Laminar Flame Speeds in Degenerate Oxygen–Neon Mixtures

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Abstract

The collapse of degenerate oxygen–neon cores (i.e., electron-capture supernovae or accretion-induced collapse) proceeds through a phase in which a deflagration wave (“flame”) forms at or near the center and propagates through the star. In models, the assumed speed of this flame influences whether this process leads to an explosion or to the formation of a neutron star. We calculate the laminar flame speeds in degenerate oxygen–neon mixtures with compositions motivated by detailed stellar evolution models. These mixtures include trace amounts of carbon and have a lower electron fraction than those considered in previous work. We find that trace carbon has little effect on the flame speeds, but that material with electron fraction $Y_e \approx 0.48 - 0.49$ has laminar flame speeds that are $\approx 2$ times faster than those at $Y_e = 0.5$. We provide tabulated flame speeds and a corresponding fitting function so that the impact of this difference can be assessed via full star hydrodynamical simulations of the collapse process.

Unified Astronomy Thesaurus concepts: White dwarf stars (1799); Degenerate matter (367); Oxygen burning (1193)

Supporting material: machine-readable table

1. Introduction

Degenerate oxygen–neon (ONe) cores with masses near the Chandrasekhar mass can form in the evolution of $\approx 8-10 M_\odot$ single stars (e.g., Miyaji et al. 1980; Miyaji & Nomoto 1987), in interacting binary systems with varying degrees of envelope stripping (e.g., Tauris et al. 2015; Poelarends et al. 2017), in binary systems with an accreting ONe white dwarf (WD; e.g., Canal & Schatzman 1976; Nomoto & Kondo 1991), or as the result of the merger of two WDs (e.g., Saio & Nomoto 1985; Brooks et al. 2017). When the core reaches a central density $\approx 10^{10} g \text{ cm}^{-3}$, exothermic electron captures on $^{20}$Ne occur and lead to the initiation of a deflagration wave (“flame”) that propagates outward. The competition between the energy release from this flame and the electron-capture reactions on its ashes determines whether this leads to an explosion (resulting in partial or total disruption of the star) or implosion (resulting in the formation of a neutron star; NS).

This situation has long been known to be finely balanced (e.g., Nomoto & Kondo 1991; Canal et al. 1992), though the general conclusion by the end of the 1990s was in favor of collapse to a NS. Recent multidimensional simulations have reiterated that the outcome is sensitive to modeling choices and opened the possibility that at least some cases may lead to a thermonuclear explosion (possibly also leaving a low-mass bound remnant) instead of collapse to an NS (Jones et al. 2016, 2019; Leung et al. 2020). One of the key ingredients in this modeling is the speed at which the flame propagates.

Timmes & Woosley (1992), hereafter TW92, calculated the physical properties of conductively propagated laminar burning fronts in high-density, degenerate carbon–oxygen (CO) and ONe mixtures. We repeat a similar set of calculations using Modules for Experiments in Stellar Astrophysics (MESA; Paxton et al. 2011, 2013, 2015, 2018, 2019), but extend these results to a wider range of compositions motivated by expectations from detailed models of the internal composition of ONe WDs (e.g., Garcia-Berro et al. 1997; Iben et al. 1997; Siess 2006). Section 2 describes the methods we use to calculate the laminar flame speeds. Section 3 reproduces the TW92 results in both CO and ONe mixtures. We then focus on the laminar flame speeds in ONe mixtures under different conditions. In Section 4 we show how the flame speeds are only mildly affected by the presence of small amounts of $^{12}$C, but in Section 5 demonstrate the significant influence of the electron fraction ($Y_e$) of the material. Section 6 provides tabulated flame speeds and a corresponding fitting function. Section 7 briefly describes the implications for models of the collapse of ONe cores.

2. Methods

We use MESA revision r12115 (Paxton 2019). The input files necessary to reproduce our work are publicly available and an illustration of this capability has been included in the test suite case conductive_flame.

We create an initial, spatially uniform MESA model with a temperature $T_0 = 3 \times 10^8$ K, specified density $\rho_0 = \rho(10^9 \text{ g cm}^{-3})$, and a specified unburned composition. These properties characterize the cold material into which the flame will propagate. So long as the upstream temperature is much less than the downstream (post-burn) temperature of $\approx 10^{10}$ K, the temperature jump across the flame is approximately the same and the initial temperature does not play an important role. The total mass $M$ defines the size of the (Lagrangian) computational domain. Because the flame width $\lambda$ varies substantially with density, our domain size must as well. We always ensure that $M/(\rho \lambda^3) \gg 1$, but typically choose this ratio to be $\sim 100$ to limit...
we demonstrate that our results are insensitive to the details of leads to an estimate of the at which nuclear reactions heat the material. This argument the diffusion timescale across it is comparable to the timescale where the computational cost. This also implies that $\lambda \ll r$, so the flame is effectively planar. In practice, $M \sim 10^{-3} \text{g}$. The small size of the domain implies that the pressure gradient due to gravity is negligible. The inner boundary is at $r = 0$. The outer boundary has a fixed temperature $T_i$ and a fixed pressure equal to the initial pressure of the material.

We then insert a hot spot at the center with fractional size in mass $q_h = 3 \times 10^{-4}$ at a temperature $T = 8 \times 10^8 \text{K}$ (for CO mixtures) or $T = 10 \times 10^8 \text{K}$ (for ONe mixtures). This hot spot should have a size of order the flame width and a temperature of order the post-burn temperature to ensure a steady-state flame. If the hot spot is too small the flame will die. If the hot spot is too large the flame will exhibit a decelerating transient. With a well chosen spot size much smaller than the domain, once the flame has propagated a few flame lengths, the initial condition will be effectively erased.

We define the location of the flame to be the location of the maximal rate of nuclear energy release in the domain (i.e., peak nuclear heating, see Figure 1). We evolve the model until the flame has propagated through 90% of the domain and then extract the steady-state flame properties. By repeating this process for different initial conditions, we calculate the laminar flame speed as a function of $\rho_0$, pressure, and composition. In Appendix, we demonstrate that our results are insensitive to the details of the initial conditions and are numerically converged.

2.1. Microphysics

As discussed in TW92, the flame will have a width such that the diffusion timescale across it is comparable to the timescale at which nuclear reactions heat the material. This argument leads to an estimate of the flame speed,

$$v_{\text{flame}} \approx \left( \frac{D_{\text{th}} \epsilon_{\text{nuc}}}{\epsilon} \right)^{1/2},$$

(1)

where $D_{\text{th}}$ is the thermal diffusion coefficient, $\epsilon_{\text{nuc}}$ is a characteristic specific rate of energy generation from nuclear reactions, and $\epsilon$ is a characteristic specific energy. Therefore, the speed of the flame is set by the energy generation rate as determined from the nuclear network and the assumed thermal transport properties of the degenerate plasma.

2.1.1. Transport Properties

In MESA the thermal conductivity of the degenerate electrons, accounted for via a conductive opacity ($\kappa_{\text{cond}} \propto D_{\text{th}}^{-1}$), comes from tables privately communicated by A.Y. Potekhin (see Section A.3 in Paxton et al. 2013). TW92 describe in detail their implementation of the transport properties; a substantially similar approach is adopted in Timmes (2000), hereafter T00. The source code for the transport properties assumed in these works is publicly available.

Figure 2 compares the conductive opacities over the range of temperatures and densities considered in this paper. The left panel shows the T00 and MESA values of $\kappa_{\text{cond}}$ and the right panel their relative difference. Both sources show similar density scalings, but the T00 values scale less steeply with temperature such that while the values agree at $10^9 \text{K}$, $\kappa_{\text{cond}}$ is higher by a factor $\approx 2$ at $10^8 \text{K}$ and lower by a factor $\approx 1.5$ at $10^{10} \text{K}$. Since these variations are not systematically in the same direction, their effect is difficult to estimate, but given the scaling in Equation (1), variations in $\kappa_{\text{cond}}$ at this level correspond to $\approx 30\%$ variations in the flame speed.

2.1.2. Nuclear Reaction Rates

The currently applicable default inputs for nuclear reaction rates are described in Appendix A.2 of Paxton et al. (2019). Rates are taken from a combination of Nuclear Astrophysics Compilation of REAction rates (NACRE; Angulo et al. 1999) and the Joint Institute for Nuclear Astrophysics REACLIB library (default version, dated 2017 October 20; Cyburt et al. 2010). The MESA screening corrections are from Chugunov et al. (2007), which includes a physical parameterization for the intermediate screening regime and reduces to the familiar weak (Dewitt et al. 1973; Graboske et al. 1973) and strong (Alastuey & Jancovici 1978; Itoh et al. 1979) limits at small and large values of the plasma coupling parameter.

Relatively large nuclear networks are required to fully capture the energy generation rate in neutron-rich compositions at these thermodynamic conditions. TW92 illustrate the increase in flame speed with increasing network size (their Table 5) and adopt a 130 isotope network. We perform a similar exercise, using three built-in MESA networks (approx21, mesa_204, mesa_495), a network constructed with the same elements as the TW92 130 isotope network (see their Table 1), and also an adaptive network that automatically adds and removes isotopes and which settles in at around 320 isotopes. Figure 3 shows the flame location as a function of time for a set of runs for a fiducial CO mixture (panel (a)) and a fiducial ONe mixture (panel (b)). Networks of more than 200 isotopes appear to be required before network size no longer makes an appreciable difference in the flame speed. This result is consistent with Chamulak et al. (2007), hereafter C07, who found that for flames in CO mixtures a 430 isotope network gave speeds up to $\approx 25\%$ greater than a 130 isotope network. We run with 495 isotopes unless otherwise stated.

The JINA REACLIB polynomial fits to the reaction rate data end at $10^{10} \text{K}$ as do the tabulated partition functions used to

6 http://cococubed.asu.edu/code_pages/kap.shtml
calculate the reverse rates and ensure detailed balance. Above \(10^{10}\) K, \textsc{Mesa} fixes the rates to be their \(T = 10^{10}\) K values. In some cases, especially for the ONe flames, the temperature exceeds \(10^{10}\) K and the peak in \(\epsilon_{\text{nuc}}\) occurs near the temperature threshold. If the \textsc{Mesa} treatment underestimates the true peak of \(\epsilon_{\text{nuc}}\), then this can lead to an underestimate of the flame speed.

For example, if we truncate the rates at \(T = 8 \times 10^9\) K, the ONe flame in Figure 3 has a speed of \(82 \pm 5\) km s\(^{-1}\), a \(\approx 10\%\) reduction. However, we have physical reasons to expect that this effect is not particularly large. By \(\approx 1.2 \times 10^{10}\) K photodisintegration is strong enough to decompose nuclei into neutrons, protons, and alpha particles. This is an endothermic process, meaning there is an upper limit to how much more positive \(\epsilon_{\text{nuc}}\) can be achieved beyond the place where \textsc{Mesa} truncates the rates.

Our results depend slightly on our adopted rate sources. If we use pure JINA REACLIB defaults (eliminating NACRE), the flame speeds increase. For the calculation shown in Figure 3, the result with the 495 isotope net and only the JINA rates is \(207 \pm 5\) km s\(^{-1}\) for the CO case and \(94.1 \pm 0.1\) km s\(^{-1}\) for the ONe case. These represent an approximately \(5\%\) speed up.

Thus, there is some systematic uncertainty from nuclear reaction rates in our results, which is difficult to characterize, but seems unlikely to be smaller than \(\approx 10\%\). We note that both the above caveats result in even faster flame speeds than the ones we will report. The open and reproducible nature of our work allows this problem to be easily revisited, enabling the impact of future experimental and theoretical progress in the relevant reaction rates to be quickly assessed.

3. Comparison with Past Work

First, we consider CO mixtures. Following \textsc{Tw92} we select a \(^{12}\text{C}\) mass fraction and put the remainder in \(^{16}\text{O}\). Figure 4 compares our results with those of \textsc{Tw92}. Qualitatively, the agreement is good, and we reproduce the trends with \(\rho_0\) and \(X_C\). Quantitatively, above \(\rho_0 = 4\), our results are \(\approx 5\%\)–\(10\%\) slower, while below \(\rho_0 = 4\), our results are faster, up to \(\approx 40\%\) at \(\rho_0 = 1\).

Figure 4 also compares the subset of our results that overlap with \textsc{C07}. We agree well at \(X_C = 0.5\). (It is difficult to see the symbols as they overlap.) We note, as do \textsc{C07}, that their fitting function does not appear to do a good job of matching their tabulated results. Our results are slower at \(X_C = 1\), though we...
note the C07 values also disagree with TW92 and that the primary focus of C07 was on $X_C = 0.3–0.7$.

Next, we consider ONe mixtures. Following TW92, we first select a mass fraction $X_O$ of $^{16}$O. When the mixture is not pure $^{16}$O, we also choose a mass fraction 0.1 of $^{24}$Mg. The remainder is $^{20}$Ne. Figure 5 compares our results with those of TW92. There is qualitative agreement, with a trend (as in the CO case) that our flame speeds are faster than TW92 below $\rho_g = 4$, up to $\approx 50\%$ at $\rho_g = 1$. Above $\rho_g = 4$ and for $X_O = 0.6$ and $X_C = 1.0$, the agreement is within $\approx 10\%$ of TW92. For $X_O = 0.8$, the agreement is somewhat worse and the speeds are systematically $\approx 15\%$ lower above $\rho_g = 6$.

This section demonstrates that our speeds are generally in good agreement ($\approx 10\%$) with the results of past work. Relative to TW92, our calculations adopt a larger nuclear network (leading to faster flames), but have slightly higher conductive opacities (leading to slower flames). In the end, these effects may offset somewhat. We have no reason to expect exact agreement with TW92 as this is not an identical calculation.

4. Influence of Trace Carbon

ONe cores are formed after off-center carbon ignition occurs and a convectively bounded carbon deflagration propagates to the center (e.g., Farmer et al. 2015). Incomplete carbon burning that occurs as the flame approaches the center can leave residual carbon mass fractions of up to a few percent. Schwab & Rocha (2019) performed calculations of accreting ONe WDs including the presence of this carbon and concluded that models are unlikely to reach carbon ignition (and subsequently oxygen ignition and the formation of the deflagration) below the threshold density for $^{24}$Mg electron captures. Here we explore whether, once the deflagration is ignited, the carbon affects the flame speed.

We select the $X_O = 0.6$, $X_Ne = 0.3$, $X_Mg = 0.1$, composition used by TW92 and add a small amount of $^{12}$C, reducing the $^{16}$O mass fraction accordingly. Figure 6 shows the ratio of this flame speed to the carbon-free speed shown in Figure 5. The flame speed increases, reflecting the additional energy release from fusion of $^{12}$C (relative to the $^{16}$O that it replaced). However, for carbon mass fractions of a few percent, the flame speed increases only by $\approx 10\%$. Therefore, we conclude that the presence of small amounts of carbon is unlikely to have a significant effect on the laminar flame speed.

5. Influence of Lower Electron Fraction

Detailed models of ONe WDs do not give compositions that are only $^{16}$O, $^{20}$Ne, and $^{24}$Mg. Several neutron-rich isotopes are typically present at mass fractions of a percent or more, meaning that material is expected to have $Y_e$ significantly below the $Y_e = 0.5$ value of a $^{16}$O/$^{20}$Ne/$^{24}$Mg mixture. Table 1 summarizes the abundances in the ONe core of a representative stellar model from Siess (2006). This mixture has $Y_e \approx 0.49$. See their Section 5 for an explanation of this core nucleosynthesis. As the core slowly grows and its density increases further, the Fermi energy of the degenerate electrons rises. Electron-capture reactions on a given isotope become energetically favored when material exceeds its threshold density. The energetics of these weak reactions are critical for understanding the thermal evolution of super-asymptotic giant branch (SAGB) cores and accreting ONe WDs (e.g., Jones et al. 2013; Schwab et al. 2015, 2017).
we indicate isotopes where these electron captures are likely to occur before the formation of the oxygen deflagration (meaning that their threshold densities are below the threshold density of $^{20}\text{Ne}$, which is $\approx 10^{10}$ g cm$^{-3}$). In what follows, we focus on the most abundant of these isotopes, $^{23}\text{Na}$ and $^{24}\text{Mg}$. The effective threshold density for $^{23}\text{Na}$ is $\approx 1.6 \times 10^9$ g cm$^{-3}$ and for $^{24}\text{Mg}$ is $\approx 4 \times 10^9$ g cm$^{-3}$. The timescales for the electron-capture reactions are typically shorter than the evolutionary timescale of the object, so they are expected to completely convert the parent isotope to its daughter.

The electron captures imply that $Y_e$ spatially varies through the core, with $Y_e$ becoming lower at higher density. By time the deflagration forms and begins to propagate, electron captures have already completely converted the $^{23}\text{Na}$ to $^{23}\text{Ne}$ and the $^{24}\text{Mg}$ to $^{24}\text{Ne}$ over the inner $\approx 0.2M_{\odot}$ of the star. For the mixture in Table 1, this further reduces $Y_e$ to $\approx 0.485$.

In a concluding comment, TW92 note that $Y_e < 0.5$ is expected and mention two calculations including reduced $Y_e$ in the form of $^{22}\text{Ne}$. They report that for a flame in CO with $Y_e \approx 0.498$ the speed decreased by $\approx 5\%$ and for a flame in ONe with $Y_e = 0.48$ the speed decreased by $\approx 30\%$. For CO flames, the effect of $^{22}\text{Ne}$ was studied by CO7. They found the opposite sign of the effect, with a $^{22}\text{Ne}$ mass fraction of 0.06 leading to a $\approx 30\%$ increase in the flame speed.

To quantify the effect of lower $Y_e$, we calculated flame speeds at $\rho_0 = 10$ with a variable amount of neutron-rich material. We performed a set of calculations using each of $^{22}\text{Ne}$, $^{23}\text{Ne}$, and $^{24}\text{Ne}$. In all cases, the mixture had a mass fraction 0.6 of $^{16}\text{O}$ with the remaining material being $^{20}\text{Ne}$. Figure 7 shows the significant impact of the neutron richness, with the flame speed relative to that at $Y_e = 0.5$ doubling by $Y_e \approx 0.488$. The sequences with the different isotopes overlap, indicating the speed up is largely independent of the neutron source.

The small change in $Y_e$ does not significantly affect the internal energy or thermal conductivity, but does lead to a significant change in $\epsilon_{\text{muc}}$ as the initial source of extra neutrons opens additional energy producing reaction channels. We examined the peak values of $\epsilon_{\text{muc}}$ in the calculations shown in Figure 7 and confirmed that the increasing flame speed is due to an increasing $\epsilon_{\text{muc}}$ at lower $Y_e$ and that it quantitatively follows the expectation from Equation (1).

To illustrate that this implies a density-dependent enhancement of the flame speed over the TW92 result, we construct two sets of models that initially have a mass fraction $X_{23} = 0.05$ of $A = 23$ elements and $X_{24} = 0.05$ of $A = 24$ elements. In one, the $A = 23$ material is always $^{23}\text{Na}$ and the $A = 24$ material is always $^{24}\text{Mg}$. In the other, the spatially uniform composition is selected differently depending on the chosen $\rho_0$. The $A = 23$ material is $^{23}\text{Na}$ if the density is below its threshold density and $^{23}\text{Ne}$ if it is above it, while the $A = 24$ material is $^{24}\text{Mg}$ if the density is below its threshold density and $^{24}\text{Ne}$ if it is above it. We then run these models and extract their flame speeds.

Figure 8 compares these two sets of calculations by showing the ratio of the flame speed in the case where the initial material has electron captured to the case where it has not. Above both threshold densities, where $Y_e$ has fallen to $\approx 0.49$, the flame is $\approx 80\%$ faster.

**6. Fitting Formula**

To allow this important effect to be incorporated in hydrodynamics calculations, we provide a simple fitting function like that of TW92, but including $Y_e$ as an additional parameter. As shown in Figure 7, the flame speed varies with $Y_e$ approximately independently of the neutron source. Therefore, we run a set of calculations for $^{16}\text{O}/^{20}\text{Ne}/^{23}\text{Ne}$ mixtures. We use densities $\rho_0 = \{1, 2, 4, 6, 8, 10, 12, 14\}$, $^{16}\text{O}$ mass fractions $X_{O} = \{0.5, 0.6, 0.75\}$, and select $^{23}\text{Ne}$ mass fractions such that $Y_e = \{0.485, 0.490, 0.495, 0.500\}$. Figure 9 plots the results.

At $Y_e = 0.5$, the flame speed always increases with increasing oxygen abundance (as found in TW92). However, in our results at lower $Y_e$, this is no longer true. Incorporating the effect of $X_{O}$ in the fit would require something beyond the power-law scaling used in the fit of TW92. Given the relatively weak dependence on $X_{O}$, we circumvent this complication and propose the following simple fitting function that includes only $\rho_0$ and $Y_e$:

$$v_{\text{flame}} = 16.0 \rho_0^{0.813} [1 + 96.8(0.5 - Y_e)] \text{ km s}^{-1}.$$ (2)

As shown in Figure 10, the fit agrees with the calculated points within 10% relative error at $\rho_0 > 4$, with the maximum error

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**Table 1**

Approximate core composition for the $Z = 0.02$ 9.5$M_{\odot}$ model of Siess (2006)

| Isotope | Mass Fraction (%) |
|---------|-------------------|
| $^{12}\text{C}$ | $\approx 1.0$ |
| $^{16}\text{O}$ | $\approx 55$ |
| $^{20}\text{Ne}$ | $\approx 30$ |
| $^{22}\text{Ne}$ | $\approx 0.8$ |
| $^{23}\text{Ne}$ | $\approx 0.7$ |
| $^{23}\text{Na}$ | $\approx 0.8$ |
| $^{24}\text{Mg}$ | $\approx 5.5$ |
| $^{24}\text{Na}$ | $\approx 3.3$ |
| $^{24}\text{Ne}$ | $\approx 1.5$ |
| $^{20}\text{Ne}$ | $\approx 0.9$ |
| $^{27}\text{Al}$ | $\approx 0.7$ |

**Note.**

* This model has a central density $\approx 7 \times 10^7$ g cm$^{-3}$. Indicated isotopes can undergo additional electron captures as the density increases towards $\approx 10^{10}$ g cm$^{-3}$, the density at which the oxygen deflagration is expected to form.

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8 A representative $Y_e$ profile as a function of mass is shown in Figure 11 of Schwab et al. (2017). One caveat is that if a large core convection zone were to develop, as happens in models adopting the Schwarzschild criterion for convection (e.g., Miyaji et al. 1980), the central region would likely be homogenized.
growing to a 30% underestimate at $\rho_g = 1$. This fitting function will also do a worse job in pure oxygen mixtures (a relative error $\approx 30\%$ for the $X_O = 1$ points shown in Figure 5), but such pure mixtures are unlikely to arise in astrophysical contexts. If a more precise reproduction of our results is desired, the flame speed values are provided in Table 2, allowing direct interpolation in our results.

7. Summary and Conclusions

Using MESA calculations that resolve the structure of conductively propagating deflagrations, we calculated laminar flame speeds in ONe mixtures. These speeds are a necessary ingredient in simulations of the final stages of electron-capture supernovae and accretion-induced collapse. We demonstrated that the values of $Y_e \approx 0.48 - 0.49$ expected in these objects lead to an increase in the flame speed by a factor of $\approx 2$ over that at $Y_e = 0.5$, the value assumed in the widely used prescription of Timmes & Woosley (1992). The low $Y_e$ is due to the nucleosynthesis during the helium and carbon burning phases that preceded the formation of the ONe core and to subsequent electron captures on isotopes initially present in the ONe core (most importantly $^{23}$Na and $^{24}$Mg). The dashed lines show the locations of these composition shifts. The electron fraction of the material is indicated in the lower panel.

**Figure 8.** Effect of the density-dependent $Y_e$ on flame speed. The composition is $X_O = 0.6$ and $X_Ne = 0.3$ ($^{20}$Ne) with 0.05 each of the $A = 23$ and $A = 24$ isotopes. The upper panel shows the ratio of the flame speed in a calculation where these were transformed based on density to the neutron-rich isotopes $^{23}$Na and $^{24}$Ne to one where they remained $^{20}$Na and $^{24}$Mg. The dashed lines show the locations of these composition shifts. The electron fraction of the material is indicated in the lower panel.

**Figure 9.** Flame speeds used to generate the fit given by Equation (2). Point shape indicates $Y_e$ and point color indicates $X_O$. The dashed black curves show the fitting function.

**Table 2** Flame Speed (in km s$^{-1}$) for Models Described in Section 6 and Shown in Figure 9

| $X_O$ | $Y_e$ | $\rho_g$ | 1  | 2  | 4  | 6  | 8  | 10 | 12 | 14 |
|------|------|-------|----|----|----|----|----|----|----|----|
| 0.50 | 0.485| 53.8  | 83.5| 128| 167| 212| 256| 296| 333|
| 0.50 | 0.490| 40.1  | 65.8| 104| 137| 170| 206| 241| 273|
| 0.50 | 0.495| 27.2  | 46.1| 75.6|102| 126| 150| 178| 205|
| 0.50 | 0.500| 15.9  | 26.3| 45.0| 62.6|79.8| 96.4|113|132|
| 0.60 | 0.485| 50.7  | 77.7| 121| 160| 208| 254| 295| 334|
| 0.60 | 0.490| 36.2  | 60.3| 98.9|133| 167| 206| 242| 277|
| 0.60 | 0.495| 25.4  | 43.6| 73.5|100| 126| 153| 182| 211|
| 0.60 | 0.500| 15.7  | 26.1| 45.3| 63.6|81.8| 99.6|117|138|
| 0.75 | 0.485| 51.2  | 75.1| 115| 154| 204| 251| 294| 334|
| 0.75 | 0.490| 35.0  | 56.2| 93.1|127| 165| 206| 245| 282|
| 0.75 | 0.495| 23.9  | 40.7| 70.6| 99.0|127| 157| 190| 221|
| 0.75 | 0.500| 15.6  | 26.1| 45.9| 65.8|86.0|106|126|150|

(This table is available in its entirety in machine-readable form.)

**Figure 10.** Relative error between calculated points and the fit. Point symbols are the same as in Figure 9.

Using MESA calculations that resolve the structure of conductively propagating deflagrations, we calculated laminar flame speeds in ONe mixtures. These speeds are a necessary ingredient in simulations of the final stages of electron-capture supernovae and accretion-induced collapse. We demonstrated that the values of $Y_e \approx 0.48 - 0.49$ expected in these objects lead to an increase in the flame speed by a factor of $\approx 2$ over that at $Y_e = 0.5$, the value assumed in the widely used prescription of Timmes & Woosley (1992). The low $Y_e$ is due to the nucleosynthesis during the helium and carbon burning phases that preceded the formation of the ONe core and to subsequent electron captures on isotopes initially present in the ONe core (most importantly $^{23}$Na and $^{24}$Mg) that occur as the core grows. As shown in Figure 8, this implies that the realized enhancement is density-dependent and most significant for $\rho \gtrsim 4 \times 10^9$ g cm$^{-3}$ (i.e., above the $^{24}$Mg threshold density).

Full star hydrodynamics simulations that follow the propagation of the deflagration through the ONe core do so by including a subgrid model for the flame. These models enhance the laminar speed by including a subgrid model of the flame-turbulence interaction (which allows for a larger, nonplanar area to undergo burning), such that the laminar speed is only a lower limit. Eventually, this speed becomes irrelevant once the turbulence is fully developed, as a turbulent deflagration no longer depends on the laminar speed. In Section 6, we provide a tabulated set of laminar flame speeds as well as a convenient fitting function. These are suitable for incorporation into subgrid flame models.

Figures 4 and 5 in Jones et al. (2016) show the laminar and turbulent flame velocities in their 3D hydrodynamic...
simulations. Typically, these flames remain laminar for \( \approx 0.4 \) s, corresponding to \( \approx 100 \) km of flame propagation. Typically, the inner \( \approx 200 \) km is above the \(^{24}\text{Mg}\) threshold density and thus at the lowest \( Y_e \). Therefore, we believe the factor of 2 speed up is representative of what will be realized in practice. The more rapid release of energy associated with a faster flame pushes models in the direction of being more likely to explode (meaning less likely to form a NS). The full implications of our results await the incorporation of this updated prescription in multidimensional models.

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Software: MESA (Paxton et al. 2011, 2013, 2015, 2018, 2019), MESASDK 20190830 (Townsend 2019), sig99 http://coccubed.asu.edu/code_pages/kap.shtml, matplotlib (Hunter 2007), NumPy (van der Walt et al. 2011), py_mesa_reader (Wolf & Schwab 2017), MesaScript (Wolf et al. 2017).

Appendix

Convergence Studies

In this Appendix, we demonstrate that the flame speeds we report are only weakly dependent on the details of the initial conditions and the spatial and temporal resolution of the MESA calculations. Figure 11 shows this for a flame in a CO mixture and Figure 12 does so for a flame in an ONe mixture. The discussion below applies equally to both figures.

Panel (a) illustrates the effect of varying our procedure for extracting the flame speed. By default, we measure the flame speed using the change in position over the second half of the time interval covered by the calculation (indicated as [0.5, 1.0] in the legend). So long as we avoid the transients during the early part (roughly the first quarter) of the calculation, the extracted velocities are consistent at the percent level.

Panel (b) and panel (c) illustrate the effect of varying the temperature of the initial hot spot \( T_i \) and its fractional size \( q_s \). As expected, so long as the hot spot causes a steady-state flame to propagate, these choices have no effect on the flame speed.

Panel (d) illustrates the effect of varying the upstream temperature \( T_u \). The flame speed increases with increasing upstream temperature, but such that a factor of \( \approx 2 \) change in \( T_u \) leads to only 1%–2% increase in the flame speed. We would expect this to remain true so long as \( T_u \) is much less than the post-burn temperature of \( \approx 10^{10} \) K.

Panel (e) illustrates the effect of varying the spatial resolution of the MESA calculation. MESA adaptively refines its mesh based on a set of mesh functions. The maximum cell-to-cell variation of these functions is maintained at around the value of the control mesh_delta_coeff which is set equal to 1 in our calculations. One of the built-in mesh functions has the form \( T_{\text{function1_weight}} \times \log(T/K) \). This function ensures that temperature gradients are resolved, placing approximately \( T_{\text{function1_weight}} \) zones per dex change in temperature. The number of zones in the calculation (which is \( \approx 1000 \) for the default) varies roughly linearly with \( T_{\text{function1_weight}} \). The results are approximately independent of the spatial resolution, with a subpercent increase between the default and higher resolution cases.

Panel (f) illustrates the effect of varying the temporal resolution of the MESA calculation. The control varcontrol_target limits the fractional step-to-step variation of quantities in the same cell. The number of timesteps in the calculation (which is \( \approx 2000 \) for the default) varies roughly linearly with the inverse of varcontrol_target. The results are approximately independent of the time resolution, with a roughly 1% increase between the default and the highest resolution.
Figure 11. Effect of modeling choices on flame speeds for the fiducial CO case ($X_C = 0.5$, $\rho_0 = 6$). Default choices are indicated with black lines. The value in parentheses in the legend is the flame speed in km s$^{-1}$. 

(a) Varying the portion of data used to measure flame speed

(b) Varying the temperature of the initial hot spot $T_s$

(c) Varying the size of the initial hotspot $q_s$

(d) Varying the upstream temperature $T_b$

(e) Varying spatial resolution

(f) Varying temporal resolution
Figure 12. Effect of modeling choices on flame speed for the fiducial ONe case ($X_O = 0.6$, $\rho_g = 10$). Default choices are indicated with black lines. The value in parentheses in the legend is the flame speed in km s$^{-1}$. 

(a) Varying the portion of data used to measure flame speed

(b) Varying the temperature of the initial hot spot $T_s$

(c) Varying the size of the initial hotspot $q_s$

(d) Varying the upstream temperature $T_b$

(e) Varying spatial resolution

(f) Varying temporal resolution
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