Modules for Experiments in Stellar Astrophysics (MESA): Convective Boundaries, Element Diffusion, and Massive Star Explosions

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Abstract

We update the capabilities of the software instrument Modules for Experiments in Stellar Astrophysics (MESA) and enhance its ease of use and availability. Our new approach to locating convective boundaries is consistent with the physics of convection, and yields reliable values of the convective-core mass during both hydrogen- and helium-burning phases. Stars with $M < 8 \, M_\odot$ become white dwarfs and cool to the point where the electrons are degenerate and the ions are strongly coupled, a realm now available to study with MESA due to improved treatments of element diffusion, latent heat release, and blending of equations of state. Studies of the final fates of massive stars are extended in MESA by our addition of an approximate Riemann solver that captures shocks and conserves energy to high accuracy during dynamic epochs. We also introduce a 1D capability for modeling the effects of Rayleigh–Taylor instabilities that, in combination with the coupling to a public version of the STELLA radiation transfer instrument, creates new avenues for exploring Type II supernova properties. These capabilities are exhibited with exploratory models of pair-instability supernovae, pulsational pair-instability supernovae, and the formation of stellar-mass black holes. The applicability of MESA is now widened by the capability to import multidimensional hydrodynamic models into MESA. We close by introducing software modules for handling floating point exceptions and stellar model optimization, as well as four new software tools—MESA – Web, MESA-Docker, pyMESA, and mesastar.org—to enhance MESA’s education and research impact.

Key words: convection – diffusion – hydrodynamics – methods: numerical – stars: evolution – supernovae: general

Supporting material: tar.gz files

1. Introduction

Over the next decade, multimessenger astronomy will probe the rich stellar astrophysics of transient phenomena in the sky, including gravitational waves from the mergers of neutron stars and black holes, light curves and spectra from core-collapse supernovae, and the oscillation modes of stars. On the observational side of this new era, the Laser Interferometer Gravitational-Wave Observatory (LIGO) has demonstrated the existence of binary stellar-mass black hole systems (Abbott et al. 2016a, 2016b, 2016c, 2017a, 2017b) and continues to monitor the sky with broadband detectors for gravitational waves from compact binary inspirals and asymmetrical exploding massive stars (Fryer et al. 2002; Gossan et al. 2016; Abbott et al. 2017a, 2016d, 2016e, 2016f). The Gaia Data Release 1, containing about one billion stars, begins the process of converting the spectrophotometric measurements to distances, proper motions, luminosities, effective temperatures, surface gravities, and elemental compositions (Gaia Collaboration et al. 2016a, 2016b). This stellar census will provide the observational data to tackle a range of questions related to the origin, structure, and evolutionary history of stars in the Milky Way (Creevey et al. 2015; Sacco et al. 2015; Lindegren et al. 2016; van Leeuwen et al. 2017). The Neutron star Interior Composition Explorer (NICER) mission, delivered to the International Space Station in 2017 June, will provide rotation-resolved spectroscopy of the thermal and nonthermal emissions of neutron stars in the soft X-ray band with over 15 million seconds of exposures (Gendreau et al. 2012; Arzoumanian et al. 2014; Gendreau et al. 2016), opening a new window into the interior structure of and the dynamics that underlie neutron stars (e.g., Miller 2016; Özel et al. 2016). With first light at Palomar Observatory in 2017, the Zwicky Transient Facility (Kulkarni 2016) will scan more than 3750 deg2 hr−1 to a depth of about 20 mag to discover young supernovae less than 24 hr after explosion each night, hunt for electromagnetic counterparts of gravitational-wave events (Ghosh et al. 2017), and search for rare and exotic transients. Repeated imaging of the northern sky, including the Galactic Plane, will produce a photometric variability catalog with nearly 300 observations each year (Laher et al. 2017) for detailed studies of variable stars and binary systems. From its unique high Earth orbit, the
Transiting Exoplanet Survey Satellite aims to survey about 200,000 nearby G-, K-, and M-type stars with apparent magnitudes brighter than about 12 mag with a 1 minute cadence across a 400 deg$^2$ area of the sky (Rickert et al. 2016; Sullivan et al. 2015, 2017) to open a new era of stellar variability studies. The Large Synoptic Survey Telescope will image the entire southern hemisphere deeply in multiple optical colors every week with a 3.5 deg$^2$, three billion pixel digital camera (LSST Science Collaboration et al. 2017) to open new perspectives on transient objects such as tidal disruption events (Bade et al. 1996; Stern et al. 2004; Arcavi et al. 2014; Komossa 2015) and interacting close binary systems (Oloffsey et al. 2012; Korol et al. 2017). The Jiangmen Underground Neutrino Observatory will usher in a new generation of multipurpose neutrino detectors (Li 2014; Brugière 2017) designed in part to open a new avenue into neutrinos from presupernova massive stars (e.g., Odrzywolek 2009; Misch & Fuller 2016; Paxton et al. 2017a, 2017b) and core-collapse supernova explosions (e.g., Hirata et al. 1987; Janka 2017).

This ongoing explosion of activity in multimessenger stellar astronomy powers theoretical and computational developments, in particular the evolution of the community software instrument Modules for Experiments in Stellar Astrophysics (MESA), for research and education. We introduced MESA in Paxton et al. (2011), hereafter Paper I and significantly expanded its range of capabilities in Paxton et al. (2013, hereafter Paper II) and Paxton et al. (2015, hereafter Paper III). These previously published papers, as well as this one, are “instrument” papers that describe the capabilities and limitations of MESA while also comparing to other available numerical or analytic results. This paper describes the major new advances in MESA for modeling convective boundaries, element diffusion, implicit shock hydrodynamics, massive star explosions and light curves, pulsational pair-instability supernovae, and black hole formation. We do not fully explore these results and their implications here. The scientific potential of these new capabilities will be unlocked in future work through the efforts of the MESA user community.

The convective regions of stars remain a rich site of fascinating challenges, including the interplay among mixing, composition gradients, and element diffusion. A convection region transports energy through the vertical exchange of matter. The location where the radial velocity of the bulk motions goes to zero is a natural way to define the edge of a convection region (Vitense 1953; Böhm-Vitense 1958). It is necessary to ensure that convective boundaries are properly positioned (e.g., Eggleton 1972; Gabriel et al. 2014), because their exact placement can have a strong influence on the evolution of the stellar model (Salaris & Cassisi 2017). An important new addition to MESA is an improved treatment of convective boundaries, allowing them to evolve toward a state where the radiative gradient equals the adiabatic gradient on the convective side of the boundary. As a consequence, the Schwarzschild and Ledoux criteria now give the same position for convective boundaries.

Gradients can drive changes in the composition profile of a star. For example, if gradients occur in the concentrations of chemical elements, then diffusion tends to smooth out the differences. Temperature gradients can push heavier species toward regions of higher temperature, while pressure gradients can propel heavier species to diffuse toward regions of higher pressure (Thoul et al. 1994; Hansen et al. 2004; Kippenhahn et al. 2012; Michaud et al. 2015). Treatments of diffusion typically assume that all diffusing species are ideal gases (e.g., Burgers 1969; Thoul et al. 1994). For white dwarf interiors and neutron star envelopes, degenerate electrons violate this assumption (Deloye & Bildsten 2002; Chang et al. 2010). In addition, strong Coulomb coupling in plasmas requires modifications to the binary scattering formalism to calculate cross-sections used to obtain diffusion coefficients (Paquette et al. 1986a; Daligault et al. 2016; Stanton & Murillo 2016; Shaffer et al. 2017). MESA’s extensions of element diffusion for degenerate and strongly coupled plasmas open a pathway into the regime relevant to the sedimentation in the interiors of white dwarfs (Iben & MacDonald 1985; Iben et al. 1992; Koester 2009; Hollands et al. 2017) and the surfaces of neutron stars (Chang & Bildsten 2003, 2004; Beznogov et al. 2016).

Massive ($M \geq 8 \, M_\odot$) stars explode when the energy from the collapse of their core into a compact object emerges as an outgoing shock wave into the outer parts of the star. The outward propagation of this shock wave generates Rayleigh–Taylor instabilities that can mix material behind the shock front (Chevalier 1976; Chevalier & Klein 1978; Weaver & Woosley 1980; Benz & Thielemann 1990; Herant & Benz 1991; Hammer et al. 2010; Wongwathanarat et al. 2015; Utrobin et al. 2017). The resulting light curves of Type II supernovae can be subdivided into multiple classes, but we focus here on Type IIP supernovae (e.g., Smartt 2009, 2015; Smith et al. 2016). Our improvements to MESA—implicit shock-capturing hydrodynamics, Rayleigh–Taylor instability modeling in 1D (Duffell 2016), and radiative transfer using the public version of the STELLA instrument (Blinnikov & Sorokina 2004; Baklanov et al. 2005; Blinnikov et al. 2006)—open up new avenues for researching the diverse set of Type II supernovae.

Pair instability leads to partial collapse, which in turn causes runaway thermonuclear burning in the carbon–oxygen core (Fowler & Hoyle 1964; Barkat et al. 1967; Rakavy & Shaviv 1967; Rakavy et al. 1967; Fraley 1968). A wide variety of outcomes is possible depending on the star’s mass and rotation. A single energetic burst from nuclear burning can disrupt the entire star without leaving a black hole remnant behind to produce a pair-instability supernova (Ober et al. 1983; Fryer et al. 2001; Scannapieco et al. 2005; Kasen et al. 2011; Chatzopoulos et al. 2013). Alternatively, a series of bursts can trigger a cyclic pattern of nuclear burning, expansion, and contraction, leading to a pulsational pair-instability supernova that leaves a black hole remnant (Barkat et al. 1967; Woosley et al. 2007a; Chatzopoulos & Wheeler 2012; Limongi 2017; Woosley 2017). Many of these variations can now be explored in MESA, as can lower-mass progenitors that do not pulse before collapsing into a black hole.

MESA is a community-driven software instrument for stellar astrophysics. New directions will be motivated by features useful to the MESA user community, advances in the physics modules, algorithmic developments, and architectural evolution. Potential examples for expanding MESA’s scientific, computational, and educational capabilities include seamlessly leveraging many-core architectures, an improved treatment of the equation of state, Jupyter/Python notebooks for education, and continued integration with software instruments useful to the astronomy and astrophysics community. Examples include ADIPLS (Christensen-Dalsgaard 2008; Christensen-Dalsgaard & Thompson 2011), CYRE (Townsend & Teitler 2013), and...
STELLA (Blinnikov et al. 1998; Blinnikov & Sorokina 2004; Balkanov et al. 2005; Blinnikov et al. 2006).

This paper is organized as follows. Section 2 introduces a new treatment of convective boundaries. In Section 3, we present an implementation of element diffusion that accounts for electron degeneracy and strongly coupled interactions. Section 4 describes the Riemann solver for shock capturing in MESA’s new implicit hydrodynamics solver, and Section 5 presents a model for approximating the 3D effects of the Rayleigh–Taylor instability. In Section 6, we introduce the coupling of MESA and an implementation of the STELLA radiative transfer instrument to explore the modeling of Type IIP supernova light curves from post-explosion to post-plateau. In Section 7, we show advances in modeling pair-instability supernovae, pulsation pair-instability supernovae, and black hole formation. Section 8 discusses energy accounting in stellar evolution.

Appendix A discusses improvements to estimating a model’s absolute magnitude in a chosen color filter, Appendix B offers guidance on importing multidimensional models into MESA, and Appendix C details the implementation of element diffusion in MESA. Appendix D introduces two new software modules for handling floating point exceptions and stellar model optimization, as well as four new software tools for education and research: MESA – Web, MESA-Docker, pyMESA, and mesestar.org.

Important symbols are defined in Table 1. Acronyms used are denoted in Table 2. We denote components of MESA, such as modules and routines, in typewriter font, e.g., "colors.

### 2. Convective Boundaries

Gabriel et al. (2014) discuss the correct positioning of convective boundaries in stellar evolution models. Following earlier work (e.g., Roxburgh 1978), they argue that a convective boundary should be defined as the point where the convective velocity vanishes. Within local mixing length theory (MLT), this condition is equivalent to the requirement \( \nabla_{\text{rad}} = \nabla_{\text{ad}} \), where \( \nabla_{\text{rad}} \) and \( \nabla_{\text{ad}} \) are the radiative and adiabatic temperature gradients, respectively. Critically, this equality must be satisfied on the **convective** side of the boundary, because the MLT convective velocity is only well-defined there. Moreover, because the fluid on the convective side is presumed to be well-mixed, the Ledoux temperature gradient \( \nabla_L = \nabla_{\text{ad}} + B \) (Equation (11) of Paper II) can play no part in setting the location of the boundary.

If the chemical composition is continuous across the convective boundary, then so too are \( \nabla_{\text{rad}} \) and \( \nabla_{\text{ad}} \), and requiring \( \nabla_{\text{rad}} = \nabla_{\text{ad}} \) on the convective side of the boundary results in the same equality on the radiative side. However, a composition discontinuity produces a jump in density and opacity, and in turn a discontinuity in \( \nabla_{\text{rad}} \) and \( \nabla_{\text{ad}} \). Hence, it is generally the case that \( \nabla_{\text{rad}} \neq \nabla_{\text{ad}} \) on the radiative side of the boundary.

In numerical codes based on discrete grids, the nuance of the foregoing discussion is often overlooked in favor of a simple approach for locating convective boundaries based on sign changes in the discriminant \( y = \nabla_{\text{rad}} - \nabla_{\text{ad}} \) (or \( y = \nabla_{\text{rad}} - \nabla_L \), if the Ledoux stability criterion is used). This approach works well when the chemical composition remains continuous, but is problematic when the composition—and hence \( y \)—is discontinuous at the boundary; it typically leads to configurations where \( \nabla_{\text{rad}} > \nabla_{\text{ad}} \) on the convective side, which is unphysical and ultimately retards the growth of the convective region. Previous versions of MESA have taken this approach; the outcome is evident in Figure 15 of Paper II, which shows the

| Name         | Description                          | First Appears |
|--------------|--------------------------------------|---------------|
| A            | Area of face                          | 4.1           |
| C            | Concentration                         | 3.1           |
| e            | Specific thermal energy               | 4.1           |
| F            | Flux across cell face                 | 4.1           |
| \(\gamma\)   | Adiabatic index                       | 4.4.1         |
| \(\Gamma\)   | Plasma coupling parameter             | 8.5           |
| K            | Resistance coefficient                | 3.1           |
| \(\lambda\)  | Screening length                      | 3.3           |
| m            | Baryonic mass coordinate              | 2.2           |
| M            | Stellar mass                          | 2.2           |
| \(\mu\)      | Chemical potential                    | 8.1           |
| \(\Phi\)     | Gravitational potential               | 8.3           |
| q            | Specific heat                         | 8.1           |
| r            | Radial coordinate                     | 3.1           |
| s            | Specific entropy                       | 8.1           |
| S            | Wave speed                            | 4.1           |
| u            | Cell-centered velocity                | 4.1           |
| w            | Diffusion velocity                    | 3.1           |
| z            | Resistance coefficient                | 3.1           |
| \(\bar{A}\)  | Average atomic number                 | 8.1           |
| \(\delta_{\text{MLT}}\) | Mixing length of MLT                  | 6.7.1         |
| \(c_p\)      | Specific heat at constant pressure    | 8.2           |
| \(c_s\)      | Sound speed                           | 4.1           |
| \(c_v\)      | Specific heat at constant volume      | 8.2           |
| \(\delta t\) | Numerical time step                   | 2.6           |
| dm           | Mass of cell                          | 4.1           |
| \(\delta m\) | Mass at cell face                     | 4.1           |
| \(D_{\text{RL}}\) | Rayleigh–Taylor decay coefficient     | 5.1           |
| \(\epsilon_{\text{ion}}\) | Specific ionization energy           | 8.4           |
| \(\epsilon_{\text{rad}}\) | Blast energy                        | 4.4.1         |
| \(\epsilon_{\text{extra}}\) | Extra specific heating/cooling rate   | 4.1           |
| \(\epsilon_{\text{grav}}\) | Gravitational heating rate           | 8.1           |
| \(\epsilon_{\nu}\) | Neutrino energy loss rate             | 4.1           |
| \(\epsilon_{\text{mc}}\) | Nuclear energy generation rate       | 4.1           |
| \(\epsilon_{\text{pot}}\) | Convective overshoot parameter       | 6.7.1         |
| \(\Gamma_1\) | First adiabatic index                 | 7             |
| \(\nabla_{\text{ad}}\) | Adiabatic temperature gradient       | 2             |
| \(\nabla_{\text{ad}}\) | Ledoux temperature gradient          | 2.1           |
| \(\nabla_{\text{rad}}\) | Radiative temperature gradient       | 2             |
| \(\nabla_T\) | Temperature gradient from MLT         | 4.1           |
| \(N_B\)      | Number of baryons                     | 8.1           |
| \(P_{\text{ad}}\) | Gas pressure                         | 8.2           |
| \(P_{\text{rad}}\) | Radiation pressure                   | 8.2           |
| \(P\)        | Pressure at cell face                 | 4.1           |
| q_e          | Electric charge                       | 3.2.2         |
| \(\rho_e\)   | Charge density                        | 3.1           |
| \(T\)        | Temperature at cell face              | 4.1           |
| \(\tau_{\text{Ros}}\) | Rosseland optical depth             | 6.5           |
| \(\tau_{\text{opt}}\) | Sobolev optical depth                | 6.5           |
| \(\lambda_c\) | (\delta log P/\delta log \rho)_{\text{tr}} | 8.2           |
| \(\chi_T\)   | (\delta log P/\delta log T)_{\text{tr}} | 8.2           |
| Z            | Average ion charge                    | 8.1           |

**Note.** Single character symbols are listed first; symbols with modifiers are listed second. Some symbols may be further subscripted by \(c\) (indicating a central quantity), by a cell index \(k\), or by an index that runs over species \(i, j, s, \) or \(t\).
convective-core mass as a function of age during the He-burning evolution of a $3 \, M_\odot$ star. In the model with no overshoot and the Schwarzschild stability criterion, the core grows only modestly in mass before reaching a plateau. Inspection of the model confirms that $\nabla_{\text{rad}} > \nabla_{\text{ad}}$ on the convective side of the core boundary, signifying that core growth is being impeded.

Gabriel et al. (2014) highlight a further issue with this simple sign-change approach, whereby the location of a convective boundary is not uniquely determined but rather depends on the mixing history near the boundary. We have confirmed that this issue is present in MESA when using the sign-change approach. This manifests itself as a lack of convergence in some models (e.g., the $3 \, M_\odot$ He-burning example) when the resolution is increased and/or the time step shortened.

To resolve these issues, we implement a new “predictive mixing” scheme in MESA. It is inspired both by the “maximal overshoot” scheme introduced by Constantino et al. (2015) and by the procedure described by Bossini et al. (2015). In the new scheme, the extent of a convection region is allowed to expand at each time step until the boundaries reach the point where $\nabla_{\text{rad}} = \nabla_{\text{ad}}$ on their convective side. We describe the new scheme in detail in the following section and then present the results obtained with this scheme in four scenarios: a growing convective core in a low-mass star on the main sequence (MS), a retreating convective core in a high-mass star on the MS, growing He-burning cores in intermediate- and low-mass stars, and a surface convective region in a low-mass star on the MS. In all cases, we assume an initial He mass fraction $Y = 0.28$ and an initial metal mass fraction $Z = 0.02$, and we ignore rotation and mass loss.

### 2.1. Predictive Mixing

The MESA predictive mixing scheme initially proceeds in the same manner as the simple sign-change approach, by finding the cells where $y > 0$ on one face (convective) and $y < 0$ on the other face (radiative). For each of these candidate boundary cells, the algorithm considers how $y$ would change if the cell were completely mixed with the rest of the adjoining convection region. This prediction involves re-evaluating the opacities, densities, and other data throughout the mixed region, under the assumption that the composition is completely uniform. If $y$ becomes positive on both faces of the candidate boundary cell, then the adjacent cell in the radiative region becomes the new candidate boundary cell, and a new round of predictive mixing begins. The process continues iteratively until the candidate cell after the predictive mixing still has a negative $y$ on the radiative face. The code reverts to the previous candidate, identifies it as the final convective boundary cell, recalculates convective diffusivities and convective velocities using MLT, and writes these into the model for use in the composition solver (see Paper I, Section 6.2). No abundances are directly modified in the model during the predictive iterations. Below, we demonstrate that this algorithm leads MESA to a solution of the stellar structure equations in which $0 \leq y \ll \nabla_{\text{ad}}$ on the convective side of each boundary cell.

The physical justification for our predictive mixing scheme can be traced back to a narrative advanced by Castellani et al. (1971). Focusing on He core burning, these authors argue that any gentle mixing outside the core boundary irreversibly alters the composition there, and the resulting increase in opacity raises the local $\nabla_{\text{rad}}$ from sub-adiabatic to super-adiabatic. The outcome is a “self-driving mechanism for the extension of the convective region,” which continues until $\nabla_{\text{rad}} = \nabla_{\text{ad}}$ on the convective side of the core boundary. Although Castellani et al. (1971) invoked overshoot as the source of the mixing outside the boundary, Michaud et al. (2007) showed that element diffusion can play this role equally well and lead to the same outcome. For MS stars with growing convective cores, the extension of the core boundary cannot be driven in exactly the same way as the He-burning case, because helium has a lower opacity than hydrogen. However, gentle mixing outside the core boundary erases any composition gradients there, and it is the loss of these gradients—and their accompanying stabilizing effect—that drives the extension of the convective region until $\nabla_{\text{rad}} = \nabla_{\text{ad}}$ on the convective side of the core boundary.

The predictive mixing scheme does not specify the nature of the gentle mixing beyond the convective boundaries, focusing instead on its effects. Tied to this agnosticism is the presumption that the mixing-driven expansion of convective boundaries is so rapid that it can be approximated as instantaneous. This is likely a reasonable approach during core H and He burning; Castellani et al. (1971) argued that the growth of the core boundary in the latter case should proceed on a timescale that is much shorter than the burning lifetime. However, there may be circumstances where the finite timescale for boundary growth cannot be ignored.

Because uniform composition is assumed during the predictive mixing iterations, there is no functional distinction between the Schwarzschild and Ledoux criteria when evaluating the discriminant $y$. However, the preliminary search for sign changes in $y$, before any predictions are made, does take into account composition gradients when the Ledoux criterion is used. As a result, the initial candidate boundary cells can differ between the two criteria. In many cases, this difference is unimportant, with the final location of the boundaries being insensitive to the choice of criterion. The one exception is when a region with $\nabla_{\text{rad}} < \nabla_{\text{ad}} < \nabla_{\text{L}}$ is bounded on both sides by...
radiative regions; then, it will be completely overlooked during a preliminary search with the Ledoux criterion. As we shall demonstrate later, such scenarios arise in our calculations outside convective cores during MS evolution.

### 2.2. Evolution of a Growing Convective Core on the Main Sequence

We evolve a 1.5 M\(_\odot\) star from the zero age main sequence (ZAMS) to the terminal age main sequence (TAMS) using the predictive mixing scheme at the convective-core boundary; this is the same mass and evolutionary stage considered in Section 5.1.2 of Gabriel et al. (2014). Figure 1 plots the mass coordinate of the convective-core boundary as a function of MS age, showing results from separate runs using the Schwarzschild and Ledoux criteria, and from additional runs with the incremental inclusion of semi-convection (in just the Ledoux case) and then element diffusion (in both cases). The semi-convection is modeled using the Langer et al. (1985) scheme with an efficiency parameter \(\alpha_{\text{sec}} = 0.1\) (see Paper II for a complete description of the semi-convection implementation in MESA). For comparison, the figure also shows the outcome of using the Ledoux criterion but no predictive mixing; in contrast to the other cases, which broadly agree with one another, core growth is inhibited and the H-burning lifetime correspondingly truncated.

Note that in Figure 13 of Paper II, the results obtained with the Ledoux criterion show a shrinking convective core; this behavior was due to a separate problem arising from the oversmoothing of the composition gradient (see Moore & Garaud 2016) and has since been rectified in MESA. For completeness, we include this case in Figure 1.

Figure 2 plots the profiles of \(\nabla_{\text{rad}}, \nabla_{\text{ad}}, \nabla_{\text{Li}}, \) and \(X_c\), in the inner part of the 1.5 M\(_\odot\) star nearing the halfway point of its MS evolution (a core H mass fraction \(X_c = 0.42\)). In the upper row, the left panel illustrates the run with the Ledoux criterion plus predictive mixing (the dotted curve in Figure 1), while the right panel shows the run with the Ledoux criterion but without predictive mixing (the black curve, ibid.). Clearly, without predictive mixing, \(\nabla_{\text{rad}}\) remains significantly larger than \(\nabla_{\text{ad}}\) on the convective side of the boundary, which as discussed previously is physically inconsistent. When using predictive mixing, however, the profiles satisfy \(\nabla_{\text{rad}} = \nabla_{\text{ad}}\) on the convective side and closely match those seen in the left panel of Figure 6 of Gabriel et al. (2014). The small bump in \(\nabla_{\text{rad}}\) just above the boundary is Schwarzschild unstable but Ledoux stable.

The middle panels of Figure 2 show the runs with the Ledoux criterion and predictive mixing, and the incremental addition of semi-convection (left) and then element diffusion (right). Inside the core boundary, the profiles are almost identical to those shown in the upper-left panel, but just outside the boundary, semi-convection converts the composition discontinuity into a steep gradient and flattens the bump in \(\nabla_{\text{rad}}\) into a neutral \(\nabla_{\text{rad}} = \nabla_{\text{ad}}\) profile. Element diffusion further softens the abundance profile, as shown in the middle-right panel. Note that element diffusion has only a small effect on the location of the convective boundary; this is barely noticeable in Figure 2, but a slight extension of the boundary can be seen in Figure 1 toward the later part of the MS, for the two cases including diffusion.

The lower panels of Figure 2 show the runs using the Schwarzschild criterion and predictive mixing, without (left) and with (right) element diffusion. In the left panel, the abundance profile shows a chaotic staircase-like profile, due to mixing by transient convective shells that appear and disappear from one time step to the next (two of these shells can be seen in the figure). The shells do not appear in the Ledoux plots (middle and upper panels) because the region outside the core is stabilized in its entirety by the abundance gradient: \(\nabla_{\text{rad}} < \nabla_{\text{Li}}\). This serves as a good illustration of the earlier discussion (Section 2.1) of how the Schwarzschild and Ledoux criteria can sometimes lead to different outcomes. It is important to note, however, that the location of the core boundary is the same in all cases with predictive mixing; the differences only appear in the inhomogeneous region beyond the boundary, which arises from slow H burning outside the core.

The lower-right panel of Figure 2 shows that adding element diffusion removes the abundance discontinuities, replacing them with a smooth gradient. The resulting profiles appear almost identical to the Ledoux case shown in the middle-right panel of the figure (compare also the curves with diffusion in Figure 1).

### 2.3. Evolution of a Retreating Convective Core on the Main Sequence

We now evolve a 16 M\(_\odot\) star from ZAMS to TAMS using the new predictive mixing scheme at the convective-core boundary; this is the same mass and evolutionary stage considered in Section 5.1.1 of Gabriel et al. (2014). Figure 3 plots the mass of the convective core as a function of MS age, showing results from separate runs using the Schwarzschild and Ledoux criteria, and with and without predictive mixing. The agreement between these four cases is very close. However, as was the case in the preceding section, there are differences outside the convective core. These can be seen in Figure 4, which plots the profiles of \(\nabla_{\text{rad}}, \nabla_{\text{ad}}, \nabla_{\text{Li}}, \) and \(X\) near the end of the star’s MS evolution (\(X_c = 0.15\)), for the two runs with predictive mixing.

Even though both runs exhibit the same core structure, with \(\nabla_{\text{rad}} = \nabla_{\text{ad}}\) at the convective side of the core boundary, the inhomogeneous region left behind by the retreating core is very different. The H abundance obtained with the Schwarzschild criterion shows the same staircase-like profile seen in the
lower-left panel of Figure 2, again due to mixing by transient convective shells. These shells are not present when the Ledoux criterion is used, with the exception of a persistent solitary shell at the top of the inhomogeneous region (corresponding to where the core boundary was located at the ZAMS); the behavior of this shell is discussed by Gabriel et al. (2014, their Section 5.5.1; compare also against their Figure 4). Between the shell and the core boundary, the abundance profile from the Ledoux run remains relatively smooth. The different abundance profiles in the two runs will have a direct influence on the Brunt–Väisälä frequency profile, and therefore on the oscillation frequencies of the stellar model.

2.4. Evolution of the Convective Core During Core He Burning

As reviewed by Salaris & Cassisi (2017), the modeling of mixing in low- and intermediate-mass stars during core He burning (CHeB) is particularly challenging. The correct treatment of convective boundaries is complicated by the fact that the $\nabla_{\text{rad}}$ profile within the core convection region develops...
a local minimum at some point during CHeB evolution (see the middle and lower panels of Figure 6). This is a consequence of the complex behavior of the physical quantities (opacity, temperature, density, etc.) involved in the expression for $\nabla_{\text{rad}}$. With further outward propagation of the convective boundary, the mixing of fresh He into the core can lower the radiative gradient throughout the core to such an extent that $\nabla_{\text{rad}} = \nabla_{\text{ad}}$ at the local minimum of $\nabla_{\text{rad}}$. When this happens, the part of the convection region interior to the minimum becomes decoupled from the part exterior to the minimum: the convection region has split. This phenomenon was first discussed by Eggleton (1972), and a variety of ad hoc approaches have been proposed to follow the subsequent evolution, mostly focused around the narrative that the exterior part undergoes partial mixing with the adjacent radiative region until it reaches convective neutrality (see, e.g., Castellani et al. 1985 and references therein).

Another problem appears near the end of CHeB. At that point, even small amounts of He added to the core (which is almost totally depleted in He) will enhance the rate of energy production and thus the luminosity, resulting in an increase in $\nabla_{\text{rad}}$. This increase leads to a sudden growth in the core boundary and a “breathing pulse.” The He is then quickly burned in the core, and the star re-adjusts itself. The existence of these breathing pulses remains controversial, and it is still unclear whether they are numerical or physical (Caputo et al. 1989; Cassisi et al. 2003; Farmer et al. 2016; Constantino et al. 2017). All of these problems are clearly described and illustrated in Salaris & Cassisi (2017). To manage these complexities, the predictive mixing scheme must be modified. When a convection region splits, it is no longer meaningful to re-evaluate $\nabla$ using opacities and other data calculated on the assumption of uniform composition throughout (Section 2.1), because the radiative region appearing at the splitting point prevents the free exchange of material between the adjacent convection regions. Although in principle we could resort to the partial mixing mentioned above, in practice it is not clear how this might be implemented within a diffusive mixing framework. Constantino et al. (2015) have developed an overshoot-like prescription that appears useful for mimicking the convective neutrality achieved by partial mixing (see their Section 2.3.3), but it involves a number of unconstrained parameters. Therefore, on the grounds of simplicity and pragmatism—and recognizing that better approaches may become apparent in the future—we modify the predictive scheme to prevent it from causing a convection region to split in the first place. This involves a new control parameter, $\text{predictive\_superad\_thresh}$; if during the predictive mixing iterations the super-adiabaticity $\nabla_{\text{rad}}/\nabla_{\text{ad}} - 1$ drops below this threshold anywhere in the mixed region, then the code backs off the mixing by one cell and updates the model convective diffusivities and convective velocities in the usual manner.

Further functionality, controlled by a new parameter $\text{predictive\_avoid\_reversal}$, also helps to prevent splitting and breathing pulses. When this parameter is set to the name of a MESA isotope, the code then monitors how the predictive mixing alters the abundance evolution of that isotope in the convection region. If it would cause this evolution to reverse (i.e., switch from decreasing to increasing, or vice versa), then the code backs off the mixing by one cell and updates the model as before. Thus, for instance, setting this parameter to “he4” during CHeB ensures that the predictive mixing scheme does not cause the core He abundance to increase across a time step.

To illustrate the preceding discussion, we evolve a 1 $M_{\odot}$ star through CHeB; this is the same mass considered by Constantino et al. (2015). Figure 5 plots the mass of the convective core as a function of CHeB age (defined as the time elapsed since the central $Y$ drops below 0.98), showing results from separate runs with and without predictive mixing, and using the Schwarzschild and Ledoux criteria. For the cases with predictive mixing, we adopt a value of 0.005 for the $\text{predictive\_superad\_thresh}$ parameter and set $\text{predictive\_avoid\_reversal}$ to “he4” to prevent any reversal in the core He abundance. Figure 5 also shows the results from an additional Ledoux/predictive mixing run where we allow the core to split by not setting the $\text{predictive\_avoid\_reversal}$ and $\text{predictive\_superad\_thresh}$ controls.

Figure 5 shows that without predictive mixing, the core is prevented from growing and the CHeB lifetime is significantly curtailed, irrespective of whether the Schwarzschild or Ledoux criteria are used (see also Figure 15 of Paper II). With predictive mixing but no splitting allowed, however, the core grows steadily until He is exhausted, and no breathing pulses are seen. There is almost no difference between the Schwarzschild and Ledoux cases. When the core is allowed to split, the evolution is much noisier. Starting at an age $\approx 25$ Myr, the core undergoes episodes of splitting and rejoicing that repeat on a short timescale. Toward the end of the evolution, as the core He abundance becomes very small, the timescale between successive splittings becomes longer, until the core finally splits without rejoicing. The overall CHeB lifetime of the model is shortened by $\approx 6$ Myr relative to the cases where splitting is avoided.

Figure 6 plots the profiles of $\nabla_{\text{rad}}$, $\nabla_{\text{ad}}$, and $Y$ for the 1 $M_{\odot}$ star at three points during its CHeB evolution, corresponding to core He mass fractions $Y_c = 0.9$, 0.6, and 0.3. The profiles are all from the run with the Ledoux criterion and predictive mixing. In the upper panel, a local minimum in $\nabla_{\text{rad}}$ has yet to develop, and the core boundary satisfies the $\nabla_{\text{rad}} = \nabla_{\text{ad}}$ equality on its convective side. In the middle and lower panels, the local minimum in $\nabla_{\text{rad}}$ can clearly be seen; in these cases, predictive mixing has extended the convection region as far as possible.
of the convective-core boundary plotted as a function of mass coordinate, in the inner part of the 16 M\(_\odot\) stellar model at X\(_c\) = 0.15. The panels show the separate runs described in the text. Gray shading indicates regions undergoing convection. Both models use predictive mixing at the convective-core boundary.

Figure 4. Profiles of \(\nabla_{\text{rad}}, \nabla_{\text{ad}}, \nabla_L\), and \(X\) as a function of mass coordinate, in the inner part of the 16 M\(_\odot\) stellar model at X\(_c\) = 0.15. The panels show the separate runs described in the text. Gray shading indicates regions undergoing convection. Both models use predictive mixing at the convective-core boundary.

Figure 5. Mass coordinate \(m_{\text{bd}}\) of the convective-core boundary plotted as a function of CHeB age, for the 1 M\(_\odot\) stellar model discussed in Section 2.4. Different line styles/colors show the separate runs described in the text. Gray shading indicates regions undergoing convection. Both models use predictive mixing at the convective-core boundary.

Without pushing the minimum \(\nabla_{\text{rad}}\) below the threshold set by the predictive superad threshold control. MESA treats the region between the \(\nabla_{\text{rad}}\) minimum and the convective boundary as fully convective. On the convective side of this boundary, \(\nabla_{\text{ad}} > \nabla_{\text{rad}}\), which is physically inconsistent but cannot be remedied with predictive mixing alone: any further extension of the boundary would cause the convection region to split. As discussed above, fixing this inconsistency requires some way of modeling the partial mixing expected to occur in the part of the convection region between the \(\nabla_{\text{rad}}\) minimum and the boundary. The abundance profiles plotted in Figure 6 show a sharp transition between the He-depleted core and the He-rich radiative region above. Although not shown, the carbon and oxygen abundance profiles exhibit corresponding jumps at the core boundary. Similar results are obtained by Constantino et al. (2015) with their “maximal overshoot” scheme (cf., their Figure 2), and those authors also find a core mass evolution during CHeB that closely resembles the outcome from predictive mixing (cf., their Figure 8 and our Figure 5). These similarities are not coincidental; although the predictive mixing and maximal overshoot schemes have different narratives and implementations, both have the effect of growing the core boundary during CHeB to the greatest extent permitted without causing the convection region to split.

The larger cores that result from this growth appear to provide a better match to Kepler asteroseismic period spacings, when compared with other mixing schemes that produce smaller cores (Constantino et al. 2015), and with certain assumptions about the post-CHeB evolution, the larger cores can also provide a satisfactory fit to observational cluster counts (Constantino et al. 2016).

To explore whether the predictive mixing performs equally well for a higher-mass star that has not passed through the He flash, we also evolve a 3 M\(_\odot\) star through CHeB; this is the same mass and evolutionary stage considered in Figure 15 of Paper II. Figure 7 plots the mass of the convective core as a function of CHeB age, showing results from separate runs with and without predictive mixing, and using the Schwarzschild and Ledoux criteria. For the cases with predictive mixing, we again adopt a value of 0.005 for the predictive_superad_thresh parameter and set predictive_avoid_reversal to “he4” to prevent any reversal in the core He abundance. As before, we find that predictive mixing allows the core to grow steadily and that the Schwarzschild and Ledoux criteria give essentially the same outcome. As a visual summary of how predictive mixing influences a star’s evolution, Figure 8 plots evolutionary tracks of the 3 M\(_\odot\) model in the Hertzsprung–Russell (HR) diagram for the same combinations of mixing and stability criteria considered in Figure 7. The left panel focuses on the MS and red giant branch phases and the right panel on the CHeB phase. In the left panel, the case with the Ledoux criterion but without predictive mixing stands out from the other three as having a slightly reduced luminosity. This behavior arises because the boundary of the hydrogen-burning convective core is incorrectly positioned during the early MS evolution, retarding the growth of the core (the same effect can be seen for the 1.5 M\(_\odot\) model in the upper panels of Figure 2). During the subsequent CHeB phase, all four tracks are similar until slightly after the luminosity minimum, when the He-burning convective core starts to grow; this growth is retarded in both cases without predictive mixing, leading to reduced luminosities and the shorter

Figure 7 plots the mass of the convective core as a function of CHeB age, showing results from separate runs with and without predictive mixing, and using the Schwarzschild and Ledoux criteria. For the cases with predictive mixing, we again adopt a value of 0.005 for the predictive_superad_thresh parameter and set predictive_avoid_reversal to “he4” to prevent any reversal in the core He abundance. As before, we find that predictive mixing allows the core to grow steadily and that the Schwarzschild and Ledoux criteria give essentially the same outcome. As a visual summary of how predictive mixing influences a star’s evolution, Figure 8 plots evolutionary tracks of the 3 M\(_\odot\) model in the Hertzsprung–Russell (HR) diagram for the same combinations of mixing and stability criteria considered in Figure 7. The left panel focuses on the MS and red giant branch phases and the right panel on the CHeB phase. In the left panel, the case with the Ledoux criterion but without predictive mixing stands out from the other three as having a slightly reduced luminosity. This behavior arises because the boundary of the hydrogen-burning convective core is incorrectly positioned during the early MS evolution, retarding the growth of the core (the same effect can be seen for the 1.5 M\(_\odot\) model in the upper panels of Figure 2). During the subsequent CHeB phase, all four tracks are similar until slightly after the luminosity minimum, when the He-burning convective core starts to grow; this growth is retarded in both cases without predictive mixing, leading to reduced luminosities and the shorter
CHeB lifetimes seen in Figure 7. For the cases with predictive mixing, there is no difference between the Schwarzschild and Ledoux criteria, either on the MS or after.

2.5. Evolution of the Bottom of the Surface Convective Region in a Low-mass Star

We now evolve a $1 M_{\odot}$ star from ZAMS to TAMS, using the predictive mixing scheme to position the lower boundary of the convective envelope. We include element diffusion in these calculations; when it is excluded, the composition remains completely uniform throughout the stellar envelope, and predictive mixing makes no difference whatsoever to the evolution. Figure 9 plots the mass coordinate of the convective boundary as a function of MS age, showing the results from separate runs with and without predictive mixing, and using the Schwarzschild and Ledoux criteria.

The four runs are in agreement until an age of $\approx 6.5$ Gyr; after this point, the downward growth of the region boundary is slower in the run that does not include predictive mixing with the Ledoux criterion. Figure 10 plots the profiles of $\nabla_{\text{rad}}$, $\nabla_{\text{ad}}$, $\nabla_{L}$, and $X$, in the outer part of the $1 M_{\odot}$ star at an age of 8.40 Gyr. The left panel illustrates the run with the Ledoux criterion plus predictive mixing, while the right panel shows the run with the Ledoux criterion but without predictive mixing. The former shows that $\nabla_{\text{rad}} \approx \nabla_{\text{ad}}$ on the convective (upper) side of the convective boundary, while the latter has $\nabla_{\text{rad}} > \nabla_{\text{ad}}$, consistent with the boundary growth being retarded.

2.6. Effect of Time Steps and Mesh Size

We now demonstrate how limiting the maximum time step $\delta t_{\text{max}}$ (set by the max_years_for_timestep control) and changing the mesh resolution parameter $\Delta$ (set by the mesh_delta_coeff control; see Section B.4 of Paper II for further details) influence the results presented in the previous sections. First, we consider the effects of changing time step and resolution on the position of the convective envelope boundary in the $1 M_{\odot}$ model considered in Section 2.5, focusing specifically on the case with the Ledoux criterion and predictive mixing. The results presented previously in Figure 9 are calculated using $\delta t_{\text{max}} = 5$ Myr and $\Delta = 0.5$. Figure 11 demonstrates that halving either $\delta t_{\text{max}}$ or $\Delta$ has little effect on these results, confirming that the calculations are converged. Such settings need to be applied when a converged result is desired from MESA for this calculation. Figures 12 and 13 repeat this exercise for the position of the core convection boundary in the $1.5 M_{\odot}$ MS model and $1 M_{\odot}$ CHeB model, respectively. The results presented previously are clearly converged, and this exercise clarifies the MESA settings that should be used for this calculation.
3. Element Diffusion

Section 9 of Paper III describes in detail the old implementation of element diffusion in MESA. Section 9.3.4 points out limitations to those methods, namely:

1. electron degeneracy was not properly accounted for in the diffusion equations and
2. strong Coulomb interaction introduced theoretical uncertainties for the diffusion coefficients.

These two issues are especially important when modeling diffusion in WDs. Here we describe the impact of degeneracy and present new methods to incorporate its effects. We also discuss recent updates to diffusion coefficients and potential approaches for further improvements.

3.1. Degeneracy and the Approach in Paper III

The approach to diffusion presented in Section 9 of Paper III assumes all particles obey the ideal gas law. Electron degeneracy pressure can significantly modify the EOS and violate this assumption.

For a plasma species $s$ (i.e., electrons and ions) with partial pressure $P_s$, mass density $\rho_s$, charge density $e_s$, number density $n_s$, and temperature $T$, the Burgers (1969) equations for diffusion are

$$\frac{dP_s}{dr} + \rho_s g - \rho_s E = \sum_{s' \neq s} K_{s's'} (w_{s'1} - w_{s1}) + \sum_{s' \neq s} K_{s's'} \frac{m_{s1} n_{s'1} - m_{s'1} n_{s1}}{m_s + m_{s'}}$$

(1)

$$\frac{5}{2} \frac{n_k k_B}{m_s} \frac{dT}{dr} = -\frac{2}{5} K_{s's'} z_{s's'}^{\mu} r_s - \frac{5}{2} \sum_{s' \neq s} K_{s's'} \frac{m_{s'1} z_{s's'}^{\mu}}{m_s + m_{s'}} (w_{s'1} - w_{s1})$$

$$- \sum_{s' \neq s} K_{s's'} \left[ \frac{3m_s^2 + m_{s'1}^2}{(m_s + m_{s'})^2} + \frac{4}{5} \frac{m_{s'1} m_s}{(m_s + m_{s'})^2} \frac{z_{s's'}^{\mu}}{r_s} \right] r_s$$

$$+ \sum_{s' \neq s} K_{s's'} \frac{m_{s'1} m_s}{(m_s + m_{s'})^2} \left[ 3 + z_{s's'}^{\mu} - \frac{4}{5} \frac{z_{s's'}^{\mu}}{r_s} \right] r_s.$$  (2)

The resistance coefficients $K_{s's'}$, $z_{s's'}$, and $z_{s's'}^{\mu}$ are defined in Equation (86) of Paper III. With $S$ representing the total number of plasma species, we must solve for $2S + 2$ unknowns: $S$ diffusion velocities $(w_s)$, $S$ heat flow vectors $(r_s)$, the electric field $(E)$, and the gravitational acceleration $(g)$. The Burgers equations above for each species provide $2S$ equations, so we can close the system with two additional constraints, which are no net flow of mass or electric current due to diffusion,

$$\sum_s \rho_s w_s = 0,$$  (3)

$$\sum_s \rho_e w_s = 0.$$  (4)

This gives a total of $2S + 2$ equations.

When electrons are degenerate, Equation (1) is difficult to apply since $dP_e/dr$ no longer takes a simple analytic
form. Moreover, the temperature term appearing on the left-hand side of Equation (2) clandestinely assumes an ideal gas law. Burgers (1969) defines the temperature for each species as 

$$T_p = \frac{P}{n_p k_B}$$

and assumes thermal equilibrium between all species so that 

$$T = T_p.$$

The quantities $P$ and $n_p$ are defined in terms of moments of a Maxwellian distribution function, but the Fermi–Dirac distribution function for electrons no longer reduces to a Maxwellian form when they are degenerate, and hence 

$$T_e = \frac{P_e}{n_e k_B}.$$ 

If the electrons remain in thermal equilibrium with their surroundings while failing to satisfy an ideal gas relation for their temperature, the Burgers treatment assigns an incorrect temperature to degenerate electrons for the $dT/dr$ term in Equation (2).

Furthermore, the approach to diffusion described in Paper III follows Thoul et al. (1994) in rearranging and rescaling all equations into one matrix system with units convenient for solving numerically,

$$\frac{P}{K_0} \left( \sum_{i=1}^{S+2} \frac{d \ln P}{dr} \right) = \sum_{j=1}^{S+2} \Delta_{ij} W_j.$$  \hspace{1cm} (5)$$

The sum on the left-hand side skips the electron index because $C_e \equiv 1$ by construction, and so we save resources by not evaluating its gradient unnecessarily. Here, indices $i = 1, 2, \ldots, S$ encode the $S$ equations given by Equation (1), indices $i = S + 1, S + 2, \ldots, 2S$ encode the $S$ equations given by Equation (2), and indices $i = 2S + 1, 2S + 2$ encode the two constraints of no current or mass flux. For the definitions of the various coefficients and matrices in Equation (5), consult Paper III and Thoul et al. (1994). We repeat a few particularly relevant definitions here. First, let $C_i = n_i / n_e$ denote the species concentration, where $n_e$ is the electron number density.
Second, define the total concentration as \( C = \sum n_i \). Then, the quantity \( \alpha_i \) appearing in Equation (5) above is defined as

\[
\alpha_i = \begin{cases} 
C_i / C & i = 1, 2, \ldots, S, \\
0 & i = S + 1, \ldots, 2S + 2.
\end{cases}
\]  

(6)

The term \( \alpha_i d \ln P / dr \) in Equation (5) is meant to capture contributions of the driving terms \( dP_i / dr \) in Equation (1). But this correspondence only holds if the ratio of the partial pressure \( P_i \) for species \( s \) to the total pressure \( P \) is given by

\[
P_i = C_i = \frac{n_i/n_e}{\sum n_i/n_e} = \frac{n_i}{n},
\]  

(7)

where \( n \) is the total number density. This holds as long as all pressures are ideal gas. However, once electron degeneracy modifies the equation of state, \( P \) does not scale linearly with \( n \), and so Equation (7) fails for all species in the plasma. This means the \( \alpha_i \) term no longer accurately represents the information in the Burgers equations for the diffusion velocity of any species.

Moreover, the prefactor \( P/K_0 \) in Equation (5) also assumes ideal gas for each species. The quantity \( K_0 = 1.144 \times 10^{-40} (T/10^7 \text{ K})^{-3/2} n_e^2 \) simply scales out some of the information common to all diffusion coefficients in the units used for Equation (5). Thoul et al. (1994) assume an ideal gas to simplify the prefactor in Equation (5) to

\[
P/K_0 = 2.00 \frac{(T/10^7 \text{ K})^{3/2}}{(\rho/100 \text{ g cm}^{-3})^5} \left( \sum_i C_i \right) \left( \sum_i A_i C_i \right),
\]  

(8)

where \( A_s \) is the mass of species \( s \) in atomic mass units. This scaling was propagated into the MESA diffusion routine described in Paper III. Since ideal gas pressure can be significantly smaller than total pressure when electrons are degenerate, this prefactor for Equation (5) is systematically too small for degenerate plasmas. This can result in diffusion velocities that are many orders of magnitude smaller than obtained by a proper solution.

We can verify that there are problems in the degenerate regime by looking at the local gravitational acceleration \( g_{\text{diff}} \), which is solved for simultaneously with the diffusion velocities in the diffusion routine described in Paper III. MESA also reports the gravitational acceleration independent of the diffusion routine, \( g_{\text{Gauss}} = Gm/r^2 \). For a MESA WD model, layers below the surface quickly become degenerate, and the difference between \( g_{\text{diff}} \) and \( g_{\text{Gauss}} \) is significant (Figure 14). This reflects the fact that the solutions given by the diffusion routine scale with a pressure that is far too small in the interior.

### 3.2. New Methods

We now describe new methods that have been introduced to avoid the limitations discussed in Section 3.1.

#### 3.2.1. Recasting the Burgers Equations

The problems with Equation (5) demonstrated in Figure 14 can be circumvented by solving the Burgers equations directly as presented in Equations (1) and (2). When avoiding the rescaling of the Burgers equations that was originally adopted from Thoul et al. (1994), no limitations on the form of total pressure are present.

To that end, we recast the diffusion solver into the form given in Appendix C. This form closely follows the general approach presented by Thoul et al. (1994) for arranging the full set of equations into a single matrix equation, but enters the Burgers equations into that matrix structure without rescaling any quantities. We therefore avoid making any additional ideal gas assumptions beyond those already present in the Burgers equations.

#### 3.2.2. Resolving the Degeneracy Problem

Electron degeneracy makes it difficult to evaluate the term \( dP_i / dr \) in Equation (1) in the case of electrons, but it is possible to form a closed set of diffusion equations that makes no explicit reference to this equation for the electrons. Even in many applications involving WDs, each ion species can be treated as approximately ideal, and hence Equation (1) remains useful for ions. We are then left with just two problematic equations out of the system of \( 2S + 2 \) equations: Equations (1) and (2) for the electrons.

---

**Figure 13.** Mass coordinate \( m_{\text{bdy}} \) of the convective-core boundary plotted as a function of CHeB age, for the 1 M_⊙ stellar model. Different line styles/colors show the separate runs with alternative time step \( (\Delta t) \) choices. The choices adopted in Section 2.4 are marked with an asterisk [*].

**Figure 14.** Gravitational acceleration reported by the diffusion routine described in Paper III compared with \( g_{\text{Gauss}} = Gm/r^2 \) for a 0.6 M_⊙ WD model.
For the $S - 1$ species of ions in the system, we can write $S - 1$ Equations (1) in the form
\[ n_i k_B \frac{d\ln T}{dr} + n_i k_B \frac{d\ln n_i}{dr} + n_A m_p g - n_i Z_i q_e E \]
\[ = \sum_{i \neq s} K_{is} (w_i - w_s) + \sum_{i \neq s} K_{is} z_{ih} \frac{A_i r_i - A_s r_s}{A_i + A_s}, \] (9)
where $Z_i$ is the average charge of species $s$ obtained using Paquette et al. (1986b). Taking this together with $S$ Equations (2) and the two constraints on the current and mass flux, we have a total of $2S + 1$ equations. If we drop $g$ as an unknown and treat it as a fixed input to the diffusion routine in MESA using $g = Gm/r^2$, we are left with $2S + 1$ unknowns. This gives a closed system of diffusion equations with no explicit reference to the problematic Equation (1) for electrons. This is the form of diffusion equations described in Appendix C.

The thermal diffusion terms (those including $dT/dr$ in Equation (2)) still contain ideal gas assumptions as described in Section 3.1. Fortunately, in WD cores where strong electron degeneracy occurs, electron conduction leads to efficient thermal transport, resulting in small temperature gradients. With $dT/dr \ll T/H$, where $H = P/\rho g$ is the local scale height, the heat flow vectors (representing the kinetic energy carried along a temperature gradient by diffusing particles) become negligible: $r_i \ll w_i$ for all $w_i$. Thus, for WD interiors, the system of diffusion equations can be simplified by dropping the $S$ heat flow terms, removing the need for the $S$ Equations (2). Indeed, according to Iben & MacDonald (1985) and Paquette et al. (1986b), thermal diffusion leads only to small corrections to the diffusion velocities for degenerate WD interiors.

Therefore, following Iben & MacDonald (1985), we provide options for neglecting thermal diffusion in electron-degenerate regions, setting $r_s = 0$ and dropping Equation (2) for each species. Equation (9) then simplifies to the following $S - 1$ equations that no longer depend on $r_s$ for the ions:
\[ \frac{1}{n_i} \sum_{i \neq s} K_{is} (w_i - w_s) + Z_i q_e E \]
\[ = A_i m_p g + k_B \frac{d\ln T}{dr} + k_B \frac{d\ln n_i}{dr}, \] (10)
which matches Equation (10) from Iben & MacDonald (1985). Together with the two constraints, this leaves a simplified set of $S + 1$ equations for $S + 1$ unknowns: $S$ diffusion velocities $w_s$ and the electric field $E$.

Thermal diffusion terms tend to enhance gravitational settling velocities (Iben et al. 1992). This can be seen in Figure 15 for a 1.25 $M_\odot$ star on the MS, where the solvers that include thermal diffusion speed the sedimentation of $^{16}$O away from the surface relative to the solver that ignores thermal diffusion. MESA also provides options for smoothly transitioning between diffusion velocities obtained with and without thermal diffusion (averaging between the two solutions in a blending region as a function of the electron degeneracy parameter). By default, this transition region occurs when the electron chemical potential is near $\mu_e \sim k_B T$, but it is left to the user to decide on an appropriate range of electron degeneracy over which thermal diffusion should be shut off, if at all. The effect of blending between solvers with and without thermal diffusion is to suppress the thermal enhancements to diffusion velocities, smoothly pushing the enhancements to zero as electrons reach a degeneracy threshold. The implementation for the simplified set of diffusion Equations (10) and the smooth turnover of thermal diffusion terms as a function of degeneracy are described in Appendix C.

In order to confirm that we recover the correct behavior on the MS, we compare results obtained with different diffusion routines for a 1.25 $M_\odot$ star in Figure 15. Here, the results based on Thoul et al. (1994) are valid, since no significant departures from ideal gas behavior are present near the surface. The results obtained with the new scheme are in agreement.

### 3.2.3. Diffusive Equilibrium

Papers II and III show abundance profiles for WDs that have reached diffusive equilibrium in their outer layers. Figure 23 of Paper II compares the diffusive tails of H and He to an analytic expression from Althaus et al. (2003) and finds good agreement. However, Althaus et al. (2003) note that their analytic expression for diffusive equilibrium follows Arcoragi & Fontaine (1980) in assuming an ideal gas, and the equilibrium abundance profiles from their evolutionary models deviate from the analytic expression due to the inclusion of electron degeneracy. Similarly, the He layer of the WD model shown in Figure 43 of Paper III is partially degenerate, and hence the driving forces for diffusion should be modified in this region.

For a fully ionized isothermal ideal gas, the electric field that serves as one of the driving forces for diffusion in Equation (9) takes the form $q_e E = (A/Z + 1) m_p g$. By contrast, in the limit of strong electron degeneracy, the electric field approaches $q_e E = (A/Z) m_p g$. When He is the background material, the electric-to-gravitational force ratio $q_e E/m_p g$ increases from $4/3$ to 2. In this limit, any trace isotopes with $A/Z = 2$ see no net sedimentation force ($Z q_e E - A m_p g = 0$), while He with $A/Z = 1$ sees a significant upward sedimentation force ($Z q_e E - A m_p g > 0$). This extra buoyant force on H in a degenerate He background pushes the diffusive tail further toward the surface relative to the ideal gas case, as shown in Figure 16. With the proper handling of electron degeneracy described in Section 3.2, our MESA models now agree with the
suggest improvements to the treatment of screening length. They provide fitting functions and tables that can be used with any choice of screening length. In \texttt{MESA}, we follow their suggested screening prescription. The electron screening length is given by a Thomas–Fermi approximation that accounts for non-relativistic degeneracy:

\[ \lambda_e = \left( \frac{4\pi q_e^2 n_e}{\sqrt{(k_B T)^2 + (\frac{2}{3} E_F)^2}} \right)^{-1/2}, \]

where \( E_F \) is the electron Fermi energy. The direct inclusion of degeneracy increases \( \lambda_e \). The ion-screening lengths are the Debye lengths for each species,

\[ \lambda_i = \left( \frac{4\pi Z_i^2 q_i^2 n_i}{k_B T} \right)^{-1/2}. \]

To prevent ions from screening below the inter-ionic spacing, \( \lambda_i \) is an ion-sphere radius, and \( \Gamma_i \) for each species, similar to the strict minimum at \( \bar{a}_i \) (by taking \( \Gamma_i \approx (3Z_i/4\pi n_e)^{1/3} \), and define an ion-sphere coupling parameter

\[ \Gamma_i = \frac{(Z_i q_i^2)}{a_i k_B T}. \]

Their net effective screening length is then

\[ \lambda_{\text{eff}} = \sqrt{\frac{1}{\lambda_e^2} + \sum_i \frac{1}{\lambda_i^2 \left(1 + 3\Gamma_i \right)}}, \]

This construction enforces a minimum on the screening length at approximately the ion-sphere radius \( a_i \) for each species, similar to the strict minimum at \( \bar{a}_i \), set by Paquette et al. (1986a), Stanton & Murillo (2016) point out that this adjustment to the ion-screening length is physically motivated by the ion pair distribution functions in a strongly coupled plasma, where the occupation probability within the ion-sphere radius is negligible, and hence no ions are present to provide screening beneath that cutoff. The proper handling of degeneracy in the electron screening length makes it unnecessary to impose any particular minimum there, so there is no longer any ad hoc appeal to a universal minimum screening length.

For repulsive Coulomb potentials of the form given in Equation (11), Stanton & Murillo (2016) provide fits and tables of collision integrals and coefficients that we now use to calculate the resistance coefficients \( K_{st} \) for inclusion in the Burgers equations in \texttt{MESA}. They do not provide fits for attractive potentials, and Paquette et al. (1986a) note that interactions with these potentials behave significantly differently from those with repulsive potentials when screened. Hence, \texttt{MESA} continues to use the Paquette et al. (1986a) coefficients for electron–ion terms, and adopts Stanton & Murillo (2016) for all ion–ion coefficients. In any case, it is evident from Equation (94) in Paper III that the resistance coefficients approximately follow \( K_{st} \propto \mu^{1/2}_s \), where \( \mu_{st} \) is the reduced mass of particles \( s \) and \( t \); so, electron–ion resistance coefficients are generally negligible compared to the ion–ion terms.

The calculations of Paquette et al. (1986a) overestimate the electron–ion resistance coefficients in the case where electrons
are degenerate. This is because diffusion and resistance coefficients are generally calculated assuming that the velocity distributions of all particles are Maxwellian, and the coefficients roughly scale as $K_{eff} \propto v^{-2}$. When the electrons become degenerate, their characteristic kinetic energies are of order $E_F \gg k_B T$, and so their velocity distribution skews toward larger velocities. This results in smaller resistance coefficients that remain negligible compared to ion–ion terms, and no attempt is made to correct it in MESA.

For repulsive potentials, the coefficients from Stanton & Murillo (2016) generally agree with those of Paquette et al. (1986a) to within a few percent. In strongly coupled WD interiors, the Stanton & Murillo (2016) coefficients lead to shorter diffusion timescales due to a screening length that is allowed to be somewhat smaller than the minimum value imposed by Paquette et al. (1986a): $\lambda_{eff} < \lambda_i$. Future prospects for further improvements to diffusion coefficients include the recent progress on effective potential methods from Daligault et al. (2016) and Shaffer et al. (2017).

3.4. Diffusion-induced Flashes on He WDs

Diffusion-induced H shell flashes on low-mass ($M \lesssim 0.4 M_\odot$) He WDs are known to alter their cooling times (Althaus & Benvenuto 2000; Althaus et al. 2001) and seismic properties (Althaus et al. 2013). Istrate et al. (2016a, 2016b) use MESA to model this process, generating tables of cooling timescales and comparing MESA models with those of Althaus et al. (2013).

Figure 17 shows an exploration of the H shell flash domain for a large grid of $Z = 0.02$ MESA models over a range of He-core and H-envelope masses. Here, the envelope mass is defined as the total mass of H-rich material ($X > 0.01$) at the surface at the beginning of the WD cooling track. Lines show the minimum envelope masses for which H shell flashes occur given various diffusion prescriptions.

For a given core mass, there is a range of envelope masses that exhibit shell flashes only if diffusion is included, but this range depends on the diffusion prescription. The two lower lines for models including diffusion in Figure 17 differ only in the handling of electron degeneracy in the diffusion scheme. This illustrates the importance of properly handling degeneracy as described in Section 3.2, since the diffusion-induced flashes are typically ignited by CNO burning in the diffusive tail of H that reaches into the partially degenerate He layers. WDs in this mass range often experience cycles of many H flashes, depleting H incrementally until insufficient H remains to ignite another flash. The disagreement between diffusion prescriptions on the minimum envelope mass for flashes is therefore significant, as this will determine the total number of flashes and final H mass that sets the ultimate cooling timescale for an object.

To explore the full range of parameters presented in Figure 17, our WD models were built by artificially stripping the H envelope down to a specific mass coordinate above the He core of a 1.0 $M_\odot$ model ascending the RGB. For a discussion of MESA models including proto-WD formation and the resulting H envelope masses, see Istrate et al. (2016b).

3.5. Heating from $^{22}$Ne Settling

In the strongly degenerate limit, $q_f E/m_p g \approx 2$ for C/O WD cores. For an isotope where $A/Z = 2$, the electric and gravitational fields result in a net force that drives diffusion. For $^{22}$Ne in cooling WD interiors, this force is $F = q_f E - A m_p g \approx -2m_p g$, causing $^{22}$Ne to sediment toward the center and deposit energy as it moves deeper into the gravitational potential (Bildsten & Hall 2001; Deloye & Bildsten 2002; García-Berro et al. 2008, 2010). This heating can prolong the WD cooling timescale, especially at late times when the WD is very dim and radiates away the energy slowly. This effect may be especially important for explaining WD luminosity functions in old and metal-rich open clusters such as NGC 6791, where abundant $^{22}$Ne is available in WD interiors to provide heating.

MESA now offers an option to include this heating term in the energy equation (see Section 8.7) when diffusion is enabled. The specific rate at which energy is deposited is

$$\epsilon_{22} = \frac{|F| v_{22}}{(A m_p)/X_{22}} = \frac{22m_p g - 10q_f E}{X_{22} v_{22}^2} \frac{X_{22} v_{22}}{22m_p} \approx \frac{X_{22} v_{22}^2}{22m_p}. \quad (16)$$

The $^{22}$Ne diffusion velocity ($v_{22}$) and electric field are calculated in the diffusion routine and then used to evaluate the above heating term. Note that the updates to diffusion described in Section 3.2 are essential for correctly calculating both the diffusion velocity and magnitude of the driving force in the degenerate interior of the WD.

Figure 18 shows the delay in WD cooling from introducing $\epsilon_{22}$ into the 0.6 $M_\odot$ models. These models turn off diffusion for $\Gamma > 175$, so $\epsilon_{22}$ is only active in material for which crystallization has not yet occurred. The time delays shown in Figure 18 are in good agreement with those shown by Deloye & Bildsten (2002) and Garcia-Berro et al. (2008) for comparable cases.

4. Implicit Hydrodynamics

In Paper III, we describe implicit shock-capturing hydrodynamics capabilities based on the use of an artificial viscosity. We now add an option for using an approximate Riemann solver, the HLLC (Harten–Lax–van Leer–Contact) solver introduced by Toro et al. (1994). (See also Batten et al. 1997 for an early implicit implementation of HLLC.) The HLLC
method provides improved shock capturing and energy conservation by avoiding the need for artificial viscosity. However, the methods presented in Paper III are still included in MESA so that users may continue to apply them.

### 4.1. Implementation of HLLC

Accurate shock-capturing methods evaluate the flux of hydrodynamical conserved quantities by extrapolating the solution on each interface between zones over the course of the time step. The different methods for projecting the solution into the future are known as different “Riemann solvers.” HLLC is designed to accurately capture the evolution of contact discontinuities. When implemented on a Lagrangian grid, HLLC is able to evolve purely advective flows without any contact smearing (Cheng & Shu 2007; Duffell & MacFadyen 2011; Cheng et al. 2012; Cheng & Shu 2014).

Paper I and Paper III discussed the evolution of a velocity variable \( v \), defined at cell faces. When using HLLC, MESA instead evolves a cell-centered velocity \( u \).

We solve a Riemann problem at the cell face with index \( k \).

The cell to the left (toward the center) is the cell with index \( k \); the cell to the right (toward the surface) is the cell with index \( k - 1 \). The cell face radius is \( r_k \). The mass contained within an individual cell is \( dm_k \). The mass enclosed from the center of the star to the cell face is \( m_k \). For convenience, we define the face area as \( A_k = 4 \pi r_k^2 \). Thermodynamic variables (e.g., \( P_k, \rho_k \)) are defined at cell centers by mass. Figure 19 shows the layout of the cells.

The calculation begins by making estimates for the density and velocity at the left and right of the face. Explicit codes sometimes use multipoint polynomial interpolation based on values in neighboring cells to improve the reconstruction of the values at the face. However, for an implicit code such as MESA, that would introduce dependencies in the partial derivatives for the Jacobian that would violate the necessary block tridiagonal structure (see Appendix B in Paper II). To avoid this problem, we use the cell center density and velocity alone to estimate the values at the edges of that cell. The variables for the left and right values are named relative to the edge rather than the cell,
and hence $u_{\text{face}} = S$. Likewise, the pressure at the cell face is the pressure at the contact wave, $P_{\text{face}} = P$, where

$$P = \frac{1}{2} \left[ (\rho_L (u_L - S_L) (u_L - S) + P_L) + (\rho_L (u_L - S_L) (u_L - S) + P_L) \right], \quad (21)$$

and $S_L$ and $S_R$ are the fastest wave speeds moving to the left and right, respectively. To evaluate these, we assume the simple and most conservative bounds on the signal velocities,

$$S_L = \min (u_L - c_{\text{L}}, u_R - c_{\text{R}}),$$
$$S_R = \max (u_L + c_{\text{L}}, u_R + c_{\text{R}}), \quad (22)$$

where $c_{\text{L}}$ and $c_{\text{R}}$ are the sound speeds on the left and right sides of the cell boundary, respectively. Having obtained values for $u_{\text{face}}$ and $P_{\text{face}}$, we now formulate the versions of the equations used when HLLC is enabled.

In the Lagrangian picture, the cell boundaries move with the fluid velocity, such that the net fluxes for mass, momentum, and energy from cell $k$ to cell $k-1$ are extremely simple (Cheng & Shu 2014) and given by

$$F_{\rho,k} = 0,$$
$$F_{P,k} = A_k P_{\text{face},k},$$
$$F_{\mu,k} = A_k P_{\text{face},k} u_{\text{face},k} + L_k. \quad (23)$$

The $L_k$ term in the energy flux does not come from the solution of the Riemann problem, but from the fact that MESA also evolves a luminosity variable that reflects the radiative or convective transport of energy.

The finite volume form of the mass conservation equation remains the same as that given in Paper I,

$$\ln r_k = \frac{1}{3} \ln \left[ \frac{3}{2} \frac{\rho_k}{\rho_k} \right]. \quad (24)$$

However, the equation for the radius (Equation (29) in Paper III) has changed. The new equation for the radius is

$$r_k = r_{\text{start},k} + u_{\text{face},k} dt,$$  

where $dt$ is the time step. For numerical precision, we rewrite this as

$$\exp(\ln r_k - \ln r_{\text{start},k}) - 1 = \frac{u_{\text{face},k} dt}{r_{\text{start},k}}, \quad (26)$$

where this recasting allows the use of the expm (de Dinechin et al. 2007; see also Paper III) function expm to evaluate the function $\exp(x) - 1$ to machine precision (as indicated by the underbrace).

The local radial momentum equation for cell $k$ is

$$\frac{u_k - u_{\text{start},k}}{dt} = -\frac{1}{2} \left( \frac{Gm_k}{r_k^2} + \frac{Gm_{k+1}}{r_{k+1}^2} \right)$$
$$+ \frac{P_k}{\rho_k} (A_k - A_{k+1})$$
$$+ \frac{1}{dm_k} (F_{P,k+1} - F_{P,k}). \quad (27)$$

On the right-hand side, the first term is gravitational, the second is a geometric source term that arises from putting the equation in conservation-law form, and the final term is the momentum flux in and out of the cell found by HLLC.

The local total energy conservation equation for cell $k$ is

$$e_k - e_{\text{start},k} + \frac{1}{2} (u_k^2 - u_{\text{start},k}^2) - Gm_k \left( \frac{1}{r_k} - \frac{1}{r_{\text{C, start}}} \right)$$
$$= \frac{dt}{dm_k} \left( F_{e,k+1} - F_{e,k} + e_{\text{nuc},k} - \epsilon_{\nu, k} + \epsilon_{\text{extra},k} \right). \quad (28)$$

(See Section 8.3 for a discussion of how this energy accounting is related to that typically used in stellar evolution calculations.) We define the cell center quantities $m_k$ and $r_k$ to be $r$ and $m$ at the center of mass of the cell. The terms on the left split the local total energy into internal, kinetic, and potential components. The right side gives the energy in and out of the cell and the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. The terms on the left split the local total energy into internal, kinetic, and potential components. The right side gives the energy in and out of the cell and the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell. Energy loss from neutrinos due to nuclear reactions is already subtracted from the energy sources and sinks in the cell.

As in Paper I, the temperature differences of interior cells $T_k$ are set by the energy transport across boundaries,

$$T_{k-1} - T_k = \frac{d m_k}{\nabla \rho_{T,k} \left( \frac{\rho}{\rho} \right)_C \frac{P}{T_k} \rho_{T,k}^2}.$$  \quad (29)

where $\nabla \rho_{T,k}$ is provided by the MESA module ml2t (see Section 5.1 in Paper I), and the overbars indicate quantities at the cell faces (see Figure 19). This equation relates the temperatures of neighboring cells; the actual temperature in each cell is then fixed by a surface boundary condition.

MESA’s HLLC includes the effects of rotation in the shellular approximation (see Paper II, Section 6.1) and can also include a post-Newtonian correction to the gravitational force. (For an example application to neutron star envelopes, see Paper III, Section 5.3). These capabilities require modifications to the momentum equation. In both cases, they can be treated as a rescaling of the local gravitational constant $G \rightarrow G \rho$. In the case of rotation, the rescaling factor is $f_P$ (Paper II, Equation (23)). In the post-Newtonian case, it is $(1 - 2GM/(r^3 \rho^2))^{-1/2}$. Therefore, when either of these is used with the hydrodynamics capabilities described in this section, the rescaling is applied to the $G$ in pressure reconstruction (Equation (18)) and separately to each $G$ (for cell $k$ and $k+1$) in the momentum equation (Equation (27)).

4.2. Mesh Refinement

During a typical stellar evolution run, MESA controls its meshing using “mesh functions” that limit the maximum allowed change of various quantities between adjacent cells (see Section 6.5 in Paper I and Section B.4 in Paper II). With HLLC, the criteria to split or merge cells are written solely in terms of the radial coordinate in order to simplify the adjustments to the mesh in response to changes in density before and after a shock. Cells split when they decompress enough that their radial extent becomes too large, and they merge with a neighbor when they compress enough that their radial extent becomes too small.

The refinement criteria can use either linear ($x = r$) or logarithmic ($x = \ln r$) radius. The user selects a target number of cells, $N_{\text{target}}$. MESA translates this into a target cell size,
\[ dx_{\text{target}} = (x_{\text{surface}} - x_{\text{center}})/N_{\text{target}}. \] A cell is considered too large if \( dx_{k}/dx_{\text{target}} > f_{\text{larg}} \) and a cell is considered too small if \( dx_{\text{target}}/dx_{k} > f_{\text{short}}. \] The refinement then proceeds iteratively. At each iteration, MESA selects the smallest and largest cells. If the largest cell is too large, it is split. If the smallest cell is too small, it is merged unless the magnitude of the difference between its velocity and that of either neighbor is a significant fraction of the local sound speed: this prevents merging in the immediate vicinity of the shock where there are sharp jumps in velocity. The refinement proceeds up to some maximum number of iterations, though in practice the procedure typically stops before then because no more cells satisfy the criteria to be split or merged.

A cell merges with its smaller neighbor, unless they have a different most-abundant chemical species, in which case the cell merges with the other neighbor instead. When a cell is split, differences in quantities such as density and chemical abundances between the two child cells are determined by interpolation from the neighboring cells.

### 4.3. Time Resolution

Since the hydrodynamics equations are being solved implicitly, MESA is not subject to the Courant–Friedrichs–Lewy (CFL) time step condition for numerical stability. The size of the MESA time step is instead limited by the restrictions on the allowed changes in the structure of the star. The usual time step controls continue to apply.

Although numerical stability does not require the restrictive CFL time step condition, the choice of time step does affect the accuracy of the solution. A CFL-like limit is often also applied because it can be a convenient additional way to restrict time steps along with the other options. Such a restriction allows for well-converged solutions. The time step can be limited by the requirement that

\[ \min \left\{ \frac{dr_{k}}{|u_{k}| + c_{s,k}} \right\} < f_{i} \times \delta t, \]  

where \( f_{i} \) is a user parameter. Unlike in an explicit code where a similar minimum must be evaluated over all cells, in MESA the minimum is taken only over cells for which

\[ \max \left\{ |u_{k} - u_{k+1}|, |u_{k} - u_{k-1}| \right\} \leq c_{s,\text{max}} \]  

where \( c_{s,\text{max}} \) is the maximum evaluated over nearby cells, and \( f_{u} \) is a user parameter.\(^{16}\) This means that only regions near the shock front limit the time step. The option for additional limitations on where this condition is evaluated (e.g., in mass) is provided.

### 4.4. Hydrodynamic Test Problems

In order to test the HLLC implementation, comparisons are now made to problems with known solutions.

\(^{15}\) By default, \( N_{\text{target}} = 1000, f_{\text{larg}} = 4.0, \) and \( f_{\text{short}} = 1.5, \) but these parameters are configurable at runtime.

\(^{16}\) The value of \( f_{u} \) is similar to the values of a CFL parameter in an explicit code, while \( f_{i} \) in the examples is typically a small value like 10\(^{-7}\). The description of these limits is schematic, and the reader is referred to the source code for the precise implementation details.

---

\( E_{\text{blast}} = 1.464 \text{ erg} \)

\( \omega = 1.0, \gamma = 7/5 \)

\( r_{\text{shock}} = 1 \text{ cm} \)

\( t = 1 \text{ s} \)

\[ \frac{E_{\text{blast}}}{P_{0}} = \frac{1.464 \text{ erg}}{0 \text{ erg cm}^{-3}} \]

\[ t = 0.06 \text{ s}, t = 0.25 \text{ s}, t = 0.50 \text{ s}, t = 1.0 \text{ s} \]

\[ \text{Density, Pressure, Velocity} \]

---

\( \text{Sedov, Spherical} \)

\( E_{\text{blast}} = 1.464 \text{ erg} \)

\( \omega = 1.0, \gamma = 7/5 \)

\( r_{\text{shock}} = 1 \text{ cm} \)

\[ \text{Density, Pressure, Velocity} \]

---

\( \text{Radius} \)

\[ 0, 0.2, 0.4, 0.6, 0.8, 1, 1.2 \]
4.4.2. Noh Problem

Noh (1987) describes a standard verification problem that tests the ability to transform kinetic energy into internal energy, and the ability to follow supersonic flows. A sphere of cold gas with an ideal gas EOS and constant adiabatic index $\gamma$, that is, $P = (\gamma - 1) \rho e$, is initialized with a uniform, radially inward speed of $1 \text{ cm s}^{-1}$. A shock forms at the origin and propagates outward as the gas stagnates. For an initial gas density $\rho_0 = 1 \text{ g cm}^{-3}$, the analytic solution in spherical geometry for $\gamma = 5/3$ predicts a density of $64 \text{ g cm}^{-3}$ in the stagnated gas.

Figure 22 shows the analytic and MESA profiles for the density, pressure, and material speed at $t = 0.3 \text{ s}$. Most implementations, including MESA’s, produce anomalous “wall-heating” near the origin (although see Gehmeyr et al. 1997). As the shock forms at the origin, the momentum equation tries to establish the correct pressure level. However, numerical dissipation generates additional entropy. The density near the origin drops below the correct value to compensate for the excess internal energy (e.g., Noh 1987; Rider 2000). Thus, the density profile is altered near the origin while the pressure profile remains at the correct constant level in the post-shock region.

Figure 22 shows the analytic solution and MESA solution for a fixed time step of $\delta t = 5 \times 10^{-6} \text{ s}$ and 10,000 cells. Deviations from the analytic solutions are $\lesssim 1\%$, except for the density near the origin and the shock front. A convergence exercise with different fixed time steps and spatial resolutions suggests that spatial resolution is relatively more important than temporal resolution in the MESA solutions for the Noh problem.

4.4.3. Supernova Shock

The problem of a supernova (SN) shock moving through a stellar envelope has been extensively studied. For a radiation-dominated strong shock, a simple analytic expression for the shock velocity is provided by Matzner & McKee (1999),

$$v_{sh} = \alpha \left( \frac{E}{m - M_{\text{center}}} \right)^{1/2} \left( \frac{m - M_{\text{center}}}{\rho r^3} \right)^{0.19},$$

where we adopt $\alpha = 0.736$ as suggested by Tan et al. (2001). The explosion has an energy $E$. The mass that enters into this expression is the mass entrained by the shock and so differs from the Lagrangian mass coordinate $(m)$ by the mass of the remnant $(M_{\text{center}})$. Since the material in the shocked envelope has an adiabatic index of $4/3$, the Matzner & McKee (1999) prediction for the post-shock velocity is $v_{\text{MM}} = 6v_{sh}/7$.

MESA defines the shock location to be the outermost point where the fluid Mach number exceeds 1, as measured in the rest frame of the star. Since the primary application of these capabilities are blast waves propagating into approximately static stellar envelopes, this shock detection criterion suffices. Figure 23 compares the velocity in a MESA model (the 19 M$_{\odot}$ model of SN 1999em; see Section 6) with $v_{\text{MM}}$. We show explosions with two different energies, $E = 0.9 \times 10^{51} \text{ erg}$ and $E = 2.7 \times 10^{51} \text{ erg}$. Both cases have $M_{\text{center}} = 1.5 M_{\odot}$. Typical differences are at the few percent level.

4.4.4. Weak Shock Propagation

We now explore weak shocks with Mach numbers $M = 1.2-2.2$ propagating outward in the hot stellar envelope of a classical nova progenitor. The model is a 0.8 M$_{\odot}$ WD. The H/He envelope extends from $r = 7.1 \times 10^8 \text{ cm}$ to $r = 7.85 \times 10^8 \text{ cm}$ with densities $\rho = 10-100 \text{ g cm}^{-3}$ and temperatures $T \approx 10^7 \text{ K}$.

After excising the core, we run the model with HLLC enabled for 100 s, corresponding to $\approx 50$ sound crossing times in the outer envelope, to allow the envelope to settle. Afterwards, the envelope has a total energy of $-9.16 \times 10^{45} \text{ erg}$, with $-9.38 \times 10^{45} \text{ erg}$ in potential energy, $2.22 \times 10^{44} \text{ erg}$ in thermal energy, and a negligible kinetic energy $1.2 \times 10^{29} \text{ erg}$. We turn off convective energy transport to study the properties of weak shocks. To create weak shocks, we inject 0.5%–5% of the total thermal energy into a single cell with mass $dm = 1.6 \times 10^{25} \text{ g}$ at $r = 7.3475 \times 10^8 \text{ cm}$ over
−4 s. Figure 24 shows the resulting upward and downward propagating shocks. We restrict our region of study to the region where the upward and downward shocks are well-separated, in the radius range of \( r = 7.47 \times 10^8 \text{ cm} \) (denoted with thin gray lines in Figure 24). We do not study the properties of the downward shock and its artificial reflection from the “floor” of our model.

We define \( u_0 = \frac{d\rho}{dt} \) as the shock velocity, where \( r_{\text{peak}} \) is defined as the radial location with maximum fluid velocity. We compare the properties of the shock to analytic expectations for cases where \( \gamma \) is identical in the pre- and post-shock material. Pre-shocked quantities carry a 0 subscript, and shocked quantities carry a 1, and we use the sound speeds, \( c_s \), and pressures, \( P \), on either side of the discontinuity. Following Zel’dovich & Raizer (1967), in the rest frame of the shock front, the pre-shock gas travels into the shock front at velocity

\[
\left( \frac{u_0}{c_{s,0}} \right)^2 = \frac{(\gamma - 1) + (\gamma + 1)P_1/P_0}{2\gamma}.
\]

The post-shock velocity \( u_1 \) has magnitude \( |u_1| = u_0 - u_{\text{peak}} \), where \( u_{\text{peak}} \) denotes the fluid velocity \( u \) at \( r_{\text{peak}} \). The analytic expression is

\[
\left( \frac{u_1}{c_{s,1}} \right)^2 = \frac{(\gamma - 1) + (\gamma + 1)P_0/P_1}{2\gamma}.
\]

Local shocked quantities are evaluated at the cell with the maximum Lagrangian fluid velocity, while pre-shocked quantities are evaluated at the cell in the initial MESA profile (before the shock has propagated) with the same mass coordinate as the shock front when it reaches \( r_{\text{peak}} \). The thin black lines in the upper and lower panels of Figure 25 are the right-hand side of Equations (33) and (34), respectively, for shocks produced by different amounts of injected energy.

Figure 23. Comparison of MESA with the analytic results of Matzner & McKee (1999) and Tan et al. (2001). This started as a 19 M\(_\odot\) model at ZAMS; at explosion it is 17.79 M\(_\odot\) with \( M_{\text{toto}} = 1.5 \text{ M}_\odot \). The upper panel shows an explosion with \( E = 0.9 \times 10^{51} \text{ erg} \); the lower panel shows \( E = 2.7 \times 10^{51} \text{ erg} \). The gray dashed curve shows the analytic prediction for the post-shock fluid velocity given the density profile of the initial model. The solid curves show the velocity profiles from the MESA calculation at specific times. The unfilled diamonds indicate where on the dashed curve the two should be compared.

Figure 24. Velocity at different times in the envelope after \( 6 \times 10^{42} \text{ erg} \) has been injected. One shock front travels upward and grows as it enters the outer atmosphere, and another pulse travels downward and reflects off of the inner boundary. The thin gray lines denote the region of study.

Figure 25. Comparison of the MESA calculation (colored lines) with analytic expressions (thin black lines) for \( (u_0/c_{s,0})^2 \) (upper) and \( (u_1/c_{s,1})^2 \) (lower) for different energies injected.
5. Rayleigh–Taylor Instabilities

The outward moving shock in a core-collapse SN explosion encounters multiple composition boundaries. Across these boundaries, the density gradient is steep, especially at the H/He boundary. Post-shock, these regions become unstable to the Rayleigh–Taylor instability (RTI). Early analytics and 2D simulations (Chevalier 1976; Chevalier & Klein 1978; Weaver & Woosley 1980; Benz & Thielemann 1990; Herant & Benz 1991) and modern 3D calculations (Hammer et al. 2010; Wongwathanarat et al. 2015; Utrobin et al. 2017) show significant impact on the density, velocity, and composition structure of the ejecta.

It has been known for decades that the resulting compositional mixing can significantly alter the photometry of the SN. This effect has been roughly included in 1D modeling of Type IIP light curves resulting from explosions deep within a red supergiant (Eastman & Pinto 1993; Utrobin 2007; Dessart & Hillier 2010, 2011). The mass densities and energy densities are also smeared out by the mixing from the RTI (see Bersten et al. 2011 for an early discussion raising this concern). In their recent modeling of the Type IIP SN 1999em, Utrobin et al. (2017) capture the impact of the RTI using a 3D model pre-breakout and connect it to observable SN properties with a 1D post-breakout radiation calculation.

To approximate the 3D effects of the RTI, we implement a scheme by Duffell (2016) that modifies the 1D spherical hydrodynamics equations. This scheme has been recently applied to the specific case of core-collapse SNe by P. Duffell et al. (2018, in preparation) and is now implemented in MESA for use along with the HLLC scheme. In this section, we describe the MESA implementation and compare it to the 3D calculations of Wongwathanarat et al. (2015). The use of the resulting RTI-mixed ejecta for SN light curves and velocities will be discussed in Section 6.

5.1. Implementation of the Duffell RTI

The Duffell (2016) scheme evolves an additional scalar quantity $\alpha_R$, representing the relative strength of turbulent fluctuations. The evolution equation for $\alpha_R$ is an advection–diffusion equation with source terms. In Eulerian form, this is

$$
\frac{\partial}{\partial t} (\rho \alpha_R) + \frac{1}{r^2} \frac{\partial}{\partial r} \left[ r^2 (\rho \alpha_R u - \eta_R \frac{\partial}{\partial r} (\rho \alpha_R)) \right] = S_\alpha^+ + S_\alpha^-,
$$

where

$$
S_\alpha^+ = (A_R + B_R \alpha_R) \max \left( 0, -\frac{\partial P}{\partial r} \right),
$$

$$
S_\alpha^- = -D_R \alpha_R \rho \gamma r^{-1},
$$

$$
\eta_R = C_R \alpha_R c_s r.
$$

The source and sink terms $S_\alpha^+$ and $S_\alpha^-$ represent the growth and decay of the turbulence, respectively. These terms along with a diffusion coefficient $\eta_R$ are determined via scaling arguments. The dimensionless coefficients in front of these quantities (growth coefficients $A_R$, $B_R$, diffusion coefficient $C_R$, and decay coefficient $D_R$) are determined by calibrating a suite of

---

17 The quantity $\alpha_R$ is denoted by $\kappa$ in Duffell (2016) and $\alpha_{RTI}$ within MESA.

---

Figure 26. Comparison of the MESA calculation (colored lines) to the expectations of shock theory (thin black lines) for the temperature increase (upper) and entropy increase (lower) for different energies injected. Colored dotted lines in the upper plot indicate the temperature change for a purely adiabatic compression.
1D models against 3D hydrodynamics simulations. The original model of Duffell (2016) calibrates against 2D simulations; see P. Duffell et al. (2018, in preparation) for the re-calibration of these constants to 3D simulations. The values of the constants found by that 3D calibration are \( A_R = 10^{-3}, B_R = 2.5, C_R = 0.2, \) and \( D_R = 2.0. \) In MESA, these constants are adjustable so that the user may explore the effect of varying them. For example, we show later the effect of \( D_R = 2.0, 3.0, \) and \( 4.0. \)

Additionally, a diffusive term (with diffusivity \( \eta_R \)) appears in each of the mass, momentum, and energy equations. For the sake of exploration in MESA, we allow each diffusivity to be scaled by an independent factor. With the diffusive term, the mass flux becomes (cf., Equation (23))

\[
F_\rho = A_\rho (u - \ddot{r}) - A_\eta_R \frac{\partial \rho}{\partial r},
\]

and the choice \( u = \ddot{r} \) (i.e., \( u_{\text{face}} = S_h \)) no longer causes this quantity to vanish. If no correction were applied, MESA would no longer preserve the mass coordinates of zone faces. In order to preserve the Lagrangian nature of the equations, we allow for an additional velocity between the cell face and the fluid.

The advective flux introduced by the relative motion of the face will then exactly cancel this diffusive flux, restoring the Lagrangian nature of the scheme. Assuming \( \ddot{r} = u + \delta u, \) the no mass flux condition can be rewritten as

\[
\delta u = \eta_R \frac{1}{\rho} \frac{\partial \rho}{\partial r} = \eta_R \frac{1}{\partial r} \frac{1}{\rho} \partial \rho.
\]

The term in parentheses is equal to \( A. \) In the finite volume form evolved by MESA, evaluating this condition at the cell face gives

\[
\delta u = A \eta_R \frac{(\rho_k - \rho_h)}{\partial m}.
\]

Therefore, we modify the HLLC equation \( u_{\text{face}} = S_h \) to

\[
u_{\text{face}} = S_h + \delta u,
\]

and proceed as in Section 4.1 (see Equation (20) and the surrounding discussion). Usually \( |\delta u| \ll S_h, \) so in practice this is a small modification and the HLLC scheme still works well.

For a scalar quantity \( f, \) the flux is the sum of the diffusive flux plus the advective flux (\( A, \rho f, \delta, u \)) created by the velocity shift \( \delta u, \) that is,

\[
F_f = A \eta_R \frac{\partial f}{\partial r} - A \eta_R \frac{\partial (\delta f)}{\partial r} = -A \eta_R \rho \frac{\partial f}{\partial r}.
\]

Rewriting the spatial derivative as a mass derivative gives

\[
F_f = -\eta_R (A \rho)^2 \frac{\partial f}{\partial m}.
\]

To evaluate the fluxes for a cell \( k, \) we define

\[
\sigma_{\eta,k} = \eta_{\eta,k} (A_k \rho_k)^2 \frac{1}{\partial m_k},
\]

where

\[
\eta_{\eta,k} = C_R \sigma_{\eta,k} \sigma_{\eta,k} \rho_k.
\]

The fluxes across faces are

\[
F_{\rho,k} = 0,
\]

\[
F_{\rho,k} = A_k \rho_{\text{face},k} - \sigma_{\eta,k} \rho_k (u_{k-1} - u_k),
\]

\[
F_{\rho,k} = A_k \rho_{\text{face},k} + L_k - \sigma_{\eta,k} \rho_k (\epsilon_{k-1} - \epsilon_k).
\]

The finite volume version of Equation (38) evolved by MESA is

\[
\alpha_R = \alpha_R \text{start}
\]

\[
= \delta t \left[ \frac{1}{\partial m_k} (F_{\rho,k+1} - F_{\rho,k}) + (A_R + B_R \alpha_R) \left[ \frac{1}{\rho_k} \max (0, -\frac{\partial P}{\partial r}) \right]_{\text{start}} \right.
\]

\[
- D_R \alpha_R \left( \frac{2 \varepsilon_{\rho,k}}{r_k + r_{k+1}} \right)_{\text{start}},
\]

where

\[
F_{\rho,k} = -\sigma_{\eta,k} (\alpha_{R,k-1} - \alpha_{R,k}).
\]

We evaluate the product of the \( P \) and \( \rho \) spatial derivatives as

\[
\frac{\partial P}{\partial r} = \frac{\rho_{k-1} - \rho_k}{d r_k},
\]

where

\[
\frac{d r_k}{4 \pi r_k^2 \rho_k}.
\]

which is numerically preferable to a subtraction of radial coordinates. At sharp jumps in density and pressure, these source terms can diverge, and therefore options to smooth \( \partial P / \partial r, \partial \rho / \partial r \) are available, though they are off by default. In practice, smoothing does not appear to be necessary in the cases we have explored, as HLLC typically smears out these sharp jumps over several cells in the model at the shock, and RTI mixing then smooths out the jumps more post-shock.

5.2. Comparing a Munich 3D Model to MESA with the Duffell RTI

We now develop a MESA analogue to a specific 3D simulation of Wongwathanarat et al. (2015). This provides a comparison of the predictions from the MESA implementation of the RTI mixing described in the previous subsection (which we refer to as the Duffell RTI) with those obtained in a 3D simulation. The Wongwathanarat et al. (2015) progenitor model we use, L15-1-cw, has a mass of \( 15 M_\odot \) based on Limongi et al. (2000). We refer to this as the Munich L15 model. Note, as made clear in Wongwathanarat et al. (2015), most prior studies simulating RTI in SN envelopes disregard early-time asymmetries, relying on explosions that are initiated assuming spherical symmetry. Since those explosion asymmetries appear to have significant consequences, it is important to start from a 3D model like L15-1-cw when evaluating the use of MESA for supernovae.

To compare with the Munich L15 model, we construct a MESA starting model with similar parameters. Future studies of a variety of 3D models will be necessary to assess the impact on our 1D results of a variety of 3D asymmetries in the initial explosion. Just as the Duffell RTI allows 1D simulations to
capture many of the effects of the 3D RTI, it may be possible to extend 1D codes in the future to include relevant effects of explosion asymmetries in a self-consistent manner rather than by expediencies such as those we describe below for initializing the $^{56}\text{Ni}$ abundance.

We now compare the 3D shell averages of Wongwathanarat et al. (2015) to \textsc{MESA} with the Duffell RTI enabled. The left panel of Figure 27 shows the resulting abundances when the shock is at $4.8 \, M_\odot$, with the thin lines from \textsc{MESA} with $D_R = 3$ and the thick lines the 3D averages from the Munich L15 model. For H, He, and O, the \textsc{MESA} lines agree with the Munich model. If nothing is done to take into account the initial asymmetry of the explosion, the radial extent of the $^{56}\text{Ni}$ in the Munich model far exceeds what can be achieved in \textsc{MESA} by the Duffell RTI mixing. Hence, at this moment in the model evolution, we use the Munich L15 results to fix the extent of the \textsc{MESA} distribution of $^{56}\text{Ni}$. Later mixing in the \textsc{MESA} run is done by the Duffell RTI. The right panel of Figure 27 shows the comparison with the Munich model just before shock breakout. For this case, we show three simulations with $D_R = 2, 3$ and 4.

In Figure 28, we show the \textsc{MESA} profiles of density (upper panel) and velocity (lower panel) at the moment when the shock is at $14.7 \, M_\odot$. The solid lines are with Duffell RTI enabled, while the dotted lines are with it turned off. As shown by Wongwathanarat et al. (2015), Utrobin et al. (2017), and P. Duffell et al. (2018, in preparation), the operation of the RTI removes the unphysical density feature produced in 1D simulations without it. Such features can be seen in Figures 2 of Eastman et al. (1994) and Dessart & Hillier (2011) and in the dotted black line in the upper panel of Figure 28. The Duffell RTI also alters the velocity structure of the material near the H/He boundary, as we will discuss more in Section 6.6. The thick gray lines in both plots show the 1D shell averages of the 3D Munich L15 model. The fainter gray lines show the density and velocity profiles for a variety of angles in the Munich model. The asymmetries of the shock in the Munich model lead to its location varying between mass coordinates $10.5 \, M_\odot$ and $14.5 \, M_\odot$. This variation with angle leads to 1D shell averages that do not show a sharp shock feature but instead have more rounded shapes. Since the 1D \textsc{MESA} results have the shock at a single mass coordinate, they are similar to the Munich profiles at a particular angle. This difference must be considered when comparing results from \textsc{MESA} to shell averages from the Munich model. It also shows that the time of shock breakout, which is well-defined in the 1D model, varies with angle in the 3D model.

6. Light Curve and Velocity Evolution of Core-collapse Supernovae

We now present \textsc{MESA} modeling of the ejecta evolution triggered by core collapse in massive stars (roughly...
The new MESA capabilities enable self-consistent calculations of the photometric evolution of core-collapse supernovae (SNe) using the STELLA code (Blinnikov et al. 1998; Blinnikov & Sorokina 2004; Baklanov et al. 2005; Blinnikov et al. 2006). A public version of STELLA is now included with the MESA distribution, and the interface from MESA to STELLA has been customized for ease of use.\footnote{When using these capabilities, one should cite this instrument paper and the following papers describing STELLA: Blinnikov & Sorokina (2004), Baklanov et al. (2005), and Blinnikov et al. (2006).}

Our main emphasis in this section is on the commonly observed Type IIP “plateau” SNe that originate from energy deposited deep in the core of an $M \approx 8$–20 $M_\odot$ red supergiant (Smartt 2009). We also exhibit how these new capabilities enable simulations of core-collapse events that occur after the star has lost the majority of its outer hydrogen envelope, the Type Ib and Ic SNe.

The new capabilities we present are provided by a powerful combination of MESA and STELLA. Post-core-collapse evolution proceeds in two distinct phases. First, we use MESA to evolve models from a few seconds after the central explosion triggered by core collapse to a time just before the outgoing shock reaches the stellar surface. These calculations make use of HLLC (Section 4) and Duffell RTI (Section 5). Subsequently, we use STELLA to evolve models through shock breakout and beyond the end of the plateau, generating light curves and velocities of the material at the photosphere and above.

Simulations using 3D models from the core-collapse event to shock breakout are computationally expensive but are now feasible (Wongwathanarat et al. 2013; Utrobin et al. 2017), and it will be a significant contribution to have more of them available in the future. To explore the subsequent \( \gtrsim \)100 days of photometric and spectroscopic evolution, 1D approximations are common. The new capabilities with MESA and STELLA also use a 1D approximation for both the pre- and post-breakout evolution. This provides a less computationally costly alternative for the initial exploration of the parameter space for potential progenitors prior to or instead of doing a more realistic but more computationally costly 3D simulation. The pair MESA and STELLA can produce useful results in a few hours running on a modern multicore desktop workstation (see Section 6.7), while the 3D pre-breakout evolution and post-breakout spectral analysis can take weeks running on a supercomputer. MESA and STELLA are not a replacement for the more computationally expensive codes but will be useful in conjunction with them.

Throughout this section, we present models that we characterize as “similar to” observed SNe. We list the properties and parameters of these models in Table 3. As we discuss in Section 6.8, where we describe the procedure by which these models were generated, they are not “best-fit” models. Rather, they simply serve as illustrative cases of these new capabilities.

6.1. From Core Collapse to Near Breakout with MESA

Models of massive stars can be evolved in MESA up to the onset of the rapid infall of the iron core (see Paper I, Paper II, and Paper III). However, MESA cannot model the core-collapse event itself. Hence, to transition from the onset of core infall to the explosion phase, we rely on a a variety of approximate procedures (Paper III).

For the current efforts, our approach is as follows. We remove the center section of the model at the location where the entropy per baryon is \( 4 \, k_B \), excising the portion of the model that will have collapsed to form a proto-neutron star. This corresponds approximately to the iron core, typically at about \( 1.5 \, M_\odot \). We allow the model to continue infall until its inner boundary (IB) reaches 200 km, near the location of the stalled shock (thanks to H.-T. Janka 2017, private communication, for suggesting this scheme). After the first few seconds, we account for further fallback by removing negative velocity material at the IB. We are not seeking a numerical model of realistic fallback since that depends on 3D details of the explosion that are beyond what MESA can simulate.

The stellar explosion is induced by injecting energy in a thin layer of approximately 0.01 $M_\odot$ at the IB for 5 ms, at a rate sufficient to raise the total energy of the model to a user-specified value. In the subsequent evolution, nuclear reactions are allowed to change abundances but not to generate energy. This choice is suitable because we are not seeking accurate nucleosynthetic yields. The explosion energy spent to photodisintegrate the core into a mix of protons, neutrons, and alpha particles is soon after roughly repaid by the energy released as those particles recombine to form products such as $^{56}$Ni.

Getting an accurate accounting of the energy balance of that complex process is beyond the scope of this paper and is not attempted in the following examples. Our choice to exclude nuclear energy generation can be seen as a simplifying assumption that the cost of photodisintegration is balanced by the return from later recombination. For users wishing to refine this, any excess change in energy from nuclear reactions can be included in the specification of the post-explosion total energy of the model.

The conservation of total energy throughout the run is estimated by summing the per-step errors from post-explosion to near breakout. At each time step, we compare the actual change in total energy between the initial and final models for the step to the change expected from surface luminosity and neutrino losses over the duration of the step. The runs for the models reported below typically show relative cumulative errors in the conservation of total energy of less than 1\%, with most of that error happening in the first few minutes post-explosion when the shock is most extreme. For later stages, the cumulative relative error is orders of magnitude smaller.

The post-explosion evolution of the MESA model is determined by the shock traversal through the star and the resulting Duffell RTI. Figure 29 illustrates the difference between models with and without the effects of the RTI by showing density and pressure profiles. They are shown when the forward shock is about halfway through the star and when the reverse shock originating at the H/He boundary has reached \( \approx 4 \, M_\odot \) on its way to the center. The reverse shock is primarily responsible for the large RTI effects evident in the plots. The online animated figure shows the time evolution of these and many other quantities of interest from seconds after explosion to near shock breakout.
### Table 3
Key Properties and Parameters Associated with the Core-collapse SN Models

| Case    | $M_{ZAMS}$ [M$_\odot$] | Z   | $(v/v_c)_ZAMS$ | $\eta_{wind}$ | $f_{ov}$ | $\alpha_{MLT,H}$ | $M_{exp}$ [M$_\odot$] | $T_{eff}$ [K] | $R_{exp}$ [R$_\odot$] | $\log(L_{exp}/L_\odot)$ | $M_{He}$ [M$_\odot$] | $M_{c,i}$ [M$_\odot$] | $E_{exp}$ [10$^{51}$ erg] | $M_{c,f}$ [M$_\odot$] | $\eta_{He}/\eta_L$ | $t_{boxcar}$ [y] | $M_{CSM}$ [M$_\odot$] | $v_{CSM}$ [km s$^{-1}$] |
|---------|------------------------|-----|----------------|---------------|---------|------------------|------------------------|---------------|------------------------|-------------------------|------------------------|-----------------------|------------------------|------------------------|------------------|----------------|------------------|--------------------------|
| std_16  | 16.0                   | 0.02| 0.2            | 0.4           | 0.01    | 3.0              | 14.5                   | 3960          | 759                    | 5.11                    | 5.58                   | 1.58                  | 0.65                   | 1.58                   | 12.9             | 0.04             | 1.0              | 0                         |
| 99em_16 | 16.0                   | 0.02| 0.2            | 0.4           | 0.01    | 3.0              | 14.5                   | 3960          | 759                    | 5.11                    | 5.58                   | 1.58                  | 0.60                   | 1.58                   | 12.9             | 0.04             | 2.0              | 3                         |
| 99em_19 | 19.0                   | 0.02| 0.2            | 0.4           | 0.00    | 3.0              | 17.8                   | 4490          | 603                    | 5.13                    | 6.58                   | 1.50                  | 0.78                   | 1.50                   | 16.3             | 0.04             | 1.0              | 3                         |
| 05cs    | 13.0                   | 0.006| 0.0           | 0.1           | 0.01    | 3.0              | 12.9                   | 4280          | 537                    | 4.95                    | 4.37                   | 1.57                  | 0.16                   | 2.51                   | 10.4             | 0.009            | 7.0              | 1                         |
| 09N     | 13.0                   | 0.006| 0.0           | 1.0           | 0.01    | 3.0              | 11.6                   | 4290          | 549                    | 4.96                    | 4.34                   | 1.67                  | 0.36                   | 1.67                   | 9.9              | 0.028            | 30.0            | 3                         |
| 12A     | 11.8                   | 0.02| 0.2           | 0.1           | 0.002   | 3.0              | 11.6                   | 4300          | 525                    | 4.94                    | 4.08                   | 1.49                  | 0.28                   | 1.49                   | 10.1             | 0.009            | 3.0              | 2                         |
| 13bvn   | 11.0                   | 0.02| 0.0           | 0.0           | 0.001   | 2.0              | 3.4                    | 26520         | 7.24                    | 4.37                    | 3.40                   | 1.57                  | 0.95                   | 1.57                   | 1.8              | 0.110            | 1.0              | 5                         |
| stripped| 17.0                   | 0.02| 0.3           | 0.0           | 0.001   | 3.0              | ...                   | ...           | ...                    | ...                     | ...                    | ...                   | 0.63                   | ...                   | 0.037            | 20.0            | 0.0              | 0                         |

**Note.** The column “Case” identifies the model. The “Progenitor Parameters” subtable lists input parameters used during the MESA evolution of the models to core infall: initial mass ($M_{ZAMS}$), initial metallicity (Z), initial rotation ($(v/v_c)_ZAMS$), overshooting parameter ($f_{ov}$), wind scaling factor ($\eta_{wind}$), and the mixing length for MLT in the H envelope ($\alpha_{MLT,H}$). The “Stellar Properties at the Time of Explosion” subtable lists the physical quantities evaluated in the MESA model at the time the Fe core begins to infall: mass ($M_{exp}$), effective temperature ($T_{eff}$), radius ($R_{exp}$), luminosity ($L_{exp}$), mass of the He core ($M_{He}$), and initial mass of the Fe core that will be excised ($M_{c,i}$). The “Explosion Properties and Parameters” subtable lists input parameters like the total energy after explosion ($E_{exp}$) and the $^{56}$Ni mass ($M_{Ni}$) as well as properties of the model including the final core mass after fallback ($M_{c,f}$) and the total ejecta mass ($M_{ej}$). This subtable also lists input parameters used in the MESA plus STELLA modeling such as the RTI parameter ($\eta_{He}/\eta_L$) and the number of boxcar smoothing passes ($t_{boxcar}$). Parameters controlling the extent of the CSM are also needed; for a wind profile, this includes the wind duration ($t_{CSM}$), mass loss rate ($M_{CSM}$), and velocity ($v_{CSM}$). Many properties are omitted for the stripped case because this is an ensemble of models with a range of envelope stripping (see Section 6.9).
6.2. From Near Breakout through the Plateau: Introducing STELLA

To follow the evolution of the model through shock breakout and beyond, we use a multigroup (i.e., frequency-dependent) radiation hydrodynamics code.\textsuperscript{19} When the shock is near breakout, we hand the \textsc{mesa} model off to \textsc{stella} in an appropriate form, which involves interpolating to the desired grid and optionally adding circumstellar material (CSM) according to user specifications. With that done, \textsc{mesa} is finished, and \textsc{stella} takes over (see Section 6.3 for a discussion of how we select when to hand off).

\textsc{stella} (Blinnikov et al. 1998; Blinnikov & Sorokina 2004; Baklanov et al. 2005; Blinnikov et al. 2006) is able to model SN evolution at early times, before the expansion is homologous. It can also handle shock breakout and interaction with CSM outside the conventional stellar photosphere. \textsc{stella} is an implicitly differenced hydrodynamics code that incorporates multigroup radiative transfer. The time-dependent equations are solved implicitly for the angular moments of intensity averaged over fixed frequency bands. \textsc{stella} takes about the same amount of time for near-breakout to post-plateau evolution as \textsc{mesa} takes to simulate from explosion to near breakout: about an hour on current workstations.

\textsc{stella} solves the radiative transfer equations in the intensity momentum approximation in each frequency bin. We use 40 to 200 frequency groups, enough to produce bolometric luminosities and broadband colors, but not sufficient to produce spectra. Better broadband light curves can be produced with a larger number of frequency groups, but 40 is sufficient for a bolometric light curve and gives faster runtimes since each group must be represented by a variable and an equation at each zone. The opacity is computed based on over 153,000 spectral lines from Kurucz & Bell (1995) and Verner et al. (1996). The expansion opacity formalism from Eastman & Pinto (1993) is used for line opacities taking high velocity gradients into account. The opacity also includes photoionization, free–free absorption, and electron scattering. LTE is assumed in the plasma, which allows the use of the Boltzmann–Saha distribution for ionization and level populations. \textsc{stella} does not include a nuclear reaction network except for the radioactive decay chain initiated from $^{56}\text{Ni}$. To calculate the overall opacity, the code uses 16 species: H, He, C, N, O, Ne, Na, Mg, Al, Si, S, Ar, Ca, a sum of stable Fe and radioactive $^{56}\text{Co}$, and stable Ni and radioactive $^{56}\text{Ni}$. Energy from nickel and cobalt radioactive decay is deposited as positrons and gamma-rays and is treated in a one-group transport approximation according to Swartz et al. (1995).

\textsc{stella} solves the conservation equations for mass, momentum, and total energy on a Lagrangian comoving grid. It employs artificial viscosity based on the standard von Neumann artificial viscous pressure used for stabilizing solutions (Von Neumann & Richtmyer 1950) and a cold artificial viscosity used to smear shocks (Blinnikov et al. 1998; Moriya et al. 2013). The coupled equations of radiation hydrodynamics are solved through an implicit high-order predictor–corrector procedure based on the methods of Gear (1971) and Brayton et al. (1972); see Blinnikov & Panov (1996) and Stabrowski (1997) for details.

We explore the sensitivity of the bolometric light curves ($L_{\text{bol}}$) reported by \textsc{stella} to the number of frequency bins, spatial zoning, and error tolerances. The result of our sensitivity study is that 40 frequency bins, 300 spatial zones, and an error tolerance of 0.001 for the Gear–Brayton method typically give a converged model. In our experience using \textsc{mesa} and \textsc{stella} for Type IIP SNe, we have not found cases that require different values for the number of frequency bins and error tolerance. Some cases may need a larger number of zones in order to minimize numerical artifacts producing spurious oscillations in the light curve. This problem can often be fixed by a relatively small increase in the number of zones; this is shown for a case similar to SN 2012A in Figure 30.

6.3. Handing Off from \textsc{mesa} to \textsc{stella}

A time must be chosen to hand off the \textsc{mesa} model to \textsc{stella}. This choice is driven by a compromise between two considerations. First, RTI modeling ceases once \textsc{stella} is running even though the effects of RTI may not be complete at that time. Therefore, one wants the model to remain in \textsc{mesa} as long as possible. But second, \textsc{stella} more accurately handles shock breakout and the outermost layers, especially if any matter is placed above the photosphere or if significant

\textsuperscript{19} \textsc{mesa} can be run through shock breakout and beyond, but we do not view gray opacity light curves as sufficient for quantitative comparisons to observed SNe.
radiation is free-streaming from just below the photosphere prior to shock breakout. Moreover, the sophisticated multigroup radiation transfer of STELLA will do a much better job than (gray) MESA at later times post-breakout. Hence, for longer-term light curve evolution, this motivates the default choice to perform this handoff just before breakout.

In order to illustrate the effects of this choice, Figure 31 shows bolometric light curves for cases where the handoffs are done at different times. Note that this plot shows MESA being forced to run post-breakout even though that is not recommended. The deviation of the light curves for later handoff is primarily the result of STELLA doing a better job because of its multigroup radiation transfer rather than any late-stage RTI effects being captured by MESA that are missed by STELLA. That is because, for this case, the H envelope is of normal thickness and the reverse shock from the H/He boundary has time to reach the center, completing essentially all of the RTI effects before breakout.

In the runs presented in the remainder of this section, we choose to do the MESA-to- STELLA handoff shortly before breakout, as determined by the outgoing shock front reaching a location 0.11 M☉ below the surface of the model (this location is a user-defined parameter). Again, we note that in some cases the reverse shock is still far from the center at this moment, and not all of the RTI mixing has completed. In particular, this is true for models with a partially stripped envelope (see Section 6.9). For now, this remains a caveat for the user; a solution would be to have the post-breakout radiation hydrodynamics code include a treatment of the effects of RTI. When presenting the results, we define t = 0 as the time of shock breakout—which we identify using the peak of the bolometric luminosity—and not the (earlier) time of the MESA-to- STELLA handoff.

Because of STELLA’s treatment of radiation hydrodynamics, we have not had to take the progress of the model toward homologous expansion into consideration in selecting a time to hand off from MESA. However, this is a consideration for doing a handoff to radiative transfer codes that assume homology. More accurate spectral and light curve modeling with full radiative transfer, such as EDDINGTON (Eastman & Pinto 1993), SEDONA (Kasen et al. 2006), and CMFGEN (Dessart & Hillier 2010), assume homologous expansion in their current applications to SNe, and this should be considered when deciding the time to hand off from another simulation. Indeed, Eastman et al. (1994) and Dessart & Hillier (2011) discuss this challenge, especially for the innermost material that has not reached a homologous stage and can still have a reverse shock running through it. Approximations made in mapping to a thereafter homologous code can impact the late-time photospheric velocity evolution and the nebular line width predictions associated with the innermost ejecta.

In contrast, STELLA does not assume homologous expansion, so early handoffs are fine; it can handle the effects of the
remaining pressure gradients as the model moves toward homologous expansion. This is important, as the time it takes to reach homology in these models can be quite long. Figure 32 shows velocity evolution results for a model similar to SN 1999em (see discussion in Section 6.8). Homologous expansion would imply that \( \frac{v}{r} \) is flat, whereas a 20% variation from simple homology is evident at 20 days. An additional way in which homology can be violated long after shock breakout is from \( ^{56}\text{Ni} \) decay, especially in Type Ia SNe (Woosley et al. 2007b). As is evident in Figure 32, the much smaller mass fractions of \( ^{56}\text{Ni} \) in Type IIP SNe do not cause such a problem. The contrasting light curves with and without \( ^{56}\text{Ni} \) are shown in Figure 33, exhibiting the prolonging of the plateau due to radioactive decay (Kasen & Woosley 2009; Sukhbold et al. 2016).

### 6.4. Connecting to Observations: Photospheric Properties from STELLA

To set the stage for the rest of this section, we describe a particular model in detail. Figure 34 shows the evolution of a model we have constructed to be similar to the Type IIP SN 1999em (99em_19 in Table 3). The quantities shown are those generated during the STELLA phase of the evolution. Panel (a) is the bolometric luminosity, while panel (b) shows the velocity at the location of the photosphere (where \( \tau_{\text{ros}} = 2/3 \)), and panels (c) and (d) show the mass and radius coordinate of this location. This illustrates the familiar result that the photosphere only reaches the deeper parts of the ejecta after about day 50. The radiation and gas temperatures at the photosphere are shown in panel (e), as is an effective temperature defined by the bolometric luminosity leaving the photosphere. Panel (f) shows the optical depth to the IB, highlighting the fact that the radiative diffusion approximation is excellent (since \( \tau_{\text{IB}} \gg 1 \)) until day 120, at which point the plateau ends and the IB temperature (panel (g)) approaches that of the photosphere.

Figure 33. Light curves and velocities for a model similar to SN 1999em with \( \approx 0.04 \text{ M}_\odot \) \( ^{56}\text{Ni} \) and without \( ^{56}\text{Ni} \). The main effect of the radioactive decay is to prolong the plateau.

Figure 34. Photosphere and IB properties of a model (99em_19 in Table 3) similar to the Type IIP SN 1999em, as a function of time. From top to bottom, the figure shows the bolometric luminosity, velocity, mass coordinate, and radius coordinate of the photosphere; three temperatures (gas, radiation, and “effective”) at the photosphere; the optical depth to the IB; and the IB temperature. The gray line in panel (c) shows the Lagrangian mass coordinate of the IB.

Figure 35. Comparison of model 99em_19 with the multicolor light curve of SN 1999em, showing colors from STELLA and blackbody colors from MESA. Circles indicate observational data. This demonstrates the effect of the number of STELLA frequency bins on the predicted colors.
Our emphasis is on bolometric luminosities, where 40 STELLA frequency bins are adequate. However, broadband light curves are also reported by STELLA. Figure 35 shows how the STELLA colors change as one goes from 40 to 200 frequency bins in a model approximately matching the bolometric luminosity of SN 1999em (99em 19 in Table 3). This reflects the fact that a given band is spanned by only a small number of frequency bins. The non-public research version of STELLA can opt to use many more frequency bins to address under-resolution issues. There are no current plans to include that capability in MESA. We also show what a blackbody would predict using the MESA colors module (see Appendix A). This makes it clear that the line-blanketing in the U band is well handled by STELLA. We do not include colors in our subsequent discussions, but we expect they may be useful to users who have access to observations in one or two bands, but not enough data to produce a bolometric light curve from observations.

6.5. Connecting to Observations: Fe II Line Velocities

It is important to be able to interpret the ejecta velocities measured by observers, which are often inferred from the absorption minimum in the Fe II 5169 Å line. Modeling these absorption features requires more detailed radiative transfer than available in STELLA. However, rather than assume that the photospheric velocity reported by STELLA is identical to that of the Fe II 5169 Å line, we have added the capability of finding the location (and hence the velocity) of the material above the photosphere where the Sobolev optical depth in the Fe II 5169 Å line is a specified value. This will prove to be most important after day 30 or so, when the photosphere has started to move inward in mass coordinate into ejecta with a shallow density profile.

The strength of a line in a homologously expanding atmosphere is quantified by the Sobolev optical depth (Sobolev 1960; Castor 1970; Mihalas 1978; Kasen et al. 2006), which for the Fe II line at any position is

\[ \tau_{\text{Sob}} = \frac{\pi q_{e}^{2}}{m_{e}c} n_{\text{Fe}} \eta_{i} f_{\text{exp}} \lambda_{0} \]

where \( \lambda_{0} = 5169 \) Å is the line center wavelength for the Fe II line, \( f = 0.023 \) is its oscillator strength, \( n_{\text{Fe}} \) is the number density of iron atoms, and \( f_{\text{exp}} \) is the time since breakout. The quantity \( \eta_{i} \) is the fraction of iron atoms that are in the lower level of the transition of interest and depends on the properties of the gas. D. Kasen (2017, private communication) provided an \( \eta_{i}(\rho, T) \) table for post-processing to produce the Fe II line velocities, calculated under the assumption of LTE and covering \( \log(\rho / \text{g cm}^{-3}) = -16 \) to \(-8\) and \( \log(T / \text{K}) = 3.3 \) to 4.3.

We use Equation (53) after the STELLA run to provide the velocity of material that satisfies a chosen value of \( \tau_{\text{Sob}} \). This yields a velocity that can be compared to the measured Fe II line velocities. Figure 36 shows the resulting comparisons for various choices of \( \tau_{\text{Sob}} \) for a model similar to the Type IIP SN 1992A found solely by matching the bolometric luminosity (upper panel). The lower panel displays the Fe II 5169 Å data and the velocities derived from the photosphere and for a range of values of \( \tau_{\text{Sob}} \). At early times, there is little difference between the photospheric velocity and that of the Fe line. However, as the photosphere moves deeper into the ejecta, the two velocities substantially diverge. The velocity inferred from the Sobolev argument gives a much better match to observations than the photospheric velocity. Motivated by this comparison, we choose \( \tau_{\text{Sob}} = 1 \) for our later plots, a parameter that the user is free to adjust.

6.6. The Impact of Pre-breakout RTI Mixing

We have previously outlined the inclusion of a method for RTI mixing in MESA (the Duffell scheme; Section 5), the use of MESA to evolve models pre-breakout (Section 6.1), and the use of STELLA to evolve models post-breakout (Section 6.2), and described how to connect the models to observations (Sections 6.4 and 6.5).

In this way, MESA plus STELLA allows users to explore the impact of RTI mixing on Type IIP light curves and velocities. Prior work in this direction (Eastman et al. 1994; Utrobin et al. 2007; Dessart & Hillier 2010, 2011; Morozova et al. 2015) focused on the impact of compositional mixing, often with averaging approaches to achieve various levels of mixing. Only the recent work of Utrobin et al. (2017) incorporated compositional mixing from a 3D model and also included the modified density and velocity structures, also seen in the 1D RTI mixing (P. Duffell et al. 2018, in preparation).

Figure 37 shows the light curves and velocities of model 99em 19. The luminosity without RTI mixing has a distinctive rise just before the plateau as shown by Eastman et al. (1994)
and Utrobin (2007). As the RTI causes many associated changes in composition, density, velocity, and energy density for the innermost material, we cannot specifically identify the immediate cause of the lengthening of the plateau phase when the RTI is incorporated without further experiments. These are now possible using MESA and STELLA but are beyond the scope of this paper. The lower panel shows the photospheric and Fe II line velocities with and without RTI mixing. The most evident change is at the end of the plateau, when the material that was near the H/He boundary in the red supergiant is approaching the SN photosphere. That material is strongly affected by RTI mixing, as shown in Figure 28 and discussed in P. Duffell et al. (2018, in preparation).

To enable the exploration of the impact of various components of the RTI mixing, we explicitly allow for the diffusion coefficients for density, momentum, energy, and composition to be scaled by independent constant factors relative to the fiducial value \( \eta_R \) given in Equation (47). We show in Figure 38 the impact of varying the coefficient in the internal energy flux in Equation (48), which we refer to as \( \eta_{R,e} \). These plots show the energy density and density of the ejecta just before shock breakout in one of our models (99em_19) in Table 3. The blue line is for the fiducial value, whereas the red line is for an extreme increase of a factor of 100. The only locations that are sensitive to these changes are the innermost mass coordinates where the RTI was most active, the same regions where the light curve and velocities seem to be sensitive to changes related to RTI mixing. The variable \( \eta_{R,e} \) was found to be a useful “knob” to vary for the modeling of specific SNe.

6.7. Exploring the Explosive Landscape

A strength of the new MESA plus STELLA capabilities is their ease of use. This enables detailed quantitative studies of large numbers of core-collapse SNe. The open source nature of MESA, the inclusion of STELLA in the MESA distribution, and the repository of examples contained within the MESAstar test suite allow a user to obtain models that can be compared directly to observations. Indeed, with minimal manual intervention, a user can take a star from the pre-MS to the SN light curve within a few hours of computer runtime. To emphasize this point, we describe here how this might be done (Section 6.7.1). To demonstrate how parameter choices affect light curves, we show a large sample of variations of a standard case for “high–middle–low” settings of some of the main parameters (Section 6.7.2). In Section 6.8, we will exhibit a few specific models created to be roughly similar to known Type IIP SNe. The potential is clear for an extensive database of such SNe models created using MESA and STELLA; its actualization is beyond the scope of this paper.

6.7.1. Generating Models with MESA plus STELLA

The first step in generating a core-collapse SN light curve is to use MESA to make a pre-SN stellar model that is undergoing core collapse. The test case example_make_pre_ccsn can serve as a useful template. As part of the required inlists, the user must select values for the main variables: initial mass \( M_{\text{ZAMS}} \), initial metallicity \( Z \), initial rotation \((\nu/\Omega)_{\text{ZAMS}}\), overshooting parameter \( f_\text{ov} \), wind scaling factor \( \eta_{\text{wind}} \), and the mixing length for MLT in the H envelope \( \alpha_{\text{MLT,H}} \). Of course, the user may tune other MESA parameters of interest. The run from pre-MS to Fe core infall runs automatically given these parameters and, depending on the case, takes roughly an hour on a modern multicore desktop workstation. Users interested in details of pre-SN models may require settings that lead to significant additional runtime (e.g., Farmer et al. 2016; Renzo et al. 2017).

The second step loads the model at core infall into MESA, emulates the core-collapse explosion by excising the core and injecting energy and Ni (as described in Section 6.1), and evolves until near shock breakout. The test case example_ccsn_IIP can serve as a useful template. Again,
the user must set the value of the various “knobs” controlling the properties of the explosion such as the total energy $E$ and the $^{56}$Ni mass $M_{\text{Ni}}$. Early ($t < 20$ days) light curves of core-collapse SNe are better fit when large amounts of CSM are placed outside the conventional photosphere (Morozova et al. 2016; Dessart et al. 2017; Morozova et al. 2017a, 2017b). We provide an option to include CSM. We also provide the option for “boxcar” smoothing of the model abundances before the handoff from MESA to STELLA (Kasen & Woosley 2009; Dessart et al. 2013; Morozova et al. 2015). The end result of this step is a model suitable for input into STELLA, so one must also indicate the number of STELLA zones to be used. This MESA phase from after explosion to near breakout typically takes about 30 minutes on a modern multicore desktop workstation.

The final step uses the results produced in the previous step as input to STELLA and evolves the model through shock breakout to the post-plateau phase. A script to execute STELLA is provided. This stage takes about an additional 30 minutes on a modern multicore desktop workstation for typical cases. When STELLA finishes, a post-processing step produces data for comparison to observational results.

6.7.2. Sensitivity to Variations in Key Parameters

Figure 39 exhibits the std_16 model light curves as progenitor parameters are varied. Many variations behave as expected from previous analytical and numerical scalings (Popov 1993; Kasen & Woosley 2009; Sukhbold et al. 2016). For example, the decrease in the plateau duration with lower ZAMS masses or higher mass loss (increased $\eta_{\text{wind}}$) is as expected. The increase in plateau luminosity with decreasing $\alpha_{\text{MLT,H}}$ is because those stars with lower $\alpha_{\text{MLT,H}}$ have a larger stellar radius at the time of explosion. However, other variations in these figures are not as easily diagnosed.

Figure 40 exhibits model light curves as explosion parameters are varied. Again, many cases lead to the expected outcomes, such as the increase in the plateau luminosity with increasing explosion energy and the increased duration of the plateau with increasing nickel mass. The changes caused by varying the RTI parameters are slight for the compositional mixing and boxcars, though, as we discussed in Section 6.6, modifying the diffusion of energy density during RTI does impact the shape at the end of the plateau. The impact of the CSM is similar to that shown by Morozova et al. (2017b) and Dessart et al. (2017).

With experience in the effects of varying the parameters (knobs) shown in Figures 39 and 40, it is sometimes possible to get a rough match between the model and observations after a dozen or so attempts. That is about the amount of effort we undertook to get the models similar to the various observed SNe presented in Section 6.8. Of course, the effects of the various knobs do not combine in any simple manner, so it can be a nontrivial challenge to find a combination that gives a good match for both velocities and light curve. Our experience suggests that it is a good strategy to match velocities before light curves since there are few ways available to shift velocities and many ways to change light curves. It is important to include velocities in judging potential matches because of the multiple degeneracies, as will be seen below where we show two models similar to SN 1999em with quite different ejecta masses and explosion energies. Even when using both velocities and light curves, it remains a challenge to find a unique “best” match.

![Figure 39. Effect of single-parameter variations of the progenitor std_16. The upper three panels vary initial properties of the star; the lower three vary modeling assumptions during evolution to core collapse.](image-url)
6.8. Applications to a Few Type IIP Supernovae

To show examples of what can be accomplished with these new capabilities, we have modeled four Type IIP SNe: 1999em, 2005cs, 2009N, and 2012A. These cover a range of luminosities, plateau durations, and nickel masses, and have readily available data (Pejcha & Prieto 2015a, 2015b) for bolometric luminosities and Fe II velocities. We follow the steps described in Section 6.7, iterating to reach the matches shown. The models are not intended to demonstrate the best matches that can be achieved using MESA and STELLA. An investment of more effort could produce better matches but is beyond the scope of this paper. The parameters we chose are shown in Table 3.

We note a few general insights gained from our modeling. We found that the radii of red supergiant models from MESA were too large for these Type IIP SNe models unless we set $3 \eta_{R,e} = \eta_{R} \times 1.0$. All models benefited at early times by having some CSM present. Figure 41 shows how the early 1999em model predictions change as CSM is added to the value shown in Table 3. The luminosity at early times is a far better match, as are the earliest velocity data. As expected, by day 50 and beyond, there is no impact of the CSM on the model predictions. Comparisons of how the luminosity collapsed at the end of the plateau drove us to prefer an enhancement in $e$, $\nu_{h}$ in several cases.

To exhibit some of the possible degeneracies, we constructed two distinct models for 1999em. As shown in Figure 42, they are both reasonable models for the bolometric luminosity and Fe II velocities. However, their ejected masses and radii differ significantly—one has 12.9 M$_{\odot}$ and 770 R$_{\odot}$, whereas the other has 15.9 M$_{\odot}$ and 600 R$_{\odot}$. Utrobin (2007) gave an ejected mass of 19.0 ± 1.2 M$_{\odot}$, a radius of 500 R$_{\odot}$, and an explosion energy of $1.3 \times 10^{51}$ erg. Bersten et al. (2011) gave an ejecta mass of 17.6 M$_{\odot}$, radius of 800 R$_{\odot}$, and explosion energy of $1.25 \times 10^{51}$ erg . Utrobin et al. (2017) model this event with a 3D simulation from explosion to shock breakout, similar to the Munich L15 model we discuss in Section 5, but with an explosion energy of about $0.5 \times 10^{51}$ erg . For comparison, the MESA models for 1999em have total energies after explosion of
0.60 \times 10^{51} \text{erg} \text{ for the case with } 12.9 \, M_\odot \text{ ejected mass, and } 0.78 \times 10^{53} \text{erg} \text{ for the case with } 16.3 \, M_\odot \text{ ejected mass.}

We previously showed 2012A in Figure 36. Our model had an ejected mass of 10.1 \, M_\odot, compared with 7.8 \, M_\odot from Morozova et al. (2017a), 12.5 \, M_\odot from Tomasella et al. (2013), and 13.1 \pm 0.7 \, M_\odot from Utrobin & Chugai (2015). Tomasella et al. (2013) also reported a progenitor luminosity of \log(L/L_\odot) = 4.73 \pm 0.13, just a bit fainter than our model’s value. Figure 43 shows our model for 2005cs. Our model has an ejected mass of 10.4 \, M_\odot, slightly higher than the 9.5 \, M_\odot reported by Spiro et al. (2014) and the 7.8 \, M_\odot reported by Morozova et al. (2017a). Figure 44 shows our model for 2009N, which has an ejected mass of 9.9 \, M_\odot, whereas Morozova et al. (2017a) found 9.3 \, M_\odot and Takáts et al. (2014) found 11.5 \, M_\odot.

6.9. Partially Stripped Core-collapse Supernovae

There is a well-defined class of core-collapse SNe where either much (Type IIb) or nearly all (Type Ib and Ic) of the H envelope was lost prior to the core-collapse event. Dessart et al. (2015) performed detailed radiative transfer models for a large set of progenitors from binary evolution, while Morozova et al. (2015) carried out diffusive calculations with varying amounts of mass loss. Yoon et al. (2017) explored MESA models constructed from binary transfer scenarios and applied them to a set of well-observed Type IIb events. We have not yet been able to deal successfully with Ic models because of numerical problems related to the extreme ejecta velocities that occur at shock breakout. However, it is possible to do both IIb and Ic models as shown here.

In Figure 45, we show the MESA plus STELLA predictions for luminosities and photospheric velocities for a range of models with varying amounts of mass stripped from a 17 \, M_\odot ZAMS model, ranging from the entire initial H envelope still remaining down to only 0.1 \, M_\odot of the H envelope left at the time of explosion. Similar to Figure 7 of Morozova et al. (2015), the plateau period becomes shorter as the residual H shell mass declines. Our smallest mass model has a H envelope mass comparable to typical models of Type IIb SNe and generates a light curve comparable to observed Type IIb SNe (Ergon et al. 2015). Figure 46 shows the interior properties of these same models near the moment of shock breakout. For models that have been stripped, the reverse shock has not reached the IB at the time the forward shock reaches the surface. Since RTI mixing does not occur in STELLA, these models would incompletely include the effects of the RTI.

Cao et al. (2013) discovered the fully stripped Type Ib SN iPTF13bvn in the nearby spiral galaxy NGC 5806 with the intermediate Palomar Transient Factory (Law et al. 2009). This is one of only a few stripped SNe with a progenitor detection. Using data from Cao et al. (2013) andFremling et al. (2014), we show in Figure 47 our model that approximately matches
the iPTF13bvn light curve. The model is derived from an
11 M\text{\odot} ZAMS model and has a remaining mass of only 3.4 M\text{\odot} at the time of explosion with total energy after explosion of $0.95 \times 10^{51}$ ergs and a $^{56}$Ni mass of 0.11 M\text{\odot} distributed throughout the remaining star (ejecta mass 1.8 M\text{\odot}). Fremling et al. (2014) also modeled this light curve, finding the total energy to be $0.85^{+0.5}_{-0.4} \times 10^{51}$ ergs, with a $^{56}$Ni mass of $0.049^{+0.012}_{-0.02}$ M\text{\odot} and total ejecta mass of $1.94^{+0.58}_{-0.50}$ M\text{\odot}.

Our parameters are similar, falling within the range of the quoted uncertainties, except for the $^{56}$Ni mass.

7. Black Hole Formation

Compact objects are a natural product of the evolution of massive stars. A broad consensus on which massive stars produce black holes (BHs) has not yet been reached (Timmes et al. 1996; Fryer & Kalogera 2001; Heger et al. 2003; Eldridge & Tout 2004; Zhang et al. 2008; Ugliano et al. 2012; Clausen et al. 2015; Müller et al. 2016; Sukhbold et al. 2016; Limongi 2017).

The lack of consensus is due to a variety of differences in the modeling, including stellar wind treatments during the pre-SN stage (Renzo et al. 2017); shellular rotation prescriptions (e.g., Limongi 2017); sensitivity to the initial metallicity (e.g., O’Connor & Ott 2011), number of isotopes in the reaction network (Farmer et al. 2016), adopted values of critical reaction rates (deBoer et al. 2017; Fields et al. 2018); and ignition of core carbon burning (Farmer et al. 2015; Cristini et al. 2018; I. Petermann et al. 2018, in preparation); variations from spatial and temporal resolution (Farmer et al. 2016); convection during core collapse (e.g., Couch et al. 2015); and effects from binary partners (e.g., Marchant et al. 2016; Batta et al. 2017). In addition, current estimates of the neutron star and BH initial mass function chiefly rely on parameterized explosion models and not on first principles calculations.

This section explores \textsc{Mesa} models that can produce BHs. First, we consider $M_{\text{ZAMS}} \lesssim 60$ M\text{\odot} models that can form a BH without encountering dynamical instability due to $e^+e^-$ pair production. Second, we survey $M_{\text{ZAMS}} \gtrsim 60$ M\text{\odot} models that encounter dynamical instability, either entering the $\Gamma_1 \lesssim 4/3$ regime once to produce a pair-instability SN (PISN; Fowler & Hoyle 1964; Barkat et al. 1967; Rakavy & Shaviv 1967; Rakavy et al. 1967; Fraley 1968; Ober et al. 1983; Fryer et al. 2001; Scannapieco et al. 2005; Kasen et al. 2011; Chatzopoulos et al. 2013), or multiple times to produce a pulsational pair-instability SN (PPISN) and a BH remnant.
neutrinos removing $\approx 10^{53}$ erg during core collapse, reducing the gravitational mass of the core by $\approx 0.3 M_\odot$. The outer part of the star responds to the sudden decrease in the gravitational field by driving a sound wave that steepens into a shock that unbinds some of the outer envelope (Coughlin et al. 2017). Mass with binding energy $\lesssim 10^{47}$ erg is likely to be ejected (Nadezhin 1980; Lovegrove & Woosley 2013) while mass that is not ejected will likely become part of the BH. Figure 48 suggests that BH masses estimated in this simple way can be significantly larger than the final He core mass and more sensitive to the assumed model parameters. For example, there is a wide variation in the expected BH mass for the 60 $M_\odot$ progenitor depending on the choice of wind scheme and scaling factor, whereas modest rotation has a smaller effect.

7.2. Pulsational Pair-instability Supernovae

Stars with $M_{\text{ZAMS}} \gtrsim 60 M_\odot$ are expected to become dynamically unstable before core O depletion as $e^+e^-$ pair production leads to regions where the adiabatic index $\Gamma_\text{ad} \lesssim 4/3$ (Fowler & Hoyle 1964; Rakavy & Shaviv 1967). The ensuing collapse results in explosive O burning, with a variety of possible outcomes. Stars can produce PISNe where the energy injected from explosive O burning completely unbinds the star without leaving a compact remnant. Alternatively, stars can undergo a cyclic pattern of entering the pair-instability region, contracting, burning, and expanding, leading to PPISNe.

Individual pulses in a PPISN can remove a large fraction of the mass of the star at velocities of several thousand km s$^{-1}$, with the remaining material settling down into hydrostatic equilibrium at a lower central temperature than before the pulse. The star then contracts as it loses energy due to radiation and neutrino emission until it undergoes an additional pulse or collapses to form a BH. Depending on its initial mass, the time between pulses varies from a fraction of a year to millennia, with the outer ejected layers expanding to very low densities and becoming optically thin.

FIGURE 48. He core mass (upper panel) and mass location where the gravitational binding energy is equal to $10^{48}$ erg (lower panel) for ZAMS masses of 30, 45, and 60 $M_\odot$. Three stellar wind treatments, two wind scaling factors, and two rotation rates are shown for each ZAMS mass. The variation, illustrated by the tan band, induced by these modeling choices increases with ZAMS masses. Also shown are models from the literature (Woosley et al. 2002; Limongi & Chieffi 2003, 2006; Sukhbold & Woosley 2014; Farmer et al. 2016; Renzo et al. 2017), although each adopts different modeling choices and definitions of the He core mass.

(Barkat et al. 1967; Woosley et al. 2007a; Chatzopoulos & Wheeler 2012; Woosley 2017).

7.1. Progenitors That Do Not Pulse

The upper panel of Figure 48 shows the He core mass from $Z = 0.02$ $M_{\text{ZAMS}} = 30$, 45, and 60 $M_\odot$ models. The lower panel shows the mass location where the binding energy is $10^{48}$ erg. Each ZAMS mass uses an exponential convective overshoot parameter $f_\text{ov} = 0.004$ applied at all convective boundaries, a mixing length $\alpha_{\text{MLT}} = 1.5$, MLT++ enabled (see Paper II), and is run to the onset of core-collapse (infall velocity $\gtrsim 1000$ km s$^{-1}$). We illustrate the variation in the He core mass and mass location where the binding energy is $10^{48}$ erg owing to the effects of rotation, wind strength, and the wind schemes of Nieuwenhuijzen & de Jager (1990), van Loon et al. (2005), and de Jager et al. (1988).

To estimate a BH mass from the structure at core collapse, we use the mass location where the binding energy integrated from the surface exceeds $10^{48}$ erg. This is motivated by...
At the onset of the first pulse, the star has a mass of 87.1 M\(_\odot\), with a He core of 45.6 M\(_\odot\) and a CO core of 41.1 M\(_\odot\). As shown in Figure 50, the first two pulses happen within two days of each other, and they remove the entire H envelope. The remaining two pulses remove almost the entire He envelope, resulting in a final mass of 41.2 M\(_\odot\) when the star collapses into a BH.

Figure 51 shows the key masses on a grid encompassing ZAMS masses for which PPISNe occur under our model assumptions. Our PPISN progenitors have He core masses in the range of 28 M\(_\odot\)–67 M\(_\odot\), and no BHs with masses above 50 M\(_\odot\) are formed. These results are in broad agreement with those of Woosley (2017). However, Figure 51 shows that the range of ZAMS masses that result in a PPISN is significantly different from the one computed by Woosley (2017). This can be attributed to a different choice of input physics, such as core overshooting, as well as a different initial metallicity.

8. Energy Accounting in Stellar Evolution

Paper I describes the stellar structure equations and their implementation in MESA. In order to provide physically and numerically accurate solutions of these equations, it is often necessary to evaluate them in different ways depending on the details of the star being simulated. In particular, there are a number of different ways to formulate and evaluate the equations solved by MESA that encode local and global energy conservation. The goal of this section is to clarify the available options, discuss when and why they are used, and describe how various forms of energy are tracked and accounted for in stellar evolution. While in places this section reads like a tutorial, it is in fact the first time we have presented a detailed description of a complex and critical aspect of how MESA works, information that is important for the intelligent use of this software tool.

In Section 8.1, we describe the fundamental equations we are solving, and in Section 8.2, we describe the choices associated with their numerical implementation. In Section 8.3, we describe the connection between the form of the energy equation typically used in stellar evolution calculations and the version used when the hydrodynamics options discussed in Section 4 are enabled. In Section 8.4, we clarify how the energy associated with ionization is included in MESA. In Section 8.5, we describe the numerical approach necessary to ensure that the latent heat associated with crystallization in a white dwarf (WD) is included in MESA. In Section 8.6, we discuss the difficulties introduced by the necessity to blend different equations of state (EOSs) as the thermodynamic conditions in the stellar interior change, and how MESA minimizes artifacts associated with these blends. In Section 8.7, we discuss the energy associated with gravitational settling.
8.1. Fundamental Equations

In the stellar structure equations (e.g., Cox & Giuli 1968; Kippenhahn et al. 2012), energy conservation is typically formulated by considering the energy flow in and out of a fluid parcel. In this Lagrangian picture, to understand how the energy of a fluid parcel is changing, we account for the specific (i.e., per unit mass) rate of energy injection into the parcel, $\epsilon$, and the specific rate of energy flow through the boundaries ($\partial L/\partial m$; $L(m)$ is the luminosity profile and $m$ the Lagrangian mass coordinate). The specific heating rate ($Dq/Dr$) for the parcel must then satisfy

$$\frac{Dq}{Dt} = \epsilon - \frac{\partial L}{\partial m}, \quad (54)$$

where $D/Dt$ is the Lagrangian time derivative. Except in the case of hydrodynamics described in Section 4 (where a total energy equation is solved; see Section 8.3), the basic equation to be solved is always some form of Equation (54). By tradition, the negative of the left-hand side of Equation (54) is called $\epsilon_{grav}$.

Thermodynamics relates the heating of material to the changes in its properties. The first law of thermodynamics states that the total heat added $bQ$ to a parcel is

$$\delta Q \equiv dE + PdV, \quad (55)$$

where $E$ is the internal energy, $P$ is the pressure, and $V$ is the volume. Let $N_i$ be the number of particles of species $i$ in the parcel. Then, expanding $E$ in terms of the independent thermodynamic basis variables ($S, V, N_i$) yields the following thermodynamic identity:

$$dE + PdV = TdS + \sum_i \mu_i dN_i, \quad (56)$$

where $S$ is the entropy and $T$ is the temperature. The sum runs over all species present, and

$$\mu_i \equiv \left( \frac{\partial E}{\partial N_i} \right)_{S,V} \quad (57)$$

is the chemical potential for species $i$.

The number abundance of every species is defined with reference to the total number of baryons $N_b$ as $Y_i \equiv N_i / N_b$. Denoting Avogadro’s number by $n_a$, the atomic mass unit is $m_{amu} = 1 / N_b$. The specific (i.e., per unit mass) form of Equation (56) is then given by multiplying by the invariant $N_A / n_b$ to find

$$\delta q \equiv de + P d\left( \frac{1}{\rho} \right) = Tds + \sum_i \left( \frac{\partial e}{\partial Y_i} \right)_{s,\rho} dY_i. \quad (58)$$

The total baryonic mass density is $\rho$, so that $1/\rho$ is the specific volume, and $e$ and $s$ are the specific energy and entropy, respectively. Local thermodynamic equilibrium (LTE) determines a unique solution for the ionization state of each isotope. Thus, the composition is completely specified by a set of number abundances $\{Y\}$ for all nuclear isotopes.

Equation (58) is relativistically correct when the rest mass is included in the energy and the chemical potentials. Therefore, in principle, changes in nuclear rest masses due to nuclear reactions could be accounted for via this equation. However, in MESA, the energetic effects associated with composition changes due to nuclear reactions are not included in $\epsilon_{grav}$. Instead, these important terms are accounted for via $\epsilon_{nuc}$ (the specific energy generation rate of nuclear reactions), which is evaluated separately and included as part of the local source term $\epsilon$ in Equation (54) (see Paper I).

It is often convenient to specify compositions in terms of the baryonic mass fractions $\{X_i\}$ via the relation $X_i = A_i Y_i$, where $A_i$ is the mass number for isotope $i$. Since the rest mass changes due to nuclear reactions are handled separately from $\epsilon_{grav}$, $\rho$ and $\{X_i\}$ can be treated as independent basis variables without introducing any ambiguity into the chemical potential term in Equation (58). Some EOS options express the composition dependence in terms of aggregate quantities; examples include hydrogen abundance $X$, helium abundance $Y$, metallicity $Z$, average mass number $\overline{A}$, and average atomic number $\overline{Z}$.

The value for $\epsilon_{grav}$ can be computed beginning from either the left- or right-hand side of the equal sign in Equation (58). Usually, some form of the left-hand side is used, but in Section 8.5 we will describe a case where it is more convenient to use the right-hand side.

8.2. Implementation

Basic variables are those quantities directly calculated by MESAstar’s solver. Examples include velocity, radius, and the thermodynamic variables. MESA offers options for selecting ($\rho, T, \{X_i\}$) or ($P_{gas}, T, \{X_i\}$) as the thermodynamic variables. The EOS routines calculate other thermodynamic quantities as a function of the chosen variables, e.g., $e = e(\rho, T, \{X_i\})$.

MESA solves the stellar structure equations implicitly, thus it is possible to approximate total time derivatives of any quantity calculated in the stellar model simply by differencing its value at the start and end of a time step. Therefore, one way to evaluate $\epsilon_{grav}$ would be to directly calculate the time derivatives in Equation (58). Two possible versions of $\epsilon_{grav}$ would then be

$$-\epsilon_{grav} = \frac{T}{v} \frac{DS}{Dt} + \sum_i \frac{\partial e}{\partial Y_i} \frac{DY_i}{Dt}, \quad (59)$$

and

$$-\epsilon_{grav} = \frac{De}{Dt} + P \left( \frac{D}{Dt} \left( \frac{1}{\rho} \right) \right). \quad (60)$$

Although simple to construct, the finite differences necessary to calculate these equations are often numerically problematic.

To see the potential numerical issues, consider the implementation of Equation (54) using Equation (60) in cell $k$ with mass $dm_k$ over a time step $\delta t$. The derivative of a quantity $DY/Dr$ is typically constructed as a finite difference of $Y$ over the time step, so after integrating over the mass of zone $k$, we have

$$0 = \left( e_k - e_{k,end} - e_{k,start} \right) / \delta t - \frac{P_k}{\delta t} \left( \frac{1}{\rho_{k,end}} - \frac{1}{\rho_{k,start}} \right) \frac{dm_k}{\delta t} - (L_k - L_{k+1}). \quad (61)$$

The implicit solver scheme in MESA attempts to reduce the residual from evaluating the right-hand side of this equation below some tolerance.

Although the implicit scheme in MESA may sometimes find acceptable results for an equation such as Equation (61), finite
numerical precision can result in troublesome behavior for the
time derivatives involving subtractions. In particular, over a
small time step where the change in $e_k$ or $\rho_k$ is small compared
to the overall magnitude of these quantities, floating point
arithmetic can suffer significant loss of precision. When energy
scales arising from these types of finite difference derivatives are
comparable to $e_k$, the implicit solver may be unable to converge to an acceptable solution.

To avoid these problems, the equations can be cast in terms of
derivatives that are not evaluated using subtractions. Such
derivatives are available only for the basic variables, since the
Jacobian matrix for an evolution step satisfying the equations
of stellar structure in MESA is written in terms of the basic variables
and their derivatives (see Paper I, Section 6.2, and
Paper II, Appendix B.2 and Figure 47). For MESA, $\rho$ and $T$
serve as default variables.

Modifying Equation (60) to take advantage of $\rho$ as a basic
variable yields

$$-\epsilon_{\text{grav}} = \frac{De}{Dt} - \frac{P}{\rho} \frac{D\ln \rho}{Dt},$$

but the change in $e$ is still evaluated using subtraction. Another
related form, obtained by application of mass continuity, is

$$-\epsilon_{\text{grav}} = \frac{De}{Dt} + P \frac{\partial}{\partial n}(\nu A),$$

where $\nu$ is the cell velocity and $A$ is the area of the cell face.
This is the form used in the artificial-viscosity-based-hydro-
dynamics options described in Paper III.

Expanding the total derivative of energy and thus eliminating
the subtraction motivates the following alternative forms.
Expanding $e$ in terms of its dependence on the basic variables $\rho$
and $T$, and dropping the dependence on composition gives

$$-\epsilon_{\text{grav}} = c_v T \frac{D\ln T}{Dt} + \left[ \rho \left( \frac{\partial e}{\partial T} - \frac{P}{\rho} \right) \frac{D\ln \rho}{Dt},
$$

where $c_v \equiv (\partial q/\partial T)_p$ and $\nabla_{\text{ad}} \equiv (\partial \ln T/\partial \ln P)_p$.

The derivation for this expression in terms of $P$ and $T$ is given in
Chapter 4 of Kippenhahn et al. (2012), from which it is
straightforward to obtain Equation (65) using $\chi_T \equiv
(\partial \ln P/\partial \ln T)_P$ and $\chi_{\rho} \equiv (\partial \ln P/\partial \ln \rho)_T$.

Since $\rho$ and $T$ are basic variables, the time derivatives
appearing in Equations (64) or (65) involve no subtractions.

Hence, solving Equation (54) with $\epsilon_{\text{grav}}$ as defined by those two
equations will not be susceptible to the same losses of
numerical precision as other forms, at the cost of dropping
the composition terms. Similarily, Equation (4.47) of Kippen-
hahn et al. (2012) will yield the same stability when $P$ and $T$
are used as the basic variables. When $P_{\text{gas}}$ and $T$ are selected as
the basic variables, the identification $P = P_{\text{gas}} + \alpha T^4/3$ allows us
to write

$$-\epsilon_{\text{grav}} = c_p T \left[ \left( 1 - 4 \nabla_{\text{ad}} \frac{P_{\text{ad}}}{P} \right) \frac{D\ln T}{Dt} - \nabla_{\text{ad}} \frac{P_{\text{gas}} D\ln P_{\text{gas}}}{P} \right].$$

Section 4.5 in Kippenhahn et al. (2012) also shows how this
local energy treatment of $\epsilon_{\text{grav}}$ results in global energy
conservation, including total gravitational potential energy from
which the name $\epsilon_{\text{grav}}$ is derived.

The superior numerical stability of Equations (64)–(66)
comes at the cost of using derivative quantities such as $c_v$ and $\chi_{\rho}$.

The Jacobian matrix of an implicit method thus requires the
partial derivatives of $c_v$ and $\chi_{\rho}$. An EOS must therefore be capable of returning the state functions $P$, $\rho$, and $s$
along with their first derivatives (e.g., $c_v$ and $\chi_{\rho}$) and second derivatives (e.g., $\partial c_v/\partial T$).

As noted above, Equations (64)–(66) drop the composition
terms, which is justifiable if the derivatives $(\partial e/\partial X)(\partial X/\partial P)$
are negligible for each $X$. Dropping composition terms is often
justified in stellar evolution scenarios where the timescales for
these changes are very slow or their associated energies are
negligible, such as MS burning where energy release from
nuclear burning dominates any small change in the internal
energy due to composition evolution over a single step
(Kippenhahn et al. 1965; García-Berro et al. 2008). Making
this assumption, MESA also offers an option for calculating $\epsilon_{\text{grav}}$
in terms of a simplified form of Equation (59),

$$-\epsilon_{\text{grav}} = \frac{TDs}{Dt},$$

which drops composition dependence to offer an expression
that is more convenient to evaluate.

However, even after composition dependence related to
nuclear burning is accounted for with a separate $e_{\text{nucl}}$ term as
discussed in Section 8.1, other processes that change abundances (e.g., mixing) may be important. In cases where
dropping these terms is not justifiable, it may be necessary to add a compensating local source term $\epsilon$ in Equation (54).

In summary, MESA currently offers options for solving
Equation (54) with $\epsilon_{\text{grav}}$ defined in any of the ways given in
Equations (60)–(67). Figure 52 schematically summarizes the
relationships between these equations, and Table 4 shows the
inlist commands necessary for invoking each of these options.

Usually, the superior numerical stability gained by using
Equation (65) is preferred, and hence it is the MESA default, but
users should be aware of the possibility that other forms may be
necessary to capture important physics. One such case for
Equation (67) is described in Section 8.5. Another is the
artificial-viscosity-based implicit hydrodynamics described in
Paper III (see Section 4, Equation (41)), where choosing
Equation (63) helps ensure intrinsic energy conservation.

8.3. Relationship to the Riemann Solver-based Hydrodynamics
Implementation

When using the Riemann solver-based hydrodynamics
capabilities described in Section 4, MESA does not cast the
stellar structure equations in terms of local heating as in
Equation (54). Instead, it combines Equation (54) with the
constraint of fluid momentum conservation to form a local total
energy equation.
We begin with the mass continuity equation,
\[ \frac{D\rho}{Dt} = -\rho \frac{\partial}{\partial r}(r^2u), \] (68)
and the momentum equation,
\[ \frac{Du}{Dt} = -\frac{1}{\rho} \frac{\partial P}{\partial r} - \frac{\partial \Phi}{\partial r}, \] (69)
written in Lagrangian form and assuming spherical symmetry. The variable \( u \) is the radial velocity and \( \Phi \) is the gravitational potential. The Lagrangian derivative operator is \( \frac{D}{Dt} = \frac{\partial}{\partial t} + u \frac{\partial}{\partial r}. \)

Multiplying Equation (69) by \( u \) gives
\[ \frac{D}{Dt} \left( \frac{1}{2} u^2 \right) = -u \frac{\partial P}{\partial r} - \frac{\partial \Phi}{\partial r}. \] (70)

The gravitational potential does not explicitly depend on time \( (\partial \Phi / \partial t = 0) \), so \( D\Phi / Dt = u \partial \Phi / \partial r. \) This implies
\[ \frac{D}{Dt} \left( \frac{1}{2} u^2 + \Phi \right) = -\frac{\partial P}{\rho} \frac{\partial \rho}{\partial r}. \] (71)

Using Equations (54) and (60), we have
\[ \frac{De}{Dt} - \frac{P}{\rho^2} \frac{D\rho}{Dt} = \epsilon - \frac{\partial L}{\partial m}. \] (72)

Adding Equations (71) and (72) gives
\[ \frac{D}{Dt} \left( \epsilon + \frac{1}{2} u^2 + \Phi \right) = \frac{P}{\rho^2} \frac{D\rho}{Dt} - u \frac{\partial P}{\rho} \frac{\partial \rho}{\partial r} + \epsilon - \frac{\partial L}{\partial m}. \] (73)

Using mass continuity (Equation (68)), this becomes
\[ \frac{D}{Dt} \left( \epsilon + \frac{1}{2} u^2 + \Phi \right) = -\frac{1}{\rho^2} \frac{\partial}{\partial r} \left( P u^2 \right) + \epsilon - \frac{\partial L}{\partial m}. \] (74)

In spherical coordinates,
\[ \frac{1}{\rho^2} \frac{\partial}{\partial r} (r^2f) = \frac{\partial (Af)}{\partial m}. \] (75)

where \( A = 4\pi r^2. \) Thus, we arrive at the equation that MESA solves,
\[ \frac{D}{Dt} \left( \epsilon + \frac{1}{2} u^2 + \Phi \right) = \epsilon - \frac{\partial}{\partial m} (L + P A u). \] (76)

### 8.4. Ionization

The internal energy reported by the EOS should include the energy associated with ionization and molecular dissociation. The assumption of LTE specifies the ionization state given \( (\rho, T, \{X_i\}) \). Since MESA does not regard a change in ionization as a change in composition, it is not necessary to include separate composition derivatives in \( \epsilon_{\text{grav}} \) in order to account for the energetic effects of changes in the ionization state.

To demonstrate a specific scenario where MESA accounts for ionization energy, we evolve a 1 M\(_{\odot}\) pre-MS model composed of pure H. We compare quantities calculated by MESA with other, simpler estimates. We calculate the thermal energy assuming a monatomic ideal gas,
\[ \epsilon_{\text{thermal}} = \frac{3N_h k_B T}{2\mu}. \] (77)

Since this energy is released upon recombination, it is also often referred to as "recombination energy."
We calculate the ionization energy for pure H as

$$e_{\text{ion}} = (1 - f_H)N_e E_{\text{H}} + \frac{N_A E_{\text{H}_2}}{2}, \quad (78)$$

where we assume the ionization fraction of H is given by the Saha equation. The variable $f_H$ represents the neutral fraction of H. The H ionization energy is $E_{\text{H}} = 13.6$ eV, and Equation (78) also includes the dissociation energy of molecular H ($E_{\text{H}_2} = 4.52$ eV) assuming that no H is in the molecular state.

During the evolution, we record the $\epsilon_{\text{grav}}$ calculated by MESA using Equation (65). We also evaluate the quantity

$$\frac{D}{Dt} \left( e_{\text{thermal}} + e_{\text{ion}} + e_{\text{PdV}} \right)$$

that separates out the thermal and ionization energies. In Figure 53, we compare these two approaches, making it clear that all three terms in the above expression play an important role. Their sum agrees with the MESA $\epsilon_{\text{grav}}$, indicating that each of these terms is accounted for in the MESA calculation.

Figure 54 shows the history of the material at the Lagrangian coordinate $(M - m)/M_0 = 10^{-5}$. We plot the $\epsilon$ reported by the MESA EOS along with $e_{\text{thermal}}$ and $e_{\text{ion}}$ (calculated in the same manner as above). At this location, the specific internal energy is dominated by the ionization energy. The lower panel of this figure shows the neutral fraction of H; toward the left of the plot, the H is fully neutral. In this region, the ionization energy plateaus at the dissociation energy of molecular H (see Equation (78)).

For a star in hydrostatic equilibrium, the virial theorem states that

$$-\frac{1}{2} \int_0^M \frac{Gm}{r} dm + \int_0^M \frac{3P}{2\rho} dm = 0. \quad (79)$$

The integrand of the right term, $3P/(2\rho)$, is the specific thermal energy of an ideal monatomic ideal gas. Figure 55 shows the total internal energy and gravitational potential energy reported by MESA for the pure-H pre-MS model. On the same scale, we show half the total potential energy plus the internal energy. This quantity is not zero; rather, by the virial theorem, it should sum to the nonthermal and non-ideal internal energy (e.g., the ionization energy). This value, recorded from the MESA model, agrees well with our estimate of the ionization energy. Also note that at early times, the total energy of the star (internal + potential, not shown) is positive. The phenomenon of positive total energy when ionization energy is included also occurs for envelopes of stars on the asymptotic giant branch (AGB; Paczynski & Ziółkowski 1968). Figure 56 shows the total energy in the envelope of a 1.0 $M_\odot$ MESA model on the AGB. This confirms that the ionization energy is included when MESA reports the total energy of a model.
8.5. Latent Heat

Paper II discusses the inclusion of the latent heat of crystallization for long-term WD cooling. Crystallization is a first-order phase transition that manifests in the PC EOS (Potekhin & Chabrier 2010) as an entropy discontinuity at a plasma coupling parameter of $\Gamma = 175$, and can be captured in stellar evolution with $\epsilon_{\text{grav}}$ in the form of Equation (67). Since the publication of Paper II, controls have been added to MESA to allow smoothing out the injection of latent heat in $\epsilon_{\text{grav}}$ over a user-specified range of $\Gamma$. By default, the range for crystallization is softened to $150 \leq \Gamma \leq 175$ to avoid numerical difficulties with the sudden energy injection associated with a sharp transition at $\Gamma = 175$. The controls allow for this range to be tightened for more precise timing on the occurrence of crystallization if necessary. Figure 57 shows the small impact on cooling time for a $0.6 \, M_\odot$ WD from spreading the latent heat over this broader range of $\Gamma$ relative to a tighter phase transition for $174 \leq \Gamma \leq 176$.

The spreading of the phase transition is accomplished by calculating both the liquid and solid solutions within the PC EOS and linearly blending the entropy $s$ and internal energy $e$ over the specified range of $\Gamma$. With $\epsilon_{\text{grav}}$ expressed in the form of Equation (67), the energy of the phase transition is captured as fluid elements smoothly traverse from the liquid phase to the solid phase. By default, MESA automatically switches to using $\epsilon_{\text{grav}}$ in the form of Equation (67) for $\Gamma > 150$. This choice ensures the capture of latent heat release.

Theoretical and observational works have suggested that crystallization in C/O mixtures may occur at higher $\Gamma$ than the classical one-component plasma value of $\Gamma = 175$ (Horowitz et al. 2007; Winget et al. 2009; Medin & Cumming 2010; Althaus et al. 2012). Our updated crystallization controls allow for the effect on stellar evolution of the crystallization at $\Gamma \approx 240$ to be investigated. Figure 57 shows the potential effects on WD cooling times of varying the $\Gamma$ for crystallization. Because the heating from crystallization is released very late in the WD evolution, its effects on cooling times are on the order of a gigayear, and variations in crystallization treatment can lead to changes that are a significant fraction of this timescale.

The composition terms in Equation (59) that were dropped to form Equation (67) are negligible as long as there is no mixing in the crystallization region. Phase separation may violate this assumption and require a modified treatment, but we do not consider this process here. Detailed phase diagrams for crystallization and the possible associated phase-separation effects are not currently supported in MESA, so our investigation here is limited to the effects of crystallization as a function of a fixed $\Gamma$ range.

8.6. EOS Blending

As shown in Figure 1 of Paper I, MESA employs a patchwork of several EOSs to provide coverage of a maximal amount of $\rho - T$ space. When blending from one EOS region into another, care is required to avoid introducing spurious contributions to $\epsilon_{\text{grav}}$. At high density, MESA blends from the Helmholtz EOS (HELM; Timmes & Swesty 2000) for $\Gamma < 10$ to the PC EOS (Potekhin & Chabrier 2010) for $\Gamma > 20$ by default. This default has been changed from the original default of $40 \leq \Gamma \leq 80$ given in Paper I due to the optimal agreement between relevant quantities shown in Figure 59, as explained below. Overall, the two EOSs agree well on thermodynamic quantities in the blending region ($\sim 1\%$ for $e$ and $s$, but Figure 58 shows that the absolute magnitude of the disagreement can still be large enough to influence the $\epsilon_{\text{grav}}$ for a cooling WD when $\epsilon_{\text{grav}}$ is expressed in the form of Equations (59)–(65).

The left panel of Figure 58 indicates that, typically, the internal energy difference is $\Delta \epsilon \sim 10^{13} \text{erg g}^{-1}$, while $c_p T \sim 10^{14} - 10^{15} \text{erg g}^{-1}$ in the region of the blend. As a WD model cools, most of its $\sim 10^{33} \text{g}$ of mass must eventually pass through this transition. If the energy equation is being solved in the form of Equation (61), $\sim 10^{48} \text{erg}$ of spurious energy would be introduced into the model by EOS blending. Since much of this blending happens after the WD model has cooled to a luminosity of $L \lesssim 0.1 \, L_\odot$, this extra energy corresponds to $\Delta T \gtrsim 100 \, \text{Myr}$ of extra WD cooling time.

The default form of $\epsilon_{\text{grav}}$ given in Equation (65) does not suffer from this spurious heating, since it is expressed in terms of thermodynamic derivatives from the EOS rather than $e$ and $s$. For this form of $\epsilon_{\text{grav}}$, the differences between $e$ or $s$ do not directly enter the equations. Instead, changes in $e$ with $D \ln \rho /Dt$ and $D \ln T /Dt$ are tracked with quantities such as $c_p$ and $\chi_T$, and Figure 59 shows that these agree well for the EOS blend region. Since the implementation of Equation (65) does not involve any derivatives constructed as finite differences, the fact that quantities such as $c_p$ agree to within a few percent guarantees that $\epsilon_{\text{grav}}$ will be consistent across the blend, with no significant spurious energy injected due to blending. Crucially, the release of latent heat described in Section 8.5 requires switching to $\epsilon_{\text{grav}}$ in the form of Equation (67) only for zones with $\Gamma > 150$, so both EOS blending and crystallization simultaneously receive appropriate treatments with different forms of $\epsilon_{\text{grav}}$ in different stellar regions.

8.7. Gravitational Settling

Equation (65) for $\epsilon_{\text{grav}}$ ignores changes in the internal energy $e$ due to composition changes. García-Berro et al. (2008) point
out that a self-consistent evolutionary approach to WD cooling including the effects of $^{22}$Ne settling requires an $\epsilon_{\text{grav}}$ that accounts for the composition changes due to element diffusion. They adopt pure $^{12}$C or $^{16}$O core compositions with trace $^{22}$Ne and no other isotopes. Although this approach is useful for a rigorous study of self-consistent WD evolution with diffusion fully coupled to evolution, it is not well-suited for a general treatment of realistic mixed core compositions.

MESA splits element diffusion into a separate step before the main structural solve, and hence diffuse effects are not included in $\epsilon_{\text{grav}}$. We ensure that the energy associated with $^{22}$Ne settling is not included in $\epsilon_{\text{grav}}$ by using Equation (65), and we compensate by including an extra heating term $\epsilon_{22}$ in Equation (54). This term is calculated using velocities saved from the element diffusion step as described in Section 3.5. Our results for the effects of $^{22}$Ne settling on WD cooling agree well with those of García-Berro et al. (2008) and Deloye & Bildsten (2002), who adopt a heating term similar to our approach.

9. Summary

We explain the significant new capabilities and improvements implemented in MESA since the publication of Paper I, Paper II, and Paper III. Progress in the treatment of convective boundaries (Section 2) and element diffusion (Section 3 and Appendix C) will improve studies of their impact on stellar evolution. Advances to MESA in implicit hydrodynamics (Section 4), approximation of 3D RTI effects (Section 5), and coupling with a public version of the STELLA radiative transfer instrument will enhance the modeling of Type IIP SN light curves from post-explosion to post-plateau (Section 6). We integrate these improvements with an exploration of PPISN and black hole formation models (Section 7). We describe energy conservation in MESA and demonstrate improvements relevant to WD cooling (Section 8). Upgrades to estimating the absolute magnitude of a model in a chosen passband (Appendix A), guidance on importing multidimensional models into MESA (Appendix B), and new MESA-based software tools (Section D) will strengthen research and education. Input files and related materials for all of the figures are available at http://mesastar.org.

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**Appendix A**

**Colors**

We describe MESA’s implementation of bolometric corrections (BCs) for use in estimating the absolute magnitude of a model in a user-chosen filter system. Note this is different from the colors reported by STELLA (Section 6), as the colors module uses pre-computed tables of BCs while STELLA solves the radiative transfer equations on the fly (Blinnikov et al. 1998).

The absolute bolometric magnitude ($M_{\text{bol}}$) of a star is defined, with reference to the solar absolute bolometric magnitude, as (Torres 2010)

$$M_{\text{bol}} = M_{\text{bol,\odot}} - 2.5 \log_10\left(\frac{L}{L_{\odot}}\right),$$  

where $M_{\text{bol,\odot}}$ is the absolute bolometric magnitude of the Sun, taken as 4.74 (2015 IAU Resolution B2). This can be transformed into the passband-dependent absolute magnitude, $M_X$, for a nominal pass band $X$ via

$$M_X = M_{\text{bol}} - BC_X,$$

where $BC_X$ is the BC for passband $X$ and accounts for the flux emitted outside of the wavelength range of the filter system.

The derivation of a BC requires an atmospheric model of a star such that a stellar spectrum can be computed over all wavelengths, a computationally costly process. To prevent the requirement of actually having to generate a spectra at each time step, we make use of pre-computed BC tables. These define the BC as a function of the stellar photosphere; $T_{\text{eff}}/K$, log ($g$/cm s$^{-2}$), and the metallicity [M/H] are derived from pre-computed grids of stellar atmosphere models (see, e.g., Kurucz 1970; Husser et al. 2013). Given the parameters at the stellar photosphere, we interpolate each set of BCs over log ($T_{\text{eff}}/K$), log ($g$/cm s$^{-2}$), and [M/H] using linear interpolation over nearest neighbors and without extrapolation for points outside of the table range.

We provide two sets of pre-processed tables of BCs, though a user may provide their own. From Lejeune et al. (1998), we provide the Johnson–Cousins–Class bands $UBVR_{\text{I}}$JHKLL/M. This table provides the BCs over the parameter range $2000 \leq T_{\text{eff}}/K \leq 50,000$, $-1.02 \leq \log (g$/cm s$^{-2}) \leq 5.0$, and $-5.0 \leq [\text{M/H}] \leq 1.0$, with a variable sampling rate. Figure 60 shows the time evolution of the absolute magnitude of a 1 M$_{\odot}$ star with the passbands defined in Lejeune et al. (1998). We also provide a set of blackbody BCs for the passbands $UBVR_{\text{I}}$, over the range $100 \leq T_{\text{eff}}/K \leq 50,000$, in steps of 100 K. As these are blackbody corrections, there is no $g$ or [M/H] dependence.

There are many other possibilities for other passbands or classes of objects (Fukugita et al. 1996; Girardi et al. 2002; Bessell 2011; Bessell & Murphy 2012). Thus, the tables we provide are not a definitive set, but merely a reasonable starting point for modeling stellar objects. Other astrophysical objects like WDs, exoplanets, or SN light curves require calculating specialized tables. Users may provide BC tables defined in terms of $T_{\text{eff}}/K$, log ($g$/cm s$^{-2}$), and [M/H].
from which code is open source and freely available at jalombar.github.io.

The hatched region in the right panel shows the previous default blending range for these quantities.

Figure 59. Percent difference between HELM and PC EOS for \(c_p\) (left) and \(\chi_T\) (right). Dashed lines show the EOS blending boundaries for \(10 \leq \Gamma \leq 20\), and the hatched region in the right panel shows the previous default blending range \(40 \leq \Gamma \leq 80\). In the new blending region, the EOSs disagree by only a few percent for these quantities.

Figure 60. Evolution of the absolute magnitude of a \(1\,M_{\odot}\) star for the bolometric magnitude and magnitude in the filter bands \(UBVR,I,JHKLL\).

### Appendix B

**Model Relaxation**

To simplify the process of importing a model into MESA, we have developed simple relaxation routines that allow the construction of a starting model in hydrostatic equilibrium with specified profiles for composition, angular momentum, and entropy. Examples that motivate importing a model into MESA include multidimensional simulations of stellar mergers, common envelope evolution, and the effects of SN explosions on nearby companions.

The relaxation process inputs include 1D profiles of the composition and angular momentum. The process also requires either an entropy profile or the profiles of pairs of values \((\rho, T)\), \((P_{\text{gas}}, T)\), or \((\rho, e)\), from which MESA extracts the entropy using the \texttt{eos} module. Note that in the case where the entropy is not provided directly, the relaxed model will match the entropy computed by the \texttt{eos} module, but not necessarily the input \((\rho, T)\), \((P_{\text{gas}}, T)\), or \((\rho, e)\) profiles. A good match for the input profiles depends on the input data corresponding to a model in hydrostatic equilibrium computed with an EOS that is consistent with MESA’s.

Relaxation is done via pseudo-evolution of a stellar model for which mixing, angular momentum transport, and changes in composition from nuclear burning are suppressed, while a quantity of interest is incrementally altered until it reaches the desired value up to a pre-defined tolerance. Throughout this relaxation process, hydrostatic equilibrium is enforced. The starting stellar model can be any MESA model with the required mass, and for most cases a ZAMS star at \(Z = 0.02\) works well. The first two steps in the relaxation of a model fix the composition and angular momentum profiles. This is done by directly adjusting the variables for the composition and angular momentum of each cell until the desired values are reached. Since the entropy is a derived quantity in MESA, the third step relaxes the entropy indirectly via the energy equation. This is achieved by adding a heating term that injects energy in regions where the entropy is below the target value and removes energy in regions where the entropy is above the target value. This specific heating rate is

\[
\epsilon_{\text{relax}}(m) = \left(1 - \frac{s(m)}{s_{\text{target}}(m)}\right) \frac{e(m)}{\tau},
\]

where \(e(m)\), \(s(m)\), and \(s_{\text{target}}(m)\) are the specific internal energy, current entropy, and target entropy, respectively, at the mass coordinate \(m\). The timescale for the relaxation process is specified by \(\tau\). The value \(\tau\) should be chosen to be small enough that energy transport is negligible during the pseudo-evolution. In practice, \(\tau\) can be chosen to be orders of magnitude smaller than the dynamical timescale of the system.

We verified that using the entropy, composition, and angular momentum profiles of a model computed with MESA as input, the relaxation procedure can reproduce the original model to within 0.1%. An example is provided in the test suite under the name relax\_composition\_j\_entropy.

We tested these relaxation routines using the outcome of a stellar merger computed with the STARSMA\textsc{sh}\textsuperscript{23} SPH code (Gaburov et al. 2010; Lombardi et al. 2011), configured to use

\[\text{The STARSMA\textsc{sh} code is open source and freely available at https://jalombar.github.io/starsmash/}.\]

\[\text{\textsuperscript{23} The STARSMA\textsc{sh} code is open source and freely available at https://jalombar.github.io/starsmash/}.\]
the MESA EOS. Two co-eval non-rotating MESA models with ZAMS masses of 20 $M_\odot$ and 15 $M_\odot$ are evolved until the 20 $M_\odot$ star reaches $X_e = 0.34$. These models are then imported into STARSMasher to simulate a head-on collision, such that the relative velocity of the two stars at infinity is zero. We find that 2.18 $M_\odot$ of material is lost from the system due to the collision.

We compute spherical mass-weighted averages of the composition, $\rho$, and $e$. These profiles are input into the MESA relaxation process, along with a zero angular momentum profile since the model is a head-on collision of non-rotating stars. Figure 61 shows that the relaxed model closely follows the input smoothed particle hydrodynamic (SPH) merger model in the central regions, though densities are $\approx 10\%$ larger throughout the inner 25 $M_\odot$. Density differences of more than an order of magnitude are present in the outer layers. This is a consequence of these layers not being in hydrostatic equilibrium in the input SPH simulation. The MESA relaxation process matches the entropy rather than the density profile of the model assuming hydrostatic equilibrium as discussed above. The relaxed model corresponds to the final configuration if it contracts adiabatically, which is a good approximation as velocities in the SPH model are well below the local sound speed (Pan et al. 2013).

### Appendix C

**Element Diffusion Implementation Details**

This appendix provides the implementation details not contained in Section 3. Equations (2)–(4) and (9) give the full set of diffusion equations that must be solved to obtain diffusion velocities. For $S$ total species in the plasma (including electrons), Equation (9) provides $S - 1$ equations (one for each ion species), Equation (2) provides $S$ equations (one for each species including electrons), and Equations (3) and (4) each provide one additional equation, for a total of $2S + 1$ independent equations. The $2S + 1$ unknowns are $S$ diffusion velocities $w_{\delta}$, $S$ heat flow vectors $r_{\delta}$, and the electric field $E$.

The inputs provided by the MESA model are the number densities $n_\delta$, temperature $T$, gradients of each of these quantities $d \ln n_\delta/dr$ and $d \ln T/dr$, species mass in atomic units $A_\delta$, species mean charge as an average ionization state $\bar{Z}_\delta$, and resistance coefficients $K_\delta$, $\bar{Z}_\delta$, and $\bar{Z}_{\delta\delta}^\prime$ (defined in Equation (86) of Paper III). The coefficients are calculated as described in Section 3.3. Together with the mean ionization states, these are the key pieces of input physics that determine the diffusion of all ions. Extra acceleration terms $g_{\text{rad}}$ for radiative levitation are either set to zero by default or calculated as in Hu et al. (2011) when the option to include radiative levitation is enabled.

In the spirit of Thoul et al. (1994), Equations (2)–(4) and (9) are grouped into a single matrix equation:

$$
\frac{\partial \bar{E}}{\partial t} + \mathbf{u} \cdot \nabla \bar{E} = \sum_{j} \Delta_{ij} W_j.
$$

The vectors capturing the driving terms are

$$
\alpha_i = \begin{cases} 
n_i A_i & i = 1, \ldots, S - 1, \\
0 & i = S, \ldots, 2S + 1,
\end{cases}
$$

$$
\nu_i = \begin{cases} 
n_i & i = 1, \ldots, S - 1, \\
S & i = S, \ldots, 2S - 1, \\
0 & i = 2S, 2S + 1,
\end{cases}
$$

$$
\gamma_j = \begin{cases} 
n_j \delta_j & i = 1, \ldots, S - 1, \\
0 & i = S, \ldots, 2S + 1,
\end{cases}
$$

$$
\beta_{\text{rad},i} = \begin{cases} 
-n_i A_i m_p g_{\text{rad},i} & i = 1, \ldots, S - 1, \\
0 & i = S, \ldots, 2S + 1.
\end{cases}
$$

The vector containing the unknowns is

$$
W_j = \begin{cases} 
w_j & j = 1, \ldots, S, \\
r_j & j = S + 1, \ldots, 2S, \\
q_{\delta E} & j = 2S + 1.
\end{cases}
$$

For $i = 1, \ldots, S - 1$, the right-hand side matrix of Equation (83) is

$$
\Delta_{ij} = \begin{cases} 
-\sum_{l=1}^{S} K_{il} & j = i, \\
K_{ij} & j = 1, \ldots, S \quad \text{and} \quad j \neq i, \\
\sum_{l=x}^{S} K_{il} A_l / (A_l + A_i) & j = i + S, \\
-K_{ij} \bar{Z}_{ij} \bar{Z}_{il} / (A_l + A_i) & j = S + 1, \ldots, 2S \quad \text{and} \\
K_{ij} & j = 2S + 1.
\end{cases}
$$

For $i = S, \ldots, 2S - 1$, the matrix terms are

For $i = 1, \ldots, S - 1$, the matrix terms are
For $i = 2S$,
\[
\Delta_y = \begin{cases} 
  n_i a_j & j = 1, \ldots, S, \\
  0 & j = S + 1, \ldots, 2S + 1. 
\end{cases} 
\] (91)

For $i = 2S + 1$,
\[
\Delta_y = \begin{cases} 
  n_i z_j & j = 1, \ldots, S, \\
  0 & j = S + 1, \ldots, 2S + 1. 
\end{cases} 
\] (92)

Indices $i = 1 \ldots S - 1$ capture the $S - 1$ Equations (9) for the ions. Indices $i = S \ldots 2S - 1$ capture the $S$ Equations (2). Indices $i = 2S, 2S + 1$ capture the two constraints in Equations (3) and (4).

For a generic driving term that takes the form of an extra force $f_i$ on ions of species $s$, a term $-n_i f_i$ appears on the left-hand side of Equation (9). This can be accounted for in the matrix setup by adding another vector $\beta_{i,j}$ to the left-hand side of Equation (83) with the form
\[
\beta_{i,j} = \begin{cases} 
  -n_i f_i & i = 1, \ldots, S - 1, \\
  0 & S, \ldots, 2S + 1. 
\end{cases} 
\] (93)

One such extra driving force that may be explored with MESA in the future is Coulomb separation in dense matter arising from non-ideal corrections for the ions (Chang et al. 2010; Beznogov & Yakovlev 2013; Diaw & Murillo 2016).

The diffusion velocities are separated into two terms capturing the distinct effects of gravitational settling and ordinary diffusion in the tradition of Equation (11) of Iben & MacDonald (1985):
\[
w_i = w_i^\alpha - \sum_j \sigma_j \frac{d \ln C_j}{dr}, 
\] (94)

where $C_j \equiv n_j / n_e$ following the notation of Thoul et al. (1994). These separate terms are constructed by inverting the matrix $\Delta_y$ and then solving Equation (83) for just one of $\alpha$, $\beta$, $\nu$, and $\gamma_{i,j}$ at a time on the left-hand side. These results can then be linearly combined to construct $w_i^\beta$ and $\sigma_j$ such that the total sum in Equation (94) gives a solution that satisfies the complete set represented by Equation (83).

When electrons become degenerate, we drop all $S$ Equations (2) and set the $S$ heat flow vectors to $r_s = 0$. Equations (83) then represents a system of just $S + 1$ equations and the vectors and matrices simplify considerably, dropping all entries for indices $i = S \ldots 2S - 1$ or $j = S + 1 \ldots 2S$ in the definitions given in Equations (84)–(92). To avoid discontinuities, we employ a blend that smoothly transitions between the diffusion velocity solutions over a range in $\eta \equiv \mu_e / k T$, where $\mu_e$ is the electron chemical potential. By default, the blend is centered around $\eta \approx 1$, with user controls available to adjust the range of this blending region.

\section*{Appendix D}
\textbf{Software Infrastructure}

Software is an integral enabler of observation, theory, and computation and a primary modality for realizing the discoveries and innovations expressed, for example, in the astronomy and astrophysics decadal surveys (e.g., National Research Council 1991, 2001, 2011). In this appendix, we describe new software stacks at a variety of scales that enhance the research and education infrastructure.

\subsection*{D.1. Not A Number}

Not a Number (NaN) is a numeric data type representing an undefined or unrepresentable value (e.g., Goldberg 1991; Hauser 1996). Examples include 0/0 and $\sqrt{-1}$ in real arithmetic. In the IEEE 754 floating-point standard (IEEE 2008), there are two types of NaNs: quiet (qNaN) and signaling (sNaN). A qNaN propagates errors resulting from invalid operations or values without triggering a floating-point exception. An sNaN precipitates an invalid operation exception whenever an attempt is made to use one as an arithmetic operand. The IEEE 754 standard requires qNaN as the default, while an sNaN can be used to support features such as filling uninitialized memory or other extensions to floating-point arithmetic.

NaN and infinity (INF) setting and testing routines are provided within the \texttt{utili.nan.f90} file. A consistent set of interfaces allows for initializing scalars/arrays to NaN values, and testing for qNaN, sNaN, or INF values. Interface overloading allows single, double, or quad precision scalars or arrays of rank between 1 and 4 to be handled. This module provides four generic interfaces. The logical function \texttt{is\_nan(x, signal)} returns true if \texttt{x} contains NaNs and false otherwise. The optional logical argument \texttt{signal} determines whether qNaN, sNaN, or both are tested for. The
logical function \texttt{is\_inf(}x\texttt{)} returns true if \( x \) contains INFs and false otherwise. The logical function \texttt{is\_bad(}x\texttt{)} returns true if \( x \) contains NaN or INF values and false otherwise. The routine \texttt{set\_nan(x, signal)} sets a scalar or array \( x \) to NaN values. The optional logical argument \texttt{signal} determines whether a qNaN or sNaN is set.

The library framework of MESA is designed to be interoperable within other software ecosystems. For example, these NaN and INF interfaces are of potential interest to users of MESA or developers of similar software instruments.

### D.2. MESA – Web

Stellar evolution software instruments can be complicated to install and use, especially when the aim is primarily pedagogical (e.g., high-school or undergraduate courses). Motivated by the community’s expressed need for a lower barrier to entry for education, a Web-based interface to MESA was developed, MESA – Web, at \url{http://mesa-web.asu.edu}. MESA – Web currently allows choices for the initial mass, metallicity, rotation, mass loss, nuclear reaction network, custom nuclear reactions rates, spatial and temporal resolution, and model output rate.

MESA – Web sends the user an email message that contains a URL of a zip file to download when their job has completed. The unzipped output directory contains a MESA history data file holding the time evolution of 57 quantities as well as a series of MESA profile data files containing information on 56 quantities in each zone of the stellar model at discrete model numbers. Also included in the output is an MP4 formatted video containing a plot dashboard of the abundance profiles, Kippenhahn diagram, Hertzsprung–Russell diagram, rotational profile, and temperature, density, and pressure profiles.

MESA – Web is currently hosted on a four-core server at Arizona State University and allows jobs to run on a single core for 4 hr of walltime or until the model reaches iron core collapse. Launched in June 2015, MESA – Web has currently served more than 3000 models to over 600 different users at over 40 academic institutions. Efforts to expand MESA – Web’s capabilities include porting the service to a host with enhanced compute resources, simulating core-collapse SN explosions (see Paper III) and light curves (see Section 6), and binary star evolution (see Paper III).

### D.3. MESA-Docker

Docker is a software technology designed to deploy and run applications by using “containers.” Containers provide much of the virtualization power of traditional virtual machines while requiring far less resource overhead. This allows the efficient packaging of an entire operating environment, with all of the necessary libraries and other dependencies for a large software tool such as MESA. The MESA-Docker package (Bauer & Farmer 2017) provides a solution that simplifies the requirements for locally running a full MESA installation with all capabilities available, with only minor overhead associated with running in a container. MESA-Docker will be useful for new users, students with educational projects, and Windows operating system users.

### D.4. pyMESA

pyMESA (Farmer 2017) allows MESA modules to be embedded into Python projects. pyMESA currently supports using the equation of state (\texttt{eos}), nuclear reaction (\texttt{rates}), neutrino (\texttt{neu}), atmosphere (\texttt{atm}), and opacity (\texttt{kap}) packages. This software infrastructure will be useful for users who want to use parts of MESA in their own Python software projects. As an example of these capabilities, Figures 58 and 59 were produced using the \texttt{pyMESA\_eos} interface to make direct calls to the MESA EOS routines.

#### D.5. MESAstar Model Optimization

The MESAstar test suite contains a sample case that shows how to use the simplex optimization algorithm (Nelder & Mead 1965) to find stellar models that minimize a specified \( \chi^2 \) by automatically adjusting a variety of control parameters.\(^{24}\) The \( \chi^2 \) to be minimized can contain both pre-supplied and user-defined terms. Pre-supplied terms include \( T_{\text{eff}}, L, R, g, \) surface \( Z, \) surface \( Y, \) and age. An easy-to-use framework allows the user to define other terms to include in the \( \chi^2 \). Control parameters include \( M, Z/X, Y, \alpha_{\text{MLT}}, \) and \( f_{\text{ox}} \). Other stellar evolution parameters can be easily added from the extensive set of controls in MESA. We provide a MESA test suite case using this new capability to calibrate a solar model. This can serve as a template for users wishing to use this method to search for models that match the observed properties of specific stars.

#### D.6. \url{http://mesa-star.org}

Reproducibility is the bedrock of scientific research. Provenance, as the term relates to software instruments (Van den Bussche & Viana 2001; Carata et al. 2014), is the ability to record the full history of a result. Scientific research is generally held to be of good provenance when it is documented in sufficient detail to allow reproducibility. The MESA project facilitates provenance by the research community in four ways. One, by curating public releases of the source code, makefiles, test suite, and how the source code was compiled—GNU compilers are redistributed in the MESA Software Development Kit (see Paper II)—at \url{http://mesa.sourceforge.net}. Two, by providing bit-for-bit consistency for all results across all the supported platforms (see Paper III). Three, by supporting a user mailing list to openly share knowledge (see the Manifesto in Paper I). Currently, over 12,000 messages are archived and searchable. Four, by hosting a Web portal at \url{http://mesa-star.org} to share MESA-oriented software contributions and reposh the MESA files (\texttt{inlist, run\_star\_extraf, etc.}) that specify all the ingredients needed to reproduce a scientific result. Currently, \url{http://mesa-star.org} offers over 120 MESA-oriented software contributions and inlist repositories.

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\(^{24}\) This is the same simplex algorithm that is used for finding matches in asteroseismology applications using MESA (see Paper III, Section 3). The code reported here is a simplified subset of that tool and is now easier to use and adapt to new problems.
Wolf, B., & Schwab, J. 2017, wmwolf/py_mesa_reader: Interact with MESA Output, zenodo, doi:10.5281/zenodo.526958
Wongwathanarat, A., Müller, E., & Janka, H.-T. 2015, A&A, 577, A48
Wood, P. R. 1974, ApJ, 190, 609
Woosley, S. E. 2017, ApJ, 836, 244
Woosley, S. E., Blinnikov, S., & Heger, A. 2007a, Natur, 450, 390
Woosley, S. E., Heger, A., & Weaver, T. A. 2002, RvMP, 74, 1015
Woosley, S. E., Kasen, D., Blinnikov, S., & Sorokina, E. 2007b, ApJ, 662, 487
Yoon, S.-C., Dessart, L., & Ciocchiatti, A. 2017, ApJ, 840, 10
Zel’dovich, Y. B., & Raizer, Y. P. 1967, Physics of Shock Waves and High-
temperature Hydrodynamic Phenomena (New York: Academic)
Zhang, W., Woosley, S. E., & Heger, A. 2008, ApJ, 679, 639