SHOULD ONE USE THE RAY-BY-RAY APPROXIMATION IN CORE-COLLAPSE SUPERNova SIMULATIONS?

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ABSTRACT

We perform the first self-consistent, time-dependent, multi-group calculations in two dimensions (2D) to address the consequences of using the ray-by-ray+ transport simplification in core-collapse supernova simulations. Such a dimensional reduction is employed by many researchers to facilitate their resource-intensive calculations. Our new code (Fornax) implements multi-D transport, and can, by zeroing out transverse flux terms, emulate the ray-by-ray+ scheme. Using the same microphysics, initial models, resolution, and code, we compare the results of simulating 12, 15, 20, and 25 M☉ progenitor models using these two transport methods. Our findings call into question the wisdom of the pervasive use of the ray-by-ray+ approach. Employing it leads to maximum post-bounce/pre-explosion shock radii that are almost universally larger by tens of kilometers than those derived using the more accurate scheme, typically leaving the post-bounce matter less bound and artificially more “explodable.” In fact, for our 25 M☉ progenitor, the ray-by-ray+ model explodes, while the corresponding multi-D transport model does not. Therefore, in two dimensions, the combination of ray-by-ray+ with the axial sloshing hydrodynamics that is a feature of 2D supernova dynamics can result in quantitatively, and perhaps qualitatively, incorrect results.

Key words: supernovae: general

1. INTRODUCTION

Despite years of tangible advance, the core-collapse supernova problem has persisted for half a century without a definitive solution. While there has been much conceptual progress during this time concerning the relevant nuclear, particle, and neutrino physics, and while the astrophysical context and goals of theory have been greatly sharpened and refined, the mechanism of explosion that leads to the measured explosion energies, nucleosynthesis, debris morphology, and residues (neutron stars or stellar-mass black holes) has not been unambiguously determined. Neutrinos are copiously produced and radiated at the high densities and temperatures achieved in collapse, and it is widely believed that neutrino heating plays a pivotal role in igniting the explosion (Colgate & White 1966; Bethe & Wilson 1985). However, the hydrodynamic instabilities and turbulence that attend collapse, bounce, and proto-neutron star mantle dynamics alter the critical condition for explosion (Burrows & Goshy 1993; Murphy & Burrows 2008; Pejcha & Thompson 2012) by introducing turbulent stress and enhancing the efficiency of the neutrino–matter coupling and must be simulated in detail to properly capture such (and related) effects. The multi-dimensional complexity of the hydrodynamics and neutrino radiative transfer seem to be central to the mechanism itself—by a wide margin, spherical models do not generally explode. This requires the solution of the coupled partial-differential equations of multi-dimensional hydrodynamics and radiative transfer, which classically is a seven-dimensional problem (time, three space, and three momentum space for each neutrino species). If performed with acceptable resolution, multi-angle transport at every three-dimensional position at every timestep cannot be done using current supercomputers. Hence, various simplifications, mostly in the radiation sector, have been introduced by the different groups attacking the core-collapse mechanism problem. In this paper, we address the accuracy of one such approximation, the so-called “ray-by-ray” transport method. Although our group does not use it (Burrows et al. 2006, 2007; Dolence et al. 2015), it is being employed by most groups currently publishing explosions, and the fidelity with which it handles multi-dimensional transport, particularly in two spatial dimensions, has been called into question (Burrows 2013; Dolence et al. 2015; Sumiyoshi et al. 2015).

Using their ChImera code, the ORNL group has obtained tepid explosions in two dimensions (2D) (Bruenn et al. 2013, 2016) and in three dimensions (3D) (Lentz et al. 2015). The Garching group, using the CoConuT hydrodynamics code (Müller et al. 2012a, 2012b; Müller & Janka 2014) in combination with the Vertex transport solver, or their earlier vertex-Prometheus code (Buras et al. 2006), has also obtained tepid explosions in 2D, and in 3D only when altering the physics (Melson et al. 2015). Suwa et al. (2014), Takiwaki et al. (2012), and Iwakami et al. (2014) neglect νμ and ντ neutrinos and use the simplifying IDSA technique (Liebendörfer et al. 2009); they also obtain explosions in 2D and have explored 3D simulations at low resolution. However, none of these groups performs multi-dimensional transport, but all use multiple one-dimensional (1D) spherical transport solves in lieu of more accurate multi-dimensional solutions in the so-called “ray-by-ray” approach (Burrows et al. 1995; Buras et al. 2003, 2006). One-dimensional transport using thermodynamic profiles along a single radial ray is calculated and coupled only locally to the spherical hydro solution of that ray. This is done for all, or a subset, of the radial rays to approximate the radiation field and its evolution. Such a dimensional reduction technique is employed to facilitate multi-processor scaling and reduce computational overhead. However, transverse radiation fluxes and lateral transport are ignored, which in reality would significantly smooth out the radiation field. The relative smoothness of a radiation field, even when the background thermal field is
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variegated and severely asymmetrical, is a consequence of the fact that radiation at a point is the composite of the contributions from many directions, with sources at a large variety of temperatures and emissivities, from both adjacent rays and distant sources at other angles. This smoothing effect was demonstrated definitively in 2D by Ott et al. (2008) and Brandt et al. (2011) using multi-dimensional/multi-angle transport, and by Dolence et al. (2015), using flux-limited diffusion.\(^3\) For instance, the magnitude of the variation with angle in either the neutrino flux or the specific neutrino heating rate from pole to equator of a 2D simulation not employing the ray-by-ray simplification is typically less than \(\sim 5\%\). However, when employing the ray-by-ray procedure, such variations are typically \(\sim 20\%\) (Dolence et al. 2015; Sumiyoshi et al. 2015). Lund et al. (2012) and Tamborra et al. (2013) perform 3D simulations using ray-by-ray+ and see variations of \(\sim 50\%\).\(^4\) Although it is often stated that small effects at the \(\sim 10\%\) level in the equation of state or neutrino–matter couplings might make a qualitative difference in the computational outcome of collapse simulations, such a large difference between ray-by-ray and real multi-dimensional transport should give one pause. This is particularly relevant for 2D simulations in which axial sloshing is so pronounced, and when such dipolar oscillations have themselves been invoked as potentially central to the mechanism of explosion (Hanke et al. 2012). It should be noted that all groups currently reporting explosions by the neutrino heating mechanism employ the ray-by-ray dimensional reduction simplification (with the “plus” extension, or otherwise).

The only self-consistent calculations that employed truly multi-dimensional transport were those of Burrows et al. (2006, 2007) and Ott et al. (2008), using the VULCAN/2D code, and Dolence et al. (2015), using the CASTRO code (Zhang et al. 2011, 2013). They did not obtain explosions in their 2D simulations. Since most other self-consistent calculations that obtained explosions in 2D used the ray-by-ray scheme, while the VULCAN/2D and CASTRO studies did not, one is tempted to suggest (as has Burrows 2013) that the ray-by-ray dimensional reduction and simplification may in 2D be yielding qualitatively incorrect results. The ray-by-ray anomalies in the angular distribution of the radiation field and corresponding neutrino heating rates may be reinforcing the axial sloshing motions and in 2D pushing the shock into explosion.\(^5\)

Hence, the current explosions in 2D may in part be numerical artifacts. Unfortunately, VULCAN/2D (Burrows et al. 2007) and CASTRO (Zhang et al. 2011, 2013) employed Cartesian grids in the central regions, and, hence, were not suitable to test in a self-consistent fashion the ray-by-ray approach to multi-dimensional collapse simulations. Surprisingly, the accuracy of the ray-by-ray+ scheme has never before been tested, neither by us nor by those who use it, and such a self-consistent head-to-head comparison in the time-dependent context is the major motivation for this paper. One of the motivations for the development of our new code FORNAX (Dolence et al. 2016) was to be able to compare directly, using the same code, the results using more realistic multi-dimensional transport against those using the ray-by-ray simplification. FORNAX, with its spherical geometry and multi-dimensional capability, can do this.

In Section 2, we briefly describe our FORNAX code and the method we use to implement the ray-by-ray+ simplification. A more complete description of FORNAX can be found in Dolence et al. (2016). Section 3 contains the results of our comparison of the ray-by-ray and multi-dimensional radiation/hydro simulations in 2D, with a particular focus on the different hydrodynamical responses, shock positions and motions, and overall outcomes. We conclude in Section 4 with a discussion of what we have found concerning the limitations of the ray-by-ray approach. We emphasize that this paper is not meant to be a full supernova mechanism study. Rather, in this paper we focus solely on the viability and accuracy of the ray-by-ray+ transport methodology and leave more comprehensive studies of the supernova phenomenon per se using FORNAX to later work.

2. METHODOLOGY

Since the speed of sound is close to the speed of light for most of the evolution of a proto-neutron star core, in the core-collapse supernova problem we can integrate the transport operator explicitly, in a manner analogous to that employed by Skinner & Ostriker (2013), but without the need for an artificially reduced speed of light in the post-bounce phase. This has also been noticed by Just et al. (2015) and O’Connor & Couch (2015) and obviates the need to do implicit global iterations, speeding up the calculations for a given core count by a factor of \(\sim 5–10\).\(^6\)

Our implementation of this is FORNAX, purpose-built by our team over the last two years. The code solves the equations of self-gravitating, compressible hydrodynamics with an arbitrary (often tabulated) equation of state, coupled to the multi-group two-moment equations for neutrino transport. We solve the equations on a fixed Eulerian grid, where the geometry and coordinates are handled in a generally-covariant way using a spatial metric and associated Christoffel symbols. Thus, FORNAX can easily incorporate a variety of geometries and coordinate choices, with the restriction that the metric and coordinate transformations remain orthogonal. For this work, we have chosen a spherical geometry and adopt a radial coordinate \(x_i\) such that \(r = x_i \sinh(x_i/x_0)\), where \(x_i\) is solved to give a specified central resolution (0.5 km) and outer radius (10\(^4\) km). Roughly, \(x_i\) is associated with the radius at which a uniform grid in \(x_i\) transitions from a uniform grid in \(r\) to a uniform grid in \(\log r\). In multiple dimensions, a spherical grid that extends down to \(r = 0\) can be restrictive as the Courant-limited timestep can become extremely small due to the convergence of radial grid lines. While others have sometimes simply evolved the inner region in 1D spherical symmetry to overcome this problem, we instead enhance our otherwise logically Cartesian mesh with a simple form of static mesh derefinement. Regardless of the resolution specified in the angular direction, the derefinen mesh has only two zones in each angular direction for those zones that reach \(r = 0\).

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\(^3\) However, Dolence et al. (2015), in particular, used multi-group flux-limited diffusion (MGFLD), which is known to smooth out radiation fields at large distances in the transparent regime.

\(^4\) A commonly used variant, “ray-by-ray+,” includes a matter velocity advection step of either the neutrino number or energy density and the radial flux component. See the Appendix for details.

\(^5\) The general absence in 3D of either an axial effect or the pronounced sloshing seen in 2D may be rendering 3D simulations performed with the ray-by-ray approach less problematic.

\(^6\) Much of that speed up is due simply to avoiding the many iterations to convergence of the spatial derivative terms required in such implicit methods, and the associated global communication overhead.
Subsequently, the number of zones in the angular direction is doubled for every doubling of radius until a specified number of cells is reached. The outcome of this process is (1) a mesh where the cell aspect ratios are never extreme, (2) a true representation of the multi-D dynamics throughout the whole domain, and (3) a drastically improved timestep limit. This type of mesh is sometimes referred to as dendritic, having nodes on zone edges in some cells instead of exclusively at corners. On our Eulerian mesh, we employ a finite-volume discretization and evolve our equations with a directionally unsplit higher-order Godunov-type scheme with Shu and Osher’s optimal second-order Runge–Kutta time stepping (Shu & Osher 1988). Hydrodynamic and compositional variables are conservatively reconstructed using a novel “geometry-aware” parabolic profile to compute edge states that define a Riemann problem on each face. With these edge states, the intercell fluxes of the conserved hydrodynamic variables are computed with the HLLC Riemann solver of Toro et al. (1994).

The multi-group two-moment equations for neutrino transport are formulated in the comoving frame and include all terms to $O(\nu/c)$. The moment hierarchy is closed with the “M1” model (Vaytet et al. 2011), though FORNAX can easily accommodate other closure choices. After core bounce, the fastest hydrodynamic signal speeds in the core-collapse supernova problem are within a factor of a few of the speed of light, so a time-explicit integration of the non-stiff transport terms is not only simpler and generally more accurate, it is also faster than globally coupled time-implicit transport solvers that are typically employed in radiation hydrodynamics methods. Radiation quantities are reconstructed with linear profiles and the resulting edge states are used to calculate fluxes via the HLLE solver, similar to Vaytet et al. (2011). As in O’Connor & Ott (2013), the HLLE fluxes are corrected to reduce numerical diffusion in the non-hyperbolic regime. The terms that transfer momentum and energy between the radiation and the gas are operator-split and treated implicitly. These terms are purely local to each cell and, therefore, do not introduce any global coupling. While in the free-streaming regime of transport the solution is agnostic to the ordering of the operator-split updates, the tightly coupled diffusive regime is sensitive to this choice. In FORNAX, the order of the operator-split updates is chosen to ensure the correct asymptotic behavior in the diffusive regime. In an earlier draft of the paper, we performed our calculations without general-relativistic (GR) corrections, but for the current version we have implemented and employed the effects of GR in the manner of Rampp & Janka (2002) and Marek et al. (2006).

For the neutrino–matter coupling, the interaction rates of Burrows et al. (2006) are used. These incorporate weak-magnetism and recoil corrections to the neutrino–nucleon interactions, screening and ion–ion correlation corrections for the neutral-current scattering off nuclei, and inelastic scattering of neutrinos off electrons. The latter is handled using the approximate scheme of Thompson et al. (2003), which treats various terms of the scattering matrix either explicitly or implicitly so that inelastic scattering can be written as a sum of effective source and sink terms in and out of an energy bin. This approach is stable, avoids the requirement of energy bin couplings, and scales linearly with energy group number.

However, the associated cross sections are low and final-state blocking (and, hence, suppression) is significant (particularly after bounce). Therefore, for this study we ignore inelastic scattering. We use the Lattimer–Swesty nuclear equation of state with $K = 220\text{ MeV}$ for all simulations (Lattimer & Swesty 1991).

For this study, comparing 2D simulation results using ray-by-ray and actual multi-dimensional transport, we employ FORNAX with 40 energy ($\varepsilon_{\nu}$) groups for each of the $\nu_e, \bar{\nu}_e,$ and $\nu_x$ species, where $\nu_x$ represents the $\nu_{\mu}, \bar{\nu}_{\mu}, \nu_{\tau},$ and $\bar{\nu}_{\tau}$ types collectively. For the $\nu_x$ types, $\varepsilon_{\nu}$ varies logarithmically from 1 to 300 MeV, while it varies from 1 to 100 MeV for the other species. For the pair processes, we assume detailed balance, but ignore the blocking of two final-state neutrino species. Spherical coordinates are used. The radial coordinate, $r$, runs from 0 to 10,000 kilometers (km) in 608 zones. The radial grid smoothly transitions from uniform spacing with $\Delta r = 0.5\text{ km}$ in the interior to logarithmic spacing, with a characteristic transition radius of $\sim 100\text{ km}$. The angular grid spacing varies smoothly from $\approx 0.95^\circ$ at the poles to $\approx 0.65^\circ$ at the equator in 256 zones, covering the full $180^\circ$. We discuss the issue of resolution in Section 3.1. The 12, 15, 20, and 25 $M_\odot$ massive-star progenitors we use as initial models are from Woosley & Heger (2007). The 2D simulations were performed on the Cray machines Blue Waters and Cori and take approximately two days each (using 1024 cores) to complete.

To perform ray-by-ray+$\pm$ calculations, we set all transverse neutrino radiation fluxes equal to zero, effectively updating only the radial components of the flux in each energy group and for each neutrino species. However, we implement the “ray-by-ray+$\pm$” variant by retaining the velocity advective terms in the zeroth-moment (energy) and first-moment (momentum) radiation equations (Buras et al. 2006). The Appendix highlights these various terms as they appear in our formulation. This also requires retaining the second- and third-moment pressure and heat-flux tensor components in the geometric source terms associated with the Christoffel terms that arise in spherical coordinates (in particular to retain transverse pressure gradients). We note that other groups using other transport implementations (e.g., multi-group flux-limited diffusion [MGFLD]; spherical Boltzmann; IDS A) will be constrained to use their own closures for these source and advective terms (if implementing the “plus” variant), but that the “transverse flux equals zero” step likely has the greatest consequences and is the central assumption of ray-by-ray.

The original version of FORNAX was built with flexibility in mind, and geometrical quantities like face areas and cell volumes were numerically computed via quadrature. Though these integrals were computed extremely precisely, a very low level of symmetry breaking was introduced. This was particularly important in the inner region where we employ static mesh refinement (the dendritic grid). The new version now eliminates this problem. For the calculations in this paper, we impose initial perturbations à la Müller & Janka (2015), with $n = 5, l = 2,$ and $v_{\text{max}} = 1000\text{ km s}^{-1}$, and do not allow the grid and computational noise to dictate initial asphericities.

One remaining source of unphysical symmetry breaking is the tabular equation of state. For the calculations presented in this work, we employed a Newton–Raphson iteration to compute thermodynamic quantities as a function of density, specific internal energy, and electron fraction from a table that provides values as a function of density, temperature, and

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9 Since cell aspect ratios are kept within a factor of 2 of unity, the explicit CFL timestep is kept within a factor of 2 of that of the innermost radial zone.
electron fraction. Part of this process relies on an initial guess for the temperature and the iteration proceeds until a relative tolerance of $10^{-11}$ is reached. Given two different initial guesses, the iteration will not result in identical thermodynamic outputs even for identical densities, internal energies, and electron fractions. These differences are small, $\sim \epsilon T \partial q / \partial T$, where $\epsilon$ is the relative tolerance and $q$ is the quantity in question. We note that for multi-dimensional models symmetry breaking is required. Most authors rely on grid level noise of some sort to achieve this. Importantly, whatever the source of noise in our calculations, it is exactly the same for both ray-by-ray and non-ray-by-ray multi-D runs.

3. RESULTS

In order to assess the differences between multi-D transport and the ray-by-ray + simplification when investigating the dynamics and outcomes of multi-D core collapse, we have simulated the evolution of the collapse, bounce, and subsequent behavior of four non-rotating progenitor models from Woosley & Heger (2007). These 12, 15, 20, and 25 $M_\odot$ models were those employed by Bruenn et al. (2013, 2016) in their recent explosion study, for which they used ray-by-ray +. Figures 1–2 depict the evolution of the average (thin line) and maximum (thick line) shock radii ($R_{\text{avg}}$ and $R_{\text{max}}$) versus time after bounce for the four models. Superposed for easier comparison are the multi-D transport FORNAX simulations (blue) and the corresponding ray-by-ray + developments (red). First, we note that the post-shock region is chaotic and turbulent due to a combination of neutrino-driven convection and the standing-accretion-shock instability (Blondin et al. 2003). Hence, the detailed hydrodynamic behavior is stochastic and will depend upon resolution, the character of perturbations, and numerical implementation. Nevertheless, Figures 1–2 show what are at times striking quantitative differences. The maximum shock radii ($R_{\text{max}}$) for the low-accretion-rate 12 $M_\odot$ and 15 $M_\odot$ models are not much different between the multi-D transport and ray-by-ray + realizations, although the average excess for the ray-by-ray + and full simulations in $R_{\text{max}}$ is $\sim 5$ km. However, for the 20 $M_\odot$ and 25 $M_\odot$ progenitors, $R_{\text{max}}$ is systematically higher for the ray-by-ray + realizations at almost all post-bounce times, with excesses ranging $\sim 10$ to $50$ km. In fact, Figures 1–2 demonstrate that the $R_{\text{max}}$ excess for the ray-by-ray + runs is an increasing function of progenitor mass, and $R_{\text{max}}$ for the full transport realizations is almost never larger than for the ray-by-ray + runs. Importantly, the 25 $M_\odot$ progenitor explodes at late times for the ray-by-ray + simulation, while it does not explode for the multi-D transport run, achieving an $R_{\text{max}}$ of $\sim 2000$ km by the end of the simulation. Figure 3 depicts the explosion on a more appropriately...
The moving frame $\nu_e$ (blue), $\bar{\nu}_e$ (red), and “$\nu_\mu$” (green) luminosities at 100 km for all eight models, integrated over solid angle (and averaged over 10 ms), are shown in Figures 4 and 5. The ray-by-ray+ values are rendered as thick solid lines. As Figures 1 through 2 indicated, there are hydrodynamic differences between the multi-D transport and ray-by-ray+ runs. However, as Figures 4 and 5 indicate, the corresponding total luminosities are generally not much different, though for the 25 $M_\odot$ model the $\nu_\mu$ luminosity can be slightly lower at later times. Since $\nu_\mu$s do not heat much, but mostly sap the core of energy, this is conducive to explosion. Also, for the 25 $M_\odot$ ray-by-ray+ model, the total luminosities are lower for all species for about $\sim$100 ms around the time the Si/O interface is accreted at $\sim$0.3 s after bounce. The overall similarities in general, particularly for the 12 $M_\odot$, 15 $M_\odot$, and 20 $M_\odot$ progenitors, are likely because the quasi-di-polar oscillation of the post-shock material and shock position average out the accretion and internal flux components.

This is not as much the case for the fluxes along the symmetry axes at $\theta = 0$ and $\theta = 180$. If we calculate the average energy flux in a 20° wedge at the northern pole (defined as the $\theta = 0°$ axis), and multiply by 4$\pi r^2$ to obtain a pseudo-luminosity, we see marked differences between the multi-D transport and the ray-by-ray+ numbers at later times. As Figures 6 and 7 indicate, the “polar” $\nu_e$ and $\bar{\nu}_e$ luminosities evolve on average to be larger using the ray-by-ray+ approach —as much as $\sim$30% larger. The $\nu_e$ and $\bar{\nu}_e$ neutrino fluxes are most strongly coupled to matter by charged-current absorption on liberated nucleons. As has been pointed out previously (Burrows 2013; Dolence et al. 2015; Sumiyoshi et al. 2015), the ray-by-ray+ scheme makes the radiation field track the matter field too closely, exaggerating its angular variation. True transport results in contributions to the neutrino energy density at a given point from the entire angular and radial domain, thereby smoothing it out. Aside from the modest effect of the advective terms, the ray-by-ray+ approach restricts the influence at a point to the characteristics of the thermal and matter profiles ($T$, $\rho$, and the electron fraction $Y_e$) along only the associated radial ray. This makes the radiation field unphysically sensitive to local and radial matter conditions, and, as we see, can alter the matter motions. Moreover, the

The minimum, average, and maximum shock radii of the 25 $M_\odot$ progenitor model using the ray-by-ray+ method that explodes at late times. This is the same as depicted in the right panel of Figure 2, but on a much expanded vertical scale in order to better show the explosion. The maximum shock radius reaches $\sim$2000 km by the end of the simulation, while the corresponding average radius achieves $\sim$1000 km, indicating the asymmetry of this 2D blast.

The expanded scale. These shock radius comparisons for a range of progenitors collectively suggest that the use of the ray-by-ray+ simplification can result in quantitatively, and perhaps qualitatively, different evolution, particularly for the more massive progenitors. The significance of the differences in the behavior of the shock radii in these 2D simulations lies in the fact that the ray-by-ray+ approach results in post-shock matter that is easier to unbind, because it resides at larger radii in the shallower depths of the gravitational potential well. Hence, we speculate that ray-by-ray+ simulations are more sensitive to the details of the numerical implementation, to the opacities, and to perturbations (perhaps due to the accretion of turbulent infalling matter). In sum, they are more “explodable,” but artificially so. In fact, for our 25 $M_\odot$ progenitor, we see an explosion only for the ray-by-ray+ variant. In general, the fact that the ray-by-ray+ method results in systematically larger excursions in the maximum shock radius suggests that its use can make explosions in 2D artificially a bit more likely.

**Figure 3.** Minimum, average, and maximum shock radii of the 25 $M_\odot$ progenitor model using the ray-by-ray+ method that explodes at late times. This is the same as depicted in the right panel of Figure 2, but on a much expanded vertical scale in order to better show the explosion. The maximum shock radius reaches $\sim$2000 km by the end of the simulation, while the corresponding average radius achieves $\sim$1000 km, indicating the asymmetry of this 2D blast.

**Figure 4.** Total solid-angle-integrated comoving-frame luminosities (in units of $10^{52}$ erg s$^{-1}$) at 100 km of the $\nu_e$ (blue), $\bar{\nu}_e$ (red), and “$\nu_\mu$” (green) neutrinos vs. time after bounce (in seconds). The luminosity is smoothed over 10 ms centered around this time. The left-hand panel is for the 12 $M_\odot$ progenitor model, and the right-hand panel is for the 15 $M_\odot$ progenitor model. The thin solid curves are for the multi-D transport scheme, and the thick solid curves are for the ray-by-ray+ approach. Note that the early bump in the $\nu_e$ luminosities results from the fact that we are calculating in the comoving frame and depicting the result at 100 km. The shock is formed deep and sweeps through the inner radii at high speed. The bump we see is due to the change in comoving-frame energy luminosity due to the abrupt change in the velocity and the consequent Doppler shifts—there would be no such bump if the calculations were in the lab frame, or the luminosity were measured much farther out. We provide the luminosities at 100 km, not at 500 km or larger radii, since this radius is in the vicinity of the gain region.
variations in the polar luminosities seen in Figures 6 and 7 are phased with the shock oscillations, thereby leading to excess heating near when the matter is less bound. Figure 8 for the 20 $M_\odot$ model comparison demonstrates this most dramatically. Here, we depict the normalized dipole component of both the shock radius and the $\nu_e$ luminosity for both the multi-D transport and ray-by-ray+ simulations. We see on both panels that the amplitudes for both quantities are roughly in phase with one another. However, the decrease seen in Figure 7 by $\sim 30\%$ in the $\nu_\mu$ luminosity along the poles for the ray-by-ray+ run of the 25 $M_\odot$ progenitor at late times may be suggestive, since $\nu_\mu$ neutrino emission is largely detrimental to explosion; such a decrease may have aided this model to explode.

Equally important, Figures 9 through 12 provide an estimate of the relative degree of neutrino heating in the gain region for this suite of four progenitors as a function of time after bounce. Here, we define the gain-region power deposition as the sum of the net $\nu_e$ and $\bar{\nu}_e$ heating rates between the shock and the 50 km...
radius when the net rate is positive. Other definitions can be envisioned, but this measure is straightforward. On each panel, this quantity is displayed for both the multi-D and ray-by-ray runs. The left panels provide the total power deposition, while the right panels are the same quantity, but for the 20° cone around the pole. We see that, for the 12M⊙ model, the differences between the multi-D and ray-by-ray are not large, although in the early post-bounce epoch the ray-by-ray numbers can be as much as \( \sim 50\% \) higher for the polar quantity. However, for the other models the differences along the poles between those two approaches are progressively more significant with increasing progenitor mass, while the total heating rates are not much different. For our 25 M⊙ model, the differences between the two transport approaches translate into a qualitatively different outcome (explosion versus no explosion).

### 3.1. Resolution Dependence

The resolution of a simulation can affect a conclusion quantitatively, and the resolution necessary to demonstrate a given behavior robustly will depend upon the algorithm employed. In addition, it has been suggested that the “explodability” of a model seems greater for lower-resolution simulations (Couch & Ott 2015). This is due predominantly to the anomalously greater power left in such runs at the larger scales at which the turbulent stress is most significant (Burrows et al. 1995; Murphy et al. 2013). Moreover, we surmise that when the initial seed perturbations are not imposed, the grid resolution will affect both the magnitude and scale of the seed perturbations for the Rayleigh–Taylor-like instabilities that grow into turbulence in the post-bounce mantles. Smaller-scale perturbations grow more slowly, and in grid codes, higher-
resolution simulations may introduce smaller amplitudes. When the initial perturbations from which the instabilities grow have not been identified and managed (as they have not been in most studies), then the strength of the resultant turbulence will be similarly contingent. Müller & Janka (2015) and Couch & Ott (2015) have started to address the importance of the initial turbulent perturbations that arise in the pre-collapse convective structures of the cores of massive stars, and we plan a separate paper on this phenomenon. However, here we compare simulations for ray-by-ray+ and multi-D transport at two different resolutions to ascertain in a preliminary way the differences.

The 2D explosion simulations of Müller et al. (2012b) employed a spherical polar grid covering 180° in θ with 64 or 128 angular zones and 10,000 km in radius with 400 radial zones. They used 12 energy groups from 0 to 380 MeV. For their 2D explosion simulations, Bruenn et al. (2016) employed 256 uniform angular zones and 512 radial zones out to 23,000 km (25 $M_\odot$) or 30,000 km (12 $M_\odot$), depending upon the progenitor. They used 20 energy groups ranging from 4 to 250 MeV. In neither of these simulations were the initial perturbations controlled for, and both simulations employed the ray-by-ray+ approach. As communicated in Section 2, the results described in Section 3 employed 256 angular zones, 608
Figure 13. Comparison of the different resolutions of the evolution of the shock radius (maximum [thick line] and average/mean [thin line]) of the 25 \( M_\odot \) progenitor ray-by-ray+ model. Low resolution is 304 (\( r \times 128 (\theta) \)) (blue), the baseline resolution is 608 \( \times \) 256 (red), and the high resolution is 1216 \( \times \) 512 (green). The baseline curves reproduce the data in Figure 2 for comparison. However, in this plot, time after bounce goes to 1.2 s and the ordinate extends to 300 km. See the text in Section 3.1 for a discussion.

radial zones (covering a radial extent of 10,000 km), and 40 energy groups, and so were rather better resolved than the two other efforts.

To gauge the potential effect of resolution, we conducted a set of simulations with 512 angular and 1216 radial zones (twice the baseline resolution), as well as with 128 angular and 304 radial zones (half the baseline resolution), attempting to maintain the aspect ratio of the grid. Figure 13 shows the comparison of the evolution of the shock positions for the ray-by-ray+ runs for the 25 \( M_\odot \) progenitor at the three different resolutions. Despite the chaos of the flow and the consequent inherent sensitivity to even small changes, all the ray-by-ray+ runs exploded at late times and manifested similar (though not exactly the same) evolutionary behaviors in \( R_{\text{max}} \) and \( R_{\text{avg}} \). In particular, though many of the oscillations in \( R_{\text{max}} \) before explosion are approximately in phase and of similar magnitude, the explosion times for the different resolutions are not the same, nor are they monotonic with resolution. Nevertheless, from this we can conclude that the resolution dependence of our runs, at least for the 25 \( M_\odot \) progenitor, is minimal and that our simulations are qualitatively converged.

4. CONCLUSIONS

We have presented the first self-consistent, time-dependent calculations to address the hydrodynamic consequences of the use in core-collapse simulations of the ray-by-ray+ transport ansatz, and our findings call into question the wisdom of its pervasive use. Such a method was introduced to simplify and speed up otherwise difficult and resource-intensive core-collapse simulations, but in doing so it may have compromised the results obtained. The combination of ray-by-ray with the axial sloshing hydrodynamics that is a feature of 2D supernova dynamics (see Burrows et al. 2012 and references therein) can result in quantitatively (and, perhaps, qualitatively) incorrect results. However, we have not here shown that the same problems arise, or arise to so dramatic a degree, in 3D. In 3D, the hydrodynamics is very much more prone to axial motions of the type that signify and define 2D simulations (see, for example, Dolence et al. 2013 and references therein). Also, we have conducted the study using only one code (Fornax), incorporating as it does its own algorithms, stencils, and term reconstructions. Moreover, there may be further angular, radial, or energy group resolution issues that are manifest to varying degrees in different codes.

However, our study merely compares the same simulations with and without the ray-by-ray+ simplification. We used the same microphysics, initial models, and hydrodynamics, as well as the same resolution (and initial perturbations), yet we find significant differences of a quantitative nature when doing so. This conclusion calls into question simulations in 2D employing the ray-by-ray+ (or simple ray-by-ray) prescription used by most groups now performing core-collapse simulations and most groups to date obtaining explosions in 2D by the neutrino mechanism. When there was strong turbulence and large shock radius excursions, we found that the ray-by-ray+ transport prescription tends to amplify them further. For one progenitor, the 25 \( M_\odot \), it resulted in explosion, while the full multi-D run did not. We note as far as “explodability” is concerned that decreasing the energy losses due to \( \nu_e \) radiation may be as important as increasing the \( \nu_x \) and \( \nu_x \) heating rates in the gain region. Such decreases could also result from legitimate alterations in the microphysics and may be one source of the differences in outcome seen in the literature.

We emphasize that a test of ray-by-ray+ does not require the code or calculations to have all the ingredients necessary for a full supernova mechanism study; this clearly was not the intention of this paper. A clean test merely requires one to compare calculations performed with exactly the same basic setup and physics, but with either the ray-by-ray+ method or an actual multi-D transport method employed. For the ray-by-ray+ method to be acceptable, such comparison calculations, performed side by side with exactly the same initial conditions, microphysics, and physics, must show little quantitative difference. However, what we have found is that in such a direct comparison there are palpable quantitative differences, and occasional qualitative ones.

However, from this we do not conclude that the neutrino mechanism itself is suspect. It is still a compelling mechanism by which to power such supernovae. Rather, aspects of the initial models and the microphysics might need closer scrutiny, and the behavior in 3D still needs to be properly addressed. The suggestion that sensitivity to given changes in the microphysics (for instance, the neutrino–matter coupling) increases with increasing dimension (from 1D to 3D) would seem to be a productive line to follow (Melson et al. 2015). Differences between the initial 1D progenitor models currently employed in core-collapse work and those that better incorporate (1) convection and semi-convection (probably fully 3D initial models; Arnett et al. 2015; Couch et al. 2015), (2) initial convective perturbations (Couch & Ott 2015; Müller & Janka 2015), and (3) rotation (Ott et al. 2006, 2008) are certainly intriguing and demand a closer look. As we have here suggested, the initial perturbation structure has not been adequately scrutinized.

Whatever the ultimate solution to the long-standing problem of the mechanism of core-collapse supernova explosions, our study serves once again to emphasize that it is crucial to ensure the physical fidelity of numerical methods that are applied to multi-dimensional problems in which turbulence and chaotic flow are of qualitative import. The next step is for other groups to check what we have found in 2D and to test this result in the 3D context. In fact, there is a rather long list of numerical
challenges and code verification issues yet to be met collectively by the world’s supernova modelers. The results of different groups are still too far apart to lend ultimate credibility to any one of them. Perhaps our study can serve to encourage the necessary global code verification initiative.

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APPENDIX

The basic comoving-frame equations of radiative transfer that we solve are the zeroth- and first-moment equations of the full equation of radiative transfer for the specific intensity of the $\nu_{ee}$, $\bar{\nu}_{ee}$ and $\nu_{\gamma}$ neutrinos, where the latter represents the $\nu_{\mu}$, $\bar{\nu}_{\mu}$, $\nu_{\tau}$ and $\bar{\nu}_{\tau}$ neutrinos collectively. The equations are

$$E_{sx,i} + (F^k_{sx} + \nu^E_{sx})_i = \nu^E_{j} \frac{\partial}{\partial \ln \varepsilon} P_{sx,i}^j = \kappa_{sx} E_{sx},$$  

$$F_{sx,j} + (c^2 P^k_{j} + \nu^F_{sx})_j = \nu^F_{k} F_{sx,k} - \nu^E_{k} \frac{\partial}{\partial \varepsilon} (\varepsilon Q^{E}_{sx,j}),$$

where differentiation is indicated with standard notation, $\varepsilon$ is the neutrino energy, $s \in \{\nu_{ee}, \bar{\nu}_{ee}, \nu_{\gamma}\}$, $E_{sx}$ is the radiation energy density spectrum (zeroth moment), $F_{sx}$ is the radiation flux spectrum (first moment), $P_{sx,i}^j$ is the radiation pressure tensor (second moment), $Q^{E}_{sx,j}$ is the heat tensor (third moment), $\sigma^E_{sx}$ is the transport scattering opacity, and the other variables have their standard meanings. We use the M1 closure to truncate the radiation moment hierarchy by specifying the second and third moments in terms of the zeroth and first moments.

For the ray-by-ray$+$ solutions, the transverse components of the radiation flux are set to zero, while we retain the velocity advection terms, even the transverse advection terms (all shown in bold above). Of course, for both schemes, we retain the advection of $Y_e$ in all directions. In our implementation of ray-by-ray$+$, the momentum exchange with the matter due to any lateral neutrino stress is not incorporated (although the corresponding radial stress is), since these lateral force terms are very small. This is particularly true in the semi-transparent region where the overall differences between ray-by-ray$+$ and 2D transport are largest and most relevant. In the Buras et al. (2006) implementation of ray-by-ray$+$, the authors include a transverse diffusive term that approximates the lateral neutrino force above densities of $10^{12}$ g cm$^{-3}$ by a gradient term in the neutrino energy density (Eddington approximation), setting it to zero below this density. We feel that including such a term in our ray-by-ray$+$ implementation is not only unnecessary, but too approximate and ad hoc. Moreover, including such an effect is not relevant when researching the differences between the two methodologies in the shock dynamics and in the hydrodynamic outcomes.

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