X-Ray Continuum as a Measure of Pressure and Fuel–Shell Mix in Compressed Isobaric Hydrogen Implosion Cores

Introduction

Measurements of continuum x-ray emission from the central hot spots of inertial confinement fusion (ICF)¹ implosions at stagnation can be directly related to hot-spot conditions using the relatively simple dependence of continuum spectral emission rates on temperature and density or pressure. Since thermonuclear ignition and high energy gain are the goals of ICF,^{2,3} one would naturally look to neutron yield as the primary measure of implosion performance. The benchmarks of progress toward ignition, or toward implosion performance that scales to ignition with higher drive energy,⁴ however, are specified in terms of core conditions at peak compression.⁵ Short of ignition, neutron yield and x-ray emission measurements can be used in similar ways to infer hot-spot conditions. The current strategy is to achieve high-temperature central hot spots within fuel shells compressed at low entropy to high areal densities. A key measure of near-ignition performance in ICF, through the Lawson criterion,⁶ is the hot-spot pressure.⁷ Consequently, a direct relationship between the hot-spot pressure and the measured hot-spot x-ray continuum emissivity, based as little as possible on prior assumptions about hot-spot temperature profiles, normalization to simulations, etc., would be very important.

At a sufficiently high spectral energy, typically $h\nu > 3$ keV for cryogenic implosions on the 60-beam OMEGA Laser System,⁸ the imploded cores are optically thin and the x-ray measurements are a direct measure of the emissivity, free of absorption and other transport effects. With instrument spectral responses narrowed to energies matched to the anticipated temperature of the hot spot, as will be described below, the free-free (FF) emissivity⁹ of hot-spot hydrogen scales as the square of the hot-spot pressure and is nearly independent of the temperature. The simple pressure dependence of the emissivity, the isobaric state of the hot spot at stagnation,^{10,11} and the known temperature-density scaling of the neutron-production rate explain a simple scaling behavior of the x-ray yield as a constant power of the neutron yield over a factor-of-10 range in neutron yield in an ensemble of similar targets imploded with a variety of laser pulses over a broad range of shell isentropes.

This is a quantitative prediction based on our understanding of isobaric hot spots that has been confirmed with measured x-ray and neutron yields. In an unstable implosion, a trace (above ~10 ng) amount of shell material mixed into the hot spot can increase the x-ray emission measurably because of the much higher emissivity per atom of carbon, without affecting the neutron yield significantly. Using the x-ray yield expected from the neutron-yield scaling as a point of reference, the excess x-ray emission and the known FF and free-bound (FB) emissivity⁹ of carbon provide a measure of this "fuel–shell" mix mass. This mix-mass estimate is similar in some respects to recent measurements of mix mass in National Ignition Facility (NIF)¹² implosions based on the ratio of the x-ray and neutron yields.¹³

The pressure profile within an imploded core at the time of peak emission can be obtained from the emissivity profile of the object, and the emissivity profile can be obtained from its projection recorded on an image plane by an imaging device. The fundamental quantity of radiation is the specific intensity $I(\vec{x}, \hat{\Omega}, \nu)$, which is the amount of radiation energy per unit of time arriving at position \vec{x} in space, per unit of area within an infinitesimal area element at this point, oriented normal to the propagation direction given by the unit normal vector $\hat{\Omega}$, per unit spectral range within an infinitesimal interval of frequency, centered at the frequency ν , and traveling within an infinitesimal cone of solid angle, per steradian, centered on the direction $\hat{\Omega}$. We will write it as I_{12} for short. The timeindependent equation of transfer governing the change dI_{ν} in the specific intensity of radiation propagating an infinitesimal distance ds along the direction $\hat{\Omega}$ is

$$\frac{\mathrm{d}I_{\nu}}{\mathrm{d}s} = \frac{\varepsilon_{\nu}}{4\pi} - \kappa_{\nu}I_{\nu},\tag{1}$$

where the interaction of radiation and matter is described entirely in terms of the plasma emissivity ε_{ν} and the opacity κ_{ν} of the matter.¹⁴ The emissivity specifies the energy per unit of time that is emitted per unit of volume isotropically into all directions within an infinitesimal interval of frequency centered at the frequency ν . The opacity is the fraction of the specific intensity absorbed per unit distance of propagation. We assume that all radiation of interest propagates at the speed of light *c* without refraction or dispersion and that any photon scattering involving a change of frequency or direction is accounted for effectively by a combination of absorption and emission included in ε_{ν} together with κ_{ν} .

In the optically thin limit, there is only emission and no attenuation, and the solution to the equation of transfer is the path integral

$$I_{\nu}(\vec{x}) = \int_{-\infty}^{x} \frac{\varepsilon_{\nu}(\vec{r})}{4\pi} \, \mathrm{d}\vec{s}. \tag{2}$$

The integration variable \vec{s} is a point along a straight-line photon path arriving at a point \vec{x} on the image plane and \vec{r} is the position of that point on the path relative to an arbitrary fixed point of reference within the emitting object. The integration path is a straight line arriving at the observer position from arbitrarily far away, indicated symbolically as "-∞," but only points along the path within the emission source contribute to the integral. The imaging device selects from the set of all paths $\hat{\Omega}$ ending at any one \vec{x} so that the image represents a useful projection of the emissivity profile. For the purposes of this discussion, we assume that the imaging device records an orthographic projection of the source, such that the direction $\hat{\Omega}$ of all paths is the same. In spherical geometry, Eq. (2) gives the specific intensity as an Abel transform¹⁵ of the emissivity profile. For example, a spherically symmetric emissivity distribution $\varepsilon_{\nu}(r)$ produces the specific intensity

$$I_{\nu}(x) = \frac{1}{2\pi} \int_{x}^{\infty} \frac{\varepsilon_{\nu}(r)r}{\sqrt{r^2 - x^2}} \,\mathrm{d}r,\tag{3}$$

and the emissivity distribution can be recovered from the inverse Abel transform of the imaged intensity,

$$\varepsilon_{\nu}(r) = -4 \int_{r}^{\infty} \frac{\mathrm{d}I_{\nu}(x)}{\mathrm{d}x} \frac{\mathrm{d}x}{\sqrt{x^2 - r^2}}.$$
(4)

The point x = 0 on the image plane is the projection of the center of the radial emissivity distribution at r = 0, and we assume that all geometrical and optical effects of an actual camera, such as magnification, etc., have already been taken into account.

If the emissivity is a known function of pressure alone, the radial pressure profile of a spherical hot spot can be inferred from the emissivity profile extracted from the imaged intensity using Eq. (4). If, in addition, the hot spot is isobaric, the pressure is constant throughout the hot spot out to its outer radius Rand the emissivity will be constant within this radius; we find using Eq. (4) that the intensity profile of its image is elliptical:

$$I_{\nu}(x) = \frac{\varepsilon_{\nu}(0)}{2\pi} \sqrt{R^2 - x^2}.$$
 (5)

The integral of the specific intensity given by Eq. (2) over the image plane gives the total emitted power per unit solid angle at the image plane:

$$\int I_{\nu}(\vec{x})d^{2}\vec{x} = \iint_{-\infty}^{\infty} \frac{\varepsilon_{\nu}(\vec{r})}{4\pi} \,\mathrm{d}\vec{s}d^{2}\vec{x} = \frac{1}{4\pi} \int \varepsilon_{\nu}(\vec{r})d^{3}\vec{r}.$$
 (6)

Assuming the hot spot radiates isotropically, the total spectral power is obtained by applying a factor of 4π . In the case of the flat emissivity profile of an isobaric core, this gives

$$P_{\nu} = \frac{4\pi}{3} R^3 \varepsilon_{\nu}(0). \tag{7}$$

The isobaric sphere, then, possesses a remarkable simplicity: All quantities pertaining to emission can be described in terms of a single radius and a single emissivity that depends on one parameter—the pressure. In the following sections, departures of the emissivity from pure pressure dependence, hot spots that are not strictly isobaric, "fuel–shell mix," and other complications will be considered.

Pressure Dependence of X-Ray Emission from Isobaric Hot Spots

The emissivity of an imploded hydrogen hot spot of a cryogenic implosion is almost entirely the result of FF emission from hydrogen. The expression for the FF emissivity of a hot plasma of fully stripped ions at photon frequency ν , temperature *T*, electron and ion densities n_e and n_i , respectively, and average nuclear charge squared $\langle Z^2 \rangle$ is

$$\varepsilon_{\nu}^{\rm FF} = \frac{32}{3} \sqrt{\frac{\pi}{3}} \alpha^3 \chi_{\rm H} a_0^3 \langle Z^2 \rangle n_{\rm e} n_{\rm i} \left(\frac{\chi_{\rm H}}{kT}\right)^{1/2} g_{\rm FF} e^{-h\nu/kT}, \quad (8)$$

where α is the fine-structure constant, $\alpha \approx 1/137$, a_0 is the Bohr radius, and $\chi_{\rm H}$ is the K-shell ionization energy of hydrogen.⁹ The units of this expression are energy per volume, per steradian, per time, and per frequency. Throughout this article, $T = T_{\rm e}$ is the electron temperature. Since the hot-spot hydrogen almost completely ionized, $\langle Z^2 \rangle = 1$ but the Z dependence in Eq. (8) will be kept, anticipating the discussion below of the contamination of the hot spot by carbon as the result of fuel-shell mix. As will be shown in **Enhanced X-Ray Emission** as a Measure of Fuel-Shell Mix (p. 176), the FB contribution to hydrogen emissivity is negligible under hot-spot conditions.

This emissivity is written in a form first obtained in a semiclassical treatment by Kramers.¹⁶ The correction factor $g_{\rm FF}$ accounts for quantum-mechanical effects in FF absorption and emission and also in other absorption and emission processes introduced by Gaunt.¹⁷ The same Gaunt factor applies to both emission and absorption, a result of the microscopic reversibility of these processes.¹⁸ Many versions of the Gaunt factor have been provided over many years by many authors who, to cite just a few, include correct isolated-ion quantum wave functions,¹⁹ relativistic effects,²⁰ collective effects,²¹ and high-density effects such as electron degeneracy²² and strong plasma coupling.^{23,24} A particularly simple and accurate Gaunt factor for hydrogen has been provided by Kulsrud:²⁵

$$g_{\rm FF} = \frac{\sqrt{3}}{\pi} \ln \left[\frac{\left(\beta^2 + u\right)^{1/2} + \beta}{\left(\beta^2 + u\right)^{1/2} - \beta} \right],\tag{9}$$

where $u = h\nu/kT$ and β^2 is an average energy parameter

$$\beta^2 = \left\langle \frac{E}{kT} \right\rangle,\tag{10}$$

representing an effective average initial electron energy *E*. Rather than averaging the FF scattering cross section over the Maxwell distribution of initial electron momenta, Kulsrud provides the Gaunt factor for a single average initial electron momentum. He finds that the value $\beta = 0.87$ provides a good fit to a more-accurate quantum result of Sommerfeld based on the Born approximation.²⁵ We verify that Eq. (9) is a good approximation to the standard results of Karzas and Latter¹⁹ (KL) in the high-temperature limit $kT \gg \chi_{\rm H}$, which is the relevant regime for implosion cores at peak conditions. We also verify that the asymptotic expression

$$g_{\rm FF} \approx \frac{2\sqrt{3}}{\pi} \frac{\beta}{u^{1/2}} \tag{11}$$

is a useful approximation to Eq. (9) at high photon energies hv > kT. This is consistent with the sensitive range of the filtered

gated monochromatic x-ray imager (GMXI) camera²⁶ ($h\nu \sim 5$ keV), described below, and the expected core temperatures ($kT \sim 2$ keV). This is also well into the optically thin spectral range where emission is directly related to the hot-spot emissivity. Using Eq. (11) in Eq. (8) provides an emissivity expression with accurate temperature and photon-energy scaling that will be used in a later discussion of the interpretation of measured energy-integrated core emission.

Some brief textbook summaries of FF and FB emission and absorption dismiss the Gaunt factor as a constant correction of order unity,³ which for our purposes would be a poor approximation. Equation (11) provides a convenient simplification and ensures correct asymptotic scaling behavior. This asymptotic expression for the Kulsrud Gaunt factor is plotted in Fig. 139.13 along with the result of the full expression, Eq. (9), and KL values.¹⁹ The Kulsrud Gaunt factor depends only on u = hv/kT, while the KL results also depend on the parameter $\gamma^2 = Z^2 \chi_{\rm H}/kT$. Here, we simply plot the KL values for hydrogen (Z = 1) at temperatures 2 keV and 6 keV to show that the departure of the KL from the Kulsrud values is small and that their temperature dependence at constant u = hv/kT



Figure 139.13

Plot of the Kulsrud²⁵ approximation to the hydrogen free-free (FF) Gaunt factor versus u = hv/kT (red line) given by Eq. (9) and its asymptotic approximation given by Eq. (11) (black line). Karzas–Latter (KL) values for kT = 2 and 6 keV are shown for comparison. The relevant range of u for the anticipated core conditions and instrumentation (GMXI B) is indicated by the blue shading. The asymptotic behavior of the Kulsrud approximation agrees with that of the KL values.¹⁹

can be cautiously disregarded for the purposes of this discussion. The plot range is extended over an order of magnitude to show that the asymptotic behavior of the Kulsrud expression is correct. The relevant range of $h\nu/kT$ for our chosen imaging instrument, the GMXI, with a response function denoted here and in Fig. 139.13 as "B," is indicated by the blue-shaded strip. Here, the agreement among the four results is particularly close. The KL and Kulsrud formulations do not consider the effects of relativistic electron motion or the effects of degeneracy and strong coupling on the energy levels of the ions and on the free-electron energy distribution. These effects will not be significant under hot-spot emission conditions, although the latter two effects must be considered in the surrounding, relatively cold compressed shell, particularly earlier in the implosion.²⁴ Evaluating the numerical coefficient in Eq. (11) gives $g_{\text{H,FF}} \approx$ 0.959 $u^{-0.5}$, which is very close to $g_{\rm H,FF} = 0.966 u^{-0.41}$, a numerical fit to the KL hydrogen FF result near $h\nu \approx 5.39$ keV and $h\nu_0/kT \approx 2$, anticipating the conditions where Eq. (11) will be applied.

Let us specialize to fully ionized hydrogen, where $\langle Z^2 \rangle = 1$ and $n_e = n_i$. We then insert the Kulsrud Gaunt factor given by Eq. (11) into Eq. (8) and obtain

$$\varepsilon_{\nu}^{\rm FF} = \frac{64}{3\sqrt{\pi}} \beta \alpha^3 \chi_{\rm H} a_0^3 n_{\rm i}^2 \left(\frac{\chi_{\rm H}}{kT}\right)^{1/2} \left(\frac{kT}{h\nu}\right)^{1/2} e^{-h\nu/kT}.$$
 (12)

Using the ideal gas equation of state $P = (n_e + n_i) kT$ to replace density with pressure, we obtain

$$\varepsilon_{\nu}^{\text{FF}} = \frac{16}{3\sqrt{\pi}} \beta \alpha^3 \chi_{\text{H}} a_0^3 \left(\frac{P}{kT}\right)^2 \left(\frac{\chi_{\text{H}}}{h\nu}\right)^{1/2} e^{-h\nu/kT}.$$
 (13)

This ideal gas equation of state is an approximation to the more general expression $P = k(n_eT_e + n_iT_i)$, recognizing that the ion temperature T_i can exceed the electron temperature during the convergence of shocks that forms the hot spot. During peak compression, however, the two temperatures equilibrate rapidly, so $T_i = T_e$ —a useful approximation that has been standard in recent discussions of hot-spot dynamics—can be applied here with caution. We will show that this emissivity is almost exclusively a function of pressure when measured with an appropriate spectral response.

Three spectral-response channels of the GMXI²⁶ x-ray camera are shown in Fig. 139.14. These response functions are nearly Gaussian in shape with spectral widths of approxi-



Figure 139.14

Plot of the GMXI²⁶ camera spectral-response functions $F(\nu)$ for three of its channels. These channels differ by the thicknesses of the Al filter layers in their respective filter packages. The emissivity of a source, integrated over one of these response functions, will be exclusively dependent on the pressure of the source when the source temperatures fall within a narrow range near the ideal source temperature kT_0 indicated for that channel.

mately $\Delta h\nu \approx 1$ keV centered at energies near $h\nu_0 \approx 5$ keV. The three channels, denoted A, B, and C, differ in the 1-, 2-, and 3-mil thicknesses of Al in their respective filter packages. These response functions are approximated adequately for our purposes by a Gaussian function

$$F(\nu) = F_0 e^{-(\nu - \nu_0)^2 / (\Delta \nu)^2}$$
(14)

with a transmission width $\Delta \nu$ centered at ν_0 . With a response function of the form of Eq. (14), the frequency-integrated emissivity expression obtained using the emissivity given by Eq. (13) is

$$E_{\nu_{0},\Delta\nu} = \frac{16}{3\sqrt{\pi}} \alpha^{3} \chi_{\mathrm{H}} a_{0}^{3} \beta \left(\frac{P}{kT}\right)^{2} \left(\frac{\chi_{\mathrm{H}}}{kT}\right)^{1/2} \times F_{0} \int_{0}^{\infty} \left(\frac{kT}{h\nu}\right)^{1/2} e^{-(\nu-\nu_{0})^{2}/(\Delta\nu)^{2}} e^{-h\nu/kT} \,\mathrm{d}\nu.$$
(15)

For the anticipated small values of the parameters $\Delta v/v_0$ and $h\Delta v/kT$, we use the leading-order approximation

$$\left(\frac{h}{kT}\right)^{1/2} \int_{0}^{\infty} e^{-\left(\nu-\nu_{0}\right)^{2}/(\Delta\nu)^{2}} e^{-h\nu/kT} \frac{d\nu}{\nu^{1/2}}$$
$$\simeq \sqrt{\pi} \frac{\Delta\nu}{\nu_{0}} \left(\frac{h\nu_{0}}{kT}\right)^{1/2} e^{-h\nu_{0}/kT},$$
(16)

and we write

$$E_{\nu_0,\Delta\nu} = \frac{16}{3} \alpha^3 \chi_{\rm H} a_0^3 \beta F_0 \Delta \nu \left(\frac{P}{kT}\right)^2 \left(\frac{\chi_{\rm H}}{h\nu_0}\right)^{1/2} e^{-h\nu_0/kT}.$$
 (17)

Near any given temperature, the temperature dependence of Eq. (17) can be treated as a power law $y \sim x^{\eta}$ fit to the actual temperature dependence of the form

$$y = \frac{e^{-1/x}}{x^n},\tag{18}$$

where $x = kT/h\nu_0$. The exponent η near a particular value of x is

$$\eta = \frac{x}{y} \frac{\mathrm{d}y}{\mathrm{d}x} = \frac{1 - nx}{x} \,. \tag{19}$$

Solving for $\eta = 0$ gives x = 1/n. The temperature dependence of Eq. (17) is the n = 2 case, which gives

$$kT_0 = \frac{h\nu_0}{2} \tag{20}$$

as the condition for stationary maximum integrated emissivity with respect to temperature. So while Eq. (17) is explicitly temperature dependent, it can be regarded as independent of temperature within a limited range of temperature centered at kT_0 . If the KL fit given above were used rather than the asymptotic Kulsrud expression, Eq. (18) would be replaced by $y = e^{-1/x}/x^{2.09}$, displacing the stationary point of the emissivity to $kT_0 = h\nu_0/2.09$, which is almost the same as Eq. (20) for the purposes of this discussion. The applicable range of temperature is easy to determine directly from Eq. (17) for a desired tolerance. For example, the integrated emissivity will be within 90% of its maximum (representing a $\pm 5\%$ minimumto-maximum variation) at $kT_0 = 2.65$ keV over the temperature range 1.95 keV < kT < 3.73 keV for the GMXI B response function with the $h\nu_0 = 5.30$ -keV center energy. This temperature range is representative of cryogenic ICF implosion hot spots.

What we now have in Eq. (17) is an expression for emissivity that is a function of pressure alone, as long as the spectral response of the detector is appropriately matched to the source temperature range. A nominal temperature value or profile must be provided to evaluate Eq. (17), but if the source–instrument matching condition given by Eq. (20) is satisfied closely enough, the emissivity can be treated as a function of pressure alone. If the source temperature profile does deviate from satisfying Eq. (20) to a degree that the emissivity temperature dependence at fixed pressure cannot be ignored, Eq. (17) is still valid, but temperature profile input must then resemble the actual temperature profile closely enough to avoid throwing off the relationship between the emissivity and the pressure profiles. A temperature profile from a simulation may suffice.

Inferring Hot-Spot Pressure from X-Ray Emission Measurements

The expression for Abel inversion given by Eq. (4) and the emissivity expression in terms of pressure given by Eq. (17) allow the pressure profile to be inferred from the imaged specific intensity $I_{\nu_0}(x)$. This will be demonstrated using a simulated image of OMEGA cryogenic implosion shot 68791 (Ref. 27). The implosion is simulated with the one-dimensional (1-D) radiation-hydrodynamics code LILAC,²⁸ and the images are calculated using the post-processor Spect3D.²⁹ The temperature, density, and pressure profiles of this implosion at 1.94 ns-the time of stagnation and peak neutron production-are shown in Fig. 139.15. A GMXI image of the emission from this configuration simulated with Spect3D, using the spectral response function B with 2-mil Al filtering shown in Fig. 139.14, is shown as the red solid line in Fig. 139.16. Spect3D uses the actual response functions specified for the GMXI [not the Gaussian approximation used in deriving Eq. (17)] and Gaunt factors based on the KL model extended to include the effects of electron degeneracy.²⁰ Spect3D calculates a full solution of the equivalent of Eq. (1), taking into account all emission and absorption effects within the entire imploded configuration.

The simulated imploded configuration in Fig. 139.15 is substantially isobaric at $P \approx 22$ Gbar out to the radius indicated by the vertical dashed line at 34 μ m. In the simulated instantaneous image at this time shown in Fig. 139.16 and in the plot of the simulated time-integrated image in Fig. 139.17, this distance represents the radius of the 17% intensity contour of the instantaneous image, which has been suggested as one criterion to use in measuring the size of a hot spot from its image.³⁰ This criterion appears to coincide with other criteria that are more physically significant, although, unfortunately, not as directly measurable, such as the inner half-peak density point of the imploding mass distribution in Fig. 139.15 or the half-emissivity radius seen in Figs. 139.16 and 139.17. The



Figure 139.15

Simulated OMEGA cryo shot 68791 (Ref. 27) profiles at 1.94 ns, the time of peak neutron production. The hot spot (kT > 1.5 keV) appears to be substantially isobaric at $P \approx 22$ Gbar out to the radius indicated by the vertical dashed line at 34 μ m. In the simulated instantaneous image at this time, this distance represents the radius of the 17% intensity contour,³⁰ which is also shown for reference on the image plots in Figs. 139.16–139.18.

temperature range within this so-defined hot spot is roughly 1.5 keV < kT < 3.5 keV, which extends only slightly below the temperature range given above for a ±5% accuracy limit on Eq. (17) with a nominal temperature of kT_0 = 2.65 keV. Therefore, the flat emissivity profile in Fig. 139.16 should track the flat pressure profile in Fig. 139.15, even as the temperature varies. This is indeed the case, as will be shown below.

The solid red curve in Fig. 139.16 is the simulated instantaneous radial intensity distribution of the image obtained using the B response function including the 2-mil Al filter layer of OMEGA cryogenic implosion shot 68791 at 1.94 ns—the time of stagnation and peak neutron production shown in Fig. 139.15. The solid green curve is the emissivity profile obtained from the inverse Abel transform of this image using Eq. (4). The image is almost indistinguishable from the dashed purple quarter-ellipse, which is exactly what Eq. (5) predicts for the image of a constant emissivity profile extending out to $R = 34 \ \mu m$, shown in Fig. 139.16 as the dashed blue profile. The relative deviations of the simulated image from the elliptical image are very small, in contrast with the larger relative deviations of the simulated emissivity from a flat profile. This



Figure 139.16

The filtered simulated GMXI image of the simulated implosion core of OMEGA shot 68791 at 1.94 ns—the time of peak neutron production. The red solid line (image) is the B-channel integrated intensity distribution of the simulated image. The green solid line (emissivity) is the emissivity profile obtained from the inverse Abel transform of the simulated image. The image is almost indistinguishable from the purple dashed line (image block), which is the image of a constant emissivity profile (emissivity block) extending out to $R = 34 \ \mu m$.

is understood as the smoothing effect of the integration in Eq. (2) on the emissivity profile.

Figure 139.17 is virtually identical to Fig. 139.16 in every respect, other than a relative factor of about 110 ps, which can be taken as the emission time scale. This simulated image intensity has been integrated over a 300-ps time window centered at 1.96 ns, long enough to include both the x-ray and neutron emission times. Again, as was the case with the results shown in Fig. 139.16, the image deviates very little from the elliptical shape, while the emissivity deviates more visibly from a flat profile near the outer radius of the hot spot.

The pressure profile of an imploded hot spot can be inferred using Eq. (17) from the emissivity profile obtained by Abel inversion of the simulated image using Eq. (4). As discussed in the previous section, this inferred pressure is insensitive to the assumed temperature profile if the camera response is centered at $hv_0 = 2kT_0$, where the hot-spot temperatures are within a limited range of T_0 . Exactly how closely the emissivity profile follows the pressure profile will depend on how far the range of the hot-spot temperature deviates from T_0 , as has been described above. In principle, any nominal *T* value or radial *T*(*r*) profile can be used with Eq. (17). For example, this nominal temperature can be the ideal temperature T_0 matched to the instrument response, or it can be the logarithmic slope of the hard end of the continuum spectrum, if it has been measured. Ignoring the Gaunt factor in Eq. (8) leaves one with the simple $\varepsilon_{\nu}^{\text{FF}} \sim e^{-h\nu/kT}$ frequency dependence. The additional factor $(kT/h\nu)^{1/2}$ in Eq. (12) bends the logarithmic slope slightly,

$$-\frac{\mathrm{d}\ln\left(\varepsilon_{\nu}^{\mathrm{FF}}\right)}{\mathrm{d}h\nu} = \frac{1}{kT_{\mathrm{inferred}}} = \frac{1}{kT} + \frac{1}{2h\nu},\qquad(21)$$

but the fractional error in the temperature inferred from measuring the spectral logarithmic slope $\delta T/T \approx kT/2h\nu$ can be minimized by measuring as high in spectral energy as possible. Using Eq. (17) with a radially dependent simulated temperature profile could provide more-accurate results than a constant nominal temperature.



Figure 139.17

The filtered simulated GMXI image of the simulated implosion core of OMEGA shot 68791 integrated over a 300-ps time window centered at 1.96 ns. This interval includes the times of stagnation, peak neutron production, and peak x-ray intensity. All plotted quantities in this figure are time integrated. The red solid line (image) is the intensity distribution of the simulated image. The green solid line (emissivity) is the emissivity profile obtained from the inverse Abel transform of the image. Out to near $R = 34 \mu$ m, the image is almost indistinguishable from the purple dashed line (image block), which is the image of a constant emissivity profile (emissivity block).

Using Eq. (17) to infer a pressure profile from an emissivity profile is demonstrated in Fig. 139.18 for the case of shot 68791 shown in Fig. 139.16. Pressure profiles inferred from the simulated GMXI B-channel image of the simulated implosion core of OMEGA shot 68791 are shown in Fig. 139.16 at 1.94 ns—the time of peak neutron production. The pressure profiles were obtained using Eq. (17) from the emissivity profile obtained from the simulated image by Abel inversion. The pressure profile plotted as the red solid line was obtained assuming the ideal $kT_0 = 2.65$ keV nominal temperature based on the



Figure 139.18

Pressure profiles inferred from the simulated GMXI image shown in Fig. 139.16 of the simulated implosion core of OMEGA shot 68791 at 1.94 ns—the time of peak neutron production. The pressure profiles were obtained using Eq. (17) from the emissivity profile obtained from the simulated image by Abel inversion. The pressure profile plotted as the red solid line was obtained assuming the ideal $kT_0 = 2.65$ keV nominal temperature based on the GMXI response centered at 5.30 keV (red dashed line), and the pressure profile plotted as the blue solid line was obtained by assuming the radius-dependent temperature profile from the *LILAC* hydrodynamic simulation (blue dashed line). The vertical dashed line indicates the 17% intensity contour radius—the nominal outer radius of the hot spot.

GMXI response centered at 5.30 keV (red dashed line), and the pressure profile plotted as the blue solid line was obtained by assuming the radius-dependent temperature profile from the *LILAC* hydrodynamic simulation (blue dashed line). The vertical dashed line indicates the 17% intensity (half-emissivity) contour radius—the nominal outer radius of the hot spot. Even with these two very different assumed temperature profiles, the two inferred pressure profiles agree with the simulated pressure profile very well, up to within a short distance of the edge of the hot spot.

Scaling Relationships of the X-Ray Yield of an Isobaric Hot Spot at Stagnation

The energy-integrated emissivity of an isobaric hot spot tracks its flat pressure profile nearly all the way to its outer radius *R*, provided that the source temperature profile does not deviate too far from the T_0 set by the condition $h\nu_0 = 2kT_0$, where $h\nu_0$ is the center of the instrument-response function. If the emissivity per volume is reasonably uniform over the entire hot spot, as is the case in the example shown in Fig. 139.16, the hot spot will produce a total filtered radiated energy or photon yield of

$$Y_{\nu_0} = 4\pi \left(\frac{4\pi}{3} R^3\right) \Delta t E_{\nu_0},$$
 (22)

where the leading factor of 4π represents integration of the isotropic emissivity over the full sphere of the solid angle. If the gate time or exposure time Δt exceeds the lifetime of the hot spot, the total effective emission time Δt must be inferred from another measurement, such as the neutron yield Y_n , assuming that both the neutron and photon emission are limited by the same hot-spot lifetime. With a radius R obtained from the GMXI image and a relatively weak temperature dependence, a useful estimate of the hot-spot pressure can be obtained from Eqs. (17) and (22). This was done for a sizable ensemble of cryogenic implosion simulation results.²⁷ The inferred pressures are compared in Fig. 139.19 with the simulated central peak pressures. The simulated photon yield was the time-integrated B-channel emission, the emission time in Eq. (22) is the quotient of the simulated neutron yield divided by the peak neutron-production rate, and the image size is the 17%-intensity contour of the simulated GMXI image. The ensemble of implosions includes shell adiabat parameters over the range $1.5 < \alpha_{\text{shell}} < 3.5$ and neutron yields over the range $1.3 \times 10^{13} < Y_n < 1.3 \times 10^{14}$. The close agreement between the pressures inferred from the simulated images and the simulated pressure values is convincing, although not entirely expected, in light of the combined systematic inaccuracy that might result from equating the photon and neutron emission lifetimes, using the reasonable but not unique choice of the 17% contour as the hot-spot radius, and from assuming that the emissivity of the entire hot spot is determined by the central pressure at stagnation alone. This effectively confirms that the agreement between the simulated and inferred central pressures in Fig. 139.18 is obtained for every implosion in this ensemble.





The hot-spot pressure inferred from *LILAC/Spect3D* simulated images shown in very good agreement with the simulated peak central pressure. Equations (17) and (22) give accurate hot-spot pressures from simulated images for an ensemble of cryogenic simulations representing a broad range of shell adiabat parameters and neutron yields.

A few interesting new expressions for the scaling of the photon yield with various parameters of the implosion can be obtained from Eqs. (17) and (22). First, we write

$$M_{\rm Sh} \frac{R}{(\Delta t)^2} = 4\pi R^2 P, \qquad (23)$$

saying that the imploding thin unablated shocked shell of mass $M_{\rm Sh}$ surrounding the hot spot of radius R experiences an outward acceleration $R/(\Delta t)^2$ at stagnation, where Δt is the scale time of the bounce of shell, by the force of the hot-spot pressure P acting on the inner surface of the shell.¹¹

Next, we can write the adiabatic scaling of the hot-spot pressure with respect to the hot-spot volume $V = 4\pi R^3/3$ as

$$P \propto \alpha_{\rm HS} V^{-5/3},\tag{24}$$

where $\alpha_{\rm HS}$, a hot-spot "adiabat" parameter, distinguishes among the hot spots of different implosions. The stagnating hot-spot material is not adiabatic in the usual sense where the pressure would scale with the hot-spot density as $P \propto \alpha_{\rm HS} \rho^{-5/3}$ because of heat flow out of and material flow into the hot spot. The hot-spot volume, however, can be treated as an adiabatic enclosure because the heat conduction out of the hot spot is exactly compensated by the heat of the material ablated off the inner surface of the shell back into the hot spot at the hot-spot boundary.¹¹ Together, Eqs. (23) and (24) give the volume–time product

$$V\Delta t \propto \frac{\left(\alpha_{\rm HS} M_{\rm Sh}\right)^{1/2}}{P}$$
(25)

needed in Eq. (22). Applying Eq. (17), subject to the conditions given above for temperature-independent emissivity, we obtain the expression

$$Y_{\nu_0} \propto \left(\alpha_{\rm HS} M_{\rm Sh}\right)^{1/2} P, \qquad (26)$$

which predicts that the photon yield will scale in direct proportion to the hot-spot pressure for an ensemble of similar cryogenic implosions. This linear scaling is verified in Fig. 139.20 for the same set of simulations used to obtain the results shown



Figure 139.20

The hot-spot pressure simulated by *LILAC* shown following the predicted linear scaling with the photon yield. The hot-spot pressure is predicted to vary in direct proportion to the x-ray yield for an ensemble of similar cryogenic implosions with shell adiabat parameters and neutron yields that vary over a wide range.

in Fig. 139.19, although with more scatter. This additional scatter may reflect the simplification of the hot-spot dynamics by Eq. (23) and the simplification of the hot-spot energy balance by Eq. (24) or by the variations in the product $\alpha_{HS}M_{Sh}$ over the ensemble. Accounting for all the details of Fig. 139.20 is beyond the scope of this article, but for now, Fig. 139.20 suffices to add validity to the scaling arguments that have been made.

The sample of 1-D OMEGA cryogenic implosion simulations illustrated in Figs. 139.19 and 139.20 displays a curious scaling of the simulated photon yield with the neutron yield, $Y_{\nu_0} \propto Y_n^{0.57}$, shown in Fig. 139.21 (Ref. 27). The value of the scaling index, as well as the fact that the photon and neutron yields appear to fall along a single curve, can be explained in terms of the dynamics of isobaric implosion cores at stagnation. We begin again with the photon yield given by Eqs. (17) and (22), this time keeping track of the precise temperature scaling of the photon- and neutron-production rates. We allow that the scaling index η of the temperature dependence of Eqs. (18) and (19), as in

$$y = \frac{e^{-1/x}}{x^2} \propto x^{\eta},$$
 (27)

may deviate from $\eta = 0$. We expect that η will be small since our instrumentation and emission source place us near $h\nu_0 =$



Figure 139.21

Photon/neutron-yield scaling in cryogenic implosion simulations.²⁷ The x-ray yields of an ensemble of cryogenic implosion simulations have been found to scale with the 0.57 power of their respective neutron yields over a broad range of shell adiabats and neutron yields.

 $2kT_0$, but the scaling calculation below does not require this. We apply Eqs. (17), (22), and (27) and obtain

$$Y_{\nu_0} \propto P^2 T^{\eta} V \Delta t. \tag{28}$$

The neutron yield can be written using the deuterium–tritium (DT) reaction rate

$$\langle \sigma \nu \rangle \propto T^{4+\varepsilon},$$
 (29)

where we obtain the small deviations $|\varepsilon| \ll 1$ from a fixed temperature scaling using the reaction rate by Bosch and Hale.³¹ The neutron yield is written as the volume and time integral

$$Y_{\rm n} = \iint n_{\rm D} n_{\rm T} \langle \sigma \nu \rangle \mathrm{d}V \mathrm{d}t \propto P^2 T^{2+\varepsilon} V \Delta t. \tag{30}$$

The neutron-production rate is a function of the ion temperature, so combining Eq. (29) with the ideal gas equation of state to form Eq. (30) is another application of the $T_e = T_i$ equilibrium assumption made earlier. While the x-ray emissivity can be approximated fairly well as uniform over the volume of an isobaric hot spot, the neutron production varies as an additional two powers of temperature and, as a result, will be more center peaked and possibly shorter in duration. Nevertheless, we proceed assuming that the usual approximation—that the source volumes and emission times of the neutrons and photons are the same—is valid to within constant factors that drop out of scaling relationships, and we apply Eqs. (24), (25), (28), and (30) to obtain

$$Y_{\nu_0} \propto \left(\frac{M_{\rm Sh}^{2/9} M_{\rm HS}^{10/9}}{\alpha_{\rm HS}^{4/9}}\right)^p Y_{\rm n}^q,$$
 (31)

where

$$p = \frac{1 + (\varepsilon - \eta)/2}{1 + 2\varepsilon/9}$$
(32)

and

$$q = \frac{5+2\eta}{9+2\varepsilon}.$$
(33)

For the nominal values $\eta = \varepsilon = 0$, this gives q = 0.56 for the neutron-yield scaling exponent, which is very close to the value obtained from the simulation ensemble, as shown in Fig. 139.21. In Eq. (31), the three constant stagnation parameters α_{HS} , M_{Sh} , and the hot-spot mass $M_{\text{HS}} \sim R^3 P/T$ combine in a product where the scaling relative to each other is fixed. We have chosen to write Eq. (31) with this three-parameter product scaled such

that p = 1 for the nominal values $\eta = \varepsilon = 0$. The effects of variations of α_{HS} , M_{Sh} , and M_{HS} throughout the simulation ensemble and the effects of the temperature-dependent scaling corrections η and ε will be considered below, but for now the successful derivation of the value of the scaling exponent q is clearly an encouraging validation of the scaling calculations so far.

The alignment of the simulated data points in Fig. 139.21 along a single curve does make sense in light of Eq. (31). The neutron yield varies over the ensemble by an order of magnitude, and the scatter about the curve, attributable to variations in $\alpha_{\rm HS}$, $M_{\rm Sh}$, and $M_{\rm HS}$, is relatively small. Extracting precise values of these three parameters from simulations depends on the time resolution of the simulation output as well as on somewhat arbitrary definitions, but it is clear that the net effect of their variations is much less than the ranges of the neutron and photon yields. It is interesting to note that the temperaturedependent scaling corrections pertain only to the microphysics of photon and neutron production, not the design or dynamics of the implosion capsules. Were this yield-yield plot generated with simulations or measurements of a different ensemble of isobaric implosions with target masses scaled up by significant factor, for example, Eqs. (31)-(33) predict that the overall proportionality constant would change with the mass and hot-spot adiabat parameter, the yield-scaling exponent would be the same for the same spectral response and source temperatures.

The result q = 0.56 quoted above is obtained only with the nominal values $\eta = \varepsilon = 0$. Their actual values are plotted as functions of temperature in Fig. 139.22 as blue and purple dashed curves, respectively, where they are read off the right-hand scale. The q value obtained from Eq. (33) is plotted on the same graph as the black curve and is read off the left-hand scale. The left-hand scale is only a fifth the numerical length of the righthand scale, indicating that q is a relatively insensitive function of temperature because of its opposite sensitivities to ε and η in Eq. (33). The range $q = 0.57 \pm 0.06$ is obtained, in reasonably good agreement with the value q = 0.57 in Fig. 139.21, for the broad range of temperature $kT = 2.21 \pm 0.43$ keV. Even though the yield-scaling exponent is relatively insensitive to variations in the hot-spot temperature, it may have diagnostic value through its temperature dependence. The red curve representing the scaling index p is to be read off the right-hand scale and is a relatively insensitive function of temperature within the 0.86-keV range indicated by the pair of vertical dashed lines. If the fit to KL results for the FF Gaunt factor described above were used in place of Eq. (11), the effect would be to reduce η by 0.09, which would reduce the predicted q values by 3.6% overall.



Figure 139.22

Yield-scaling indices as functions of temperature. The yield-scaling indices p and q are functions of temperature through the temperature-dependent scaling indices η and ε of the photon- and neutron-production rates, respectively. The q curve shows that the q = 0.57 value from Fig. 139.21 is at the center of the range $q = 0.57\pm0.06$, corresponding to the temperature range $kT = 2.21\pm0.43$ keV.

Equation (31) with its scaling index given by Eq. (32) accounts for the yield-scaling behavior shown in Fig. 139.21 because the scaling parameter agrees with the fit to the simulation ensemble and because the scaled product $M_{\rm Sh}^{2/9} M_{\rm HS}^{10/9} \alpha_{\rm HS}^{-4/9}$ of stagnation parameters has negligible correlations with both the photon and neutron yields. The simulation ensemble was constructed to follow a long series of cryogenic implosion experiments as closely as possible; the experiments were not designed to control this product. If this product did have a significant correlation with either the photon or neutron yields, then Eq. (32) would not agree with the yield scaling exhibited by the simulation ensemble. For example, a slightly modified version of Eq. (31) can be derived by using the scaling expression

$$M_{\rm HS} \sim M_{\rm Sh}^{1/7} P^{4/7} R^{16/7}$$
(34)

for the hot-spot mass given by Zhou and Betti³² to remove $M_{\rm HS}$ from the calculation. This gives the results

$$Y_{\nu_0} \propto \left(M_{\rm Sh}^{4/11}\right)^{p'} Y_{\rm n}^{q'},$$
 (35)

where

$$p' = \frac{1 + \left(\varepsilon - \eta\right)/2}{1 + 2\varepsilon/11} \tag{36}$$

and

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$$q' = \frac{7+2\eta}{11+2\varepsilon}.$$
(37)

These results are only a slight departure from Eqs. (31)–(33). The yield-scaling parameter q' does not agree as well as q with the fit to the ensemble, but, remarkably, the only stagnation parameter appearing is M_{Sh} , or, in other words, the normalized photon yield

$$Y_{\nu_0} / Y_n^{q'} \sim M_{\rm Sh}^{4p'/11}$$

is a function of the shell mass alone. Both Eqs. (31) and (35) are correct expressions, but if Eq. (33) agrees more with the simulation ensemble than Eq. (37), it is because $M_{\rm Sh}$ has a less negligible correlation with the photon and neutron yields. Another reason to regard Eqs. (35)–(37) with more caution than Eqs. (31)–(33) is that introducing Eq. (34) brings additional approximations into the calculation that were made to evaluate thermal transport in evolving inner-shell density and temperature profiles,¹¹ although these same approximations were made in establishing the hot-spot adiabatic behavior expressed as Eq. (24); the hot-spot pressure varies adiabatically with respect to the hot-spot volume during the approach to stagnation.

Enhanced X-Ray Emission as a Measure of Fuel–Shell Mix

The yield-scaling results in Fig. 139.21 were obtained from $LILAC^{28}$ 1-D hydrodynamic simulations, which exhibit no unstable hydrodynamic behavior. Implosions with shell adiabat parameters in the lower end of the range $1.5 < \alpha_{shell} < 3.5$ have thinner shells that are more susceptible to the Rayleigh-Taylor instability³³ during the deceleration phase of the implosion. Breakup of the unstable shell would leave some amount of the shell carbon mixed into the hot spot at the time of stagnation. Since carbon is much more emissive than hydrogen, very small concentrations of carbon can significantly increase the x-ray emission. Since the photon and neutron yields in Fig. 139.21 exhibit a $Y_{\nu_0} \propto Y_n^{0.57}$ behavior, the normalized yield quotient $Y_{\nu_0}/Y_n^{0.57}$ forms a constant normalized photon yield when plotted as a function of any quantity, such as the shell adiabat parameter, as shown by the blue squares in Fig. 139.23. The same quantity derived from actual measured yields is plotted as red circles. These values are generally constant, in agreement with the 1-D simulated results, for shell adiabat parameter values above the $\alpha_{shell} \approx 2.3$ value, but they show a distinct excess photon yield below this point. This has been attributed to the expected contamination of the hot spot by shell carbon. We extend this analysis to infer the mass of shell material mixed into the hot spot from measurements of this excess emission.

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Figure 139.23

Excess photon yield as a measure of fuel–shell mix. The normalized yield quotient $Y_{\nu_0}/Y_n^{0.57}$ is nearly constant for the simulated implosions, according to the results plotted here and in Fig. 139.21. This quotient is plotted versus the shell adiabat parameter here as blue squares. The same quotient calculated from measured yields is plotted as red circles. These measurements conform to the 0.57 power scaling for higher shell adiabat parameters $\alpha_{\rm shell} > 2.3$. As unstable shells break up during the deceleration phase of the implosion, carbon can be mixed into the hot spot from the shell, which accounts for the excess x-ray emission above the amount expected from the yield scaling. This twofold to threefold enhancement can be attributed to masses of 125 to 250 ng of shell CH mixed uniformly into the hot spot. (Figure from Ref. 27.)

Using the FF emission of hydrogen as a pressure diagnostic assumes a pure hydrogen core. Any mix of non-hydrogen into the hot spot will raise the per-ion emissivity of the hot spot. The twofold-to-threefold enhancement of the hot-spot emission shown in Fig. 139.23 for low-adiabat implosions can be interpreted in terms of contamination of the hot spot by shell carbon. The emission from a carbon-contaminated hot spot will have a strong contribution from radiative recombination, which is FB emission, the time inverse of photoionization or bound-free (BF) absorption. The carbon emission will have comparable FF and FB contributions with much stronger total emission per ion than hydrogen.

Following a conventional approach, we discuss the combined FF and FB emissivity in parallel by considering their respective absorption cross sections. The Kramers photon absorption cross sections are

$$\sigma_{\rm FF}(\nu) = \frac{256\pi^{5/2}}{3\sqrt{3}} \alpha a_0^5 n_{\rm e} Z^2 \left(\frac{\chi_{\rm H}}{kT}\right)^{1/2} \left(\frac{\chi_{\rm H}}{h\nu}\right)^3 g_{\rm FF} \quad (38)$$

for FF absorption by an ion of total charge Z and

$$\sigma_{\rm BF}(\nu) = \begin{cases} \frac{64\pi}{3\sqrt{3}} \alpha a_0^2 \frac{Z^4}{p^5} \left(\frac{\chi_{\rm H}}{h\nu}\right)^3 g_{\rm BF}; h\nu < \chi_p \\ 0; h\nu < \chi_p \end{cases}$$
(39)

for BF absorption by the photoionization of the single electron bound to a nucleus of charge Z in principal quantum level p (Ref. 9). We keep the explicit dependence of $\sigma_{BF}(\nu)$ on p for now, but only the ground-state p = 1 photoionization contribution will be significant. The Gaunt factors $g_{\rm FF}$ and $g_{\rm BF}$ account for departures from the Kramers semi-classical approximation. Based on the micro-reversibility of each process, the Gaunt factor for emission also applies to absorption. The ionizationedge cutoff energy χ_p of the BF cross section is the ionization energy of the bound electron in the p shell. This can be expressed using the Bohr formula $\chi_p = Z^2 \chi_H / p^2 - \Delta \chi$. We mention continuum lowering³⁴ $\Delta \chi$ only for completeness, showing only its effect on χ . Since x-ray measurements are made far above this $h\nu = \chi_p$ cutoff, we will not consider it further. Further discussion beyond the scope of this article should consider continuum lowering and other high-density effects in more detail.

The hot-spot plasma is accurately described as nearly completely ionized, with the fully ionized atoms accounting for essentially all the FF and FB emission and with only a trace of the H-like species remaining to provide BF absorption. We neglect excited states and consider only the density $n_{i,1}$ of ground-state (p = 1) H-like ions and the density $n_{i,2}$ of the fully stripped species, so that $n_{i,1} + n_{i,2} = n_i$, where n_i is the total ion density of element *i*. Certainly for hydrogen and carbon, $n_{i,1}$ will be very small, and all other ionization species can be neglected. The hot spot is dense enough to maintain collision-dominated, detailed-balance local thermodynamic equilibrium (LTE) ionization-species population ratios given by the Saha equation⁹

$$\frac{n_{i,2}n_{\rm e}}{n_{i,1}} = \frac{2}{a_0^3} \frac{g_2}{g_1} \left(\frac{kT}{4\pi\chi_{\rm H}}\right)^{3/2} e^{-\chi_i/kT},\tag{40}$$

where g_1 and g_2 are the statistical weights of the bound and stripped states, respectively. The statistical weights for H-like and stripped ions are $g_1 = 2p^2$ and $g_2 = 1$, respectively.

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The FF opacity κ_{ν}^{FF} for the fully ionized species with nuclear charge Z,

$$\kappa_{\nu}^{\text{FF}} = \frac{256\pi^{5/2}}{3\sqrt{3}} \alpha a_0^5 n_{\text{e}} n_{i,2} Z^2 g_{i,\text{FF}}$$
$$\times \left(\frac{\chi_{\text{H}}}{kT}\right)^{1/2} \left(\frac{\chi_{\text{H}}}{h\nu}\right)^3 \left(1 - e^{-h\nu/kT}\right), \tag{41}$$

accounts for the entire FF contribution from that element since lower degrees of ion ionization contribute less, in proportion to their fractional populations. We can be more precise now by stipulating that the Gaunt factor $g_{i,FF}$ pertains to the fully stripped species of element *i*. The correction for stimulated emission is included. Considering a mixture of elements *i*, using $n_2 \approx n_i$, we have

$$\sum_i n_i Z_i^2 g_{i,\rm FF} = n_i \left< Z^2 g_{Z,\rm FF} \right>, \label{eq:relation}$$

where n_i is the total ion density, and

$$\kappa_{\nu}^{\rm FF} = \frac{256\pi^{5/2}}{3\sqrt{3}} \alpha a_0^5 n_i n_e \left\langle Z^2 g_{\rm FF} \right\rangle$$
$$\times \left(\frac{\chi_{\rm H}}{kT} \right)^{1/2} \left(\frac{\chi_{\rm H}}{h\nu} \right)^3 \left(1 - e^{-h\nu/kT} \right). \tag{42}$$

The BF opacity for a single element is written in similar terms. Here, only the H-like species population $n_{i,1}$ is relevant in the fully ionized limit, giving simply

$$\kappa_{\nu}^{\rm BF} = n_{i,1} \frac{64\pi}{3\sqrt{3}} \alpha a_0^2 \frac{Z^4}{p^5} \left(\frac{\chi_{\rm H}}{h\nu}\right)^3 \times \left(1 - e^{-h\nu/kT}\right) g_{i,\rm BF}$$
(43)

for the spectral range above the ionization edge. The BF opacity written in this way has strong hidden temperature and density dependence through $n_{i,1}$. The more-stationary product $n_{i,2}n_e$ is substituted using the Saha equation [Eq. (40)] to obtain

$$\kappa_{\nu}^{\rm BF} = \frac{256\pi^{5/2}}{3\sqrt{3}} \alpha a_0^5 n_2 n_{\rm e} \left(\frac{\chi_{\rm H}}{h\nu}\right)^{3/2} \frac{g_1}{g_2} \frac{Z^4}{p^5} \left(\frac{\chi_{\rm H}}{h\nu}\right)^3 \times \left(1 - e^{-h\nu/kT}\right) e^{(\chi - \Delta\chi)/kT} g_{i,\rm FF}.$$
(44)

Again, considering a mixture of elements *i*, using $n_{i,2} \approx n_i$, we can write

$$\sum_{i} n_{i} Z_{i}^{4} e^{\left(\chi - \Delta\chi\right)/kT} g_{i,\text{BF}} = n_{i} \left\langle Z^{4} e^{\left(\chi - \Delta\chi\right)/kT} g_{\text{BF}} \right\rangle, \tag{45}$$

giving

$$\kappa_{\nu}^{\rm BF} = \frac{256\pi^{5/2}}{3\sqrt{3}} \alpha a_0^5 n_i n_e \left(\frac{\chi_{\rm H}}{kT}\right)^{3/2} \frac{g_1}{g_2} \frac{1}{p^5} \left(\frac{\chi_{\rm H}}{h\nu}\right)^3 \\ \times \left\langle Z^4 e^{(\chi - \Delta \chi)/kT} g_{Z,\rm BF} \right\rangle \left(1 - e^{-h\nu/kT}\right). \tag{46}$$

Applying the appropriate values for g_1 , g_2 , and p given above for the H-like species, we have

$$\kappa_{\nu}^{\mathrm{BF}} = \frac{512\pi^{5/2}}{3\sqrt{3}} \alpha a_{0}^{5} n_{i} n_{e} \left(\frac{\chi_{\mathrm{H}}}{kT}\right)^{3/2} \left(\frac{\chi_{\mathrm{H}}}{h\nu}\right)^{3} \times \left\langle Z^{4} e^{\left(\chi - \Delta\chi\right)/kT} g_{Z,\mathrm{BF}} \right\rangle \left(1 - e^{-h\nu/kT}\right).$$
(47)

In LTE, the Kirchhoff relationship¹⁴ $\varepsilon_{\nu} = \kappa_{\nu}B_{\nu}(T)$ is the statistical detailed-balance relationship between emissivity and opacity, or between any one absorption process and its time-inverse emission process. Therefore,

$$\frac{\varepsilon_{\nu}^{\rm BF}}{\varepsilon_{\nu}^{\rm FF}} = \frac{\kappa_{\nu}^{\rm BF}}{\kappa_{\nu}^{\rm FF}} \tag{48}$$

and, applying Eqs. (42) and (47),

$$\frac{\varepsilon_{\nu}^{\rm BF}}{\varepsilon_{\nu}^{\rm FF}} = \left(\frac{2\chi_{\rm H}}{kT}\right) \frac{\left\langle Z^4 e^{(\chi - \Delta\chi)/kT} g_{\rm BF} \right\rangle}{\left\langle Z^2 g_{\rm FF} \right\rangle}.$$
(49)

This ratio is independent of density, except at higher densities where continuum lowering $\Delta \chi \sim n_i^{1/3}$ becomes important. To gauge the relative importance of the participating processes, we set aside the Gaunt factors temporarily and write

$$\left\langle Z^4 e^{\left(\chi - \Delta \chi\right)/kT} g_{\rm BF} \right\rangle \approx \left\langle Z^4 \right\rangle,$$
 (50)

which is a valid simplification at very high temperatures, $kT \gg \chi \ge \chi_{\text{H}}$, and

$$\langle Z^2 g_{\rm FF} \rangle \approx \langle Z^2 \rangle.$$
 (51)

It is easy to see from Eqs. (49)–(51) that we have been correct in neglecting FB emission in hydrogen, relative to FF emission, but that the $\langle Z^4 \rangle / \langle Z^2 \rangle$ charge scaling of the FB-to-FF ratio predicts significant FB emission from mid-Z contaminants, in addition to their FF emission, even if they are present in trace amounts relative to hydrogen.

It may appear paradoxical that the BF-to-FF ratio given by Eq. (49) is independent of density, considering that BF absorption is initiated by a two-body (photon-ion) interaction, implying that its opacity is linear in density, while FF absorption is initiated by a three-body (photon-electron-ion) interaction, implying that its opacity is second order in density. In general, the ratio of densities in the BF-to-FF opacity ratio is density dependent, but since we are working in the near-complete ionization regime where all the bound-electron species are H-like, the Saha equation [Eq. (40)] removes this density dependence.

The Planck function in the Kirchhoff relationship can be written as

$$B_{\nu}(T) = \frac{\chi_{\rm H} \alpha^2 \left(\frac{h\nu}{\chi_{\rm H}}\right)^3}{8\pi^2 a_0^2} \frac{1}{e^{h\nu/kT} - 1}.$$
 (52)

The individual emissivity contributions are then

$$\varepsilon_{\nu}^{\rm FF} = \frac{32}{3} \sqrt{\frac{\pi}{3}} \alpha^3 \chi_{\rm H} a_0^3 n_{\rm i} n_{\rm e} \left\langle Z^2 g_{\rm FF} \right\rangle \left(\frac{\chi_{\rm H}}{kT} \right)^{1/2} e^{-h\nu/kT}, \quad (53)$$

repeating Eq. (8), and

$$\varepsilon_{\nu}^{\rm BF} = \frac{64}{3} \sqrt{\frac{\pi}{3}} \alpha^3 \chi_{\rm H} a_0^3 n_{\rm i} n_{\rm e} \left(\frac{\chi_{\rm H}}{kT}\right)^{1/2} \\ \times \left\langle Z^4 e^{(\chi - \Delta \chi)/kT} g_{\rm BF} \right\rangle e^{-h\nu/kT}, \qquad (54)$$

with a total emissivity

$$\varepsilon_{\nu} = \frac{32}{3} \sqrt{\frac{\pi}{3}} \alpha^{3} \chi_{\mathrm{H}} a_{0}^{3} n_{\mathrm{i}} n_{\mathrm{e}} \left(\frac{\chi_{\mathrm{H}}}{kT}\right)^{1/2} \\ \times \left[\left\langle Z^{2} g_{\mathrm{FF}} \right\rangle + 2 \left(\frac{\chi_{\mathrm{H}}}{kT}\right) \left\langle Z^{4} e^{(\chi - \Delta \chi)/kT} g_{\mathrm{BF}} \right\rangle \right] \\ \times e^{-h\nu/kT}.$$
(55)

This expression can be evaluated to obtain the enhancement of the photon yield resulting from the increase in both $\langle Z^2 g_{FF} \rangle$ and $\langle Z^4 e^{(\chi - \Delta \chi)/kT} g_{BF} \rangle$ from CH shell–mix contamination. The CH mix does not affect the neutron yield, except possibly through increased radiative cooling since the $n_D n_T$ product of fuel ion densities product in Eq. (30) is only minutely affected by a trace contamination.

As was noted above, the Kulsrud Gaunt factor for hydrogen differs only slightly from the more reliable KL values for hydrogen in our relevant temperature range, but it is not applicable to carbon under ICF hot-spot conditions or to FB emission. Power-law fits to KL results have been obtained near $hv_0 \approx 5.39$ keV and $hv_0/kT \approx 2$ for the Gaunt factors needed to evaluate Eq. (55) (Ref. 19). These are

$$g_{\rm H,FF} \approx 0.966 (h\nu/kT)^{-0.41},$$

$$g_{\rm H,FB} \approx 0.299 (h\nu/5.39 \,\rm{keV})^{-0.43},$$

$$g_{\rm C,FF} \approx 1.28 (h\nu/kT)^{-0.23},$$

$$g_{\rm C,FB} \approx 0.926 (h\nu/5.39 \,\rm{keV})^{-0.141}.$$
(56)

The FF Gaunt factors depend on temperature because the FF absorption cross section is an average over the Maxwell– Boltzmann distribution of initial free-electron states. The FB Gaunt factors, on the other hand, do not depend on temperature because all properties of the single-electron bound states are attributes of the ion, not the free-electron plasma. We find that adding a fraction $f_{\rm CH} = 2.4\%$ by atom of CH to a pure hydrogen core, as in the uniform mix of shell polymer into the hot spot, doubles the emissivity at $h\nu_0 \approx 5.39$ keV. This represents a mix mass of $\Delta M_{\rm CH} = 125$ ng in the chosen example above where the hot-spot mass is $M_{\rm HS} = 2.1 \ \mu g$, which happens to be the mean hot-spot mass of the entire ensemble of simulations considered above. The standard deviation of hot-spot masses in this ensemble is only 17%, so the emission-doubling mix

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mass is roughly the same for all the implosions. The measured emission enhancements of the low-adiabat implosions plotted in Fig. 139.23 range over factors from 2 to 3, which correspond to a range of mix mass from 125 ng to 250 ng. The enhancement factor corresponding to a 2.4%-by-atom contamination level is shown plotted as the black curve in Fig. 139.24 as a function of temperature. Within the range of plausible hot-spot temperatures indicated by the blue-shaded region, 2 keV < kT <3 keV, the enhancement factor is a weak function of temperature, primarily the result of the temperature dependences of the relevant Gaunt factors listed in Eq. (56). The yield-ratio scaling index q, plotted as the red curve, varies with temperature in the contaminated core, as it does in the pure hydrogen core in Fig. 139.22, although much less so, because the Gaunt factors for the FB emission, which counts for very roughly half the emission in Fig. 139.24 and none of the emission in Fig. 139.22, do not depend on temperature. The black and red horizontal dashed lines show the enhancement factor (unity) and the index value (0.56) expected for zero contamination.



Figure 139.24

Photon-yield enhancement resulting from a contamination of the hot spot, as a function of temperature. The photon emissivity of a hot spot is expected to double when contaminated at a level of 2.4% by atoms of the CH shell material. This enhancement factor varies with hot-spot temperature, as shown by the black curve and the left-hand scale. The blue shading emphasizes the range of temperature from 2 to 3 keV—the plausible range of hot-spot temperatures. The red curve and right-hand scale show the scaling index q of the yield quotient Y_{ν_0} / Y_n^q as a function of temperature. The black and red horizontal dashed lines show the enhancement factor (unity) and the index value (0.56), respectively, expected for zero contamination.

A set of similar mix-mass measurements has been reported for cryogenic implosions on the NIF based on the ratio of the x-ray and neutron yields.¹³ To describe this technique in terms of the discussion above, we write Eq. (28) once again, but with the mix effect included, as in Eq. (55),

$$Y_{\nu_0} \propto P^2 \frac{e^{-h\nu/kT}}{T^{2-\eta}} \left(\frac{\chi_{\rm H}}{kT}\right)^{1/2} \left(1 + \langle Z \rangle\right) \\ \times \left[\langle Z^2 g_{\rm FF} \rangle + 2 \left(\frac{\chi_{\rm H}}{kT}\right) \langle Z^4 e^{(\chi - \Delta \chi)/kT} g_{\rm BF} \rangle \right] V \Delta t. \quad (57)$$

Our discussion parallels the discussion in Ref. 13 to some extent, although the mix-mass estimates presented in Ref. 13 are based ultimately on emissivity values obtained from detailed atomic calculations, rather than the Kramers–Gaunt formulation underlying Eq. (57). The ratio of the photon to neutron yield can be constructed from Eq. (30) and Eq. (57) with the result being

$$\frac{Y_{\nu_0}}{Y_{\rm n}} \propto \frac{e^{-h\nu/kT}}{T^{4+\varepsilon-\eta}} \left(\frac{\chi_{\rm H}}{kT}\right)^{1/2} \left(1 + \langle Z \rangle\right) \\ \times \left[\langle Z^2 g_{\rm FF} \rangle + 2 \left(\frac{\chi_{\rm H}}{kT}\right) \langle Z^4 e^{(\chi - \Delta \chi)/kT} g_{\rm BF} \rangle \right]$$
(58)

to within fixed constants. This is the quantity that was measured in the NIF experiments. In forming this ratio, the pressure factors, the hot-spot volumes, and the emission times cancel. It was assumed here as well that the temperature and neutron emission rate were spatially uniform, the electron and ion temperatures were equal, and the photon and neutron emission volumes and times were equal. Since OMEGA and non-igniting NIF implosions stagnate in comparable temperature ranges, we can examine this yield ratio in an approximate fashion by allowing $\eta = \varepsilon = 0$, and applying the Kulsrud limit $g_{\text{FF}} \propto T^{0.5}$ for both hydrogen and carbon and the same temperature independence for g_{FB} , which is a better approximation near $h\nu_0 = 10.85$ keV used in the NIF measurements than near $h\nu_0 = 5.30$ keV used in the OMEGA experiments. This gives

$$\frac{Y_{\nu_0}}{Y_{\rm n}} \propto \frac{e^{-h\nu/kT}}{T^4} \left(1 + \langle Z \rangle\right) \left[\langle Z^2 \rangle + 2\left(\frac{\chi_{\rm H}}{kT}\right) \frac{g_{\rm BF}}{g_{\rm FF}} \langle Z^4 \rangle \right].$$
(59)

The argument supporting the mix-mass estimates based on measurements of this yield ratio in the NIF experiments is that, according to an equation very much like Eq. (59), the yield ratio is related directly to the charge averages and consequently to the mix atomic fractions:

$$\langle Z^n \rangle = 1 + (6^n - 1) \frac{f_{\rm CH}}{2}.$$
 (60)

Since the hydrogen and carbon are nearly fully ionized, the ion charges are effectively fixed, and everything else is a weak function of temperature.¹³ Based on the condition given by Eq. (19) for n = 4, the leading factor of Eq. (59) is independent of temperature for a range of temperatures centered at $kT_0 = h\nu_0/4$ or $kT_0 = 2.71$ keV, which is almost exactly the center of the range of temperatures, 1.7 keV < kT < 3.9 keV given in Ref. 13 as the range of applicable source temperatures. These NIF mix-mass measurements are another example of how taking x-ray measurements at an appropriate spectral energy simplifies their analysis in a very important way.

The contamination fractions measured on OMEGA at the level of $f_{CH} = 2.4\%$ to 4.8% are similar to the range of mix fractions measured on the NIF.¹³ Given the obvious dissimilarities between NIF indirect-drive implosions driven by two orders of magnitude more laser energy than the OMEGA direct-drive implosions, the similar mix fractions become an interesting point of comparison for future consideration. A few remarks will suffice for now. We note that the NIF capsules are roughly twice the diameter of the OMEGA capsules.³⁵ Assuming naively that all characteristic lengths and times of an implosion scale in direct proportion to the initial capsule radius, i.e., if the experiments on the two platforms were self-similar versions of each other, then the material composition profiles should be self-similar as well. Comparable mix fractions would result from shell perturbation amplitudes growing to comparable fractions of the capsule radius. For spherical-harmonic perturbation modes of the same harmonic order, the perturbation wavelengths scale with radius. The saturation amplitude, which is the point where linear perturbation growth transitions to nonlinear growth and, perhaps, turbulent mix, occurs at a specific ratio of amplitude to wavelength.³⁶ Since both the wavelengths and the amplitudes scale in direct proportion to the capsule radius, the transition to turbulence should be self-similar in both series of implosions. No doubt, this self-similarity does not apply to all aspects of an implosion.⁴ If mix is regarded as a surface phenomenon, comparable mix masses per volume indicate roughly twice the mix mass per shell area in the NIF capsules than in the OMEGA targets since the former have roughly twice the volume-to-surface-area ratio as the latter. These tentative remarks do not identify which series of implosions was more unstable or which of the two mix fractions is larger, relative to

expected performance, but it is clear that mix measurements are now possible and experimental progress in addressing these and other questions can be expected in the future.

Conclusions

The scaling behavior of the x-ray emissivity of hydrogen hot spots in ICF implosions has been examined. Using the pressure- and temperature-dependence of x-ray continuum emission, we have shown that the pressure can be inferred from the emissivity measured within a specific spectral energy range without requiring accurate prior knowledge of the source temperature. This is significant because the pressure is the single most important parameter that characterizes an isobaric hot spot; it is a key measure of the scaled-equivalent ignition performance of an implosion; and it is an example of how quantities can be measured in direct or advantageous ways with spectral responses that have been optimized to the task.

The scaling relationship between the photon and neutron yields of OMEGA cryogenic implosions that has been observed in simulation results and used to isolate excess x-ray emission from low-adiabat implosions has been explained. We have brought together x-ray emission and neutron yield scaling results to produce the first estimates of the fuel-shell mix mass in OMEGA implosions. The scaling properties of both the x-ray emissivity at the chosen spectral energy and the neutron yield allow one to normalize the x-ray emission with respect to the scaled neutron yield so that excess normalized emission becomes a measure of the CH polymer mass from the shell that has mixed into the hot spot during the implosion, up to the time of stagnation. It is a matter of some interest that this neutronphoton scaling, which has gone unexplained in the past, is now understood. It is of particular interest to the progress of cryogenic implosion research on the OMEGA Laser System that we have estimates of the mix mass. In a number of important respects, this approach to mix-mass estimation is similar to that of the x-ray continuum-based mix-mass measurements on the NIF. It is interesting that both sets of measurements describe similar levels of mix contamination, the implications of which will be interesting to consider in the future.

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