A flexible, on-line magnetic spectrometer for ultra-intense laser produced fast electron measurement

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Abstract

We have developed an on-line magnetic spectrometer to measure energy distributions of fast electrons generated from ultra-intense laser-solid interactions. The spectrometer consists of a sheet of plastic scintillator, a bundle of non-scintillating plastic fibers, and an sCMOS camera recording system. The design advantages include on-line capturing ability, versatility of detection arrangement, and resistance to harsh in-chamber environment. The validity of the instrument was tested experimentally. This spectrometer can be applied to the characterization of fast electron source for understanding fundamental laser-plasma interaction physics and to the optimization of high-repetition-rate laser-driven applications.

Keywords: electron spectrometer; on-line spectrometer; ultra-intense laser-solid interaction

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1. Introduction

The interaction of ultra-intense laser with overdense plasma produces large-current relativistic (fast) electron beams. Knowing the electron energy distribution and its dependence on laser and target parameters are important to the understanding of fundamental laser-plasma physics, and to the optimization of electron sources for applications, e.g., the fast ignition (FI)\(^1\) approach to inertial confinement fusion (ICF), laser-driven ion acceleration\(^2, 3\) and bright radiation sources development\(^4, 5\).

The electron energy spectrum is normally measured with a dipole magnet based electron spectrometer. The electrons are first dispersed according to the different kinetic energies in the magnetic field and then recorded by the detector(s). Three major detection methods previously reported are image plates (IPs), scintillating fibers and scintillator screen both imaged directly with C-CD/sCOMS camera. The major drawback of IPs\(^6, 7\) is the need to vent the vacuum chamber to read the stored signals. Therefore IPs are not appropriate for high-repetition-rate experiments. The scintillator fibers\(^8–11\) are normally placed parallel to the magnetic lines along the sides of both plates. The effects of fringe field on electron deflection trajectory are inevitable\(^12\). In addition, the effective fiber length which produces scintillating light is only a small portion of the fiber. The longer redundant fibers beyond the magnetic field region will produce background scintillating light when exposed to optical light or particles. To eliminate the undesired background, Chen \textit{et al}\(^10\) has put the CCD camera inside the vacuum chamber to shorten the fiber length. However, the CCD suffered from the strong electromagnetic pulses (EMP)\(^13\) in the harsh environment and may not work properly\(^14\). When scintillator screens\(^15–18\) were used, they were normally imaged directly with an optical lens coupled CCD camera. Extensive shielding is an essential requirement to avoid scattered optical light entering the imaging system. Due to the large aspect ratio of screen (> 10) and considerable length along the dispersion direction (normally tens of centimeters), effectively only a very small portion of CCD chip was used,
limiting the spatial resolution on the scintillator detector by the imaging system to only several hundred microns[2]. Another important issue is that the imaging system needs be realigned when changing the positions of the electron spectrometer to measure electron spectra along other directions.

In this paper, we present a novel design of a flexible, on-line magnetic spectrometer, which utilizes the scintillator screen to produce scintillating light and employs the non-scintillating fibers to guide light to high frame-rate sCMOS camera imaging system. This instrument combines the advantages of the on-line capability of scintillator, flexibility and versatile transforming configuration of the fibers, and resistance to harsh environment by moving the recording electronics out of the vacuum chamber.

2. Spectrometer design

The schematic of the spectrometer is shown in Fig. 1(a) (the imaging system is not presented). A pair of parallel NdFeB permanent magnets, shown in light blue, deflects the electrons according to their different kinetic energies (representative lines in black). A thin sheet of plastic scintillator (in dark red) is inserted inside the magnetic spectrometer to convert the electron signal into visible optical light. The scintillator screen can facilitate an on-line detection capability since the emitted light from the scintillator can be captured by the imaging system without disturbing the detector. This would attract more interests especially in high-repetition-rate single-shot evaluations for statistical analysis.

A bundle of fibers, arranged in multiple rows of linear arrays, are placed with one end in close contact with the scintillator sheet. The light signals from the scintillator are then relayed by the fibers to the imaging system placed outside the vacuum chamber[19]. A high frame-rate sCMOS camera located outside the vacuum chamber is used to record the optical light transmitted through a glass window. Considering the aspect ratio of sCMOS chip size, the other end of the fiber bundle is rearranged to an areal configuration. The insets in Fig. 1
(a) illustrate the linear and areal interfaces of the fiber bundle, respectively. A photo of the assembly is shown in Fig. 1 (b).

Trapezoidal-shaped (truncated square) magnets (marked by dashed-line in Fig. 2 (a)) are used in this work. The magnet plates have an overall side-length of 200 mm and a separation gap of 10 mm. They are enclosed with a 10-mm-thick iron yoke. The spatial distribution of the magnetic field strength in the mid-plane was mapped out with a Hall probe, and is shown as the grayed area in Fig. 2(a). The peak field strength is 0.12 T. The fringed field decreases from 0.10 T at the edge to 0.05 T 10 mm away from the magnet plates. If the detector is not inside the uniform region and there is no collimator at the entrance, the electron trajectories and energy dispersions need to be precisely simulated using a particle tracing program\[12\] in order to include the effects of fringe fields. Simplified calculations using simple lorentz force consideration can be obtained by minimizing fringe field effects with a collimator placed inside a central region of 160 mm with field strength variation of only 1.26% (standard deviation).

An ultralow-carbon stainless-steel collimator was chosen as the entrance. The collimator is 30 mm long with a 3.5 mm hole in the center (yellow in Fig. 2(a)). The magnetic field increases from 0 to 0.12 Tesla within 13 mm beyond the collimator hole. The scintillator screen was also inserted into the near-uniform magnetic field region further reducing the effect of fringe field. A visible CW laser (red dash line) was used to align the spectrometer.

The plastic scintillator is a DRZ fluorescent screen (Gd$_2$O$_2$S:Tb, type Pl-200) having dimensions of 200 mm (length) × 10 mm (width) × 630 μm (thickness) and emits light at 546 nm when impacted by energetic electrons. The scintillator screen itself has a higher spatial resolution (a few 10’s of microns) than the large-diameter scintillator fibers (typical 500 μm)[10] and it is not the limiting factor for the energy resolution. A 2 μm-thick aluminum-anodized mylar film was placed before the scintillator to block scattered light. This type of scintillator has been calibrated using ultrashort electron beam from a radio frequency linear electron accelerator[20]. The 20 cm-long plastic scintillator sheet (dark-red line in Fig. 1(a)) was positioned at 45° with respect to the entrance axis. In this case,
Figure 1: (color online) The schematic (a) and photograph (b) of the magnetic spectrometer. The regions in light blue and in green are magnet plates and the yoke, respectively. A collimator was inserted into the region of near-uniform magnetic field strength. The red lines indicate the fiber bundles. Two insets in (a) show the linear and areal interfaces of the fiber bundle.
the spectrometer works in the focusing mode, i.e. finite-diameter collimated
electron beam is bent by the magnet and overlaps on the detector plane[11]. The
corresponding trajectories of the electrons with different energies were computed
by calculating the deflection angle according to the Lorentz force, which are
shown in Fig. 2(a).

A total number of 875 equally spaced 1-meter-long plastic fibers are placed in
close contact with the scintillator, to sample and guide the optical light signals
out of the spectrometer. The fiber bundle was further wrapped with aluminum
to avoid stray light. The cross-talk between channels was also checked using
a point laser source to be <4%. Although the spatial resolution of the non-
scentillating fibers is not as high as that of IPs, they were chosen due to several
advantages. Firstly, it is easier to shield the imaging system from scattered light-
s, and no optics are required in the crowded chamber, saving space for other
diagnostics. Secondly, the fibers can be easily bent. Therefore, the imaging sys-
tem can be setup using chamber windows away from the influences of radiations
(eespecially EMPs) and also to avoid conflicts with other diagnostics. Thirdly,
the plastic fibers can be rearranged into a matrix to fit the sCMOS/CCD camera
sensor for better imaging. Fourthly, the imaging component does not require
alignment, when re-positioning of the main body of the spectrometer becomes
necessary to measure the electron spectra along other directions. Lastly, the
fibers are insensitive to the radiations, hence no extensive shieldings are need-
ed.

Each fiber has a diameter of 0.75 mm. The separation of the centers of
adjacent fibers is 1 mm. In the scintillator end, five fibers were employed in
the non-dispersion direction for each specific electron energy, thus forming a
matrix of 5 × 175 fibers. This gives rise to 175 energy points in the dispersion
direction. Figure 2(b) shows the energy dispersion and the resolving power
\((E/\Delta E)\) with respect to the electron energy \((E)\), where \(\Delta E\) is the energy reso-
lution at \(E\). The electron traces in the magnetic field could be calculated using
equation \(d(\gamma m \vec{v})/dt = -e \cdot \vec{v} \times \vec{B}/c\). Where \(\gamma\) is the relativity factor, \(m\)
is the electron mass, \(e\) is the electronic charge, \(\vec{B}\) is the magnetic field, \(\vec{v}\) is the
speed of electrons and \( c \) is the speed of light. In this case, \( E \) is calculated to be ranging from 0.5 MeV to 4.8 MeV. The energy resolutions of the spectrometer are determined by diameter of fibers and the beam sizes on scintillator. The typical electron trajectories in the energy range of 1-4.5 MeV for an acceptance angle of \( 2.4 \times 10^{-4} \) sr are shown in Fig. 2(a). The trajectories of mono-energetic electrons entrancing the magnetic at two edges of the collimator will crossed on the scintillator at two different positions as shown in Fig. 2(a). The distance between these two positions would be the beam size on the scintillator and it is marked by the size of the black dots in Fig. 2(a). The beam sizes on the scintillator (black dots) are determined by the electron divergence, the acceptance angle of the collimator, electron energies, magnetic field and the position of scintillator. It is less than 0.75 mm for energies lower than 1.4 MeV. For electrons with energies higher than 1.4 MeV, the beam size is larger than the fiber diameter, which becomes a predominant factor limiting the energy resolution. As shown in Fig. 2(b), the resolving power first increases from 30 to 77 for the electron energies of 0.5-1.4 MeV due to the fiber diameter and then decreases to 57 from 1.4 MeV to 4.8 MeV when the beam sizes are bigger than fiber diameter. Since smaller diameter optical fibers (a few 10's of microns) are available, decreasing the acceptance angle or the diameter of the fibers will improve the energy resolution.

The other end of the fiber bundle is in close contact with a thin glass window. Due to the fiber flexibility, the window could be selected from any flange on the chamber, preferably away from the laser incident plane. A 16-bit sCMOS camera (ORCA-Flash4.0 LT, from Hamamatsu Photonics) coupled with a f=6-12mm/F1.6 lens was used to image the fibers. The sCMOS camera was chosen with consideration of several superior advantages: small pixel size (6.5 \( \mu m \)), high frame rate and low cost while remaining comparable in quantum efficiency to EMCCD. The fiber bundle was re-organised to an areal configuration of 25\( \times \) 35 fibers to maximize the usage of the sCMOS chip such that the resolution of the imaging system is \( \sim \)50 \( \mu m \)/pixel. Similar work was reported by Kodama et al that two-dimensional fibers were rearranged to a line to fit a
streak camera[21]. The light transport efficiency of the fiber bundle and imaging system was relatively calibrated and the variation between energy channels was 4.5% (standard deviation). Since the camera was detached from the target chamber, the shielding was much easier.

3. Experiment and results

The experiment was carried out using the 200 TW Ti: sapphire laser at Shanghai Jiao Tong University. The setup is schematically shown in Fig. 3. After reflected from the high reflection mirror (M), the 25 fs p-polarized laser beam was focused by an f/4 off-axis parabola mirror (OAP) at 36° to the target normal on 2-μm-thick iron foil targets. The focal spot is 6 μm in diameter enclosing 25% of total energy. In this experiment, the energy on target is 1 J, giving a peak intensity of $\sim 4 \times 10^{19} W/cm^2$. The electron spectrometer was positioned 17 cm away from the target in the front side at an angle 12° to the target surface for measuring the spectra of electrons emitted along the target surface. We tested the flexibility of the spectrometer by quickly changing the detection direction from 12° to 45° in less than 5 minutes. The energy spectra of electrons emitted at 45° was much weaker than that at 12°. The acceptance solid angle is $2.4 \times 10^{-4}$ sr.

An example raw data measured along an angle of 12° to the target surface is shown in Fig. 4(a). The light intensities from the fibers have been normalized and are comparable with each other. Each of the five columns (for I, II, III, IV and V sub-groups) consists of 35 rows and each row corresponds to one energy channel. The electron energy increases with the increment of channel number from top left to bottom right. The overall trend is that the measured light intensities decrease gradually with electron energies, indicating a decreasing continuous energy spectrum (assuming a a flat calibration curve with electron energies). A null test was made by blocking the entrance collimator with a piece of 5-mm-thick lead, resulting in a background count of <10. This is more than two orders of magnitude lower than the case when the block was removed.
Figure 2: (color online) (a) Trajectories of electrons in magnetic field (gray area). Two colorbars depict the magnetic field distribution and the electron trajectories, respectively. Black circles show the electron beam sizes on the scintillator. (b) Dispersion and energy resolving power with respect to the electron energies.
Figure 3: (color online) Schematic of the experimental setup. After reflected from the high reflective mirror (M), the laser beam was focused by an off-axis parabola mirror (OAP) to the solid target. The electron spectrometer was placed in the target front side to measure the energy distribution of the fast electrons emitted close to the surface direction. The fluorescent light from the scintillator screen were guided by the fiber bundle and detected with a lens-coupled sCMOS imaging system. A silver mirror (Ag) was used to avoid the γ-ray photons or high-energy particles directly bombarding the sCMOS sensor.
Figure 4: (color online) (a) Experimental raw data. Each colored circle spot corresponds to one fiber channel. The red numbers and arrow indicate the energy channel and energy increasing direction, respectively. (b) The measured light intensities (red) of the scintillator-based detector and error bars (gray) for the fluctuations of different fibers in one energy channel. A fit of the experiment data was also plotted by considering a two-temperature distribution of electrons.
The arrangement of the fiber bundle can cause fluctuations of coupling efficiency between scintillator and fibers. To reduce the fluctuation in the data, the maximum and minimum values for each energy channel were removed. Taking into account the energy resolutions, we calculated the electron spectra using the remaining three values of the fibers in each energy channel. The spectral distribution of the sampled fast electrons showing a two-temperature distribution is presented in Fig. 4(b) where the error bars are the data fluctuations from the remaining three values in the same energy channel.

The reliability and the on-line capability were also tested experimentally. 10 consecutive shots were taken in 3 minutes for the same laser and target parameters. The shot frequency was only determined by the time of replacing targets, which could be as short as a few 10's of seconds. Compared with the use of IPs, which need to vent the chamber to read the stored signals, the use of scintillator coupling with imaging system is much more attractive in high-repetition-rate experiments. One could obtain the data on electron energy spectrum shortly after each laser shot, which is the prerequisite to source optimization for applications. The shot-to-shot fluctuations of the spectra was measured to be less than 5%, which is within the error bars shown in Fig. 4.

4. summary

In summary, we have designed and tested a flexible, on-line electron spectrometer using a combination of plastic scintillator, plastic fiber array and sCMOS camera. This design would allow on-line measurement of fast electrons produced by high-repetition laser pulses. Better energy resolution can be obtained by using optical fibers with smaller diameters.

This spectrometer can also provide, though relatively modest, one-dimensional angular resolution from the five fiber channels perpendicular to the dispersion direction. The angular resolution could be improved by enlarging the entrance pinhole with a long slit and increasing the number of fiber channels in the plane perpendicular to the dispersion direction. With further revision, this novel
design could be employed to measure the angular and energy distributions of
low-divergence laser wakefield accelerated electron beams. Continuous improve-
ment and development could facilitate many high-power laser-plasma research
projects with a versatile diagnostic for fast electron generation.

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