 pump, P40PFT01, Scientific Systems, Inc., USA) at 100 mL/min, which produced an effective pressure of 800 psi. The belt was then dried with He that was forced through small diameter nozzles (Atto super air nozzle, model 1108SS, EXAIR Corporation, USA). To minimize He consumption, the gas was continuously circulated using a compressor comprising a diaphragm pump (KNF Neuberger, Inc., type N145.1.2AN.18) and a small reservoir. This mechanism was necessary to have enough backing pressure to purge water from the belt surface. Effective drying is also important for the operation of the
rollers to maintain a constant tape speed and was achieved by blowing He at a flow rate of 100 standard cubic feet (SCF) per hour with a backing pressure of 20 psi.

**Sample delivery system**—The tape drive instrumentation included a 15 MHz spot-focused piezoelectric immersion transducer (Part # V319, Olympus NDT) with a focal length of 1.0 in. A waveform generator (model 33612A, Keysight Technologies) triggered by the XFEL master clock was used to produce short bursts of RF energy which were amplified by a 500-W RF amplifier (Model 500A250C, Amplifier Research, Souderton, PA, USA) and used to drive the transducer for droplet ejection. The transducer was immersed in degassed water within a 3D-printed plastic sleeve with a sample reservoir at the end. A motorized stage was used to adjust the up and down position of the transducer such that the acoustic energy was focused at the liquid-gas interface. To change the position of the droplet deposition parallel to the beam, a second motorized stage was used to move the transducer and sleeve together. The sample reservoir was a 3D printed small plastic well (6 μL volume) with a thin polypropylene window to minimize the attenuation of the acoustic ejection pulse. Sample was continuously fed to the reservoir, using a syringe pump, through a 250 μm ID glass capillary. To avoid crystal settling, the syringe pump was mounted onto a shaker that rotated the syringe 180°. The 3D files/schematics of custom-built parts and more details of the sample delivery system are included in the Supplementary Protocol.

**Laser excitation system**—A Coherent Evolution frequency doubled Nd:YLF laser provided 100 ns pulses of 527 nm light which were fiber coupled into the multi-flash excitation system depicted in Supplemental Fig. 1. An identical and independent Evolution provided the free-space excitation pulse near the X-ray interaction region for studies of PS II. For all excitations in PS II studies a fluence of 0.12 J/cm² was used, with a focal diameter of 400 μm. The penetration depth of PS II at 527 nm is about 40 μm in PS II crystals and 400 μm in PS II solution at the concentrations we employ. Thus, we are close to the limit in crystal size that can be excited from a single side. For solution samples, the situation is better (as the concentration is about 10× lower).

**Gas activation system**—The oxygen incubation chamber (Fig. 3a) used for solution studies of RNR was 3D printed using ABS-like resin and has 5 internal chambers. The central compartment, where O₂ gas is pumped, is a 60 mm long, 14 mm diameter cylinder. The flanking chambers, two on each side, are each 15 mm long and 14 mm in diameter. To mount the conveyor belt on the drive, the top half of the chamber was made to be removable and after mounting the belt the chamber top was sealed to the bottom half with the aid of vacuum grease. Entrance holes into the sides of the O₂ activation chamber and between internal chambers measured 0.9 mm tall by 3 mm wide. The vacuum pressure on the chambers immediately flanking the central O₂ chamber was set to −0.5 psig through a variable vacuum regulator. More details of the gas activation system are included in the Supplementary Protocol.

**Data collection**—The DOT setup was employed at two end stations in Linac Coherent Light Source (LCLS): X-ray Pump Probe (XPP) and Macromolecular Femtosecond Crystallography (MFX). Photoactivated studies began at XPP and continued at MFX over
several beamtimes. Chemically activated studies were performed at MFX. The X-ray beam with a pulse width of ~50 fs was focused at either end station to 5×5 μm² (FWHM) using Be lenses. A pulse energy of 2.5 mJ at 9.5 keV, corresponding to ~1.6 × 10¹² photons/pulse, was used at a repetition rate of 10 Hz for crystal samples. For XES, solution samples of PSII were excited at 7.5 keV and at 30 Hz rep rate. Fe XES of RNR solution was collected at 60 Hz using 9.5 keV excitation and 2.5 mJ pulse energy. To minimize the belt diffraction background, the tails of the beam were obscured with a pinhole lead collimator. XRD data were collected in the forward direction using a Rayonix MX 170 HS detector at a distance of 58 to 108 mm from the interaction point allowing for detection of a maximum inscribed resolution of 1.4 or 2.0 Å, respectively. To calibrate the detector distance, the XRD pattern of silver behenate, enclosed in a quartz capillary (Hampton Research, 10 μm wall thickness) was recorded with an attenuated beam (≤1%). Mn XES spectra were recorded perpendicular to the beam on a Cornell-SLAC pixel-array detector (CSPAD-140k) using a multi-crystal wavelength-dispersive hard X-ray spectrometer based on the von Hamos geometry. Studies on Fe Kα XES employed the newly available ePIX-100 detector in the same geometry. Mn XES was analyzed by an array of 16 Si (440) crystal analyzers, mounted at a distance of 500 mm from the interaction point, with the center of the array at an angle of 81° with respect to the interaction point. The Bragg angle range covered by these analyzer crystals was 85.8–83.4° which corresponds to a Mn Kβ₁,₃ energy range from 6474.6–6499.4 eV. To calibrate the emission energy, the XES spectrum of MnCl₂ solution (500 mM), in a quartz capillary, was collected using an unfocused beam. The emission energy was then calibrated to the published value of 6493.3 eV of MnCl₂ spectrum peak position. Fe Kα XES was analyzed by a single Ge (440) crystal placed 250 mm from the interaction point, with the center of the crystal at 75.41° with respect to the interaction point and resolved both Kα lines from 6370 to 6420 eV. Calibration of the Fe spectrometer was similar to the method for Mn, using as the reference Fe(III)(NO₃)₃ instead.

**Processing of Data of Raw XRD images**—Analysis of the diffraction data was performed using cctbx.xfel. Diffraction images were indexed using LABELIT with a target cell to guide the choice of basis vectors. For PS II samples, two populations of unit cells in the P2₁2₁2₁ space group were identified (isoform A: a=117.5 Å, b=223.6 Å, c=329.5 Å; isoform B: a=117.7 Å, b=223.2 Å, c=310.5 Å). PS II diffraction images were re-indexed twice, imposing either isoform A or isoform B unit cell parameters as constraints during unit cell refinement, to generate two disjoint sub-datasets for each illuminated state. For other samples, the single predominant unit cell identified in the first indexing step was imposed as a basis set target and a unit cell constraint in a second round of indexing. For all samples, resolution limits were determined independently for each image in order to ensure that the weak signal at high resolution in the highest-resolution images was not diluted by noise contributions from lower-resolution images. To determine resolution limits on each image, reflections were sorted into bins of equal size by resolution, and all bins beyond and including the lowest-resolution bin in which the average signal to uncertainty ratio <I/σ(I)> was below 0.1 were discarded. This approach permits the inclusion of a small number of negative intensities while in practice eliminating most nonphysical negative intensity reflections. Signal was integrated to these resolution limits and the resulting data were merged using cxi.merge. The resolution cut-off for the final merged data was determined.
based on a combination of several criteria, including where the data falls below 10 fold multiplicity, where the CC_{1/2} becomes ‘unstable’, meaning it begins to oscillate at higher resolution instead of uniformly decreasing, and where the values of I/σ(I) do not uniformly decrease any more.

Roughly half of the forward scattering signal passes through the PI belt, slightly lowering the Bragg spot intensities in this region. We developed a ray-tracing model to predict the absorption by the belt at a given point on the detector and used this to correct Bragg intensities during integration. Our model uses three fit parameters: distance of the crystal from the edge of the belt f, crystal height h, and the angle Θ between the vertical axis of the detector and the conveyor belt. Belt thickness and material absorption cross section are treated as known constants. These various parameters are illustrated in Supplemental Fig. 2c with the overall problem depicted in Supplemental Fig. 2b. Parameters f, h, and Θ are obtained indirectly through geometric relations to two boundary lines that are observed on the detector. The boundary lines (corresponding to maximum and minimum belt absorption) are highlighted in Supplemental Fig. 2a and c in red and blue, respectively. An estimate for the uncertainty of the Bragg spot intensity is needed in subsequent processing steps to weight observations. The uncertainty of the absorption correction to the spot intensity is estimated by the change in spot intensity upon perturbing the fit parameters. We typically find that the perturbations in f produce the largest effect and that perturbations on the order of a droplet diameter dominate other uncertainty estimates. Consequently, Bragg spots, which are shadowed by the belt, are down-weighted relative to those that are un-shadowed during merging. Separate resolution cut-offs are used for Bragg spots that are affected and unaffected by Kapton belt absorption.

**XES data reduction and analysis**—Supplemental Fig. 6 panel a shows the raw data of the averaged XES spectrum collected with the pixel array detector. To subtract the background signal, a fifth order two-dimensional polynomial background was fit to the image using information outside of the Region of Interest (ROI), shown in bounded by red lines. The fit of the background was extrapolated into the ROI (panel b, with the intensity multiplied by 4) to estimate its intensity there. The estimated background is shown as the red line in panel Supplemental Fig. 6 c. The final XES spectrum is obtained by integrating a tight ROI around the background subtracted signal (depicted in Supplemental Fig. 6 d). The resulting spectrum is shown in panel e. Difference spectra are computed normalizing the component spectra by their respective sums and then subtracting the components.

**Synchrotron XES data collection**

Room temperature difference Mn Kβ_{1,3} XES shown in Figure 2 are shown in reference to cryogenic difference spectra of PS II S-states collected at BL 6–2 in SSRL (Stanford Synchrotron Radiation Lightsource). For these synchrotron measurements, the incidence energy was set to 10.4 keV with the beamline monochromator, which uses two cryogenically cooled Si crystals in (111) reflection. The X-ray beam was focused to 0.45 (V) × 0.45 (H) mm (fwhm) by means of vertical and horizontal focusing mirrors. The X-ray flux at 10.4 keV was ~ 1 × 10^{13} photons/s/mm^2. Samples were prepared in 0, 2, and 3 flash states by a freeze quench method, wherein a sample holder containing a 300 µm thick layer of PS II
solution from T. elongatus (8.5 mM chlorophyll) is flashed from two sides with a Spectra-Physics Quanta Ray Pro 230-10 Nd:YAG laser (0.3 J/cm2, 9 ns pulse duration) with a 1 s flashing interval and subsequently plunged (within 1 second) into liquid nitrogen. During the measurement, samples were kept at 10 K in a continuous flow liquid helium cryostat (Oxford Instruments CF1208) under helium exchange gas atmosphere. Emission spectra were recorded by means of a high-resolution crystal-array spectrometer, using the 440 reflection of 7 spherically bent Si(110) crystals (100 mm diameter, 1 m radius of curvature), aligned on intersecting Rowland circles. An energy-resolving Si drift detector (Vortex) was positioned at the focus of the 7 diffracting elements. A helium-filled polyethylene bag was placed between the cryostat and the spectrometer to minimize signal attenuation due to air absorption. Each energy point in the spectra was collected at a fresh sample spot. The maximum exposure time at each spot was 2.5 s and the signal was read out in bins of 50 ms duration. To determine an appropriate exposure time to avoid radiation-induced effects, the signal intensity was measured as a function of exposure time at a single emission energy for each S-state. No significant changes were observed within the first 1.5 s of exposure, and so 20 time bins (equivalent to 1 s) were averaged for the final spectra. The signal intensity from each sample spot was normalized by the emission signal intensity recorded at 6491.5 eV within 7 s from the same sample spot, after going through all the fresh spots.

**Phytochrome Structure determination, model building and refinement**—Initial phases for the PAS-GAF and PSM datasets were calculated by PHASER\textsuperscript{55} (using the 1.45-Å DtBphP PAS-GAF structure (PDB ID code 2O9C) generated by Wagner \textit{et al}\textsuperscript{45} as the search model for PAS and GAF domains, and residues 325–501 of PDB ID code 4Q0J as the PHY domain search model for the PSM\textsuperscript{34}. To minimize model bias at the chromophore, the biliverdin and side chains of the following residues were omitted during molecular replacement and initial structural refinements: Cys24, Ile29, Arg70, Met174, Tyr176, Glu193, His196, Phe198, Phe203, Asp207, Ile208, Pro209, Tyr216, Arg222, Arg254, Ser257, Met259, His260, Tyr263, Leu264, Met267, Ser272, Ser274, Leu286, His290, and Arg310. Manual model building was conducted with COOT\textsuperscript{56}, and structures were refined and simulated-annealing omit maps were calculated with PHENIX\textsuperscript{57}. The initial refinement for each structural model included simulated annealing to reduce model bias. Model validation was conducted with MOLPROBITY\textsuperscript{58}, and superpositions and figure preparation were conducted with the PyMOL Molecular Graphics System.

**RNR Structure determination, model building and refinement**—The RNR structure was solved using PHASER\textsuperscript{55} by molecular replacement using the atomic coordinates of the R2 subunit of ribonucleotide reductase from \textit{Chlamydia trachomatis} (PDB ID code: 1SYY\textsuperscript{59}) as starting model. A well-contrasted solution was obtained with 1 molecule per asymmetric unit (space group \textit{P}4\textsubscript{1}2\textsubscript{1}2). A preliminary crystallographic refinement was performed using PHENIX\textsuperscript{57} and the three-dimensional model was examined and modified using the program COOT\textsuperscript{56}. Supplemental Table 2 lists the crystallographic statistics in which the test set represents 5% of the reflections. The current Ramachandran plot indicates that 93.0% and 0.3% of the residues are in the favored and disallowed regions, respectively. Figure 4e was prepared using the PyMOL Molecular Graphics System.
Data availability

**Accession codes**—Coordinates and structure factors of the phytochrome PAS-GAF and PSM structures have been deposited in the RCSB Protein Data Bank under accession codes 5MG0 (doi:10.2210/pdb5mg0/pdb) and 5MG1 (doi:10.2210/pdb5mg1/pdb), respectively.

**Code Availability**—Source code for the cctbx.xfel software package, used for diffraction data processing is freely available and is distributed via the Phenix software package (http://phenix-online.org/). Further information can be obtained via the cctbx.xfel homepage (http://cci.lbl.gov/xfel/).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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References


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Figure 1. Experimental Setup

a. A comparison of sample consumption rate (assuming 120 Hz operation) and pathlength accessible with the replenishing methods used at XFELs. b. The experimental setup of DOT. The conveyor belt delivers droplets of sample at a high rate (up to 120 Hz). A large (283 liter) gas-tight chamber houses the entire instrument and is maintained at 95–97% Helium via a 30 cubic feet per hour purge flow. Droplets are deposited onto a polyimide belt using an ADE, and ejected from an open 2.5 mm diameter reservoir that is continuously resupplied via a capillary feed line attached to a syringe pump (top left b). In the interaction region, the belt was run at a small angle with respect to the z-axis (vertical) in the xz-plane (zoomed in view of interaction point). Positioning the droplet in the X-ray focus is accomplished by moving the entire system in the horizontal plane, while the droplet z-position (vertical) on
the tape at the intersection is adjusted by changing the deposition delay. The belt is cleaned and dried in situ, enabling continuous use for days. The reaction initiation point for longer time delays is shown in the top right of panel b. **Bottom right b.** The XFEL beam passes parallel to the belt surface, striking the droplet atop the belt. c. The data collection geometry for XRD and XES. An inflatable (and X-ray transparent) plastic film door with a 160° aperture to the X-ray interaction region allows both XES and XRD to be collected simultaneously.
Figure 2. Photo-initiated XES of PS II

a. The photoexcitation setup employed for PS II, comprised of a precision-machined grid of fixed excitation positions with 60 mm spacing, is shown. Optical gates, which measure droplet arrival times on the grid, are realized by two low-power (<0.5 mW) continuous-wave near infrared (NIR, 850 nm) point sources, delivered via optical fiber. NIR light scatters from the droplets as they pass over the gates and is collected onto high-speed Si-photodiodes for readout.

b. A schematic of the mechanism depicting S-state advancement using periodic laser flashes. The interval between laser flashes for both DOT and freeze-quench methods was 1.0 s. Flash states (denoted as 0F, 1F, 2F, or 3F) are highly enriched in the pure reaction intermediates S$_1$, S$_2$, S$_3$, and S$_0$, respectively, but are not completely pure due to small back reaction rates and photon misses (see ref. 23).

c. A comparison of Mn K$\beta_{1,3}$ XES difference spectra of PS II solution collected with the DOT instrument at room temperature using an XFEL (red) and the same state differences collected using a freeze-quench method at 8 K with a synchrotron source (black). The XFEL spectra contain about 120,000 shots per difference spectrum. For details of the collection conditions and difference analysis see Online Methods and Supplementary Fig. 7.
Figure 3. XES of gas-activated RNR
a. A schematic of the differentially pumped oxygen gas activation setup is shown. b. The known reaction scheme of $CrRNR^{17-19}$ is shown. Oxygen is activated at the Mn$^{II}$/Fe$^{II}$ cluster to produce a high valent Mn$^{IV}$Fe$^{IV}$ intermediate in a biomolecular reaction ($k_{\text{form}} = 13\pm3 \text{ mM}^{-1}\text{sec}^{-1}$ at 5 °C from the literature$^{18}$), which then decays to the active Mn$^{IV}$Fe$^{III}$ state (first-order $k_{\text{decay}} = 0.02 \text{ sec}^{-1}$ at 5 °C also from literature$^{18}$). c. RNR solution was monitored at 25 °C with Fe K$\alpha$ emission for various $O_2$ exposure times. The inset shows the K$\alpha_1$ FWHM as a function of exposure time relative to an 8 s exposure. Error bars are computed via bootstrap residual sampling (1000 samples per data point). Difference spectra with respect to an 8 s exposure (smoothed by wavelet denoising) are shown in the main plot.
Figure 4. XRD of various enzymes

a. The PAS-GAF (blue and green, respectively) and the PSM (white) constructs from *D. radiodurans* BphP in their dark-adapted Pr states are shown superimposed. b. Superposition of atomic models of the PSM in the room-temperature Pr state (colored) with that derived from diffraction data collected at 100 K (white, PDB ID 4Q0J). The β-sheets of the GAF domains were superposed, allowing the respective positions of the PAS and PHY domains of the two models to be identified. The largest differences between the models were found at the PHY domain. For the room temperature model the domains were colored blue, green, and orange for the PAS, GAF, and PHY domains, respectively. Several important features are labeled: HP, hairpin; NTE, amino-terminal extension; BV, biliverdin. BV from the room temperature structure is shown for orientation. c and d. Composite simulated-annealing omit
maps ($2F_o-F_c$) contoured at 1 $\sigma$ were superimposed with the corresponding PAS-GAF (e) or PSM (d) model of DrBphP$^{22}$. For clarity only electron density around Cys24 (gray), biliverdin (cyan) and for (c) the pyrrole water is shown. e. Metal site electron density ($2F_o-F_c$) at the heterodinuclear Mn/Fe site in aerobic class Ic RNR, metal ions and protein ligands are indicated (contoured to 1.3 $\sigma$) in blue. Residual positive difference electron density map ($F_o-F_c$) representing non-protein ligands is indicated in green (contoured to 3.5 $\sigma$). The manganese and iron atoms are depicted as purple and orange spheres, respectively. $K_{\beta_{1,3}}$ XES of oxidized RNR in crystal and in solution collected at room temperature with an XFEL is shown to the left. A Mn(II)Cl$_2$ calibration standard is also shown to illustrate the absolute oxidation state of the solution and crystal spectra. Fe $K\alpha$ XES Data collected from oxidized RNR crystals is shown at the right.