

Compact laser-driven plasma X-ray source for time-resolved diffraction, spectroscopy and imaging experiments at ELI Beamlines

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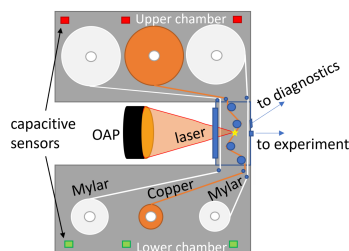
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In this work, experimentally measured characteristics of a kilohertz laser-driven Cu plasma X-ray source that was recently commissioned at the ELI Beamlines facility are reported. The source can be driven either by an in-house developed high-contrast sub-20 fs near-infrared terawatt laser based on optical parametric chirped-pulse amplification technology or by a more conventional Ti:sapphire laser delivering 12 mJ and 45 fs pulses. The X-ray source parameters obtained with the two driving lasers are compared. A measured photon flux of the order up to 10^{12} $K\alpha$ photons s^{-1} $(4\pi)^{-1}$ is reported. Furthermore, experimental platforms for ultrafast X-ray diffraction and X-ray absorption and emission spectroscopy based on the reported source are described.

1. Introduction

Laser-driven plasma X-ray sources (PXSs) hold a niche in studying ultrafast phenomena thanks to their ability to provide pulses as short as hundreds of femtoseconds. Moreover, their compact size makes them favorable for small-scale laboratories, which enhances their availability to a broad scientific community. X-ray pulses of such short duration are otherwise achievable only by the synchrotron slicing technique (Prigent *et al.*, 2013), LINAC sources (Enquist *et al.*, 2018) or at XFEL facilities (Altarelli, 2011; Schietinger, 2018), and allow time-resolved studies in material science (Mattern *et al.*, 2021), molecular dynamics (Freyer *et al.*, 2013) and structural biology (Khakurel *et al.*, 2024). The two main application branches are ultrafast time-resolved X-ray diffraction, using the characteristic line emission of the target material, and X-ray absorption and emission spectroscopy (XAES) that benefits mainly from the continuous *Bremsstrahlung* emission of hot electrons' collisions in the target.

Laser-driven plasma sources for X-ray generation have been demonstrated across solid, liquid and gas targets with a diverse range of flux and source sizes. Sources with solid targets, such as Cu, Mo and Ti (Holtz *et al.*, 2017; Afshari *et al.*, 2020; Li *et al.*, 2017; Azamoum *et al.*, 2018; Rousse, Audebert *et al.*, 1994), reach photon fluxes up to 10^{12} photons s^{-1} $(4\pi)^{-1}$ with source sizes varying from a few micrometres to several tens of micrometres. Solid target sources, used together with high-power lasers, presently produce the highest brightness among PXSs but are prone to long-term flux decline due to target deformation and damage, as well as higher operational



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costs due to the consumable target. Liquid sources, such as Ga and Hg jets (Reich *et al.*, 2007; Zhavoronkov *et al.*, 2004; Ivanov *et al.*, 2011), show photon fluxes of around 10^9 photons s^{-1} $(4\pi)^{-1}$ and are mostly used with lower peak-power lasers. These sources hold the potential of long-term low-maintenance operation by recycling the target metal. However, technical challenges, such as maintaining a homogeneous target temperature and effective debris mitigation, are yet to be solved, in particular when increasing the driving-laser power. Gas-based sources, such as argon (Chen *et al.*, 2007, 2010), allow the use of much more powerful laser pulses due to inherent debris-free operation, allowing for fluxes of up to 10^{11} photons s^{-1} $(4\pi)^{-1}$. Nevertheless, they come with a much larger source size, up to 100 μm . This prevents their use for applications requiring some degree of spatial coherence.

For solid-target PXSs, the main challenge is maintaining their operation stability. The shot-to-shot instabilities are mainly caused by mechanical vibrations of the target and fluctuations in the laser pointing. A decrease in X-ray flux on a timescale of several hours is usually observed. This is due to the debris accumulation affecting the mechanical stability of dynamical parts (*e.g.* guiding bearings) or, in some setups, coating the infrared focusing optics and reducing their efficiency. Several normalization strategies for data processing applicable to both types of instability have been suggested (Schick *et al.*, 2012; Zhang *et al.*, 2014), commonly requiring the introduction of a second X-ray detector. Further efforts are focused on automation control (Zhao *et al.*, 2022) and mechanical design improvements.

The choice of driving laser also affects the performance of the X-ray source. Plasma physics models and experimental results indicate that increasing the driver wavelength leads to higher X-ray photon flux (Weisshaupt *et al.*, 2014; Koç *et al.*, 2021), accompanied by a relative increase of the X-ray source size. The presence of the nanosecond pre-pulse decreases the X-ray yield (Eder *et al.*, 2000), whereas the pulse duration determines the maximum temporal resolution for pump-probe experiments. PXSs require a stroboscopic approach for achieving appropriate signal-to-noise ratio. The total exposure time is then highly dependent on the available repetition rate. For solid-target PXSs, it ranges from 10 Hz to nowadays established maximum of 10 kHz. Target Z number also affects the X-ray yield of a source with given driving-laser parameters (Rousse, Audebert *et al.*, 1994).

ELI Beamlines is a part of The Extreme Light Infrastructure ERIC that offers a wide range of high-power lasers, laser-driven sources of short-wavelength radiation and beams of accelerated particles (Rus *et al.*, 2013). A variety of end-stations enable numerous ultrafast experiments relevant to many fields, such as atomic, molecular and optical physics; material and life sciences; plasma physics; *etc.* A copper tape PXS is offered to ELI users as a source with a unique set of parameters for time-resolved X-ray diffraction and spectroscopy experiments, as well as for X-ray imaging applications.

In this article, we report on the Cu tape target PXS at ELI Beamlines, together with two setups for applications in time-

resolved X-ray diffraction and spectroscopy. The article is organized in the following way. Section 2 is devoted to the physics of X-ray generation in a PXS. Section 3 describes the laser systems available for driving our X-ray source and reports on measurement of their characteristics. Section 4.1 describes the target system that provides a renewable solid Cu target. In Section 4.2, we present the measured parameters of the PXS. In Section 5.1, the time-resolved experimental station for ultrafast diffraction (time-resolved experiments with X-rays – TREX) is presented, and Section 5.2 offers an outlook for the X-ray absorption and emission (XAES) end-station. Finally, we provide a discussion and summary in Section 6.

2. Theoretical background: X-ray generation mechanisms in laser plasma

An important milestone for compact laser-driven PXSs was enabled by invention of the chirped-pulse amplification method that introduced high-peak power sub-picosecond laser technology (Strickland & Mourou, 1985). Consequently, the dominant laser absorption mechanism shifted from resonant absorption with longer pulses (Forslund *et al.*, 1977) to vacuum heating (Brunel, 1987) with short ones, provided that the laser pulse has sufficient contrast.

In the long-pulse regime, the heated part of the target undergoes expansion and the region with most absorption occurs at lower plasma density n . The plasma frequency in this region given by $\omega_p = (e^2 n / m_e \epsilon_0)^{1/2}$, where e and m_e are the electron charge and mass, respectively, and ϵ_0 is vacuum permittivity, is close to the laser frequency. The laser energy is absorbed by collisions of plasma electrons that oscillate in the electric field of the laser. Given that plasma expands with the speed of sound c_s , this happens when the laser pulse length is longer than 100 fs (Murnane *et al.*, 1991).

For short pulses interacting with solid targets, the created plasma density profile is much steeper. The laser pulse terminates before the required density for laser propagation is achieved through plasma expansion. Moreover, during laser illumination, the applied electric field can pull electrons off the surface of the thin plasma layer out of the target. Following a laser half-cycle, electrons are re-accelerated back to the target, delivering the acquired energy to the bulk through collisions with neutral atoms.

In both cases, X-ray radiation is produced by energetic electrons, which collide with ions and neutral atoms. These collisions can be either inelastic, ionizing the target atoms by creating vacancies in the inner electronic shells, which allows electrons from upper shells to decay and emit characteristic X-ray fluorescence photons, or they can be elastic and produce continuous *Bremsstrahlung* radiation.

3. Driving kilohertz lasers available at ELI Beamlines

The Cu tape X-ray source at ELI Beamlines can be driven by one of the kilohertz laser systems operated in experimental

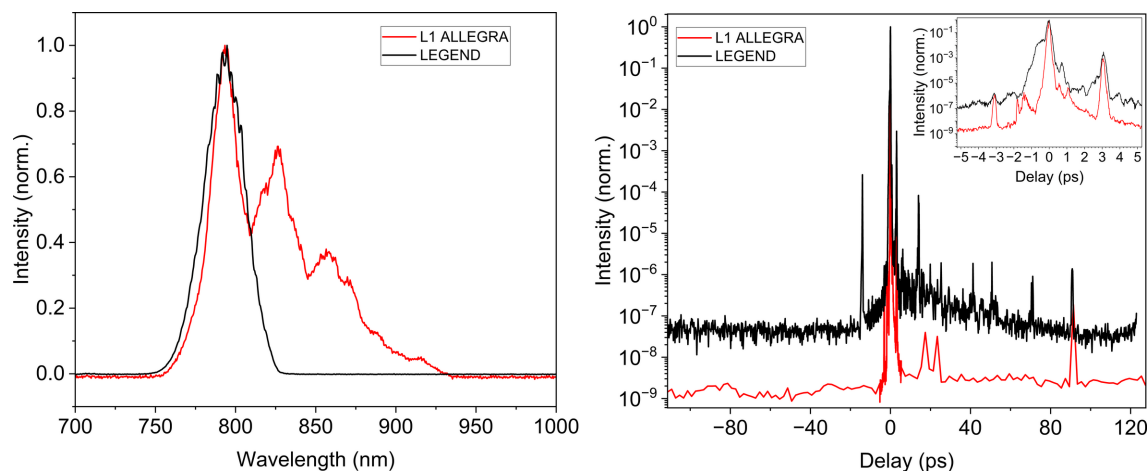


Figure 1

(Left) Measured spectra of 1 kHz lasers available for driving the Cu tape PXS: L1 ALLEGRA with a central wavelength of 820 nm and LEGEND with a central wavelength of 795 nm. (Right) Picosecond temporal contrast of the driving lasers L1 ALLEGRA and LEGEND, highlighting a pre-pulse with intensity 10^{-4} at -14 ps for LEGEND and a ghost peak at -3.2 ps for both lasers.

hall E1: the in-house developed laser L1 ALLEGRA or the commercial LEGEND Elite Duo laser (from Coherent). While LEGEND (12 mJ, 40 fs) is a chirped-pulse amplification system with a Ti:sapphire oscillator and a regenerative amplifier followed by a one-pass booster amplifier, L1 ALLEGRA utilizes optical parametric chirped-pulse amplification (OPCPA) for achieving ultrashort pulses with 15 fs duration and energy up to 50 mJ in a pulse (Antipenkov *et al.*, 2021). The L1 ALLEGRA nanosecond contrast corresponds to the total gain of all amplification stages, which is more than 10^7 , whereas for LEGEND it is as low as 10^3 , as seen at the oscilloscope. Both systems operate at similar central wavelengths (see Fig. 1, left), allowing for convenient switching between them using the same beam-transport optics.

The picosecond temporal contrast of both driving lasers (L1 ALLEGRA and LEGEND) is shown in Fig. 1 (right). Both measurements were performed using a SEQUOIA 800 cross-correlator from Amplitude Systems. The background noise and the limit of detection were determined to be around the 10^{-10} level by blocking each arm of the cross-correlator separately. For L1 ALLEGRA, the contrast was measured using sub-20 fs pulse duration with $\sim 500 \mu\text{J}$ of energy, revealing a 10 ps contrast base level of $\sim 10^{-9}$. Presumably, there are two weak pre-pulses as close as 2 ps and 1.5 ps before the main pulse with relative intensity 10^{-6} . A peak observed at -3.2 ps is identified as a ghost peak, because its amplitude is approximately equal to the square of the amplitude of the post-pulse peak at +3.2 ps. In third-order correlation measurements, such ghost pre-pulse peaks arise due to the mixing of the second harmonic of the post-pulse with the main pulse at the fundamental frequency (Tavella *et al.*, 2005; Chen *et al.*, 2024). For LEGEND, the contrast was measured using a 45 fs pulse with $\sim 500 \mu\text{J}$ of energy, revealing a 10 ps contrast base level of $\sim 10^{-7}$. However, there was a pre-pulse observed at -14 ps with intensity of 10^{-4} and a picosecond-long pedestal with intensity of 10^{-2} .

4. Plasma X-ray source

4.1. Technical description

The PXS is situated in a radiation-shielded hutch, with a footprint of $2.4 \text{ m} \times 3.6 \text{ m}$. Its walls have a thickness of 6 mm and consist of 2 mm of lead sealed from both sides with 2 mm of steel. The layout of the interior of the Cu tape PXS is illustrated in Fig. 2. The chamber is shielded with a lead-steel housing and it consists of three connected vacuum chambers operating at a pressure of 10^{-4} mbar. The upper and lower chambers contain three spools each that regulate the motion of the copper tape and two Mylar bands. Spools are connected with driving motors outside of the chamber with 90° joints.

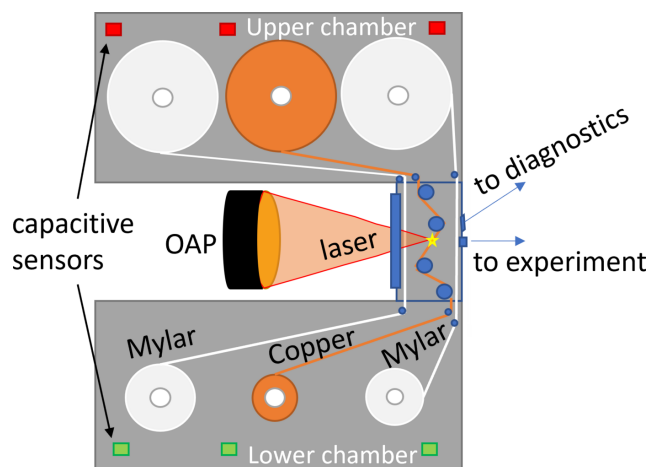


Figure 2

The layout of the PXS. The U-shaped vacuum chamber contains the spools of the copper tape, as well as shielding Mylar tapes that protect the entrance and exit windows from copper dust. The OAP focuses the laser onto the copper tape through the AR-coated vacuum window. The laser interaction point is marked by a star. The X-rays are emitted at a 4π solid angle. Radiation exits through the two Kapton-sealed slits, towards the experiment and diagnostics. The capacitive sensors are marked with red rectangles, if switched by the spool size, and green rectangles, if not activated.

Table 1

A summary of the X-ray source characteristics including driving-laser parameters.

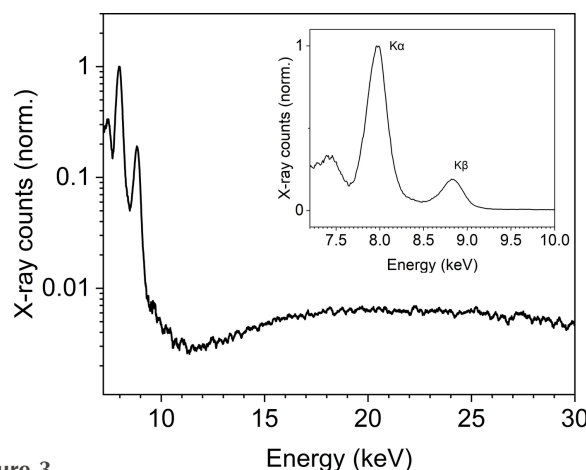
		LEGEND	L1 ALLEGRA
Driving laser	Power†	7.2 W	14.3 W
	Nanosecond contrast	3×10^{-3}	$< 1 \times 10^{-7}$
	10 ps contrast	3×10^{-4}	$< 1 \times 10^{-9}$
	Intensity‡	$1 \times 10^{17} \text{ W cm}^{-2}$	$5 \times 10^{17} \text{ W cm}^{-2}$
X-ray source	Repetition rate	1 kHz	1 kHz
	Plasma source size	$\sim 30 \mu\text{m}$	$\sim 30 \mu\text{m}$
	Average radiant intensity§	$10^{11} \text{ photons s}^{-1} (4\pi)^{-1}$	$10^{12} \text{ photons s}^{-1} (4\pi)^{-1}$
	Montel optics focus size	100 μm	100 μm
	Average photon flux in the X-ray focus§	$2 \times 10^6 \text{ photons s}^{-1}$	$2 \times 10^7 \text{ photons s}^{-1}$

† After the beam transport and beam splitters for the pump arm in the PXS hutch. ‡ Considering 10 μm (at $1/e^2$) focal spot diameter, which was the same for both lasers. § Only $K\alpha$ and $K\beta$ radiation.

The interaction chamber, the smallest one, is where the laser is focused onto the Cu tape target using a 2-inch gold-coated 90° off-axis parabola (OAP) with a focal length of 76.2 mm. The OAP is installed on a motorized kinematic mount for convenient alignment. The copper tape is guided by a set of bearings that define the laser incidence angle of 23° from the target normal. The p-polarization of the laser pulse ensures efficient X-ray generation. The laser is focused onto the target through a 1 mm-thick anti-reflective (AR)-coated vacuum window. The X-ray burst is emitted isotropically but it exits the interaction chamber in beams that are defined by two exit slits enclosed by a 50 μm -thick polyimide tape. The main beam dedicated to applications is situated along the laser-propagation axis, while the second beam points upwards at a 45° angle and is used for diagnostic measurements and source alignment (see Section 4.2 for details).

The PXS at ELI Beamlines uses a 20 mm-wide 20 μm -thick copper tape target. The velocity of the tape can be adjusted based on the plasma spot's size so that the surface damage of neighboring interaction spots does not overlap, and each laser pulse impacts the renewed target. The turning points of the copper tape are monitored and controlled by capacity sensors that are activated when the growing spool reaches its expected maximum size. The motion of the Cu tape is stopped at this point. For starting the new line, the PXS chamber shifts horizontally, perpendicular to the laser axis, while the OAP is decoupled from this motion, therefore the focal spot remains fixed in space. This approach ensures that the X-ray source stays at the same position with respect to the downstream X-ray optics and no realignment is needed when the new line is started. An ultrafast Uniblitz laser shutter (80 ms full cycle) is automatically closed during the line-changing sequence to prevent the tape from being cut by the laser during the chamber shift.

As a large amount of copper dust debris is produced from the laser impact, two Mylar foils are used to protect the vacuum windows. Moving tapes continuously remove the debris and their motion is also regulated by the capacitive sensors. On activation of the capacitive sensors, the Mylar tape automatically reverses its spooling direction. The interaction chamber is separated from the rest of the vacuum system only by thin slits for the tapes, to reduce the debris in the upper and lower vacuum chambers and protect the spooling system.


Figure 3

An emission spectrum of the Cu PXS driven by the LEGEND laser, depicting characteristic copper lines together with *Bremsstrahlung* radiation, measured with a 360 μm -thick aluminium filter inserted in front of a single-photon-counting silicon detector (AMPTEK Si-PIN detector with thickness of 500 μm). The inset displays the $K\alpha$ and $K\beta$ emission lines in higher resolution. The resulting spectrum is corrected for air absorption and aluminium attenuation, as well as absorption in silicon (Henke *et al.*, 1993). Energy calibration of the spectrometer was performed using lines of ^{55}Fe and ^{241}Am radioactive sources. The spectrum was measured at a distance of 1.6 m with an exposure time of 22 min.

4.2. X-ray source parameters

A typical X-ray spectrum of the Cu PXS (shown in Fig. 3) consists of two pronounced characteristic lines, $K\alpha$ at 8.04 keV and $K\beta$ at 8.91 keV, and a broadband continuous *Bremsstrahlung*. The drop of spectral brightness around 11 keV might be caused by reabsorption of Cu atoms and ions.

In Table 1 we summarize basic characteristics of the driving lasers and key properties of the X-ray sources driven by each laser. The individual measurements of the X-ray beam are described in the following subsections.

4.2.1. X-ray source size

The size of the X-ray source was measured from a radiograph of perpendicular steel blades with a two-dimensional knife-edge method. An image of the blades with a magnification of 7 was recorded onto a back-illuminated X-ray CCD with deep depletion sensor technology with 13.5 μm pixel size

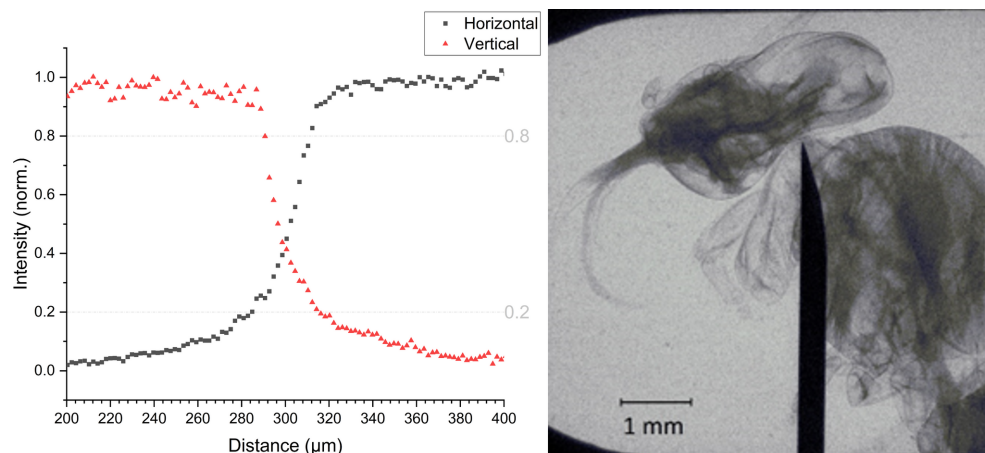


Figure 4

(Left) Knife-edge test results measured with the LEGEND laser. A normalized intensity profile of a sharp edge of a steel razor blade placed between the source and the detector with ratio of distances being 7. (Right) A radiography image of a bee acquired with 60 s exposure time. The black pointy object represents a needle fixing the sample.

(Andor ikon-L 936) providing an effective spatial resolution of $\sim 2 \mu\text{m}$. The results showed that the source is symmetric in horizontal and vertical dimension. To assess the X-ray source size, we applied the ‘20–80% intensity’ criterium and calculated the diameter of the X-ray source to be $30 \mu\text{m}$ (Fig. 4, left). Assuming a Gaussian source, the corresponding full width at half-maximum source size would be $20 \mu\text{m}$.

Such a small size enables the PXS to be utilized for X-ray imaging applications. The imaging capabilities of the PXS are illustrated in Fig. 4 (right). For the proof-of-concept demonstration, a bee was placed in front of the exit slit and the transmission image was captured onto a CCD camera with a magnification of 4.3 and an exposure time of 60 s. Among other possible applications, near-field ptychography was successfully demonstrated with a compact laboratory plasma source for the first time at the PXS beamline (ELI Beamlines) (Fardin *et al.*, 2024).

The post-shot traces on the tape in Fig. 5 reveal significantly different interaction regimes of the two driving lasers. While L1 ALLEGRA produced clean holes of $65 \mu\text{m}$ diameter, Fig. 5(a), LEGEND created crater-like traces with a notably asymmetric halo of comparable size, Fig. 5(b). The burnt-through part is only $20 \mu\text{m}$ in diameter. We believe this is due to different peak intensity and contrast of the two lasers that transfer into different laser–matter interaction scenarios.

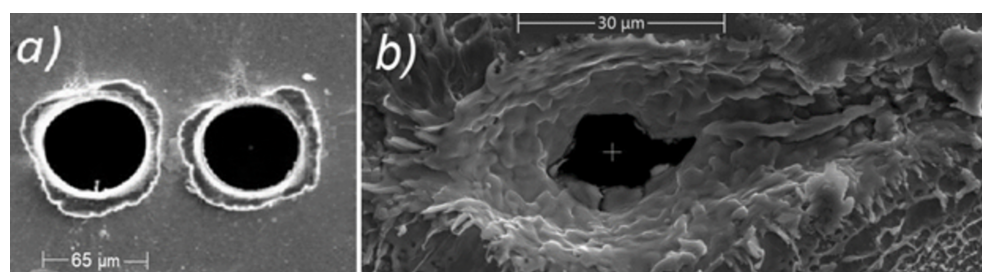


Figure 5

Comparison of laser impact for (a) the L1 ALLEGRA laser and (b) the LEGEND laser. The corresponding scale is inserted in each image.

4.2.2. Radiant intensity

For estimating the number of photons in each X-ray pulse, listed in Table 1, we utilized an X-ray detector based on a PIN diode (xPIN by Rigaku Innovative Technologies Europe). The silicon diode was covered by a $25 \mu\text{m}$ -thick black Kapton window, which blocks all visible light but transmits X-rays. The sensitivity of the diode, with thickness of $300 \mu\text{m}$ and an active area of $7 \text{ mm} \times 7 \text{ mm}$, ranges from 3 to 30 keV, peaking at 8 keV. The diode was biased by 50 V to increase its sensitivity and the signal-to-noise ratio of the oscilloscope readout. Moreover, the diode can capture each shot at a 1 kHz repetition rate to provide reference data for application experiments. For conversion of the photo-charge measured by the diode to the number of photons, we used the responsivity of the diode at 8 keV, as most of the photons are emitted in this spectral line (Fig. 3).

The radiant intensity of the *Bremsstrahlung* was estimated from the spectrum at 20 keV. The flux per 1 keV is two orders of magnitude lower in this spectral region compared with the $K\alpha$ flux.

4.2.3. X-ray pulse duration estimate

The laser-driven PXS occupies its niche among other X-ray sources mainly for its short pulse duration. The theoretical

model predicts sub-picosecond pulses (Reich *et al.*, 2000), which was strongly supported by indirect measurements (Linde *et al.*, 2001; Zamponi *et al.*, 2009; Zhavoronkov *et al.*, 2005). In these studies, the rapid change in physical properties of the sample was monitored via an ultrafast pump–probe scheme. These methods are usually based on monitoring a drop of intensity or shift of the Bragg peak due to the atomic lattice oscillation or observing a drop of reflectivity due to the lattice melting. Successful resolution of the femtosecond dynamics gives an upper limit for the probe pulse duration and is estimated to be 300–500 fs. The first cross-correlation measurement of X-ray pulse duration from the laser-driven PXS indicated the pulse length to be ~ 100 fs (Iqbal *et al.*, 2015).

The current understanding of the emission process suggests that the $K\alpha$ radiation is limited by the laser pulse duration and electron thermalization time (Rousse, Antonetti *et al.*, 1994; Kieffer *et al.*, 1996; Teubner *et al.*, 1996). To the best of the authors' knowledge, direct measurements of sub-picosecond pulses in the hard X-ray range have not yet been performed. However, continued advancements in ultrafast X-ray detectors are driving progress toward achieving 100 fs streak camera technology (Qiang *et al.*, 2009; Liu *et al.*, 2003; Toufexis & Dolgashev, 2019; Li *et al.*, 2015; Feng *et al.*, 2010), bringing us closer to the PXS pulse-duration direct measurements. Recently, an indirect measurement technique was suggested, predicting the achievable temporal resolution to be a few tens of femtoseconds (Nazarkin *et al.*, 2004).

The time-resolved structure of the accompanying *Bremsstrahlung* radiation is an important parameter for the time-resolved absorption spectroscopy. The pulse durations of both *Bremsstrahlung* and $K\alpha$ emission were modeled suggesting similar pulse duration (Reich *et al.*, 2007).

5. Experimental end-stations

In addition to its compact footprint, a key feature of our PXS is its ability to produce pulses with durations of the order of hundreds of femtoseconds. To fully utilize this capability, we have identified two time-resolved applications – X-ray diffraction and X-ray spectroscopy – that are ideal for studying ultrafast sample dynamics initiated by photoexcitation in a pump–probe setup. For this arrangement, a pump beam is generated by splitting 10% of the main driving-laser beam earlier in the beam-transport section and synchronized with the X-ray pulse using a delay line. The pump pulse, typically at 800 nm, can be converted to higher harmonics or used to generate a supercontinuum to optimize the sample response.

5.1. The X-ray diffraction end-station

The TREX end-station at ELI Beamlines is a state-of-the-art instrument designed for time-resolved X-ray diffraction and scattering experiments to study fast processes in crystallographic samples. The end-station is a versatile diffract-

ometer designed to investigate electronic and structural dynamics in a wide range of scientific fields, including chemical reactions, biomolecular dynamics and material science. The setup features a Eulerian cradle diffractometer (STADIVARI from Stoe & Cie GmbH) that enables precise sample positioning through three-axis orthogonal translations and three rotations (SmarAct, GmbH). This configuration supports a wide range of experiments, including powder diffraction, wide-angle scattering and studies on thin solid film samples. The main detector is a Jungfrau detector (three panels, total 1.5 megapixel with 75 μm pixel size) from PSI, Switzerland (Mozzanica *et al.*, 2018; Leonarski *et al.*, 2018), which can be translated 40–400 mm from the sample and rotated in the θ – 2θ mode. An additional X-ray back-illuminated CCD with deep depletion chip technology (Andor ikon-L 936) is available for alignment purposes. Some experiments were performed using a hybrid photon-counting detector, Eiger X 1M (DECTRIS) (Fig. 6). For single-crystal protein diffraction experiments,

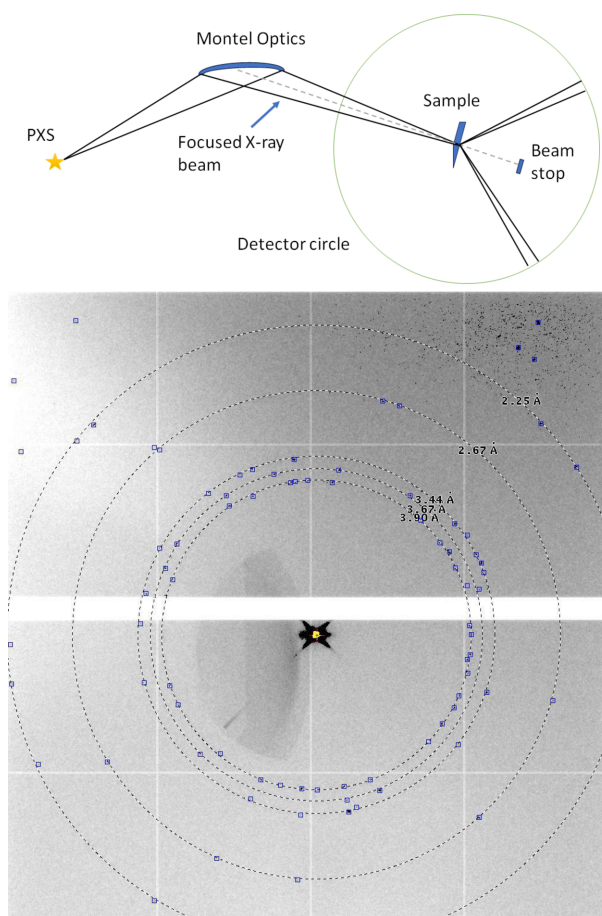


Figure 6
(Top) A scheme of the TREX end-station. A scheme of 1D parallel beam optics geometry with fixed entrance and exit apertures of Montel optics ASTIX++ from AXO Dresden. (Bottom) Diffraction from hexagonal and cubic ice formed at 100 K, measured with DECTRIS Eiger X 1M at a distance of 46.4 mm from the sample. Localization of diffraction peaks was performed using ADXV (Arva, 2015) with a signal-to-noise ratio threshold of 7. The shadow of the direct beam is visible on the left side of the image owing to the absence of the collimator. Diffraction rings of the ice are at interplanar spacings of 2.25, 2.67, 3.44, 3.67 and 3.90 Å.

sample cooling can be provided by a cryogenic gas stream (Oxford Cryosystems).

Femtosecond X-ray pulses generated by the PXS are conditioned using motorized Montel optics (Montel, 1957; AXO Dresden GmbH) for monochromatization, focusing and collimation (Shymanovich *et al.*, 2008). A 15 cm-long Montel mirror with an acceptance angle of 16.8 mrad is placed 11.5 cm from the source delivering $K\alpha$ radiation onto a probed sample with enhanced reflectivity thanks to an Ni/C multilayer coating. The focus is 54 cm from the mirror's exit edge. Considering a primary focal length of 190 mm and a secondary one of 615 mm, the magnification of the optics is ~ 3.2 , resulting in a divergence of the focused beam of 5 mrad. The beam intensity depends on the driving laser and for L1 ALLEGRA laser operation was measured to be $\sim 2 \times 10^7$ photons s^{-1} with a focal spot size of $\sim 100 \mu m$.

The Jungfrau detector, a key component of the TREX end-station, operates at a high repetition rate of 1 kHz, generating a substantial amount of data that can fill several terabytes of disk space daily. To tackle this data challenge, ELI Beamlines is developing customized software solutions for calibrating and converting the raw data into a common photon-per-pixel format. Simultaneously, progress is being made in standardizing 2D image formats based on the HDF5 specification, which will promote harmonization with other large-scale facilities and enable more efficient data sharing and collaboration among researchers.

The station's modular design also allows for adaptations to accommodate user-initiated experiments, providing researchers with the flexibility to install their own equipment. The combination of femtosecond X-ray pulses and advanced detectors at the TREX end-station will enable high-resolution time-resolved studies of various phenomena, such as phase transitions, chemical reactions and protein dynamics (Schmidt, 2019; Rischel *et al.*, 1997). Future plans include the implementation of wide-angle X-ray scattering, thin-film reflectometry and surface diffraction techniques.

5.2. The X-ray spectroscopy end-station

X-ray spectroscopy is an experimental tool that gives access to information about the electronic structure of materials. X-ray emission spectroscopy allows one to probe occupied states while X-ray absorption spectroscopy traces the difference between the incoming and outgoing X-ray signal, revealing the photon fraction absorbed in the sample and thus exploring unoccupied states. Fundamentally, the two techniques provide complementary information. Thus, the goal of the X-ray spectroscopy station at ELI Beamlines is simultaneous acquisition of emission and absorption data via development of parallel measurements accommodated around the Cu tape PXS with sub-picosecond temporal resolution.

Typically, an X-ray spectroscopy experiment consists of an X-ray source, the sample under study (with corresponding environment if required) and a spectrometer. The latter can be a single-body detector, *e.g.* of a microcalorimeter type

(Doriese *et al.*, 2017), in a very compact setup, or a crystal-based solution (Sokaras *et al.*, 2013) that can achieve much higher spectral resolution. In the case of a crystal-based design, there are two alternative approaches to spectrum collection. In Johann and Johansson-type spectrometers, a 2D focusing crystal selects a single energy and focuses the X-rays down to a point. To acquire the full spectrum an energy scan is needed, which requires moving the crystal and the detector. Alternatively, in von Hamos geometry (Szlachetko *et al.*, 2012), a crystal bent in one dimension and energy resolving in another allows for collection of full spectra simultaneously, eliminating the need for scanning but reducing the solid angle of collection for a given photon energy. For a focusing crystal, a few alternatives exist, such as segmented (Németh *et al.*, 2016) or continuous Si wafers glued to a cylindrically bent substrate, highly oriented pyrolytic graphite (Legall *et al.*, 2006) and highly annealed pyrolytic graphite (Malzer *et al.*, 2021).

A requirement to resolve chemical changes of the oxidation state (*i.e.* well below 1 eV) has imposed the choice of a crystal-based spectrometer. However, the space constraints in the experimental hutch called for strategic selection of a compact design. Thus, we have selected a von Hamos type spectrometer (Zymaková *et al.*, 2023). An additional benefit from this choice is that it enables us to use the spectrometer solutions developed for the PXS at other X-ray sources available at ELI Beamlines, such as the plasma betatron X-ray source (Chaulagain *et al.*, 2022), providing a user with a wide choice of source parameters.

Polycapillary X-ray focusing optics have been incorporated (Zymaková *et al.*, 2020) for experiments with inhomogeneous samples and lower requirements on temporal resolution. The system can be used for solid, powder or liquid samples. The ability to handle liquid samples paves the way for biological sample studies. A few different liquid sample delivery systems, such as a wire-guided jet (Picchiotti *et al.*, 2023), a colliding jet (Koralek *et al.*, 2018) and a microlitre stirred cell (Fanselow *et al.*, 2022) are available for users at ELI Beamlines.

6. Discussion and outlook

6.1. Comparison with other laser-driven sources and outlook

Among established PXS beamlines, our source radiant intensity reaches the current best-efforts flux for such sources, ranging up to 10^{11} – 10^{12} photons s^{-1} $(4\pi)^{-1}$ (Zhao *et al.*, 2022; Holtz *et al.*, 2017). An exceptional flux of 10^{12} photons s^{-1} is reached by a driving laser with pulse energy as low as 3 mJ thanks to the use of a long 5 μm wavelength (Koç *et al.*, 2021). An additional X-ray yield can be reached by optimizing a prepulse (Lu *et al.*, 2024), achieving 10^5 photons per shot on the sample. Similar flux values were achieved by Afshari *et al.* (2020), utilizing high-energy laser pulses of 100 mJ.

Depending on the choice of focusing optics, the average loss at monochromatization is four (Zhao *et al.*, 2022), five (Koç *et al.*, 2021) or six orders of magnitude (Holtz *et al.*, 2017; Schick *et al.*, 2012). For our Montel optics, which belong to the

longer-focus category, the flux loss is 10^6 compared with the full solid angle.

Our radiant intensity will be further enhanced through improved stability achieved by a redesigned interaction chamber. The current distance between the guiding bearings and the plasma is insufficient, leading to gradual damage of the bearings and consequent tape quivering. Furthermore, active beam stabilization that utilizes a quadrupole diode for monitoring the laser beam position and pointing will prevent the source from drifting due to temperature drift of the system.

6.2. The PXS as an alternative to large-scale facilities

Time-resolved diffraction as well as X-ray spectroscopy are well established techniques in large-scale facilities such as synchrotrons and XFELs. While ultrafast high-intensity pulses from XFELs ensure exceptionally fast data collection, ‘diffraction before destruction’ is assumed (Chapman *et al.*, 2014). Nevertheless, as 10^{12} photons are delivered to the sample in just 1–100 fs (Abela *et al.*, 2017), it raises the moderate concern of the possibility of rapid changes in protein structure in response to extreme irradiation, especially where pulse durations are greater than a few tens of femtoseconds (Nass, 2019; Nass *et al.*, 2020; Garman & Weik, 2023; Brinkmann & Hub, 2016; Arnlund *et al.*, 2014; Ansari *et al.*, 1985). In addition, the destruction of a crystal after a single exposure from a high-intensity XFEL pulse results in high rates of sample consumption.

Synchrotrons provide flux up to 10^{15} photons s^{-1} (Garman & Weik, 2023) but usually operate in a pseudo-continuous wave mode, limiting the achievable temporal resolution when in normal operation. Detector gating (Donath *et al.*, 2023), femtoslicing (Labat *et al.*, 2018) and rotating choppers may be used (Meents *et al.*, 2009; Cammarata *et al.*, 2009) but radiation damage remains an important consideration.

However, both XFELs and synchrotrons demand large infrastructure for their operation. Thus, PXSs emerge as a promising alternative beamline for time-resolved experiments. PXSs combine the advantages of a pulsed source and short pulse duration. The PXS is a tabletop and potentially could fit into a small university laboratory. Their relatively low cost makes them an affordable option. However, the downside of the PXS is its unstable flux and comparatively low brilliance. Nevertheless, the latter can usually be overcome by a stroboscopic (Kim *et al.*, 2002) or multiplexing (Klureza *et al.*, 2024) approach, where an ensemble of the sample responses is gathered and merged together. This makes PXSs a suitable tool for studying ultrafast reversible processes, such as superlattice oscillations, X-ray absorption spectroscopy, *etc.*

7. Conclusions

In this article, we have introduced the capabilities of a newly commissioned 1 kHz laser-driven Cu tape PXS, showcasing its versatility and potential for a range of scientific applications. The source can be driven either by a sub-20 fs near-infrared

OPCPA system or by a commercial Ti:sapphire laser. Our findings highlight the differences in X-ray beam characteristics and photon flux achievable with these different laser systems. A significantly higher X-ray yield with the OPCPA driver is given by the contribution of several factors. First, higher laser-pulse energy and shorter pulse duration lead to higher laser intensity on the target. In addition, better nanosecond contrast further enhances the $K\alpha$ yield compared with the case driven by the conventional Ti:sapphire laser.

The ability to deliver photon fluxes up to the order of 10^{12} $K\alpha$ photons s^{-1} $(4\pi)^{-1}$ underlines the high performance of our X-ray source. Additionally, we have outlined the potential of this system for ultrafast X-ray diffraction and X-ray absorption and emission spectroscopy, paving the way for innovative research studying ultrafast structural dynamics. As the system is now available for user-based access, we anticipate it will serve as a powerful tool for the scientific community, enabling a wide range of experiments in the fields of material science, chemistry, biology and physics.

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Conflict of interest

Authors declare that there are no conflicts of interest.

Data availability

Data supporting this publication are available upon reasonable request to the corresponding author.

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