Turbulent Mixing and Nuclear Burning in Stellar Interiors

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ABSTRACT

The turbulent burning of nuclei is a common phenomenon in the evolution of stars. Here we examine a challenging case: the merging of the neon and oxygen burning shells in a 23 M⊙ star. A previously unknown quasi-steady state is established by the interplay between mixing, turbulent transport, and nuclear burning. The resulting stellar structure has two burning shells within a single convection zone. We find that the new neon burning layer covers an extended region of the convection zone, with the burning peak occurring substantially below where the Damköhler number first becomes equal to unity. These characteristics differ from those predicted by 1D stellar evolution models of similar ingestion events. We develop the mean-field turbulence equations that govern compositional evolution, and use them to interpret our data set. An important byproduct is a means to quantify sub-grid-scale effects intrinsic to the numerical hydrodynamic scheme. For implicit large eddy simulations, the analysis method is particularly powerful because it can reveal where and how simulated flows are modified by resolution, and provide straightforward physical interpretations of the effects of dissipation or induced transport. Focusing on the mean-field composition variance equations for our analysis, we recover a Kolmogorov rate of turbulent dissipation without it being imposed, in agreement with previous results which used the turbulent kinetic energy equation.

Key words: turbulence – mixing – nuclear burning – stellar evolution

1 INTRODUCTION

It is now feasible to simulate stellar convection in three-dimensions (3D), with realistic microphysics, multiple species of nuclei, and sufficient resolution in space and time to represent turbulent flow (Meakin & Arnett 2007; Mocá 2011; Woodward et al. 2013). Historical work on stellar convection (Böhm-Vitense 1958) and 3D simulations of stellar atmospheres (Stein & Nordlund 1998) have generally focused on flows having uniform composition, a case which is usually appropriate for the outer layers of stars.

By contrast convection in stellar interiors is generally characterized by nuclear burning and nonuniform composition. Here we examine the interaction between turbulent convection, thermonuclear burning, and entrainment at boundaries. Simulations of convective shells, driven by nuclear burning, show entrainment of material from surrounding stable layers. Erosion at boundaries introduces inhomogeneities in composition, entropy, and buoyancy into the convective flow. This can be viewed as a multi-stage process of entrainment, transport, dispersion (or stirring) induced from the largest to smallest eddies, as well as diffusion, spanning the full spectrum of space-time scales of the flow (Dimotakis 2005). The feedback of such mixing on nuclear burning, convection, and its impact on the evolution of the star remains largely unexplored: an ad hoc diffusion operator is almost universally used in stellar evolution. This paper begins to analyze these stages.

We focus on the oxygen burning shell in a massive supernova progenitor. Oxygen burning and neon burning occur at sufficiently similar temperatures that these burning shells may interact (Arnett 1974a; Arnett 1974b). Interaction was indeed found in the 3D simulations of Meakin (2006) and Meakin & Arnett (2007), but was not analyzed in detail there. Here we present a detailed account of the compositional mixing and modified nuclear burning, which occurs as the convective oxygen-burning shell merges with the (initially) stable overlying neon shell.

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We use 3D numerical hydrodynamics in the implicit large eddy simulation framework ILES (Grinstein et al. 2007), which means we solve the Euler equations with a non-oscillatory finite volume numerical fluid solver. In the current study we use the PPM method (Colella & Woodward 1984). The dissipation in such solvers comes from solving the Riemann shock problem over each zone \( \Delta s \), giving a dissipation rate \( \approx v^3/\Delta s \) as a shock of speed \( v \) traverses a zone. In a turbulent cascade the mean damping is \( D \sim v^3/\Delta s \sim v^3/\ell \), which is determined by the rms velocity \( v \) and dimension \( \ell \) of the turbulent region. Use of such solvers introduces an implicit sub-grid model which corresponds to a Kolmogorov turbulent cascade, freeing computational resources to capture the large scales relevant to astrophysics. See Grinstein et al. (2007) for references and a more rigorous discussion.

This is in contrast to the direct numerical simulation approach DNS (Pope 2000), which solves the Navier-Stokes difference equations on the grid, all the way down to the dissipation scale. In this approach most of the computational effort is spent on these small scales, which are buried in the turbulent cascade. Sylte et al. (2000) showed that both methods converge to the same result, but that ILES is more efficient for highly turbulent flows. We have confirmed that our simulations extend from the integral (large) scale down into the inertial range of the turbulent cascade (e.g. Cristini et al. 2017).

In this paper, we extend our analysis by using an approach inspired by Reynolds-averaged Navier-Stokes (RANS) methods (Viallet et al. 2013; Mocák et al. 2014; Arnett et al. 2015). Our approach differs from traditional RANS (Besnard et al. 1992; Chassaing et al. 2010) in two fundamental ways: (1) the fluctuations are taken from our simulations, and hence are dynamically constrained, and (2) we solve the ILES Euler equations. A more accurate acronym than “RANS” (which we have used previously) is needed for clarity; we choose “Reynolds averaged ILES” (RA-ILES), to distinguish our approach. Unlike unconstrained RANS analysis, our RA-ILES equations are complete, exact to the accuracy of our grid, and require little added computational cost. There is no closure issue.

Previous papers (Meakin & Arnett 2007; Viallet et al. 2013; Arnett et al. 2015; Cristini et al. 2017) have focused on the turbulent kinetic energy equation (TKE); here our analysis shifts to mean-field transport equations for the density \( \rho \), momentum \( \rho \mathbf{v} \), and their turbulent fluxes and variances, complemented by analysis of relevant timescales and nuclear burning processes. These transport equations are the ones which deal with changes in composition variance: i.e., mixing.

The paper is organized as follows: In §2 we describe the initial conditions and the 3D stellar model that we investi–

\footnote{The change in specific kinetic energy over the shock traversal time gives the dissipation rate. PPM approximates sub-grid structure as piece-wise parabolic, smoothing higher-order terms and decreasing information (complexity). Variances in velocity and in scalar variables dissipate/diffuse at this same rate at the sub-grid scale.}

\footnote{There is no explicit Navier-Stokes viscosity term to generate higher-order moments in the conventional way (Tritton 1988); the implicit turbulent cascade gives closure.}

\begin{table}
\begin{tabular}{|c|c|}
\hline
Grid dimensions \((\Delta r, \Delta \theta, \Delta \phi)\) & \( 6 \times 10^8 \text{cm} \times 27.5^\circ \times 27.5^\circ \) \\
Grid zoning & \( 400 \times 100 \times 100 \) \\
\( \Delta \rho_{\text{av}} \) (s) & \( 300 \) s \\
\( v_{\text{rms}} \) & \( \sim 1.4 \times 10^7 \text{cm/s} \) \\
\( \tau_{\text{conv}} \) & \( \sim 65 \) s \\
\hline
\end{tabular}
\caption{3D oxygen burning simulation properties (model ob.3d.B). \( \Delta \rho_{\text{av}} \) is the averaging timescale for the mean field analysis; \( v_{\text{rms}} = \sqrt{2K_{\text{tot}}/M} \) is the approximate global rms velocity (where \( K_{\text{tot}} \) is the total turbulent kinetic energy in the convection zone and \( M \) mass contained in the convection zone), and \( \tau_{\text{conv}} = 2l_{\text{conv}}/v_{\text{rms}} \) is the convective turnover timescale (where \( l_{\text{conv}} \) is size of the convection zone, \( \sim 4.3 \times 10^9 \) cm). All values were obtained at the central simulation time 1060 seconds around which we perform all subsequent time-averaging.}
\end{table}

\section{2 Initial Model and Simulation Method}

The initial 1D model for our 3D simulation is a 23 \( M_\odot \) supernova progenitor. It was evolved with the 1D TYCHO stellar evolution code (Young & Arnett 2005; Arnett et al. 2010) using mixing-length theory (MLT) to a point just following core oxygen burning where oxygen, neon, carbon, helium, and hydrogen are burning in concentric shells outside a degenerate core of silicon and sulfur. The structure of the 1D initial model is shown in the panels of the top row of Figure 1. It has a single convective oxygen shell enclosed by two stable layers. The initial convective region is driven primarily by nuclear burning of oxygen and extends from its base around \( 4.3 \times 10^8 \) cm up to a radius around \( 7.2 \times 10^8 \) cm, where the Ledoux criterion indicates a stable boundary. The stable layers below and above are composed primarily of silicon and oxygen; in the top layer the dominant nuclear burning is that of neon, and nonconvective. Such stellar structures are quite common and can be found, e.g., in cores of low-mass red giants during the core helium flash, or during core carbon flashes of “super-AGB” stars (Mocák et al. 2012).

Reactive-hydrodynamic evolution in 3D was computed with the PROMPI code, a version of the legacy PROMETHEUS code (Fryxell et al. 1991) adapted to parallel computing via the Message Passing Interface (MPI). This code is an Eulerian implementation of the piecewise parabolic method (PPM) of Colella & Woodward (1984) updated with a Riemann solver for real gases according to Colella & Glaz (1985). Additional added physics include self gravity in the spherically symmetric approximation, a realistic equation of state to handle the semi-degenerate stellar plasma (Timmes & Swesty 2000), and a general nuclear
reaction network. For the current simulation we use a 25-isotope network that includes neutrons, protons, $^4$He, $^{12}$C, $^{16}$O, $^{20}$Ne, $^{23}$Na, $^{24}$Mg, $^{28}$Si, $^{31}$P, $^{32}$S, $^{34}$S, $^{35}$Cl, $^{36}$Ar, $^{38}$Ar, $^{39}$K, $^{40}$Ca, $^{42}$Ca, $^{44}$Ti, $^{46}$Ti, $^{48}$Cr, $^{50}$Cr, $^{52}$Fe, $^{54}$Fe, $^{56}$Ni. All of the important strong and weak reactions are included. It is conceptually convenient to decompose the network results into separate processes. In Appendix §A we give simpler approximations which are used to identify the different stages of burning.

In the simulation neither Si nor C burning are ever the primary nuclear process. The dominant energy release is due to burning of $^{16}$O and $^{20}$Ne. The dominance of neon burning is the result of entrainment as it mixes $^{20}$Ne into the deeper, hotter layers of the oxygen burning shell. Oxygen burning is invigorated by mixing in new $^{16}$O fuel from the previously non-convective region (Meakin & Arnett 2007).

We focus on the region encompassing the oxygen and neon burning shells and their interactions with each other and with the adjacent stably stratified layers. The computational grid is defined in a spherical coordinate system with periodic boundary conditions in the angular directions and reflecting (non-transmitting) boundaries in the radial directions. The opening angle of the grid is $27.5^\circ \times 27.5^\circ$ in $\theta$ and $\phi$. A short summary of the simulation properties, including zoning, is presented in Table 1.

Due to its lower computational cost, this medium resolution simulation could be extended over a longer time span than higher resolution simulations, and attain a new and reliable quasi-steady state.

### 3 MEAN-FIELD EVOLUTION EQUATIONS

In this section we describe and develop the Reynolds-Averaged ILES (RA-ILES) analysis for composition i.e. mean-field equations describing the evolution of mean fields...
related to composition on spherically averaged shells. The evolution equations for mean composition (§3.3), turbulent composition flux (§3.4), and composition variance (§3.5) are then considered in turn.

3.1 Reynolds-Averaged ILES Analysis

The RA-ILES equations derived here are exact, and do not employ approximations, at least to the extent that the continuum approximation is appropriate and flow features are resolved. This contrasts with methods using closure relationships and truncations of the RANS equations to construct approximate models of turbulence. The RANS equations, when closed by 3D numerical simulations to become “RA-ILES” equations, are able to represent the full range of hydrodynamical behavior present in a stellar interior.

As shown in Hinze (1975), Besnard et al. (1992), and Chassaing et al. (2010), the RANS framework provides a rational approach to interpreting complex 3D fluid dynamical data. The extension of the methodology to “RA-ILES” (see §1) for stellar interiors is discussed in Viallet et al. (2013), Mocák et al. (2014), and Arnett et al. (2015).

3.2 The Averaging and Decomposition Procedure

In this section we define the averaging rules needed to obtain 1D mean and fluctuation fields from 3D flow data. This decomposition makes precise the relationship between the 1D mean fields evolved by a stellar evolution code and 3D hydrodynamical simulations (Viallet et al. 2013). Two types of averaging are used: in time, and in space. In practice, both types of averaging are combined and contribute to a meaningful average of the flow within a spherical layer by virtue of the ergodic hypothesis\(^3\) and spherical symmetry\(^4\).

The average of a quantity \(q\) on a spherical shell at radius \(r\) (i.e. a mean field) is defined by

\[
\overline{q}(r, t_c) = \frac{1}{T} \Delta \Omega \int_{t_c-T/2}^{t_c+T/2} \int_{\Delta \Omega} q(r, \theta, \phi, t') \, d\Omega \, dt'
\]

where \(d\Omega = \sin \theta d\theta d\phi\) is the solid angle in spherical coordinates, \(T\) is the averaging time period, \(t\) is time and \(\Delta \Omega\) is the solid angle being averaged over. The time coordinate \(t_c\) represents the center of the time-averaging window used (central time).

Fluctuations, which retain the full time and space dependence as the original, self-consistent 3D flow field, are defined according to the decomposition \(q(\vec{x}, t) = \overline{q}(r, t) + q''(\vec{x}, t)\), noting that \(\overline{q}(r, t) = 0\) by construction. Similarly, a Favre (or density weighted) average is given by

\[
\tilde{q} = \frac{\overline{pq}}{\overline{p}}
\]

which defines a complimentary decomposition of the flow according to \(q = \tilde{q} + q''\) where \(q''(r, \theta, \phi, t)\) is referred to as the Favrian fluctuation and its mean is zero when Favre averaged: \(q''(r, t) = 0\). A more complete elaboration on the algebra of these averaging procedures can be found in Chassaing et al. (2010).

The mean fields presented in this paper were calculated

\(^3\) An average of a physical quantity over a statistical ensemble is equivalent to its average over time.

\(^4\) If spherical symmetry were broken, as it would be in the presence of a magnetic field or a mean stellar rotation field, spherical averaging would not be appropriate. In the case of a mild rotation field, for example, mean fields could not be restricted to less than two dimensions and would be represented most naturally in a meridional plane.
by post-processing snapshots (which were written to disk every 0.5 s of simulated time, for a period from 888 s up to 1211 s), by following these steps:

(i) Calculate Reynolds fluctuations for each snapshot, at time \( t \), from the raw 3D simulation data, as defined above:

\[
q_{\text{inst}}^i(r, \theta, \phi, t) = q(r, \theta, \phi, t) - \langle q \rangle(r, t)
\]

where the dependence on space and time variables are shown for each term. \( \langle q \rangle(r, t) = 1/\Delta t \int_{i\Delta t} q(r, \theta, \phi, t) \, d\Omega \) gives spatial average of \( q \) for a given time \( t \) and radius \( r \).

(ii) Calculate needed products of any required thermodynamic quantities \( q_1, q_2, q_3 \), etc. (for example \( q_1 q_2 \) or \( q_1 q_2 q_3 \)) from 3D fields. Favrian fluctuation can be obtained from

\[
q_{\text{inst}}^i = q_{\text{inst}} - \langle q_{\text{inst}} \rangle (\rho).
\]

(iii) space and time average the products calculated in the prior step around central time \( t_p \), as defined in Eq. 1 above.

We find that the turbulent flow can be very well sampled when using an averaging time window of width \( T \) around two convective turnover times (Mocák et al. 2014). Figure 2 shows an example of the averaging window effect on our flow for the mean specific turbulent kinetic energy (TKE) defined as \( \tilde{k} = \frac{1}{3} \left( \rho \left( \tilde{u}_i \tilde{u}_i + \tilde{u}_j \tilde{u}_j + \tilde{u}_k \tilde{u}_k \right) \right) \) where \( u_i \) are components of the velocity field. Small timescale features are absorbed into the mean-field when the averaging window exceeds around 150 seconds (~2 turnover times). In order to present a robust statistical analysis we use an averaging window of 300 seconds throughout this paper, which is ~4 turnover times.

### 3.3 Evolution Equation for Mean Composition

The instantaneous evolution equation for mass fraction of element \( i \) in spherical geometry is (Arnett 1996), Eq.4.97,

\[
\partial_t (\rho \xi) = -\nabla \cdot (\rho \xi \mathbf{u}) + \rho \dot{X}^{\text{nuc}}.
\]

Applying our decomposition and averaging procedure (Sect.3.2), we obtain the following 1D transport equation

\[
\tilde{p} \tilde{D}_i \tilde{X}_i = -\nabla \cdot \tilde{f}_i + \tilde{p} \cdot \dot{X}^{\text{nuc}} + \tilde{N}_i,
\]

where \( X_i \) is mass fraction of chemical element \( i \), \( \rho \) is density, \( \mathbf{u} = [u_r, u_\theta, u_\phi] \) is the velocity vector, \( \nabla \) is the diverger operator, \( \dot{X}^{\text{nuc}} \) is the rate of nuclear burning of \( i \), and \( \tilde{f}_i = \tilde{p} \dot{X}^{\text{nuc}} \tilde{u}_i \tilde{u}_j \) is the turbulent flux of element \( i \). \( \nabla \cdot \tilde{f}_i \) is the radial divergence operator and the \( \tilde{D}_i \) is the mean-flow Lagrangian derivative \( \tilde{D}_i(\cdot) = \partial_t (\cdot) + \tilde{u}_r \partial_{r}\tilde{X}_i \).

The mean-field transport equation (Eq. 6) states that the temporal change of mass fraction of an element \( i \) in the Lagrangian frame of reference, \( \tilde{p} \tilde{D}_i \tilde{X}_i \), is caused by either a spatial redistribution by the turbulent flux, \( -\nabla \cdot \tilde{f}_i \); or by nuclear burning, \( \tilde{p} \dot{X}^{\text{nuc}} \).

We define the numerical residual in these equations by \( \tilde{N}_i \), which represents the implicit action of the numerical simulation algorithm. These terms are discussed in more detail in the following sections and again later in §5.

#### 3.4 Evolution Equation for Turbulent Composition Flux

The transport equation for the turbulent flux of an arbitrary chemical element can be obtained by using the following general formula for second-order moments (Mocák et al. 2014),

\[
\tilde{p} \tilde{D}_i \tilde{u}_j \tilde{u}_k = \left[ \frac{\partial \tilde{f}_j}{\partial \xi} \right] + \tilde{p} \cdot \dot{X}^{\text{nuc}} \tilde{u}_i \tilde{u}_j \tilde{u}_k + \tilde{d}_i \tilde{d}_j \tilde{d}_k \right]
\]

\[
+ \tilde{d}_i \tilde{d}_j \tilde{d}_k \tilde{u}_l \tilde{u}_m \tilde{u}_n \tilde{u}_p \tilde{u}_q \tilde{u}_r \tilde{u}_s \tilde{u}_t
\]

by substituting \( X_i \) for \( c \) and \( u_r \) for \( d \) and using the radial momentum equation

\[
\tilde{p} \tilde{D}_i (u_r) = \nabla \cdot (\tau_{rr} - G_{\text{M}}) - \partial_r P + \rho g_r
\]

where \( \tau_{rr} = [\tau_{rr}, \tau_{r\theta}, \tau_{r\phi}] \) contains the radial components of the viscous stress tensor (not explicitly included in our simulation model), \( G_{\text{M}} = -\rho v_r^2 / \tau_{rr} \) is a geometric term, \( P \) is the pressure, and \( g_r \) is the gravitational acceleration in the radial direction.

After averaging, we arrive at the flux evolution equation,

\[
\tilde{p} \tilde{D}_i (f_i/P) = -\nabla \cdot \tilde{f}_i - f_i \tilde{u}_r \tilde{\partial}_r \tilde{X}_i - \tilde{X}_i \partial_r \tilde{P} - \tilde{X}_i \partial_r \tilde{\partial}_r \tilde{P} + \tilde{N}_{\text{f}_i}
\]

where \( f_i = \tilde{p} \tilde{X}_i \tilde{u}_i \tilde{u}_j \tilde{u}_k \) is the radial component of the “flux of the turbulent flux” of element \( i \). Here \( -f_i \tilde{u}_r \) is a production term due to velocity effects controlled by the flux itself. The \(-\tilde{X}_i \partial_r \tilde{X}_i \) term is a production term which transports the flux from regions with higher \( X_i \) to regions with lower \( X_i \), or vice versa and is controlled by the Reynolds stress \( \tilde{R}_{ir} = \rho \tilde{u}_i \tilde{u}_j \). The terms \(-\tilde{X}_i \partial_r \tilde{P} \) and \(-\tilde{X}_i \partial_r \tilde{\partial}_r \tilde{P} \) drive evolution of the flux in the presence of a pressure gradient and pressure fluctuations. The term \(+\tilde{p} \tilde{X}_i \tilde{u}_i \tilde{u}_j \tilde{u}_k \tilde{u}_l \tilde{u}_m \tilde{u}_n \tilde{u}_p \tilde{u}_q \tilde{u}_r \tilde{u}_s \tilde{u}_t \) drives evolution of the flux by the net nuclear burning of element \( i \). Finally, the term \( \tilde{G}_i = G_{\text{M}} - \tilde{X}_i \tilde{G}_{\text{M}} \), where \( G_{\text{M}} = -\rho \tilde{u}_i \tilde{u}_j \tilde{u}_k \tilde{u}_l / r - \rho \tilde{u}_i \tilde{u}_j \tilde{u}_k \tilde{u}_l / r \) mediates the production of \( f_i \) due to centrifugal forces caused by horizontal (non-radial) motion of the flow. The equation contains additionally a term \( \tilde{X}_i \tilde{g}_r \tilde{N}_{\text{f}_i} \), which we neglect due to gravitational acceleration \( g_r \) being constant in the simulation.

The residuals of all these mean fields in the flux transport equation, \( \tilde{N}_{\text{f}_i} \), represent numerical effects which we do not calculate explicitly in RA-ILES, which has no explicit viscous (Navier-Stokes) term. Their magnitude is determined solely by the implicit action of our numerical scheme at the sub-grid scale. The precise mathematical formulation of the term is following:

\[
\tilde{N}_{\text{f}_i} = -\nabla \cdot (\tilde{X}_i \tau_{rr}) + \varepsilon_i^r + \tilde{G}_i^r
\]

where

\[
\varepsilon_i^r = -\tau_{rr} \partial_r \tilde{X}_i - \tau_{r\theta} \partial_\theta \tilde{X}_i^\theta - \tau_{r\phi} \partial_\phi \tilde{X}_i^\phi
\]

\[
\tilde{G}_i^r = \tilde{X}_i \tau_{r\theta} \partial_\theta \tilde{X}_i^\theta + \tilde{X}_i \tau_{r\phi} \partial_\phi \tilde{X}_i^\phi
\]

The term \(-\nabla \cdot (\tilde{X}_i \tau_{rr}) \) is a flux of a composition flux controlled by viscosity; \( \varepsilon_i^r \) is a viscous dissipation of the turbulent flux and \( \tilde{G}_i^r \) is of geometric origin and contributes to flux production due to action of viscosity.
3.5 Evolution Equation for Composition Variance

To derive a transport equation for the variance of composition fluctuations of an arbitrary element $i$, we use Eq. 7 and substitute $c$ for $d$:

$$\bar{p}\bar{D}_i\bar{c}'\bar{c}'' = +2\bar{c}'\bar{p}\bar{D}_i\bar{c} - 2\bar{p}\bar{D}_i\bar{c}'\bar{c}''\bar{c}'\bar{c}'' - \bar{c}'\bar{p}\bar{D}_i\bar{c}''\bar{c}'\bar{c}'' + \bar{c}'\bar{p}\bar{D}_i\bar{c}''\bar{c}'\bar{c}''.$$  (11)

Next we substitute a mass fraction of a given element $X_i$ for $c$ and find

$$\bar{p}\bar{D}_i\sigma_i = -\nabla f_i^* - 2f_i\partial_i\tilde{X}_i + 2\bar{X}_i\rho\tilde{X}_i^{\text{nuc}} + \mathcal{N}_{c,i}$$  (12)

where $\sigma_i = X_i^fX_i^p$ is the variance of the mass fraction of species $i$; $f_i^* = \rho X_i^fX_i^p u_i^f$ is the turbulent flux of this variance; $2f_i\partial_i\tilde{X}_i$ is the down-gradient production/destroyal; and $2\tilde{X}_i\rho\tilde{X}_i^{\text{nuc}}$ is a variance source/sink due to nuclear burning.

The term $\mathcal{N}_{c,i}$ represents the dissipation of composition variance due to the numerical scheme, which is discussed further in §5.

4 RELEVANT TIMESCALES

The simulations are strongly dynamic, and may be better understood by comparison of several timescales, which we define here. The convective turnover timescale is

$$\tau_{\text{conv}} = 2(r_i^t - r_i^b)/v_{\text{rms}},$$  (13)

where $r_i^t$ and $r_i^b$ are the radii of the bottom and top convection boundaries, and $v_{\text{rms}}$ is the rms of the velocity field in the convection zone. The net nuclear (e-folding) burning timescale for element $i$ is

$$\tau_{\text{nuc}} = \frac{\tilde{X}_i}{\tilde{X}_i^{\text{nuc}}}.$$  (14)

The nuclear (e-folding) burning timescale due to photodisintegration of element $i$ is

$$\tau_{\text{nuc-phot}} = 1/\lambda_i,$$  (15)

and the nuclear (e-folding) timescale for burning element $i$ due to two-body reaction of $j$ and $k$ is

$$\tau_{\text{nuc-two}}^{i,j,k} = 1/\left(\rho/\lambda_{jk}Y_jY_k/\tilde{Y}_i\right),$$  (16)

where $Y_i = X_i/A_i$ are molar abundances of species $i$, and $A_i$ is mean number of nucleons per isotope $i$. $\lambda_i$ and $\lambda_{jk}$ are nuclear reaction rates expressed as $N_A\langle\sigma v\rangle$, where $N_A$ is Avogadro’s number, $\sigma$ is the reaction cross section, and $v$ the relative velocity of reactants (Clayton 1983; Arnett 1996).

The (e-folding) transport timescale of element $i$ is

$$\tau_{\text{tran}}^{i} = \frac{\tilde{X}_i}{\nabla \cdot f_i/\bar{p}};$$  (17)

and finally, the (e-folding) dissipation timescale of element $i$ is

$$\tau_{\text{dis}}^{i} = \sigma_i/\varepsilon_i,$$  (18)

where $\varepsilon_i$ is the dissipation of variance $\sigma_i$, which we identify with the residual term $\mathcal{N}_{\sigma_i}$ defined in Eq. 12 above and discussed further in §5.3.3.

5 RESULTS

To initiate convection in our 3D simulation we seed the initial hydrostatic model with small (10$^{-3}$) random perturbations in density and temperature (Meakin & Arnett 2007) in the unstable regions. Convection starts with an increase of turbulent kinetic energy near the base of the convection zone, similar to other qualitatively comparable cases, e.g. Mocák et al. (2009); Stancliffe et al. (2011); Woodward et al. (2015). The velocities eventually grow from zero to 10$^{-2}$ of sound speed, to become a self-consistent turbulent cascade, with final amplitudes that happen to be of the same order as MLT velocities because they involve the same buoyancy.

Since the 1D initial model does not contain sufficient dynamic information for self-consistent quasi-static 3D convection, a readjustment occurs as a self-consistent convective flow forms. This is usually referred to as the “initial transient” phase, and is a common feature of 3D stellar hydrodynamics simulations; see §5.1. Even with perfect hydrostatic matching of the initial model, a readjustment is required – 3D convection is different from 1D stellar convection theory (e.g. Meakin & Arnett 2007; Arnett et al. 2015).

5.1 Initial transients

For neutrino-cooled stages such as oxygen burning, convection is vigorous and turbulent (Meakin & Arnett 2007). One-dimensional (1D) stellar evolutionary sequences use mixing-length theory (MLT), which assumes a specific average correlation between fluctuations in velocity and entropy; these correlations drive the fluxes crucial for thermal balance. In contrast, 3D simulations must develop such correlations in a dynamically self-consistent way, to obtain the appropriate average. This takes a turnover time, after which the turbulent cascade can remove the excess entropy to obtain a balance on average. Because mean velocities are dominated by fluctuations (Meakin & Arnett 2007), turbulent convection is made up of alternate bursts and pauses, occurring throughout the convection zone. It is intermittent; only the average is quasi-steady. The 1D models cannot provide the dynamically consistent phases of the fluctuations which are crucial to accurately determine the net effect of cancellations, and for setting up a consistent 3D initial model.

In 3D stellar interior simulations, this fact, combined with the (generally negligible) errors introduced by mapping a 1D model to a 3D space, leads to an “initial transient” – even if the most exacting hydrostatic balance is enforced in the mean.

5.2 Multi-layered convective-reactive burning

During the initial transient phase (see §5.1 above), the convective plumes emerging from the oxygen burning shell quickly extend into the neon burning region, which was previously separated by a very narrow region of stability. The two shells merge within the first 100 seconds (~1 turnover) of the simulated star time. Soon a quasi steady-state is established, when the total turbulent kinetic energy density of the flow in the convection zone reaches roughly 10$^{14}$ erg g$^{-1}$, at around 300 seconds (~3 turnovers) of the simulation.

In a local 1D treatment of convection, a linear criterion
such as the Ledoux or Schwarzschild defines a region of stability which is capable of preventing mixing, regardless of how weak the stable layer may be. In reality, the ability of a stable layer to survive against turbulent convection depends on both the strength of the stabilizing gradient and the strength of the adjacent turbulence (the Richardson criterion; Meakin & Arnett (2007); Cristini et al. (2017)). In this particular model, the upper convective boundary was calculated by the Ledoux criteria (at a radius around $7.2 \times 10^8$ cm), and was quickly overwhelmed by the adjacent turbulent flow. The oxygen and neon-burning shells are rapidly but gently\(^5\) mixed together.

The onset of convection, the subsequent merging of the burning shells, and the establishment of a quasi steady-state, is depicted in Figure 3 as a group of “Kippenhahn plots”, which are space-time diagrams. Shown are the time evolution of (1) the turbulent kinetic energy, (2) mean and variance of \(^{16}\text{O}\) and \(^{20}\text{Ne}\) mass fractions, and (3) nuclear energy generation rates. The top row shows the time evolution of the turbulent kinetic energy (TKE), including, from left to right, the radial, horizontal, and total values. The kinetic energy at a given radius is calculated as half of the variance of the velocity component in the horizontal plane, e.g., the radial TKE \(k_r\) is defined by \(k_r(j) = \sigma_{u_r}^2 = (u_r - \bar{u_r})^2\). The horizontal TKE is defined by \(k_h = \sigma_{u_\theta}^2 + \sigma_{u_\phi}^2\), and the total TKE by \(k_{\text{tot}} = k_r + k_h\).

The most prominent features of the quasi-steady state include a single convection zone with two internal burning layers, as well as the interplay of turbulent mixing and material entrainment. We describe these features later for \(^{16}\text{O}\) and \(^{20}\text{Ne}\) nuclear burning\(^6\). The net nuclear burning of other prominent species in the convection zone (i.e., \(^{12}\text{C}\) or \(^{28}\text{Si}\)) are several orders of magnitude smaller and are not discussed further here.

\(^5\) No shocks or even strong pressure waves could be seen, only steady growth of turbulence.

\(^6\) A more comprehensive presentation of the chemical element transport for all the remaining nuclear species in the model is presented in Mocák et al. (2014).
gen burning shell from the upper boundary. Figure 4 shows both (left) a cross sectional slice through the 3D simulation; as well as (right) the radial profile of the Ne\textsuperscript{20} mass fraction in both the initial model and after a quasi-steady state has been achieved. The reason this phenomena has been missed in 1D models is that those models use MLT, which is a local theory, and must be supplemented by ad hoc boundary conditions (Schwarzschild or Ledoux). The correct boundary conditions are nonlinear, dynamic and complex i.e. not linear, static, and simple, as we shall see below.

The nuclear energy production increases in the vicinity of the temperature maximum by a factor of 5. We find a secondary peak in the energy release at \(r \approx 5.7 \times 10^8\) cm, where the main source of nuclear energy is neon burning (Eq. A2). The contributions to the net nuclear energy generation by different reactions are presented in the left panel of Figure 5.

In the region of this secondary energy generation peak, the mass fraction of Ne\textsuperscript{20} is maintained in a quasi-steady state by a balance between a net inflow from the overlying neon-rich stable layer and depletion by nuclear burning. This balance can be seen by comparing the timescales for the different processes involved, as shown in the right panel of Figure 5: the timescales for destruction of Ne\textsuperscript{20} by nuclear burning (blue curve) and inflow by transport (green curve) match in the burning region. In contrast, the overlying layers are dominated by neon transport where its timescale is \(\sim 3\) orders of magnitude shorter than that for depletion by nuclear burning.

A Damköhler number, \(Da\), may be defined as the ratio of transport to nuclear burning timescales. Since we have more than one burning process, multiple Damköhler numbers should be defined, one for each reaction process \(i\):

\[
Da^i = \tau_{\text{tran}}^i / \tau_{\text{nuc}}^i
\]

For \(Da \sim 1\), such that the transport timescale is comparable to the burning timescale, we have the regime of convective-reactive mixing (Herwig et al. 2011). In this regime it is important to model the convection well, in order to predict the coupled mixing and burning correctly. Such convective-reactive situations are likely to be common in stellar evolution — if allowed by the evolutionary algorithm. They might be common, for instance, in the early Universe in low-metallicity stars (Fujimoto et al. 2000; Schlattl et al. 2002), or during dredge-up in AGB stars (Herwig 2004; Goriely & Siess 2004).

The Damköhler number varies across a convective region because of the local variation in transport and nuclear timescales. The nuclear timescale depends on burning rates which vary with fuel abundance, density, and temperature, while the transport timescale is determined by the topology of the velocity field. In the upper layers of the convection zone in our simulation, the transport timescale is much shorter than the nuclear burning timescales, so \(Da^i_{\text{nuc}} \ll 1\). There is much more neon entrained and transported than nuclear burning is able to destroy. These timescales become close only at around \(7 \times 10^8\) cm, where effective nuclear destruction of entrained Ne\textsuperscript{20} begins (see right-hand panel of Fig. 5).

It is instructive to compare this 3D result to 1D simulations of similar convective-reactive events. One such class of calculations is the ingestion of protons into He-burning convective regions (“proton ingestion episodes”). These calculations result in a splitting of the He convection zone into two parts (see e.g. Herwig et al. 1999; Campbell & Lattanzio 2008). The split occurs because the large energy release from hydrogen burning creates a temperature inversion at the point of peak H-burning luminosity, creating a (formally) stable region just below. If time-dependent mixing is used in the 1D calculation, this peak occurs at the point where the lifetime of a proton against capture by \(^{12}\text{C}\) is just equal to the timescale of convective mixing, i.e., \(Da = 1\). In some...
In contrast, our 3D calculation shows that, in the neon ingestion case, the $Da \sim 1$ condition defines a substantial burning region – rather than a single radial location. This suggests that a split of the convective zone is less likely, as the burning is distributed rather than sharply peaked. Such splitting is likely to be an artifact of the 1D formulation. In order for a radial velocity to go to zero in a quasi-steady state, baryon conservation requires a finite horizontal velocity (Arnett et al. 2015), which in turn implies a shear instability and probably mixing instead of splitting. This may be considered a boundary issue, or “overshooting”.

The (smooth) peak of nuclear energy release from neon burning is actually located near the middle of the burning region, at around $5.7 \times 10^8$ cm (Fig. 5). Neon mixes down still further, to $5 \times 10^8$ cm. As suggested by Herwig et al. (2011), the width of these regions must be due to a combination of factors, such as the temperature sensitivity of the particular nuclear reactions, the range of plume velocities, and the various mixing ratios in the plumes of (in our case) neon-enriched material. In summary, the 3D model shows that the energy release from Ne burning dominates that of O in the top half of the convection zone.

Regardless of whatever the final answer may be for the evolution of Ne and O burning shells, the inconsistency of 1D and 3D models is a cause to question present 1D evolutionary scenarios.

5.3 Mean-Field Analysis of 3D Simulation Data
We use the three composition related mean-field RA-ILES equations to interpret our 3D simulation data, which we depict graphically in Figs. 6, 7, and 8. Each of these balance equations is discussed in the following subsection in turn, with a focus on a physical interpretation of each term in the equations.

5.3.1 Mean Composition
The mean profiles of $^{16}$O and $^{20}$Ne mass fraction are presented in the top row of Fig. 6; while the bottom row of the same figure shows the individual terms in the mean-field balance equation given by Eq. 6 which underly these profiles.
This figure shows that the composition profiles $p\bar{X}$ are established as an interplay between the material entrainment, turbulent mixing, and nuclear burning.

**Time Dependence:** $pD_t\bar{X} = \partial_t(p\bar{X}) + \nabla_r(p\bar{X}\bar{u}_r)$. While the convective layer is in a quasi-steady state, ongoing mixing at the upper boundary increases the depth of the convection zone over time. The signature of this mass entrainment appears in the Eulerian component of the time derivative, which is plotted as the red curve in Figure 6. By contrast, the change in the composition profile due to an overall expansion or contraction is negligible, as revealed by the turquoise curve.

**Turbulent Transport:** $-\nabla rf_i = -\nabla r(p\bar{u}_r\bar{X}_i')$. The stable layer material which is mixed into the convective layer is transported by turbulent velocity fluctuations. This spatial redistribution of material is shown by the blue curve in Figure 6 where we see that $^{16}\text{O}$ and $^{20}\text{Ne}$ are transported out of the boundary layer ($r \sim 9 \times 10^8$ cm) and into the lower nuclear burning regions of the convection zone ($r < 7 \times 10^8$ cm). A simple balance is found: the transport out of the stable layer is closely matched by the time rate of change there; and at the other end, the rate at which material is being brought into the nuclear burning zone is balanced by the rate it is being destroyed by nuclear reactions.

In the case of $^{20}\text{Ne}$, the turbulent transport at the outer convective boundary shows depletion by downdrafts – there is more $^{20}\text{Ne}$ going out from the region than is going in. In contrast, the transport is positive in the neon burning region below. This indicates that there is more neon going into this region than out, and the net input is equal to the $^{20}\text{Ne}$ burned by nuclear reactions. The situation is almost identical for $^{16}\text{O}$, except there is additional transport of freshly produced oxygen in the neon burning layer due to its photodisintegration at $r \sim 5.7 \times 10^8$ cm.

**Nuclear Burning:** $p\bar{X}_{\text{nuc}}$. The rate at which the mean abundance of $^{20}\text{Ne}$ is changed due to nuclear burning is shown by the green curve in Figure 6. It reveals a region of
depletion near the lower part of the convection zone where the temperature and densities rise above $\sim 1.9 \times 10^5$ K and $\sim 7 \times 10^9$ g cm$^{-3}$, respectively. The $^{20}$Ne burning rate is governed by the rate at which entrained material is transported into the burning region.

The width of the neon burning region is set at the upper end by the reaction rates which are a function of the density and temperature, so that burning does not begin until the material is transported below a radius of $7 \times 10^8$ cm. At lower radii, the burning region is truncated by the complete depletion of the $^{20}$Ne as it is being transported downwards. The nearly symmetrical, gaussian shape is therefore a coincidence of the near balance in timescales between these two competing processes: a more vigorous turbulent transport, and hence a shorter convective mixing timescale through the burning region, would result in a deeper lower boundary consisting of fuel which was able to penetrate deeper before being completely consumed. The higher temperatures present near the base of the oxygen-burning shell, however, ensures that the bulk of any entrained $^{20}$Ne is consumed before traversing the entire convective layer, resulting in a structure with “stacked” nuclear burning zones and a stratified composition. The plateau in $\overline{p}X_{\text{NeO}}$, at $r \sim 6.5 \times 10^8$ cm (see Fig. 6), reflects this balance between entrainment and burning.

The transport of $^{16}$O in the upper layers of the convection zone shows similar properties to that of $^{20}$Ne, with oxygen being pulled down into the convection zone from the upper stable layer by turbulent velocity fluctuations. However, in layers where Ne burning dominates, $^{16}$O is produced through photo-disintegration: $^{20}$Ne($\gamma$, $^{16}$O)$^{16}$Mg (see lower left panel in Fig. 6). Toward smaller radii, oxygen depletion continues to increase due to nuclear fusion.

**Numerical Residual:** $\nabla \cdot \mathbf{u}$. Unlike traditional solvers of the Navier-Stokes equations, a conservative hydrodynamics algorithm such as PPM results in errors in conservation laws with magnitudes comparable to machine precision. Therefore, the residuals found in the balance equations are due to the much larger effects – and can be understood as loss of information at the grid level (dissipation and diffusion).

A quantitative measure of this numerical residual is shown by the dashed black curve in Figure 6. The impact that it has on the mean flow can be quantified by comparing the amplitude of this curve with the other dominant physical processes controlling the time evolution. In comparison to nuclear burning and turbulent transport, which are the dominant processes, the numerical flux is relatively small over the majority of the domain. However, it does make an important contribution in narrow regions around both the upper and lower convective boundaries (Fig. 6). Here turbulent boundary layers develop (Landau & Lifshitz (1959), §44). As expected, this flux is present exactly where the mean composition fields possess steep gradients. The steady presence of internal wave fields in and around the convective boundaries ensures that the material fields are constantly in motion against the computational mesh, thereby weakening the gradients. The strength of this numerical flux is seen to diminish with finer resolution as gradients are better resolved (Arnett et al. 2015; Cristini et al. 2017).

### 5.3.2 Turbulent Composition Flux

The turbulent composition flux $f_i$ represents the rate at which matter of species $i$ is transported across a given spherical shell. Flux profiles and balance equation terms for the flux evolution equations for $^{16}$O and $^{20}$Ne are presented in Figure 7. The negative values of the fluxes indicate that they are oriented towards the stellar center, which is consistent with the picture of depletion from upper stable layers dominating the mixing of new material into the convection zone.

**Time Dependence:** $\overline{p \mathbf{D}_i(f_i/\rho)} = \partial_i (f_i) + \nabla \cdot (f_i \mathbf{u})$. The rate of change of the flux, at a given spatial location and within a fixed mass shell (i.e., at a fixed Lagrangian coordinate) – shown by the orange and brown curves in Fig. 7 – are both found to be small compared to the other terms determining the flux profile, indicating a quasi-steady balance. The expansion velocities in the model are only $\sim 10^2$ cm s$^{-1}$, which is 5 orders of magnitude smaller than the r.m.s. of the turbulent velocity field ($\sim 10^7$ cm s$^{-1}$). The quasi-steady flux of $^{16}$O and $^{20}$Ne is here due to a balance between nuclear burning, turbulent transport, and pressure-force and centrifugal-force coupling.

**Turbulent Production:** $-\overline{R_e \partial_i \overline{X_i}}$. The presence of radial velocity fluctuations in regions where there is a mean radial composition gradient leads to the production of a turbulent mass flux. This process involves the radial component of the turbulent kinetic energy $k_r = \overline{R_e}/2$ and the mean composition gradient for isotope $i$, given by $\partial_i \overline{X_i}$. In the case of $^{20}$Ne, where the mean abundance gradient is everywhere positive, the presence of turbulent velocity fluctuations leads to a down directed directed mass flux, and hence has a negative sign. This term is shown by the red curve in Figure 7. The situation is similar for $^{16}$O with the exception of (1) the neon burning region – where the oxygen abundance locally increases due to photo-disintegration of neon – giving rise to a contribution to the outward-directed composition flux; and (2) the region around the lower boundary, where oxygen burning produces a very steep gradient and hence a large contribution to the downward-directed flux component.

**Pressure-force Coupling:** $-\left( \nabla P \frac{\partial_i}{\overline{X}} + \nabla \left( \frac{\partial_i}{\overline{P}} \right) \right)$. This process is localized near the upper and lower convective boundaries where it peaks and drops off on either side of the interface. These terms are shown by the turquoise colored curve in Figure 7. On the convection zone side of each boundary, the overall strength of this effect drops as the buoyancy diminishes relative to the velocity fluctuations; on the stable side of each boundary, the strength of buoyancy is set by the entropy stratification while the turbulent velocity fluctuations decay with distance from the boundary.

Material which resides in the stable layer is buoyant relative to material in the adjacent convection zones by virtue of its relative mean molecular weight and entropy: in the upper stable layer, the material is lighter per ion and is in a higher entropy configuration. Therefore, as material is drawn down into the convection zone by turbulent velocity fluctuations its downward motion is resisted by this buoyancy. As the freshly entrained stable layer material is dredged deeper into the convection zone, it is continually being mixed with its surroundings by turbulence which reduces its relative buoyancy. By the time this newly ingested ma-
material reaches the far end of the convection zone the relative buoyancy affecting its flux is negligible.

The term $-\nabla r f_r$ can be rewritten to highlight its relationship to buoyancy: using the identity $\nabla \rho = -\rho \nabla X/\rho$ and hydrostatic equilibrium $\partial_r \rho = -\hat{\rho} \hat{g}$ we find

$$-\nabla r \partial_r \rho = \nabla \rho \nabla \hat{X}.$$  

The term $-\nabla \rho \nabla \hat{X}$ is related to pressure fluctuations and is of a dynamic rather than a thermodynamic origin. In the anelastic approximation, the pressure fluctuations are governed by a Poisson equation

$$\Delta \rho' = -\nabla : (\rho a' \nabla \nabla) - \nabla \cdot (\hat{\rho} \hat{g}).$$

In stably stratified layers the buoyancy component dominates; in the well-mixed turbulent layer the advective term dominates (Viallet et al. 2013); and the partially mixed boundary layer is a region of transition between the two.

**Centrifugal-force Coupling:** $\varpi$. This term is driven by horizontal flow produced by the deflection of plumes at the inner boundaries of the convection zones and provides a radially directed acceleration to the matter with which it is coupled. At the upper boundary, the plumes which ultimately become horizontal flows carry material which is primarily representative of the mean composition of the convection zones, and hence will be depleted in those isotopes which are being actively entrained at the upper boundary. Therefore, the net effect of this process will be a net negative contribution to the entrained mass flux. Since we are considering downward directed fluxes of material being entrained at the upper boundary, this term will therefore act as a source for this entrained matter flux. This term is plotted as the yellow curve in Figure 7.

A similar, but mirrored effect, takes place at the lower boundary for $^{16}$O.

**Nuclear burning:** $u_i' \rho X_i$. As the $^{20}$Ne approaches the bottom of the convection zone it begins to undergo nuclear burning which diminish fluctuations in its abundance, and
therefore reduces the downward-directed mass flux. This process is represented by the magenta curve in Figure 7 where it can be seen to operate in the neon burning region. This same process also operates for the $^{16}$O flux, but with the opposite sign.

**Turbulent Transport:** $-\nabla \cdot \vec{f}_i = -\nabla \cdot \left( \overline{\rho u_i u_i} \vec{X}_i \right)$.

In the same manner in which velocity fluctuations act to redistribute Reynolds stress, or its trace, the turbulent kinetic energy $k = \text{Tr}(\overline{\rho u_i u_i})$; velocity fluctuations also redistribute compositional flux $(f_i = \overline{\rho u_i X_i})$ in space. Although not a dominant effect in controlling the radial profile of $^{20}$Ne flux, it does transport some of the downwardly directed flux from the middle of the convection zone to the upper stable layer and the region just below the $^{20}$Ne burning zone. This spatial redistribution can be seen in Figure 7 where it is represented by the darker blue curve.

**Numerical Residual:** $N_{fi}$. The implicit action of turbulent cascade on the turbulent composition fluxes is apparent at the convective boundaries.

### 5.3.3 Composition Variance and its Dissipation Rate

Composition variance $\sigma_i$ is a measure of the amplitudes of the composition fluctuations at a given radial location in the stellar interior. It is typically largest at convective boundaries (where the gradients are steepest), due to the oscillatory distortion of the boundary by gravity waves as well as the presence of turbulent entrainment of material into the convection zone. This $\sigma_i$ and the terms in the balance equation for this mean-field variable is shown in Figure 8. The $^{16}$O variance distribution shows maximum values at both convective boundaries, whereas $^{20}$Ne peaks only at the upper boundary due to a lack of neon at deeper layers due to depletion by nuclear burning and turbulent mixing.

Dissipation is an important feature of turbulence. In the ILES approach, the Euler equations are solved so that dissipation is implicitly included; in our case, the numerical algorithm provides an effective damping at the dissipation (sub-grid) scale. The rate at which composition fluctuations are destroyed by this process can be expressed by a dissipation timescale, as in Eq. 18. We associate this timescale with the residual of the transport equation of variance (Eq. 12) since it characterizes the magnitude of numerical dissipation. We discuss this important feature further in the paragraph below, under the subheading **Numerical Residual**.

**Time Dependence:** $\vec{p} \vec{D}_i \sigma_i = \partial_t (\overline{\rho \sigma_i}) + \nabla \cdot (\overline{\rho u_i \sigma_i})$.

The variance is in a steady state throughout most of the computational domain but does show a sign of the convective boundary mixing at the upper boundary. The signature of this mass entrainment appears in the Eulerian time derivative which is shown by the turquoise curve in Figure 8 – while the overall effect of background expansion on the mean variance field is negligible, as shown by the magenta curve.

The sine wave-like shape of the time dependence is due to the outward migration of the variance peak following the mass entrainment (note the sign of the terms plotted as indicated in the figure legend).

**Turbulent Production:** $-2 f_i \partial_i \vec{X}_i$. The primary source of variance in the composition field is turbulent production, which is analogous to the source of turbulent flux discussed above. In this case, variance is produced when a turbulent mass flux is present in regions where there is a mean convection gradient. Physically, the growth of variance can be thought of as the coupling of the composition fluctuation associated with the turbulent flux to the newly created composition fluctuation which is generated by the associated velocity fluctuation.

In the case of the $^{20}$Ne, the net downward directed composition flux works against a positive gradient in the $^{20}$Ne abundance (see Fig. 4), thus amplifying the local fluctuations and resulting in an overall positive increase in the mean variance. Production is seen to dominate in and around the upper boundary, then drops off with depth into the convection zone, until a small peak in the $^{20}$Ne burning region where again the mean composition gradients are very steep – in this case due to the nuclear depletion of $^{20}$Ne. This term is shown by the green curve in Figure 8. The situation for $^{16}$O is similar, but has significant contributions at both boundaries.

**Turbulent Transport:** $-\nabla \cdot \vec{f}_i = -\nabla \cdot \left( \overline{\rho u_i X'_i} \right)$.

Variance is transported by the variance flux $f'_i$ which is the correlation of a velocity fluctuation with the square of the composition fluctuation. As with all turbulent transport, it is a conservative process and neither creates nor destroys variance but simply redistributes it spatially. The redistribution of $^{20}$Ne has a relatively simple behavior in our 3D models: the variance that is generated within the upper boundary by turbulent production is moved both outward, to a region lying just above the main region of turbulent production; as well as being moved downward to the bottom of the $^{20}$Ne burning region to the depth at which $^{20}$Ne is completely consumed by nuclear burning. The case for $^{16}$O is similar, but can be seen operating at the lower convective boundary as well.

**Nuclear burning:** $2X'_i \rho X'_i$. Nuclear burning results in the growth or decay of fluctuations due to nuclear reactions. In the case of $^{20}$Ne, the abundance is depleted through nuclear burning and so this term acts as a sink for the variance that was transported or produced in the $^{20}$Ne burning zone.

**Numerical Residual (dissipation):** $N_{\sigma_i}$. The dashed black curve in Figure 8 shows the residual term. The thin solid black curves show a model of the scalar dissipation which assumes that the dissipation timescale is equal to the Kolmogorov damping timescale, defined by

$$\tau_L \equiv \left( \frac{\nu}{\epsilon} \right)^{1/2} \equiv \frac{\hat{k}}{\epsilon_k}$$

where $\nu$ is viscosity, $\epsilon$ is the turbulent kinetic energy transfer rate, and $\hat{k} = 1/2u''_{rms}^3$ is the Favrian specific turbulent kinetic energy mentioned also in Section 5.2. Here, $\epsilon_k$ is the dissipation rate of $\hat{k}$, which can be approximated by $\epsilon_k \sim u''_{rms}/L$, where $L$ is the scale of the flow i.e. essentially the size of convection zone (Arnett et al. 2009; Cristini et al. 2017).

These timescales allow us to test if our simulation results agree with Kolmogorov’s first similarity hypothesis, which states that in turbulent flows with high Reynolds numbers, the statistics of the motion at the dissipation scale has a universal form that is uniquely determined by $\nu$ and $\epsilon$ (Pope 2000). We do indeed find that the scalar dissipation rates in our simulation, calculated using $\tau_L$ as defined above,
Figure 8. Top: Profiles of \(^{16}\text{O}\) and \(^{20}\text{Ne}\) variances. Bottom: \(^{16}\text{O}\) and \(^{20}\text{Ne}\) variance transport equations. The shaded vertical region and vertical line as defined in Figure 5.

provides a good fit to our data (Fig. 8), with

\[
N_{\text{ri}} \sim \frac{\nabla \sigma_i}{\tau_L} \approx \frac{\rho u_i \varepsilon_k}{k} \approx \frac{\rho U_{\text{rms}}}{L} \frac{1}{k}. \tag{23}
\]

This result supports the physical consistency of the sub-grid scale numerics of our simulation and the ILES approach, even at rather low spatial resolution and effective Reynolds numbers, \(Re^{\text{eff}}\). In our simulation, \(Re^{\text{eff}} \sim N_c^{4/3} \sim 1440\), where \(N_c \approx 234\) is the number of radial grid points across the convection zone. That the composition variance dissipation rate \(N_{\text{ri}}\) is directly proportional to \(\nabla \sigma_i / \tau_L\) (Eq. 23) is consistent with mixing models of molecular diffusion. This further confirms the assumption that the rate of scalar mixing is determined by scalar variance production at large scales (Fox 1995).

Despite the inviscid assumption of the Euler equations, scalar dissipation in the composition variance will occur because of the presence of a numerical mass flux \(\mathcal{J}\). This will modify all continuity equations such that they have a form:

\[
\partial_t (\rho X_i) = \nabla \cdot (\rho u_i X_i + \mathcal{J}) + \rho X_{i,\text{mix}}. \tag{24}
\]

In these equations \(\nabla \cdot \mathcal{J}\) is therefore a conservative redistribution term that will lead to smearing effects, i.e. numerical diffusion. Mass conservation is however still maintained across the domain. We use these equations to represent the modified equations being solved by the numerical algorithm.

5.4 Effects of Resolution

As mentioned above, the effective Reynolds number of the flow in our simulation is \(Re^{\text{eff}} \sim 1440\) and it is thus expected to be in the turbulent regime. This is supported by our finding that the variance dissipation is proportional to the variance over the large eddy timescale – a phenomenology expected when turbulence is developed and has an inertial range.
Our previous work utilising RA-ILES (“RANS”) analysis shows that the qualitative features of the interior of convection zones is not strongly affected by resolution, for a wide range of effective Reynolds numbers (Re$_{\text{eff}}$ ∼ 500 to 3300; Viallet et al. 2013; Mocák et al. 2014; Cristini et al. 2017). This shows that our RA-ILES technique is useful even at relatively low resolution – at least for PPM hydrodynamics. Moreover, the RA-ILES approach can be used to assess resolution dependence and convergence by analyzing the residuals defined in by the mean-field equations.

The boundaries of convection zones, on the other hand, are generally under-resolved in stellar interior simulations. This is readily identifiable – and quantified – in the RA-ILES residuals (eg, $N$ in Fig. 6, 7). In the current simulation, the lower boundary has a large ‘spike’ in residuals. This is a turbulent boundary layer, and is relatively thin, with steep gradients in various quantities. We have previously shown that these residual spikes diminish with increasing resolution (Cristini et al. 2017). The top boundary also shows a spike in the residuals, but is much less pronounced because of the shallower gradients present there – it is a ‘softer’ boundary.

While these residuals must affect the quantitative rate of entrainment, we do not expect the qualitative features of the convective-reactive flow to be strongly impacted, especially since the residuals become negligible away from the convective boundaries.

6 SUMMARY

We have analyzed coupled turbulence and nuclear burning in a 3D hydrodynamic implicit large eddy simulation (ILES) of an oxygen burning shell in the core of a 23 M$_\odot$ supernova progenitor star.

6.1 Hydrodynamic simulation

Our initial stellar structure encompassed a single oxygen-burning convection zone with a stable Ne-burning layer above. After an initial transient episode which led to a gentle merging of the O-burning convective shell with the Ne-burning shell, a quasi-steady layered-shell state was formed. Two layers of stable nuclear burning were set up in the joined velocity field of a single convection zone, with $^{16}$O dominating the nuclear energy production at the bottom of the convection zone, but $^{20}$Ne dominating in the upper layers.

The quasi-static state showed a Ne-burning rate in the convection zone up to seven orders of magnitude higher than in the initial 1D model. This was due to a steady entrainment of $^{20}$Ne into the hotter, oxygen burning shell.

Further, neon burning was found to occur over a wide region in the convection zone. This is in contrast to the situation found in 1D codes for similar convective-reactive cases, for example during proton ingestion into He burning regions in low-mass stars. As a consequence of limitations in their mixing algorithms, 1D codes generally show burning in a very thin layer, at a depth where the ratio of the burning timescale equals that of the convective turnover timescale, i.e. where the Damköhler number equals one. In our simulation this defines only the top of a wider burning region, and the peak of the burning is well below this point (Fig. 5).

6.2 RA-ILES analysis

The key results presented in this paper are based on a Reynolds-Averaged ILES (RA-ILES) analysis of the composition-related mean fields found for our 3D ILES simulation data, and extend our previous investigations which used the turbulent kinetic energy (TKE) equation.

We develop RA-ILES equations (§3) which describe the time evolution of mean composition, turbulent composition flux, and composition variance for each isotope present in the stellar plasma. The equations derived are exact and do not employ approximations to the extent that the continuum approximation is appropriate and flow features are resolved. This contrasts with use of closure relationships and truncations of the RANS equations to construct models of turbulence. The RANS equations, when closed by 3D ILES numerical simulations (RA-ILES), are able to represent the full range of hydrodynamical behavior that is present in a stellar interior simulation.

Our detailed RA-ILES analysis revealed, over the timescale of the simulation (~4 turnovers):

- A simple balance between turbulent transport and nuclear burning in the convection zone is maintained, whereby any amount of material entrained into the convection zone is counterbalanced by nuclear burning (§5.3.1, Fig. 6).
- The turbulent composition flux appears to be quasi-static due to a complex balance between turbulent production, nuclear burning, turbulent transport, pressure force coupling, and centrifugal forces (§5.3.2, Fig. 7).
- Composition variance provides information about composition fluctuation amplitudes, and the residual of the composition variance mean field equation then gives direct information about numerical dissipation timescales (§5.3.3).

6.3 Dissipation and ILES

The residual of the composition variance equation implies a match with the Kolmogorov damping timescale (§5.3.3, Fig. 8). This independently confirms a similar result found using the turbulent kinetic energy equation (Meakin & Arnett 2007; Arnett et al. 2010), and indicates a deep consistency in our ILES approach. In the simulation, the dissipation of both turbulent kinetic energy and composition variance occurs at the sub-grid level, as desired. This should be true of the entropy and temperature variances as well, because ILES joins onto the inertial range of the turbulent cascade.

6.4 Implications for Stellar Evolution Theory

As discussed above (§6.1) we have identified very substantial differences between 1D stellar evolution modelling and our 3D modelling. These findings indicate that if shell mergers and entrainment do happen in real stars, 1D evolutionary models that do not include these phenomena are likely to be substantially in error. Moreover, based on our finding of the way in which ingested fuel burns in the convective zone, it appears that even if 1D models do incorporate shell mergers and entrainment, current 1D theory would be unable to model them reliably.

The differences identified here will need to be addressed.
to improve modeling of convective-reactive phases. If not addressed the errors from these events will accumulate through the subsequent evolutionary phases and will cause uncertainty in, for example, pre-supernova evolution and the 3D explosion models that rely on the 1D models for their initial states.

Further 3D modeling – and, just as importantly, detailed analysis such as that presented in this paper – is vital to guide these improvements by illuminating the physical processes at play.

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APPENDIX A: APPROXIMATE RATES OF ENERGY GENERATION

The nuclear burning of carbon, neon, oxygen and silicon during advanced evolutionary stages of massive stars involve increasingly extensive nuclear reaction networks, which have been analyzed and reduced to much simpler equivalent forms (Arnett 1996). These equivalent nuclear reactions (see Arnett 1972a,b, 1974a; Arnett 1974b for original work), allow calculation of approximate formulas for nuclear energy production rates coming from each of these equivalent reaction links, and are useful for understanding the energy generation. Our simulations actually use the 25-species network in each zone. Here we give simpler approximations which are useful for analysis; see Fig. 5.

Nuclear energy production due to $^{12}$C burning goes primarily to unstable highly excited state of $^{23}$Mg which decays further through neutron, $\alpha$ and proton channel to $^{24}$Mg, $^{20}$Ne and $^{23}$Na and its rate (in erg g$^{-1}$ s$^{-1}$) can be approx-
imated by (Arnett 1972b):

$$
\dot{\epsilon}_{\text{c12}} \approx 4.8 \times 10^{18} Y_{\text{c12}}^2 \rho \lambda_{\text{c12,c12}}.
$$  \hspace{2cm} (A1)

where $\lambda_{\text{c12,c12}}$ is reaction rate for $^{12}\text{C}(^{12}\text{C},\gamma)^{24}\text{Mg}$ Caughlan & Fowler (1988).

Neon burns in massive stars primarily by reaction $2\text{Ne}^{20} \rightarrow \text{He}^4 + \text{O}^{16}$, assuming approximate equilibrium between one photo-disintegrated $^{20}\text{Ne}$ into $^{16}\text{O}$ and $^4\text{He}$ and one $\alpha$ capture on the new oxygen restoring $^{20}\text{Ne}$. At this point, one $^4\text{He}$ starts to add to $^{20}\text{Ne}$ and produces $^{24}\text{Mg}$. The net effect is that for each 2 neon nuclei there is one new nucleus of $^{16}\text{O}$ and one of $^{24}\text{Mg}$ produced. Its nuclear production rate (in erg g$^{-1}$ s$^{-1}$) can be approximated by (Arnett 1974a):

$$
\dot{\epsilon}_{\text{nuc,\text{ne}}}^{\text{20}} \approx 2.5 \times 10^{29} T_{9}^{5/2} \frac{Y_{\text{ne}^{20}}^2}{Y_{\text{o}^{16}}} \lambda_{\text{ne}^{20}}^{\alpha\gamma} e^{-\left(54.89/T_{9}\right)}
$$  \hspace{2cm} (A2)

where $\lambda_{\text{ne}^{20}}^{\alpha\gamma}$ is the reaction rate for $^{20}\text{Ne}(\alpha,\gamma)^{24}\text{Mg}$ Caughlan & Fowler (1988) and $T_{9}$ is temperature $T$ divided by 10$^9$K. The equilibrium in the neon burning holds in our network, as indicated by a proximity of the neon nuclear timescales for $\text{Ne}^{20} \rightarrow \text{He}^4 + \text{O}^{16}$ and $\text{He}^4 + \text{O}^{16} \rightarrow \text{Ne}^{20}$ shown in Figure 5.

Oxygen burning produces compound nuclear states of $^{32}\text{S}$ which may decay by neutron, proton, deuteron or $\alpha$ channel into $^{31}\text{S}$, $^{31}\text{P}$, $^{30}\text{P}$ or $^{28}\text{Si}$ but the $n$, $p$ and $\alpha$ are recaptured, giving a distribution of nuclei from $^{28}\text{Si}$ to Ca isotopes (Truran & Arnett 1970; Arnett 1974b). The energy production rate (in erg g$^{-1}$ s$^{-1}$) can be approximated by:

$$
\dot{\epsilon}_{\text{nuc,\text{o}}}^{\text{16}} \approx 8 \times 10^{18} Y_{\text{o}^{16}}^2 \rho \lambda_{\text{o}^{16},\text{o}^{16}}
$$  \hspace{2cm} (A3)

$\lambda_{\text{o}^{16},\text{o}^{16}}$ is reaction rate for $^{16}\text{O}(^{16}\text{O},\gamma)^{32}\text{S}$ Caughlan & Fowler (1988).

Simplified view of silicon burning involves 2 nuclei of $^{28}\text{Si}$ producing one $^{56}\text{Ni}$ nucleus via specific sequence of photo-disintegration and fusion reactions. The critical reaction that allows the photodisintegration of $^{28}\text{Si}$ is $^{24}\text{Mg}(\gamma,\alpha)^{28}\text{Ne}$. The energy production rate can during such silicon burning be approximated by:

$$
\dot{\epsilon}_{\text{nuc,\text{si}}}^{\text{28}} \approx 1.8 \times 10^{28} T_{9}^{3/2} X_{\text{si}^{28}} \lambda_{\text{ni}^{20}}^{\alpha\gamma} e^{-142.07/T_{9}}
$$  \hspace{2cm} (A4)

(Clayton 1983; Woosley et al. 2002).