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Tracer spectroscopy diagnostics of doped ablators in inertial confinement fusion experiments on OMEGA

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I. INTRODUCTION

The efficient compression of an indirect-drive inertial confinement fusion (ICF) fuel capsule depends sensitively on the coupling of the hohlraum x-ray drive to the capsule ablator. In order to control the deposition of energy in the ablator, and thus the shock compression of the fuel, mid-Z ablator dopants such as germanium, bromine, or copper are often added to the low-Z ablator material. The increased opacity provided by ablator dopants serves to change the radiation–matter coupling as well as to shield the fuel from preheat. It is desirable for experimentalists to be able to monitor the effects of ablator dopants on the physical conditions within a capsule ablator exposed to a hohlraum x-ray drive. In the past, the effects of ablator dopants have been studied via imaging of imploded capsules and via shock breakout or burnthrough and foil motion measurements. In these cases, the integrated effects of the radiation hydrodynamics over the course of a capsule implosion, burnthrough, etc., are diagnosed. In this paper we report on experiments and associated modeling undertaken to study the effects of dopants at a specific location in the interior of an ablator sample. This technique, which involves point-projection spectroscopy of a thin tracer layer, could be generally applied to the study of material samples exposed to strong radiation fields.

It has previously been shown that absorption spectroscopy can be used to study energy transport and material properties such as opacities, and from witness plates mounted on hohlraums has also been employed as a diagnostic of physical conditions in hohlraum plasmas. Additionally, Chenais-Popovics et al. have shown that chlorine absorption spectroscopy, backlit by bismuth M-shell emission, is a useful diagnostic of laser-produced plasmas.

We have combined these ideas in developing a program to study the time-dependent conditions within ablator samples mounted on halfraums at the OMEGA laser facility. We use thin (submicron) chlorine-bearing tracer layers buried at known depths within these ablator samples to diagnose the ionization state (and thus localized plasma conditions) as a function of time at depths of several microns in these samples. Ultimately, these data reflect the Marshak wave velocity in the ablator. Modeling of the experiments using view factor, radiation hydrodynamics, and spectral synthesis codes is an important component of the program, allowing us to infer physical conditions from spectral measurements and giving us greater confidence in making comparisons between doped and undoped ablator samples.

In the experiments we mounted planar ablator samples
on the back ends of halfraums, which we shot on OMEGA in a standard configuration. We used planar samples for the sake of simplified hydrodynamics, and model the radiation drive onto the samples, which, in any case, is very similar to what a spherical capsule in a hohlraum sees. At this point we do not have plans to attempt this measurement in an actual spherical pellet, although this would be possible as long as the backlighter x rays were not too attenuated by the material in the pellet. The drive modeling can, in principle, be constrained by absolutely calibrated DANTE measurements of the halfraum wall reemission. However, we have found that in general, the radiation field inside the halfraum must be modeled, as it varies spectrally with location as well as time.

The planar ablator samples are specially fabricated to have a thin (roughly 0.4 μm) NaCl layer about 5 μm below the surface. This chlorine-bearing tracer gives rise to a Kα absorption signal just below 2800 eV seen in absorption against a laser-produced bismuth plasma generated by diverting several beams onto a Bi foil hung off the halfraum in front of the laser entrance hole (LEH). See Fig. 2 for an image and model of a target. We used the LLE x-ray spectrometer (LXS) coupled to a streak camera to make time-dependent measurements of the tracer spectral signal.

Our modeling, which is described in detail in Sec. III, shows that the Cl Kα features turn on when the Marshak wave reaches the tracer layer and heats the tracer to roughly 100 eV. The shock wave driven in front of the Marshak wave does not heat the tracer enough to ionize up to the L-shell, which is required for the production of Kα absorption. If the effect of ablator dopants is to reduce the Marshak wave velocity, as is expected, then one should see the tracer signal turn on earlier in an undoped ablator sample compared to a doped sample (in cases where the tracer depth is the same in the two samples). This is demonstrated by the model spectra time series shown in Fig. 3. A second-order effect that one can also observe is the time-dependent ionization balance in the tracer, which reflects changing temperature and density conditions at the location of the tracer as the Marshak wave passes through.

II. EXPERIMENTS AND DATA

We performed a series of shots on the OMEGA laser as part of the National Laser Users Facility (NLUF) program, comparing doped and undoped ablator samples on scale-1 halfraums mounted on the P6-P7 axis. We used 15 cone 2 and cone 3 beams, pointed at the center of the LEH (giving laser spot positions on the wall ranging from 480 to 890 μm from the LEH plane), for a total of approximately 6.7 kJ delivered in a 1 ns square pulse. Unfolded DANTE measurements indicated peak wall reemission temperatures of 182 eV, with the two shots we compare here both peaking at this
temperature, and having very similar time histories. Note that in the next section we discuss the halfraum radiation field modeling and show that the calculated drive temperatures are within the DANTE error bars. We diverted six additional beams onto the bismuth backlighter foil, with staggered timing, to produce a relatively temporally smooth backlighter spectrum.

The ablator samples were manufactured at General Atomics using glow discharge polymerization (GDP) to lay down a layer of plastic (doped or undoped) several tens of microns thick. This wafer of plastic was then coated with sodium chloride (NaCl) to a thickness of approximately 4000 Å. On top of this, another, thinner layer of plastic was deposited in the same GDP chamber as the first layer. The sample was then cut into roughly ten pieces and each was characterized in terms of its density and composition. The undoped samples were typically 54.6% H, 42.2% C, and 3.3% O with a density of 1.11 g cm$^{-3}$. The germanium-doped samples had 55.4% H, 38.6% C, 4.2% O, and 1.8% Ge with a density of 1.28 g cm$^{-3}$. These are the compositions used to calculate the opacities shown in Fig. 1. We discuss two different shots in this paper. In shot 19 526 the tracer layer was 6.3 μm from the surface of the undoped sample and in shot 19 528 the tracer layer was 4.1 μm from the surface of the doped sample.

In Fig. 4 we show time-dependent spectra measured on these two shots. Although there is a fair amount of emission structure in the backlighter spectrum, absorption signals from at least three ionization stages of chlorine can be seen turning on after the backlighter signal itself, and with the lower ionization stages appearing before the higher ones (this last effect is more apparent in the undoped sample, shot 19 526). The turn-on time is slightly earlier in the undoped sample than in the doped sample, which is consistent with the qualitative expectations of the effect of the dopant on the Marshak wave velocity. Note the Be-like feature may appear weakly in the 300 ps lineout in the undoped sample (top panel), and by 400 ps the Be-like and Li-like features are strongly present.

In the doped sample (lower panel of Fig. 4), the spectral signal becomes visible at 400 ps, and the ionization balance
in this sample is weighted toward higher ion stages than in the undoped sample. The spectral signal is weaker overall in the doped sample because of the effect of the added opacity from the germanium dopant. It should be borne in mind that the tracer in the doped sample in the experiments is 2 µm closer to the surface than the tracer in the undoped sample, so this modest time lag between the doped and undoped samples represents a significantly lower Marshak wave velocity in the germanium-doped plastic than in the undoped plastic. In other words, the tracer spectral signal turns on later in the doped sample, despite the fact that the tracer is located more than 30 percent closer to the surface in that sample.

III. MODELING

To understand the experimental results, and to infer the physical conditions in the halfraum and the ablator, we perform three stages of numerical modeling: (1) View-factor calculations, which provide the temperature structure (both spatial and temporal) within the halfraum as well as the incident radiation flux as a function of time on all surfaces, including the ablator sample; (2) hydrodynamics simulations of the interaction between the time-dependent radiation field in the halfraum and the ablator sample; and (3) spectral post-processing of the hydrodynamics calculations. Associated atomic modeling is carried out to provide inputs at stages (2) and (3).

We used the VISRAD view-factor code\textsuperscript{13} to do the first step in the modeling. This code has been used to model both laser hohlraum and Z-pinch experiments, and accurately reproduced data from similar, previously published OMEGA hohlraum experiments.\textsuperscript{14,15} VISRAD is a time-independent code that assumes albedos and x-ray conversion efficiencies for each surface in a simulation, which are assumed to emit as blackbodies. Temperature-dependent albedos are calculated for the gold halfraums separately in a hydrodynamics code, using detailed opacities, the recorded beam power profiles are used, and with a value for the (time-dependent) x-ray conversion efficiency (XCE) assumed, this modeling is fully constrained, with no adjustable parameters. This modeling procedure has been quite successful in reproducing the observed conditions in laser hohlraums.\textsuperscript{15}

In Fig. 5 we show the good match obtained between VISRAD view-factor modeling of the wall reemission in earlier OMEGA hohlraum experiments and DANTE measurements. We also show in that same figure the midplane drive temperature from the same modeling and note that it is significantly different than the wall reemission profile. This difference in the DANTE and drive temperatures makes sense because DANTE sees only hot spots and wall reemission, while a package at the midplane of a hohlraum sees hot spots and wall but also sees the cold LEHs as well. The same will be true for a package on the end of a halfraum.

The percentage reduction in temperature at the package versus that seen by DANTE will depend on the relative distances of spots and LEH from the package, which differ between hohlraum and halfraum geometry. These results demonstrate the need for modeling of the drive conditions on an experimental package, even when DANTE or other wall reemission measurements are made. And they also demonstrate, through the good agreement between the DANTE measurements and the view-factor modeling, the accuracy of the modeling technique.

In Fig. 6 we show the results of similar view-factor modeling for the OMEGA experiments we report on in this paper. This modeling was carried out in exactly the same way as that described above, using the same temperature-dependent gold albedo and time-dependent XCE (0.55 after 200 ps, ramping up from 0.2 initially) as was used to successfully
match the data in the earlier hohlraum experiments. Our calculations also predict the wall reemission temperatures seen by DANTE, which are within the error bars of the absolutely calibrated DANTE measurements.

The time-dependent x-ray spectrum incident on the ablator sample calculated from the view-factor modeling was next used as input to a hydrodynamics simulation performed with HELIOS. This one-dimensional, planar, Lagrangian simulation used multigroup opacities for both the plastic and the tracer calculated for the purpose. SESAME equations of state were used, and scattered laser light onto the ablator sample was also included. The scattered laser intensity was calculated in the VISRAD simulations to be roughly 100 TW cm$^{-2}$ for the first 200 ps, falling off rapidly afterward. In Fig. 7 we show representative temperature profiles from these simulations. The shock wave and radiation wave can easily be seen, with the slower velocity in the doped sample apparent in these time series. We note here that we shifted the radiation drive calculated with VISRAD by 100 ps to account for uncertainties in the relative zero points of the DANTE timing and the LXS timing. The apparent turn-on of the LXS-measured spectra should be somewhat later than the DANTE turn-on time, as the LXS is sensitive only to hard x rays (between 2 and 3 keV).

Ultimately, we synthesized the time-dependent chlorine absorption spectra via a series of time-independent calculations of the excitation/ionization and radiation transport through the hot (and cold) plasma derived from the HELIOS simulations. We custom compute atomic models for the ablator and tracer materials, which are used as inputs to the spectral calculations, which we carried out with the Spect3d code. We use over 1100 levels in our chlorine model, and assume local thermodynamic equilibrium (LTE) for the calculations we present here [using a time-independent collisional-radiative equilibrium (CRE) model gave very similar results]. We present spectral simulations, in Figs. 8 and 9, for the two samples shot. We stress that the earlier turn-on times seen in the doped sample in these simulations are entirely due to the smaller tracer depths. In these figures, we again show the data lineouts for comparison.

These spectral simulations show that the Be-like and higher ion stages become visible slightly earlier in the doped sample than in the undoped sample (400 ps versus 500 ps). In both simulations a progression from lower to higher ion...
IV. DISCUSSION

Despite the noisiness of the spectral data (due to emission spectral structure in the Bi backlighter), we have detected the tracer absorption signals in the two samples. The detection is weaker in the doped sample, due to the lower signal-to-noise in the spectrum transmitted through the higher-opacity material. The tracer turn-on times observed in the undoped sample are slightly earlier than in the doped sample, especially if one looks at the He-like feature, which is relatively robust in the 400 ps lineout for the doped sample, but only just beginning to appear in the 500 ps lineout in the undoped sample.

It seems likely that our nondetection of ion stages lower than Be-like can be understood in terms of the shape of the underlying backlighter pseudocontinuum, which is significantly weaker at energies below about 2750 eV, as has been seen in other experiments. The tracer turn-on time observed in the undoped sample is somewhat earlier than our modeling predicts, even under the most conservative assumptions. The appearance of Be-like absorption at 300 ps is roughly 200 ps earlier than in the simulations. Taken at face value, the effect of the germanium dopant in the experiments is thus even more dramatic than what is expected from our calculations.

It is conceivable that the chlorine tracer was closer to the surface of the ablator samples than we assumed, either due to uncertainty in the fabrication and characterization process or due to the possible migration of chlorine from the tracer salt into the plastic after fabrication. If so, this would seem to have affected the undoped sample more than the doped sample, based on the fact that it was the undoped sample for which there was a discrepancy in the calculated and measured tracer turn-on times. In the future, fabrication of tracer layers using chlorinated plastic rather than salt could be attempted. It is also possible the spatial nonuniformity in the tracer salts (due, for example, to crystallization) could have caused the tracer absorption signal to be weaker than it might otherwise have been. Using chlorinated plastic tracer layers, rather than salt, in the future might therefore also improve the signal strength.

In any case, if we infer the Marshak wave velocity from the experimental results we can tentatively conclude the following: If we take the turn-on time of the tracer signal to be 300 ps in the undoped ablator and 400 ps in the doped ablator (for the appearance of Be-like absorption), then the Marshak wave moves through an areal mass of $7 \times 10^{-4} \text{g cm}^{-2}$ in 300 ps for the undoped ablator and $5 \times 10^{-4} \text{g cm}^{-2}$ in 400 ps for the germanium-doped ablator. Expressed in units of areal mass per time, the effect of the ablator dopant is thus to reduce the Marshak wave velocity by almost 50% ($2.3 \times 10^{-3} \text{g cm}^{-2} \text{ns}^{-1}$ versus $1.3 \times 10^{-3} \text{g cm}^{-2} \text{ns}^{-1}$).

Finally, it is clear that the detailed experimental modeling is crucial for interpreting the relatively complex data from these experiments. And, that in conjunction with such modeling, point-projection absorption spectroscopy of buried tracer layers is a promising—but challenging—diagnostic technique for inferring the localized properties of plasmas in ICF environments.

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15. J. J. MacFarlane, astro.swarthmore.edu/prism/visrad-benchmark.pdf; see EPAPS Document No. E-PHPAEN-11-911405 for “Visrad benchmark calculation: OMEGA Hollraum radiation temperatures.” A direct link to this document may be found in the online article’s HTML reference section. The document may also be reached via the EPAPS homepage (http://www.aip.org/pubservs/epaps.html) or from ftp.aip.org in the directory /epaps/. See the EPAPS homepage for more information.


18. See EPAPS Document No. E-PHPAEN-11-911405 for a color version of the lower panel of Fig. 2. A direct link to this document may be found in the online article’s HTML reference section. The document may also be reached via the EPAPS homepage (http://www.aip.org/pubservs/epaps.html) or from ftp.aip.org in the directory /epaps/. See the EPAPS homepage for more information.