Demonstration of a sub-picosecond x-ray streak camera

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A novel design, magnetically focused, x-ray streak camera was designed and tested using sub-20 fs soft-x-ray pulses generated by high harmonic emission in a gas. The temporal resolution of the camera was demonstrated to be under 0.9 ps throughout the ultraviolet to soft-x-ray wavelength region. Our streak camera represents the fastest x-ray detector developed to date. © 1996 American Institute of Physics. [S0003-6951(96)03327-X]

During the past decade, the development of ultrafast x-ray sources based on laser-produced plasmas, high harmonic emission, and synchrotrons, has advanced rapidly. 1–3 It is now possible to generate sub-picosecond pulses throughout the vacuum ultraviolet and x-ray region of the spectrum, and sub-10 fs pulses have been generated using high harmonic emission. 4 However, progress in the development of ultrafast sub-picosecond x-ray detectors has been relatively slower. 5,6 Although cross-correlation techniques of ultrafast sub-picosecond x-ray detectors has been tally demonstrated to be 0.88 ps. 4,8 The resolution of our camera was experimentally demonstrated to be 2 ps. This measurement was limited both by the time response of the streak camera itself, and by the laser-plasma-based x-ray source. 5,6 In this letter we describe the design and implementation of a novel x-ray streak camera, which exhibits sub-picosecond time resolution. The time response of the streak camera was measured using ultrashort sub-20 fs high-order harmonics produced by a 25 fs laser. 4,5 The resolution of our camera was experimentally demonstrated to be 0.88 ps.

It is well known9,10 that the temporal resolution of streak cameras is limited mainly by the transit time dispersion of the photoelectrons as they travel from the photocathode to the deflection plates. It is also limited by the spatial resolution, and the deflection speed of the streak plates. For sub-picosecond time resolution, space-charge effects may also limit the time resolution, and thus limit the dynamic range. For our work, we designed and tested a novel streak camera design to reduce the limitations on temporal resolution as much as possible. The configuration of the new x-ray streak camera is shown schematically in Fig. 1. In our camera, a pair of meander-type deflection plates is located before a magnetic focusing lens. This has several advantages: first the electron transit time from the anode to the deflection plates is minimized, as is the transit-time dispersion; second, the fast time response and high deflection sensitivity (8 cm/kV) of the meander-type deflection plates provides the possibility of high sweep speeds on the exit phosphor screen; finally, the short transit time also reduces space-charge effects.

The electron transit time dispersion from the photocathode to the deflection plates in the camera can be evaluated analytically. In the photocathode to anode region, it is straightforward to show that the transit time difference between an electron liberated with energy $eV_0$, and an electron liberated with zero energy is given by

$$t_{pa} = \sqrt{\frac{2mV_0}{e} \frac{1}{E}}. \quad (1)$$

where $m$ and $e$ are the charge and mass of the electron, respectively, $eV_0$ is the initial energy of an emitted photoelectron in the axial direction, and $E$ is the extraction field.

For x-ray photocathodes in the 100 eV to 10 KeV region, the distribution of the initial energies of the emitted photoelectrons can be expressed as

$$N(eV_0) = \frac{eV_0}{(eV_0 + W)^2}, \quad (2)$$

where $W$ is related to the photocathode material ($W = 1$ eV for KBr). The photocathode material also determines the full width at half-maximum (FWHM) of the energy distribution ($\delta e$). From Eqs. (1) and (2), we obtain the transit time distribution of the electrons. The FWHM of this distribution is defined as the time dispersion, and can be shown to be

$$\delta t_{pa} = \frac{2.63 \sqrt{\delta e}}{E} \text{ (ps)}, \quad (3)$$

where $\delta e$ is in eV, and $E$ is in kV/mm. For our camera, $\delta e = 1.1$ eV (KBr photocathode), and $E = 10$ kV/mm, resulting in a calculated time dispersion of about 276 fs.

![FIG. 1. Configuration of the sub-picosecond x-ray streak camera.](image_url)
In the nearly field free region between the anode and the entry to the deflection plates, we can again calculate the transit time distribution of the electrons. The distribution FWHM is then given by

\[ \delta t_{ad} = \frac{t_{ad} \delta \phi}{eV_a}, \]

where \( t_{ad} \) is the transit time for electrons with \( eV_a = 0 \). For our streak camera, \( t_{ad} = 500 \) ps, so that \( \delta t_{ad} \) is 25 fs. It is clear that the time dispersion in the anode-to-deflection plates region is much smaller than in the photocathode-to-anode region. The above analysis does not take into account the angular distribution of the photoelectrons emitted from the photocathode. We have, however, calculated the transit time dispersion from the photocathode to the deflection plates, assuming the angular distribution is Lambertian. The calculated dispersion is 250 fs.

The spatial resolution of the streak tube was simulated by tracing the trajectories of the electrons from the photocathode to the phosphor screen. This was done by solving the dynamic equations of motion for the electrons in an electromagnetic field, using the Runge–Kutta method. The magnetic field distribution was calculated by the finite difference method, assuming that the shield metal has an infinitely high magnetic permeability, and is not saturated. The calculated image width of the slit on the phosphor screen is \( \sim 60 \mu \text{m} \).

In our camera, the photoelectrons are multiplied by a microchannel plate (MCP) detector placed in front of the output phosphor screen, as shown in Fig. 1. The output phosphor screen is then fiberoptically coupled to a proximity focused second generation image intensifier. The image is then lens coupled to a low light change coupled device (CCD) camera, which is connected to a frame grabber. The experimentally measured width of the slit image on the CCD camera is \( \sim 80 \mu \text{m} \). The high voltage on the streak camera photocathode was applied by superimposing a \(-5 \text{ kV}\) pulse to a \(-5 \text{ kV}\) dc voltage. Use of a pulsed extraction field prevents electrical breakdown between the photocathode and the anode. A GaAs photoconductive switch\(^{12,13}\) was used to obtain a fast ramp voltage for driving the deflection plates. This switch also exhibits a small relative time jitter of about 5 ps. The resulting sweep speed is \( 2 \times 10^3 \text{ m/s} \), so that the camera time resolution is limited at 400 fs by the spatial resolution and sweep speed. Also taking into account the time dispersion, the total estimated time resolution is \( \sim 0.5 \) ps.

Preliminary tests of our camera were performed using the third harmonic of a Ti:sapphire laser.\(^4\) The 265 nm light was generated by focusing an 800 nm, 26 fs, 3 mJ laser pulse in air. They demonstrate a time resolution of 880 fs in the x-ray region for our streak camera, as shown in Fig. 3. This is, to our knowledge, the first demonstration of a sub-ps response x-ray streak camera. Our results are somewhat longer than the expected 0.6 ps response calculated from a convolution of the dispersed x-ray pulse duration (\( \sim 0.3 \) ps) and estimated camera resolution (\( \sim 0.5 \) ps). We believe that the difference is due to the decrease of spatial resolution in the dynamic or streaked mode. Further work is in progress to improve our results, by designing better electrically matched streak plates.

In conclusion, we have demonstrated a novel design x-ray streak camera with sub-picosecond time resolution. Sub-20 fs high harmonics were used to calibrate our streak camera. We believe that the time resolution our camera can
be improved further by redesigning the deflection plates to achieve higher sweep speeds. Our work demonstrates that sub-picosecond time resolution experiments can now be performed using currently available synchrotron or laser–plasma sources.

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