Technical Report

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\textbf{ABSTRACT}

Recent experiments using Terawatt lasers to accelerate protons deposited on thin wire targets are modelled with a new type of gridless plasma simulation code. In contrast to conventional mesh-based methods, this technique offers a unique capability in emulating the complex geometry and open-ended boundary conditions characteristic of contemporary experimental conditions. The simulations shed new light on a number of experimentally observed features, including the hitherto unexplained ‘double-disc’ emission pattern of the MeV protons accelerated away from the wire. These discs appear to be formed by the combined action of target normal sheath acceleration and resistive hot electron transport effects.

\textbf{1. Introduction}

Since the first experiments measuring ion emission from multi-Terawatt laser-solid interactions \cite{1, 2}, laser-induced acceleration of MeV protons (fast ions) has become one of the most contentious issues in the field. Such protons originate from water vapour or other impurities adsorbed onto the target surface prior to laser irradiation, and by virtue of their lower mass, are preferentially accelerated over heavier constituent plasma ions when the laser creates a charge separation either inside or outside the target. The ability to create multi-MeV protons in a relatively cheap and compact manner has generated widespread interest because of its potential in a number of emerging fields, such as hadron therapy \cite{3}, novel neutron sources \cite{4} and advanced fusion concepts \cite{5}. Experimental campaigns begun by the Livermore and Imperial College groups some four years ago resulted in two apparently irreconcilable pictures of proton acceleration \cite{6, 7, 8}.

The first interpretation, proposed by the Livermore team \cite{6, 9}, supposes that protons will be primarily accelerated from the \textit{rear} surface of thin (1–100 µm) foil targets by the space charge set up by the laser-generated hot electron cloud. This intuitive scenario, dubbed ‘target normal sheath acceleration’, or TNSA, has since been strongly supported by 2- and 3-dimensional particle-in-cell (PIC) simulations performed by various authors over the last 3 years \cite{9, 10, 11}. These simulations, based on a self-consistent solution of the Lorentz-Maxwell equations for the electromagnetic fields and plasma electrons and ions, all show an efficient initial transfer of laser energy to MeV electrons, which proceed virtually unhindered through the target and beyond. A large charge separation is thus rapidly created on the rear side, which then tugs ions away from this surface.

An alternative school of thought argues that most of energetic protons in high intensity interactions must come from the \textit{front} side of the target, a viewpoint supported by experiments performed by the Imperial College group at the Rutherford-Appleton Laboratory \cite{12, 13, 14} and by the Michigan group \cite{15}. The details of the mechanism for the ‘front-side’ scenario are still unclear however: ponderomotively driven charge separation and the associated ion shock formation appears – ac-
According to PIC simulations – to be insufficient by itself to account for the high number, energies and angular distribution of protons observed.

It is generally acknowledged that both mechanisms probably play a role: the real bone of contention is which one dominates for a particular laser-target configuration. In order to probe the physics of proton acceleration further, recent campaigns by the ICL and Darmstadt groups have been carried out using different target geometries [16, 17]. In particular, a series of experiments with the VULCAN laser using wire targets has added fuel to this debate, as well as throwing up new questions concerning the role of ‘spectator’ targets which appear to radically alter the field distribution in the vicinity of the laser-irradiated region [18].

The purpose of this paper is to report on simulations of ion acceleration from wire targets using the new parallel tree code PEPC (Pretty Efficient Parallel Coulomb-solver). Like the Particle-in-Cell method, this technique also follows the motion of charged particles in self-consistent electric (and in principle magnetic) fields. In contrast to PIC, however, the tree code computes inter-particle potentials and forces directly rather than by employing a grid to mediate the fields via charge and current densities. As will become apparent shortly, this mesh-free, Lagrangian approach lends itself rather well to the kind of open-ended, complex geometry typical of contemporary high intensity laser-matter interactions. After an introductory description of the tree-code-based model in Sections 2–4, simulations of proton acceleration from laser-irradiated wire targets are presented for parameters close to conditions in the recent ICL-RAL experiments.

2. Finite-Size Particle Kinetics with a parallel tree code

The hierarchical tree method on which PEPC is based actually has more in common with molecular dynamics than with particle-in-cell simulation. Briefly, this technique makes systematic use of multipole expansions to reduce the computational effort expended in the force-summation to a time $O(N \log N)$, which for large systems of charges ($N > 10^4$), leads to substantial speed-ups over the conventional $O(N^2)$ algorithm, independently of machine architecture. The technical details of the parallel algorithm used here have been documented elsewhere [19], and we will concentrate on the main components of the physical model in what follows. An earlier plasma tree code (in many respects a sequential forerunner to PEPC) has previously been used to perform microscopic MD simulations of dense, strongly coupled plasmas [20].

In the laser-plasma context of interest here, we use the tree algorithm to model ‘macroscopic’ plasma behaviour in the same spirit as PIC or fluid simulation. This model is based on the ‘Finite-Size-Particle’ (FSP) approach, in which point particles are replaced by spherical clouds, and are allowed to interpenetrate or cross each other. A detailed theoretical basis for this approach was actually laid down over 30 years ago by Langdon, Okuda and Birdsall [21, 22]. An important outcome of their work was to show that the collisionality of FSP plasmas is reduced by orders of magnitude compared to a plasma comprising point particles, so that the plasma parameter $n \lambda_D^2$ is effectively replaced by $n \varepsilon^3$, where $n, \lambda_D$ are the number density and Debye length respectively, and $\varepsilon$ is a measure of the particle size, or cloud radius. This property is implicitly and deliberately exploited in PIC codes, where the smoothing arises automatically by the imposition of a spatial grid, with the result that the above parameters are typically restricted to values: $\varepsilon \approx \Delta x \approx \lambda_D$. Henceforth, we will use the term FSP to mean gridless particle simulation.

The pure FSP method has two immediate advantages over PIC in kinetic plasma simulation: i) collisions are in principle included naturally through the choice of $\varepsilon/a$, where $a = n^{-1/3}$ is the average interparticle spacing, and do not need to be patched back into the code in an ad hoc (and usually expensive) fashion [23]; ii) there are no geometrical restrictions on the simulation region: fast (laser-accelerated) particles do not have to be artificially absorbed or recycled, and may fly as far as they wish away from the interaction region. This does not preclude the application of periodic or
reflective boundary conditions for special geometries: a fully periodic system for strongly coupled plasmas was developed, for example, in Ref. [24].

The drawback of the model is that it is, for the time-being, purely electrostatic: induced magnetic fields are neglected and no electromagnetic wave propagation is supported. At first sight, this may seem too simplistic to describe the kind of highly relativistic, nonlinear phenomena which prevail in high-energy-density laser-matter interactions. As we shall see, however, this ansatz does in fact allow us to capture the salient features of ion acceleration, including important collisional physics which has evidently been missing from the vast majority of PIC simulations of laser-solid interactions to date.

We now proceed with a ‘formal’ description of the electrostatic FSP model as currently implemented in PEPC. The choice of units is somewhat subtle for macroscopic mesh-free plasma simulation, and contrasts with the microscopic ‘Debye’ system used, for example in Ref.[20]. The base normalizations for time, space, velocity, charge and mass respectively are as follows:

\[
\begin{align*}
  t &= \omega_p^{-1} \tilde{t}, \\
  v &= c \tilde{v}, \\
  r &= c \omega_p^{-1} \tilde{r}, \\
  q &= N_p \tilde{q}, \\
  m &= N_p m_e \tilde{m}.
\end{align*}
\] (1)

The constant \(N_p\) represents the number of physical charges contained within a simulation (macro-) particle, to be determined through the equation of motion, which for a given particle \(i\) with charge \(q_i\) and mass \(m_i\) is given (in cgs units) by:

\[
\begin{align*}
  m_i \frac{du_i}{dt} &= q_i E_j \\
  &= q_i \sum_{i \neq j} \tilde{q}_j \tilde{r}_{ij} \tilde{r}_{ij}^3,
\end{align*}
\] (2)

where \(r_{ij} = r_i - r_j\) is the separation between particles \(i\) and \(j\), and \(u_i = \gamma v_i\) is its proper velocity; \(\gamma = (1 + |u|^2/c^2)^{1/2}\) the relativistic factor. In a tree code, the \(O(N)\) sum over all other particles is replaced by a sum over multipole expansions (expanded here up to quadrupole) of groups of particles, whose size increases with distance from particle \(i\). The number of terms in this sum is \(O(\log N)\), which even after the additional overhead in computing the multipoles, results in a substantial saving in effort for large \(N\).

Rewriting Eq. 2 in terms of the normalized variables (1), we find:

\[
\begin{align*}
  \tilde{m}_i \frac{d\tilde{u}_i}{dt} &= \frac{N_p e^2 \omega_p}{m_e c^3} \tilde{q}_i \sum_{i \neq j} \tilde{q}_j \tilde{r}_{ij} \tilde{r}_{ij}^3, \\
  \end{align*}
\]

which, after adding an external field \(E^p\), and making use of the plasma frequency definition, \(\omega_p^2 = 4\pi e^2 n_e/m_e\) for electron density \(n_e\), reduces to:

\[
\begin{align*}
  \tilde{m}_i \frac{d\tilde{u}_i}{dt} &= \frac{1}{3} \tilde{q}_i \sum_{i \neq j} \tilde{q}_j \tilde{r}_{ij} \tilde{r}_{ij}^3 + \tilde{q}_i E^p(r_i),
\end{align*}
\] (3)

provided we take:

\[
N_p = \frac{4\pi}{3} n_e \left( \frac{c}{\omega_p} \right)^3.
\] (4)

Physically, the constant \(N_p\) is just the number of electrons in a sphere with radius \(c/\omega_p\). Since it has been normalized out, we do not actually need to know \(N_p\) in order to carry out a simulation, although it does provide a convenient conversion factor.
As in classical MD simulation, we cannot use the pure Coulomb law for point charges because of the finite timestep, which will cause some particles to experience large, stochastic jumps in their acceleration, eventually destroying the energy conservation. We therefore modify the force-law in Eq. 2 to include a softening parameter \( \varepsilon \), so that the electric field looks like:

\[
E(r) = \frac{q r}{(r^2 + \varepsilon^2)^{3/2}}, \tag{5}
\]

The effect of the softening parameter is to introduce a cutoff into the potential, and to ensure that \( E(r) \to 0 \) as \( r \to 0 \), which greatly assists numerical stability in the time-integration (or particle-pusher) scheme. Physically, we no longer have point charges, but rather charge clouds with a smooth charge density. It is instructive to compute the latter by applying Gauss’ law to (5), giving:

\[
\rho(r) = \frac{3q\varepsilon^2}{4\pi(r^2 + \varepsilon^2)^{5/2}}. \tag{6}
\]

Using the same normalisations as before, and taking \( \rho = en_0\tilde{\rho} \), where \( n_0 \) is some number density to be determined, we find:

\[
n_0\tilde{\rho}\tilde{r} = n_e\frac{\tilde{q}\varepsilon^2}{(\tilde{r}^2 + \varepsilon^2)^{5/2}}. \tag{7}
\]

To simplify this expression, we choose \( n_0 = n_e \), or \( \tilde{n}_e = 1 \). Charge assignment is then straightforward: the total charge contained within a cuboid volume \( V = x_L \times y_L \times z_L \) (in normalized units) is

\[
Q = \sum_i q_i = \tilde{\rho}_0 V = N_e Q_s,
\]

where \( N_e \) is the total number of simulation electrons and \( Q_s \) is the macro-charge carried by them. Since the initial density \( \tilde{\rho}_0 = -\tilde{n}_e = -1 \), we simply have

\[
Q_s = -\frac{V}{N_e}. \tag{8}
\]

3. Target preparation (particle loading)

Assigning charges \( Q_s \) and \(-Q_sZ\) to the electrons and ions respectively, and masses \( M_e^s = |Q_s|, M_i^s = A|Q_s| \), where \( Z \) and \( A \) are the atomic number and mass, sets up a macroscopic plasma system whose internal dynamics is governed solely by Equation 3. Before we can proceed, however, we must pay some attention to its initial spatial and thermal configuration. Whereas a PIC code can be fairly easily initialised through a ‘quiet start’ – an orderly placement of particles in phase space – the FSP model suffers the same kind of pitfalls encountered in classical MD simulation, such as: i) strong initial heating resulting from the system being out of equilibrium at \( t = 0 \), and/or ii) persistent drift currents and oscillations due to localised random concentrations of ion charge.

In the present work, these problems are resolved by a two-step ‘target preparation’ phase. First, ions are forced into a quasi-crystalline structure bounded by the target geometry (which could be, for example: cuboid, wedge-shaped or cylindrical). This is efficiently achieved by allowing the ions to interact via an artificial Lennard-Jones-type potential (the Coulomb interaction having been switched off), thus collectively seeking out a spatial configuration such that the mean distance to each nearest-neighbor is maximised [25]. The wire targets of the present investigation are constructed from cylinders of length \( H \) and radius \( R \), as depicted in Fig.1. The laser is focussed either at the midpoint along the \( z \)-axis or with some offset \( z_0 \).
Next, electrons are placed close to the ions (assuming $Z = 1$) with a velocity randomly selected from a Maxwellian distribution with temperature $T_e$. The whole system is then allowed to relax with the Coulomb force-law reinstated and with the additional thermodynamic constraint that $T_e = \text{const.}$ [20]. This allows the system to seek out its own minimum potential energy while maintaining the temperature desired for the actual simulation.

The end result, arrived at after a few plasma periods, is a configuration with well-defined boundaries, quasi-uniform initial density and minimum potential energy. The same potential energy $U_P$ can also be reached by forcing total energy conservation ($U_P + U_K$), but only at the expense of increasing the electron temperature to some unpredictable value $\gg T_e = 100\,\text{eV}$, as demonstrated in Fig.2a).

It is important to note that unlike in conventional explicit PIC codes, the FSP model does not suffer from numerical heating associated with the grid instability (after all, there is no grid here!). The initial heating seen in Fig.2a) is physical, not numerical: the total energy (central line) is conserved – relative to the kinetic or potential energy values – to better than 1%.

The temperature-clamped system ($\omega_p t = 0 \rightarrow 20$) in Fig.2b) remains in thermal equilibrium when allowed to evolve in the absence of external fields. Note that in this case the potential energy ends up over 8 times larger than the kinetic energy ($\omega_p t = 20 \rightarrow 40$), a situation normally associated with strongly coupled plasmas. For a charge-cloud plasma, however, the relevant parameter is $N_c = 4\pi/3(\varepsilon/a)^3$, rather than $N_D = 4\pi/3(\lambda_D/a)^3$, where $a$ is the interparticle spacing. Although we still have to take some care over the choice of these parameters, the FSP model provides an effective means of modelling plasmas with finite, variable collisionality.
4. Laser model

Because wave propagation within the plasma is not yet supported by this model, the laser is incorporated by a ponderomotive source term, phase-matched to the instantaneous critical density surface at the plasma edge – Fig.3. The appropriate amplitude and phase of the standing wave set up at the interface $x = x_c$ is determined by the solution of the Helmholtz equation for a normally incident, s-polarized plane wave on a step profile. Assuming zero absorption, this solution yields the following electric field:

$$E_z = 2 E_L \cos \omega t \begin{cases} \sin(kx' + \phi), & x' < 0 \\ \sin \phi \exp(-x'/l_s), & x' \geq 0 \end{cases}$$

where $\tan \phi = -kl_s$, $x' = x - x_c$ and $l_s = c/\omega_p$ is the collisionless skin depth. This field is assumed to maintain the above time dependence $\sim \cos \omega t$, which, after dropping the prime from the variable $x$ (henceforth taken relative to the vacuum-plasma boundary) leads to the following expression for the $x$-component of the $v \times B$ force:

$$f_p^x = v_z B_y = -E_z \frac{\partial E_z}{\partial x} = 2E_L^2 \sin^2 \omega t \begin{cases} k \sin[2(kx' + \phi)], & x' < 0 \\ -\frac{2}{l_s} \sin^2 \phi \exp(-2x'/l_s), & x' \geq 0 \end{cases}$$

Figure 3: Ponderomotive laser model. A standing wave solution for the laser is applied at the plasma-vacuum interface, giving rise to the intensity pattern $E_z^2$ (dotted curve) and a ponderomotive force (solid curve).

Note that unlike $E_z$, the ponderomotive force changes sign with $x$ but not $t$: here it comprises an oscillating component at $2\omega$ plus a DC component (the actual ponderomotive part), both of which always point in either the positive or negative $x$-direction, as depicted in Fig. 3.

To make this laser model viable for relativistic interactions, it needs two further modifications: a correction for large quiver amplitudes, $a_0 = eE_L/m\omega c > 1$, and a radial dependence to allow for a finite focal spot. The expression used in the code therefore takes on the following form:

$$f^p = -\nabla \gamma,$$

where

$$\gamma = \left(1 + \frac{\Psi}{2}\right)^{1/2},$$
\[
\Psi = 4a_0^2 X^2(x) R(r) T(t),
\]

\[
X(x) = \begin{cases} 
\sin \chi, & x < 0 \\
\sin \phi \exp(-x), & x \geq 0 
\end{cases}
\]

\[
R(r) = \begin{cases} 
\cos^2 \left( \frac{\pi r}{4\sigma} \right), & r \leq 2\sigma \\
0, & r > 2\sigma 
\end{cases}
\]

\[
T(t) = \sin^2 \left( \frac{\omega t}{\omega_p} \right).
\]

The above expressions are written in terms of normalized variables, hence the skin depth, \( \tilde{l}_s = 1 \). For readability, we have retained an explicit frequency ratio \( \omega/\omega_p \), so that the phase factors become: \( \phi = -\tan^{-1}(\omega/\omega_p), \chi = \omega_p x + \phi \). The radial coordinate \( r = (y^2 + z^2)^{1/2} \) is taken relative to the center of the focal spot. The latter has a \( \sin^2 \) form rather than a Gaussian one in order to create a sharp radial cutoff at \( 2\sigma \) (\( \sigma \) is the FWHM). This is found to give a more physically reasonable modelling of profile deformation, avoiding penetration of the low-intensity wings in the overdense plasma, which would tend to occur for a Gaussian focal spot.

The longitudinal and radial field components are finally given by:

\[
E^p_x = \frac{\partial \gamma}{\partial x} = \frac{a_0^2}{\gamma} R(r) T(t) \begin{cases} 
\frac{\omega}{\omega_p} \sin 2\chi, & x < 0 \\
-2 \sin^2 \phi \exp(-2x), & x \geq 0 
\end{cases}
\]

\[
E^p_y = \frac{\partial \gamma}{\partial y} = \frac{a_0^2}{\gamma} T(t) X^2(x) \begin{cases} 
-\frac{\pi y}{4\sigma r} \sin^2 \theta, & r < 2\sigma \\
0, & r \geq 2\sigma 
\end{cases}
\]

where \( \theta = \pi r/4\sigma \).

This obviously simplistic model cannot hope to match the rich array of physical phenomena accessible through a full solution of Maxwell’s equations. Nonetheless, when combined with a rudimentary density-tracking algorithm to monitor the position of the critical surface, it does serve rather well in reproducing some of the main features of hot electron generation and pondermotive ion dynamics. We illustrate this with a test problem in slab geometry, namely collisionless shock-formation through pressure imbalance: a hole-boring simulation. Balancing continuity and momentum at the critical surface (laser reflection point) gives the well-known formula for the recession velocity [26, 27]:

\[
\frac{u_h}{c} = \left( \frac{Zm_e}{m_i} \frac{n_e}{n_c} \left( \frac{2 - \eta}{4} \right) a_0^2 \cos \theta \right)^{1/2},
\]

where \( a_0 \) is the normalised laser amplitude or quiver velocity, \( \eta \) is the absorption fraction of laser energy coupled to the plasma and \( \theta \) is the angle of incidence.

A simulation to verify this behaviour was set up using a plasma block with dimensions \((60 \, c/\omega_p \times 150 \, c/\omega_p \times 150 \, c/\omega_p)\) and initial electron and ion temperatures of \( T_e = 5 \, keV \) and \( T_i = 0 \) respectively. The other simulation parameters were: \( a_0 = 2.7, m_i/Zm_e = 1836, \eta = \theta = 0, n_e/n_c = 4 \). A total of \( 1.44 \times 10^6 \) particles were used with effective size \( \varepsilon = 2 \) and average (ion) spacing \( a = 0.23 \), giving a smear factor \( N_c = 4\pi/3(\varepsilon/a)^3 = 2700 \), placing the simulation well into the collisionless limit. Figure 4 shows successive lineouts of the ion density along the laser axis, from which we deduce a hole-boring velocity \( u_h/c = \Delta x_c/\Delta t = 15/700 \approx 0.022 \). This is in good
agreement with the theoretical value given by (13) of $u_h/c = 0.02$, giving us some confidence in the ponderomotive laser model described above.

![Figure 4](image)

**Figure 4:** Ion density profile sequence along laser axis as a result of ponderomotive laser pressure.

### 5. Wire simulations

The laser-wire simulations were set up with a quasi-neutral plasma with $n_i = n_e = 4n_c$ and initial electron and ion temperatures $T_e = 200 \text{ eV} - 1 \text{ keV}$ and $T_i = 0$ respectively, configured in a cylinder with radius $R \simeq 1 - 4 \mu m$ and height $H \simeq 10 - 16 \mu m$, as depicted in Fig. 1. Up to $3.2 \times 10^6$ simulation particles were used, with a mass ratio $m_i/m_e = 1836$ and ion charge state $Z = 1$. The laser wavelength is assumed to be $1 \mu m$ the spot size is $0.5-1 \mu m$ FWHM, or $12-24c/\omega_p$ generally chosen so that the focal spot just fits within the wire diameter ($25c/\omega_p$). The pulse is turned on over 5 laser cycles and then kept at constant intensity for around 300 fs, or until the wire is burned through, at which point the standing wave ansatz is no longer reasonable. These parameters are still some way short of the experimental conditions, in which wires with diameters of $20 \mu m$ were irradiated by a 1 ps laser focused to $20 \mu m$. The total laser energy converted into hot electrons is therefore 100-1000 times less in the simulations than in the experiment, so that we concentrate on identifying trends in the interaction behaviour rather than attempting a 1:1 quantitative comparison.

Scaling up the simulations is non trivial because the statistics deteriorate rapidly: doubling the wire radius alone results in a $4 \times$ larger plasma volume $V = \pi R^2 H$, and therefore requires $4 \times$ the number of particles to maintain the same particle macro-charge $Q_s$ (keeping $n_e/n_c$ constant) and inter-particle spacing (or collisionality, unless $\varepsilon$ is adjusted as well). These parameters determine the maximum timestep permitted for numerical stability and hence the total simulation time required. A ‘minimal’ simulation with $N_e + N_i = 1.44 \times 10^6$ particles took 50 hours on 16 CPUs of the Jülich IBM p690+ Regatta. The largest simulation considered in the present work, a $4\mu \times 16\mu$ wire with $3.2 \times 10^6$ particles, took over 100 hours on 32 CPUs. For convenience we include a summary of the simulations referred to here in Table 1.

We begin our study by examining some general aspects of the laser-wire interaction for Run C; the $2 \mu m (50 c/\omega_p)$-radius wire in the table. The large-scale electron and ion dynamics can be traced in Fig.5, which shows a sequence of ion density slices in the $x - z$ plane while the laser is incident. A number of features in a) and b) are immediately apparent: the strong bow-shock structure resulting from the ponderomotive push of the laser; the characteristic low-density ion blowoff back towards the laser; the hot-electron current into the target, and ion layers starting to peel off the rear-side
due to hot electrons circulating behind and around the wire. This last effect is the familiar target-normal sheath acceleration (TNSA) mechanism, but in cylindrical geometry, ultimately leading to a disc-like fast ion emission.

From Fig.5 one might conclude that rear-surface protons will dominate the emission spectrum here, yet this is only part of the picture. Inspection of the ion phase space \( p_z - x \) for Run C in Fig.6a) indicates that front-side ions are also accelerated significantly via the ponderomotive shock, some of which have already emerged from the rear surface (at \( x = 50 \)) as a beamlet in the forward direction. The onset of a double-disc structure is apparent in the \( p_z - x \) plot of Fig.6b): the TNSA-ions \((x > 100)\) are beginning to fork at an angle of \(5–10^\circ\) to the laser axis. At this point these ions have energies of \(> 6\) MeV, and are still being accelerated. Also evident from Fig.6b) are the significant blowoff components at \( p_z \approx \pm 0.05 m_c \) from the wire tips, reflecting the fact that the hot electrons have formed a large plume around the wire. Indeed, the electron phase space shows that this plume extends more-or-less symmetrically with a radius of \(\sim 1200 \ c/\omega_p\), or 50 times the initial wire radius. This corresponds to an effective simulation volume of almost \(10^7 \ \mu\text{m}^3\) - a feat which would be difficult to match with a grid-based particle code.

The far-field structure of the ion emission in a more appropriate form for comparison with exper-
Figure 6: Ion phase space at the end of Run C in the laser direction (x-axis): a) longitudinal and b) vertical (p_z - along wire axis) momentum components. The wire is initially located between x = 0 and x = 50.

Figure 7: Angular MeV ion emission at in a) forward and b) backward directions.

To get a feel for how these results scale with laser and target parameters, and to make a connection with the PIC simulations in Refs.([9]-[11]), Table 2 provides a summary of the energy balance statistics for the runs listed in Table 1.

Although this sample of the available parameter space is too small to draw definitive conclusions, some general trends are worth pointing out. First, the maximum ion energy U_{i,max} is clearly correlated to the laser intensity (or I\lambda^2) rather than the total energy. The lower value for the 4 \mu m wire reflects the fact that most of the absorbed energy is either still carried by hot electrons, or has gone into heating a larger bulk of wire material at this time. This is in contrast to the 1 \mu m wires, for which even after 700 \omega_p^{-1} (200 fs), around 3x as much energy has been transferred to the ions than is carried by hot electrons.
Table 2: Energy balance for the runs listed in Table 1. The wire dimensions are expressed in microns to aid identification. $U_a^e$ and $U_a^i$ are the total energies absorbed by electrons and ions respectively at the end of the run; $U_{\text{max}}^i$ is the maximum ion energy; $U_{\text{ave}}^i$ the median ion energy (peak in spectrum).

<table>
<thead>
<tr>
<th>RUN #</th>
<th>Dimensions $R \times H$ (µm)</th>
<th>$I_{19}$</th>
<th>Laser energy $(\text{mJ})$</th>
<th>$U_a^e$ $(\text{mJ})$</th>
<th>$U_a^i$ $(\text{mJ})$</th>
<th>Total absorption $(%)$</th>
<th>$T_h$ (MeV)</th>
<th>$U_{\text{max}}^i$ (MeV)</th>
<th>$U_{\text{ave}}^i$ (MeV)</th>
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<td>21</td>
<td>1.5</td>
<td>4.8</td>
<td>30</td>
<td>0.33</td>
<td>2.5</td>
<td>1</td>
</tr>
</tbody>
</table>

The reason for this enhanced transfer efficiency is not clear at present. Normally, one would expect a smaller-radius wire to favour the TNSA mechanism because the hot electrons have less material to pass through. However, runs A, B and E have a far higher collisionality than C and D, implying a lower mean-free-path for the cold electrons. This in turn leads to inhibition of hot electron transport [29] and correspondingly more pronounced front-side ion acceleration. This can be clearly observed in the ion phase-space of run B in Fig. 8, which shows the front-side ions emerging from the rear side with more than twice the energy than the TNSA-accelerated ions.

Figure 8: Ion phase space for the 1 µm wire in Run B at times a) $t = 600$ and b) $t = 1080$ showing enhanced front-side ion acceleration. The wire is initially located between $x = 0$ and $x = 25$.

A detailed analysis of the physics behind this effect will be presented elsewhere [30]: for the time-being, we compare the ion dynamics in the 1 µm -radius wire with that observed in Fig. 5 for the 2 µm wire of Run C. As before, we show a sequence of ion density isovolumes, but this time consisting of a 1/2-wire vertical slice – Fig. 9. Superimposed on these plots are slices of the instantaneous electron temperature in MeV, showing that while the laser is incident, the hottest electrons are actually confined to the shock region, yet there is also a strong circulation of hot electrons around the wire.

The most striking feature of this simulation is that the entire mid-section of the wire is pushed out by the laser: the beamlet visible in Fig. 9d) has detached itself completely from the wire and continues to propagate away, spreading as it does so. This is reminiscent of 3D PIC simulations double-layer targets in which a proton beam was created from the low-Z coating on the rear-side [31]. By contrast, the main thrust in this case comes unmistakably from the target frontside, even though the beamlet comprises ions which originate from the across the whole wire.

6. Discussion and Conclusion

The simulations presented here demonstrate that high-intensity laser-wire interactions can be effectively modelled with a 3D electrostatic tree code, despite simplifications to the absorption physics and the neglect of self-generated magnetic fields. The disc-like ion emission pattern appears to
Figure 9: Time-sequence of ion density isovolume $n_i/n_e \geq 0.25$ and electron temperature $T_e$ slice in plane of laser incidence for a 1/2 wire-section sliced along the wire $z$-axis – Run B. Times shown are a) $200/\omega_p$, b) $400/\omega_p$, c) $600/\omega_p$, and d) $800/\omega_p$.

originates initially from the cylindrically symmetric charge separation caused by hot electrons circulating around the wire. However, it is not yet clear from the present analysis why two discs (or a double-stripe in the far-field image) emerge, both in forward and backward directions, as in the RAL experiments. One possible explanation is that a large number of hot electrons tend to arc back towards the wire (which gets positively charged during laser irradiation) thus setting up a return current along the wire ($z$) axis from the tips to the focus. Ions exiting the wire surface will therefore be pulled at a slight angle to the target normal, in the $\pm z$-direction for ions above and below the laser focal plane respectively.

Return current effects also appear to be responsible for the development of disc-like emission in the small-radius wire simulations. In Run B for example, radial components develop in the aftermath of the burn-through phase, albeit at somewhat lower energies (0.5-1.5 MeV) in this case. Whether

Figure 10: Electron circulation along the wire axis. The arrow length is proportional to the electron momentum.
this effect persists as the wire radius and laser energy is scaled up will be addressed by future work.

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