Pre-explosive Accretion and Simmering Phases of SNe Ia

Luciano Piersanti1,4, Eduardo Bravo2, Oscar Straniero1,5, Sergio Cristallo1,4, and Immaculada Domínguez3

1 INAF-Osservatorio Astronomico d’Abruzzo, via Montore Maggini, snc, I-64100, Teramo, Italy; luciano.piersanti@inaf.it, oscar.straniero@inaf.it, sergio.cristallo@inaf.it
2 E.T.S. Arquitectura del Vallés, Universitat Politècnica de Catalunya, Carrer Pere Serra 1-15, E-08173 Sant Cugat del Vallès, Spain; eduardo.bravo@upc.es
3 Universidad de Granada, E-18071 Granada, Spain; inma@ugr.es

Abstract

In accreting white dwarfs (WDs) approaching the Chandrasekhar limit, hydrostatic carbon burning precedes the dynamical breakout. During this simmering phase, e-captures are energetically favored in the central region of the star, while β-decay are favored more outside, and the two zones are connected by a growing convective instability. We analyze the interplay between weak interactions and convection, the so-called convective URCA process, during the simmering phase of Type Ia supernovae (SNe Ia) progenitors and its effects on the physical and chemical properties at the explosion epoch. At variance with previous studies, we find that the convective core powered by the carbon burning remains confined within the 21Ne,F URCA shell. As a result, a much larger amount of carbon has to be consumed before the explosion that eventually occurs at larger density than previously estimated. In addition, we find that the extension of the convective core and its average neutronization depend on the WD progenitor’s initial metallicity. For the average neutronization in the convective core at the explosion epoch, we obtain \( n_{\text{exp}} = (1.094 \pm 0.143) \times 10^{-3} + (9.168 \pm 0.677) \times 10^{-2} \times Z \). Outside the convective core, the neutronization is instead determined by the initial amount of \( C + N + O \) in the progenitor star. Since \( S, Ca, Cr, \) and \( Mn \), the elements usually exploited to evaluate the pre-explosive neutronization, are mainly produced outside the heavily neutronized core, the problem of too high metallicity estimated for the progenitors of the historical Tycho and Kepler SNe Ia remains unsolved.

Unified Astronomy Thesaurus concepts: Stellar accretion (1578); Stellar nucleosynthesis (1616)

1. Introduction

Type Ia supernovae (SNe Ia) are among the most reliable extragalactic distance indicators and, in turn, they play a fundamental role in our understanding of the universe. Hubble diagrams based on SNe Ia have shown the recent acceleration of the expansion rate of the universe (Riess et al. 1998; Perlmutter et al. 1999), as well as the previous deceleration (Riess et al. 2004). While high-redshift SNe Ia are fundamental tools to probe dark energy (see, for example, Betoule et al. 2014; Jones et al. 2018), those at lower redshifts constrain the progenitors of the historical Tycho and Kepler SNe Ia.

The precision of SNe Ia as distance indicators relies on empirical relations, such as, for example, the maximum-decline relation (Phillips 1993; Phillips et al. 1999), which link their maximum absolute magnitudes with light-curve shape and color. Thanks to dedicated surveys collecting thousands of SNe Ia, it has been proved that these relations show a quite small dispersion, i.e., \( \sim 0.1 \) mag (Scoulnic et al. 2018). This occurrence demonstrates that SNe Ia are a very homogeneous class of events, which is the key feature for being a reliable tool for cosmology.

On the other hand, the homogeneity of this class of supernovae is supported by their most popular model. It consists of a thermonuclear explosion of a carbon-oxygen (CO) white dwarf (WD) with a mass of the order of the Chandrasekhar limit (Hoyle & Fowler 1960). According to this scenario, the WD is destabilized by mass accretion from a close stellar companion. The supernova engine is a thermonuclear explosion that starts with the carbon fusion. As a result, the whole WD is incinerated, and its ashes contain intermediate-mass elements and a substantial amount of iron-group isotopes. SNe Ia are indeed the main contributors to the iron pollution of the interstellar gas (see, for example, Matteucci & Tornambe 1985; Bravo et al. 1993; Timmes et al. 1995; Prantzos et al. 2018). Among these nuclei, \( ^{56}\text{Ni} \) is mostly produced, and it is the radioactive energy from the cascade \( ^{56}\text{Ni} \rightarrow ^{56}\text{Co} \rightarrow ^{56}\text{Fe} \) that powers SNe Ia light curves (LCs). This simple scenario implies that the memory of the properties of the WD progenitor, such as the initial mass or the original chemical composition, are lost in the supernova outcomes, i.e., LCs and spectra. This occurrence is often referred to as “stellar amnesia” ( Höflich 2006).

SNe Ia still remain an intriguing mystery. First of all, no clear consensus exists concerning the nature of their stellar progenitors, the evolutionary paths up to the explosion and the explosion mechanisms. Moreover, several questions have been raised about the actual diversity of the SNe Ia. It is well known that the total amount of \( ^{56}\text{Ni} \) determines the maximum brightness and other features of the LCs. Therefore, the rather large range of values of the maximum brightness would imply that different amounts of \( ^{56}\text{Ni} \) should be produced by different supernovae. What determines these differences? Moreover, various features of the LCs correlate with the properties of the host galaxies (Hamuy et al. 1996, 2000; Gallagher et al. 2005; Mannucci et al. 2005; Sullivan et al. 2006; Rigaut et al. 2013;
Several progenitor systems have been proposed (see Branch & Wheeler 2017, for a more detailed review). A critical property is the mass of the WD at the explosion time, which is strictly connected to the explosion mechanism. In the classical model, when the WD mass attains the Chandrasekhar limit, the compressional heating of the whole star induces a central carbon ignition in highly degenerate conditions (Chandrasekhar-mass scenario). Basing on a 1D model, the outward burning front initially moves with subsonic velocity (deflagration). However, in order to obtain a good reproduction of the observed spectral evolution, a transition to a detonation should occur at a certain point (delayed detonation model, Khokhlov 1991). The physical mechanism that induces this transition is, however, unknown. Recently, explosions of sub-Chandrasekhar mass WDs, initially proposed to explain some subluminous events (Livne 1990; Woosley & Weaver 1994), have been suggested to represent a large part of the normal SNe Ia (Flörs et al. 2020). In these sub-Chandrasekhar WDs, the explosion is a consequence of He accretion onto a CO WD. It starts with a detonation of the external He-rich layer. Then, an inward shock moves toward the center triggering a carbon detonation. At the opposite, rotating WDs with a mass exceeding the nonrotating Chandrasekhar limit have been proposed to explain a few superluminous SNe Ia that imply a large amount of $^{56}$Ni (Howell et al. 2006; Hicken et al. 2007; Brown et al. 2014). However, rotation is a very common feature for WDs belonging to close-binary systems, and in turn, it should play an important role for the majority of the SNe Ia, rather than for just a minority.

Moreover, the nature of the companion object is still under debate: broadly speaking, it could be a normal star (Single Degenerate, or SD, scenario; Whelan & Iben 1973) or another WD (Double Degenerate, or DD, scenario; Iben & Tutukov 1984; Webbink 1984). In the SD scenario, the companion could have a H-rich envelope (classical SD), or it may be a He star (He-donor channel); while in the DD scenario, the companion could be another CO WD (classical DD) or a He WD (He-donor channel; Tutukov & Yungelson 1996; Yoon & Langer 2004; Solheim & Yungelson 2005; Wang et al. 2009). Other scenarios have been proposed. Examples are the merging of a CO WD with the CO core of an AGB companion star, during a common envelope episode (Core-Degenerate scenario; Iikov & Soker 2012), or violent mergers of two WDs (Pakmor et al. 2012, 2013).

In principle, the composition of the progenitors at the time of explosion, specifically the C/O ratio and the degree of neutronization, influences the explosive nucleosynthesis (Höflich et al. 1998; Hamuy et al. 2000; Domínguez et al. 2001; Timmes et al. 2003; Moreno-Raya et al. 2016; Piersanti et al. 2017). For instance, abundance ratios of intermediate-mass elements, like Si, and iron-peak elements in the material ejected by a SN may provide some hints on the neutronization degree. In this context, Badenes et al. (2008; see also Park et al. 2013) used the Mn to Cr abundance ratio to derived the average pre-explosive neutronization of the Tycho and Kepler SNe Ia, namely the following: $\tilde{\eta}_{\text{exp}} = 4.36 \times 10^{-3}$ and $4.55 \times 10^{-3}$, respectively. In principle, the neutronization of a WD should depend on the progenitor’s metallicity. Indeed, the composition of the C-O core of an intermediate-mass star is the result of both the H burning and the subsequent He burning. In the first evolutionary phase, the original CNO material is mainly converted into $^{14}$N. Later on, during He burning, $^{22}$Ne is produced through the chain $^{14}$N (α, γ)$^{18}$F (β+), $^{18}$O (α, γ)$^{22}$Ne. In practice, the final abundance (by number) of $^{22}$Ne is equal to the original C + N + O abundance (by number). Since for the major constituents, i.e., $^{12}$C and $^{16}$O, the contribution to $\eta$ is 0, the neutronization is mainly due to the final $^{22}$Ne abundance and, in turn, to the original C + N + O. On this base, the initial metallicity of the Tycho and the Kepler progenitors would have been as large as $Z = 3.5 \pm 4Z_{\odot}$, values that are much larger than the average metallicity of the galactic disk. This occurrence suggests that the average neutronization of the exploding WD should increase during the accretion phase. Indeed, due to the mass deposition, the WD density increases up to a few $10^{9}$ g cm$^{-3}$. At such a high density, $e$-captures are energetically favored with respect to the reverse processes ($/β$-decays), and for this reason, the neutronization is expected to rise up.

Piro & Bildsten (2008) and Chamulak et al. (2008) investigated such a possibility, showing that the average neutronization increases during the so-called simmering phase, i.e., the hydrostatic C-burning phase that precedes the dynamical breakout. In particular, they found that the larger the carbon consumption during the simmering, the higher the pre-explosive neutronization. Nevertheless, for $Z > Z_{\odot}$, they found that the $e$-capture contribution to the final $\eta_{\text{exp}}$ is small compared to the $Z$-dependent contribution, so that the high metallicity problem for the Tycho and Kepler supernova progenitors remains unsolved. Similar conclusion was also derived by Martínez-Rodríguez et al. (2016; hereinafter MR2016), while Piersanti et al. (2017; hereinafter P2017) found that the variation of the average neutronization during the accretion and the simmering phases increases as the initial metallicity of the progenitor increases, thus showing that the estimated $\eta_{\text{exp}}$ for the Tycho and Kepler SNe Ia are compatible with a progenitor’s metallicity of $Z \sim Z_{\odot}$, close to the average metallicity of the thin disk.

Schwab et al. (2017; hereinafter S2017) suggest that these “differences ... could arise due to differences in the net effect of the convective Urca process.” Actually, from a careful reading of all these theoretical studies, it appears that in addition to the URCA process; the various computations were carried out under quite different assumptions about the treatments of relevant physical processes, such as the convective mixing, the heat and the transports, and the nuclear network. As a matter of fact, a self-consistent treatment of all the physical processes operating during the last few thousand years prior to the explosion is still missing. The aim of the present work is to extend previous studies, by reviewing the treatment of all the relevant physical processes that may influence the final neutronization of the exploding WD. Then, in Section 2, we critically analyze the evolution of an accreting WD before and after the development of the convective core. We review, in particular, the properties of the convective URCA processes, paying attention to identify all the possible URCA pairs that may affect the thermal budget of the CO core and the extension of the convective region. Moreover, we investigate the effects of changing the numerical treatment of the convective

---

As usual, for a mixture of $^N$ isotopes, the neutronization is defined as $\eta = \sum_{A} \frac{Z_i}{A_i} (X_i - Z_i)$, where $X_i$, $A_i$, and $Z_i$ are the mass fraction, the atomic number, and the charge number, respectively, and $i = 1, ..., N$. 

---

Piersanti et al. 2020; Hakobyan et al. 2021; Kelsey et al. 2021). These occurrences appear in contrast with the supposed stellar amnesia.

---

THE ASTROPHYSICAL JOURNAL, 926:103 (18pp), 2022 February 10

---

---

2
instabilities, in particular, the mixing scheme and the criterion adopted to identify unstable zones. Eventually, we check the completeness of the adopted nuclear network. Basing on this analysis, in Section 3, we show that all the models currently available for the SNe Ia progenitors are inaccurate, and their predictions are not reliable. After discussing in Section 4 the effects of each energy contribution associated with URCA processes and their connection with convection, in Section 5, we present new models of accreting CO WDs up to the explosion, and in Section 6, we review the effects of the often neglected \(^{23}\)F(\(\alpha\),\(\gamma\)) URCA shell. Eventually, in Section 7, we discuss our updated predictions for the pre-explosive structures, and we present the new relation between progenitor metallicity and pre-explosive neutronization. Finally, in Section 8, we explore the dependence of the simmering phase on the initial mass of the accreting WD and of its cooling age as well, on the adopted accretion rate. Conclusions follow in Section 9.

In Appendix, we summarize all the models discussed in the present work and the setup adopted for their computation.

### 2. The Overall Evolutionary Scenario

According to the classical Chandrasekhar-mass scenario for SNe Ia progenitors, two distinct phases precede the explosion, namely the following: the accretion phase and the simmering phase. In the former, owing to the inverse relationship between mass and radius of a degenerate star, the density of the accreting WD increases until conditions for carbon ignition are attained at the center. The latter is instead characterized by a thermonuclear runaway, as usual for a nuclear burning in degenerate conditions, and the consequent development of an extended convective core, whose external border moves progressively outward.

A key process occurring, since the accretion phase, is the activation of cyclic weak interactions, in particular \(\beta\)-captures followed by \(\beta\)-decays. The former start at the center, when the density attains a critical value, typically, of a few \(10^9\) g cm\(^{-3}\). Then the reverse process, the \(\beta\)-decay, may eventually close the cycle. Each cycle, which is called URCA process, usually involves a pair of isobars. At first, a mother isotope, IS\(_1\), evolves into the daughter, IS\(_2\), according to the weak reaction IS\(_1')(A, Z) + \nu -\rightarrow IS\(_2')(A, Z - 1) + \nu^\prime\). Then, the reaction IS\(_2')(A, Z - 1) -\rightarrow IS\(_1')(A, Z) + e^- + \gamma\) transforms the daughter into the mother’s nucleus. The density for which the \(e^-\) capture on the mother nucleus and the \(\beta\)-decay on the daughter having the same probability to occur defines the location of an URCA shell. In practice, for each isobar pair, this threshold density \(\rho_{th}\) separates the more internal zone, where the \(e^-\) captures are favored, from the external one, where \(\beta\)-decays dominate. Indeed, the energy released by an \(e^-\) capture is as follows:

\[
E_{ec} = E_M - E_v + E_k
\]

where \(E_M = (M(IS_1) + m_e - M(IS_2)) \cdot c^2\) is the mass excess between reactants and products, \(E_v\) is the energy carried away by neutrinos, and \(E_k\) is the average thermal energy of the captured electrons.\(^7\) The latter is of the order of the electron Fermi energy that scales as \(\rho^{\alpha}\), with \(1/3 \leq \alpha \leq 2/3\). Since the first term is usually negative, the process becomes energetically feasible \((E_{ec} > 0)\) only if the Fermi energy is sufficiently large.

This occurrence implies high density. On the contrary, \(\beta\)-decays are hampered at high density, because of the Pauli suppression. More outside, where the density is lower, \(\beta\)-decays can occur, while \(e^-\) captures are suppressed, because of the negative \(E_{ec}\). In this case, the energy released by the \(\beta\)-decay is as follows:

\[
E_\beta = E_M - E_\nu - E_k
\]

where \(E_M = (M(IS_1) - m_e - M(IS_2)) \cdot c^2\), and the other terms have the same meaning as in Equation (1). Close to the Urca shell, both reactions are active, and the rate of energy production/subtraction is as follows:

\[
e_{urca} = N_\alpha \lambda_{ec} E_{ec} + N_\beta \lambda_{\beta} E_\beta
\]

where \(\lambda_{ec}\) and \(\lambda_{\beta}\) are the \(e^-\) capture and the \(\beta\)-decay rate, respectively, and \(N_\alpha\) is the Avogadro number.

During the accretion phase, the activation of these weak interactions has two major consequences. First of all, because of the electron captures, the average neutronization of the WD increases. On the other hand, at the URCA shell, there is a net energy loss due to the emission of \(\nu\)\(\gamma\) pairs. When an Urca shell first forms at the center, the neutrino/antineutrino energy loss causes a sharp decrease of the central temperature (see, e.g., Figure 2 in MR2016, Figure 1 in P2017 and Figure 1 in S2017, where the effects of the activations of the \(e^-\) capture on \(^{22}\)Na and \(^{25}\)Mg are shown). Then, as matter continues to be accreted, the mass coordinate of the URCA shell, as defined by the condition \(\rho = \rho_{th}\), moves outward. This progressive displacement of the URCA shell produces a uniform cooling of the core. Below the URCA shell, the abundance of the daughter nuclei increases, because of the \(e^-\) capture, until all the mother nuclei have been consumed. As a result, the matter inside a given URCA shell becomes \(n\)-rich. Once the maximum abundance of the \(n\)-rich isobar is attained, the \(e^-\) captures cease, and the core heating, as due to the WD compression, restarts.

As discussed in the extant literature, the most important isotopes involved in URCA processes during the accretion phase of SN Ia progenitors are \(^{23}\)Na, \(^{25}\)Mg,\(^{26}\)Ne, and the triplet \(^{26}\)Fe,\(^{26}\)Mn,\(^{26}\)Cr.\(^8\) Indeed, the abundances of the mother nuclei, in these cases, large enough to produce sizable effects on the WD thermal content (see, e.g., P2017).

Later on, as the accreting WD approaches the Chandrasekhar mass, it undergoes a rapid compression, until physical conditions for C-burning are attained. The C-ignition occurs near the center and proceeds mainly through the \(p\)-channel \(^{12}\)C\(^{(12}\)C,\(p\))\(^{23}\)Na and the \(\alpha\)-channel \(^{12}\)C\(^{(12}\)C,\(\alpha\))\(^{26}\)Ne, the relative contribution being \(0.44-0.45\) and \(0.56-0.55\), respectively (see, for instance, Caughlan & Fowler 1988). As the \(p\)-channel starts to release \(^{23}\)Na, it immediately undergoes an \(e^-\) capture, thus producing a further increase of the local neutronization. The contribution of these \(e^-\) captures to the total nuclear energy budget can be estimated according to Equation (1). Note that, since the Fermi energy of the electrons is quite large, the \(^{23}\)Na daughters are produced in excited states, and the energy released after the radiative decay exceeds that of the emitted neutrino. As a result, this process now produces a local heating. This is clearly shown in Figure 1, where we report for the \(^{23}\)Na\(^{26}\)Ne URCA shell the three contributions in Equation (3).

\(^7\) As usual, the difference of the thermal energy of the two ions is neglected.

\(^8\) In this case, two consecutive \(e^-\) captures convert the mother \(^{56}\)Fe into the daughter \(^{56}\)Cr, through \(^{56}\)Mn.
energy of the electrons should increase up to the local value of the Fermi energy. Therefore, the electron excess causes a cooling of the convective core. As already noted in Iben (1978), the electron abundance profile within the convective core critically depends on the mixing timescale, as compared to the local $e$-capture timescale. In order to take into account this energy contribution, Equation (3) should be modified as follows:

$$
\varepsilon_{urca} = \mathcal{N}_i \lambda_{ec} E_{ec} + \mathcal{N}_i \lambda_{\beta} E_{\beta} + \varepsilon_w, 
$$

where $\varepsilon_w$ is the negative contribution representing the amount of thermal energy absorbed when electrons are dredged down (see Section 5.1).

Last, but not least important, is the criterion adopted to define if a layer is stable or unstable against convection. In general, convection arises when the temperature gradient drops below a critical value. However, the molecular weight gradient may prevent convection (if negative) or may enhance the instability (if positive). According to the Ledoux criterion, a layer is stable against convection if the following is true:

$$
\nabla_{rad} < \nabla_{ad} + \delta \nabla_{\mu},
$$

where $\nabla_{rad} = (\partial \ln T / \partial \ln \rho)$ is the radiative gradient, $\nabla_{ad} = (\partial \ln T / \partial \ln \rho)_{\delta}$ is the adiabatic gradient, and $\nabla_{\mu} = (\partial \ln \rho / \partial \ln \rho)_{\mu}$ is the molecular weight gradient, $\phi = (\partial \ln \rho / \partial \ln T)_{\mu}$, and $\delta = (\partial \ln \rho / \partial \ln T)_{\delta}$. In case of a very efficient mixing, $\nabla_{\mu}$ vanishes, so that the Ledoux criterion reduces to the Schwarzschild criterion, i.e., $\nabla_{rad} < \nabla_{ad}$. During the simmering phase, this condition is satisfied when the border of the convective core remains well inside the $^{23}(\text{Na,Ne})$ URCA shell. On the contrary, when the convective-core border reaches the URCA shell, the rapid decay of the $n$-rich isobars conserves and reinforces the $\mu$-gradient, which represents a barrier to a further outward mixing. In practice, the growth in mass of the convective core stops when the convective border attains the location of the URCA shell (see Lesaffre et al. 2006, for a detailed and complete analysis of this process).

3. Critical Analysis of the Extant Literature

The analysis performed in the previous section gives us the possibility to critically review models for the simmering phase presented up to now (see Section 1), explaining the origin of the differences in the estimated neutronization at the explosion and evaluating their reliability. Piro & Bildsten (2008) provided the first evaluation of the neutronization evolution during the simmering phase. They started their computation from the onset of the simmering phase, ignoring the effects of URCA pairs/triplets in determining the physical conditions at the onset of central convection. However, they tried to account for such an effect by varying the ($\rho, T$) starting values of their simulation. Moreover, even if they consider the WD structure, they practically used a one-zone model, evaluating nuclear burning at the center of the star and using the Schwarzschild criterion to define the temperature gradient (assumed to be adiabatic). In this way, they were able to estimate the amount of $^{12}$C consumed during the evolution up to the explosion and, hence, the produced neutronization. In this regard, they assume...
that convective mixing is instantaneous. The adopted nuclear network is very short, including 9 isotopes linked by 7 reactions only (see their Table 1); in particular they include one \( p \)-capture \((^{12}\text{C}(p, \gamma)^{13}\text{N})\), one \( \alpha \)-capture \((^{13}\text{C}(\alpha, n)^{16}\text{O})\), and one \( n \)-capture \((^{12}\text{C}(n, \gamma)^{13}\text{C})\) only, which are relevant to evaluate the neutronization produced during the simmering phase but underestimate their contribution to the total energy budget. At the end, they assume that \( e \)-captures onto \(^{14}\text{N}\) and \(^{23}\text{Na}\) do not contribute to the energy budget. In particular, this assumption is equivalent to set \( \varepsilon_{\text{URCA}} = 0 \), because at the center of the CO WD during the simmering phase, density is definitely larger than the threshold value for the \(^{23}\text{(Na,Ne)}\) URCA shell so that the contribution from the \( \beta \)-decay of \(^{23}\text{Ne}\) could be neglected. As a matter of fact, Piro & Bildsten (2008) ignore any energy contribution from convective URCA processes, only suggesting that if these processes are at work they should produce a cooling of the convective core so that a larger C-consumption is expected (see their footnote 1).

On a general ground, the assumption that URCA processes do not contribute to the total energy budget during the simmering phase could be realistic, even if it cannot be proved a priori. Notwithstanding, the circulation via convective mixing of \(^{23}\text{Na}\) and \(^{23}\text{Ne}\) could produce local cooling and/or heating in the innermost zones where burning occurs, so that the C-consumption estimated by Piro & Bildsten (2008) could be completely unrealistic and, consequently, the neutronization at the explosion could be wrong.

Chamulak et al. (2008) used an approach similar to that of Piro & Bildsten (2008), even if they used a full nuclear network including 430 isotopes, and they computed the energy released by nuclear processes by including the mass term \( \varepsilon_{\text{M}} \), the neutrino term \( \varepsilon_{\nu} \), and the electron kinetic energy variation \( \varepsilon_{K} \) (see their Equation (2)). As a consequence, they found that the \(^{23}\text{(Na,Ne)}\) URCA process heats up the burning zones, because, as illustrated in Figure 1, for density values above \( \sim 2 \times 10^{9} \text{g} \cdot \text{cm}^{-3} \), the kinetic term dominates over the sum of the mass and neutrino terms \( \varepsilon_{\text{M,\nu}} \sim 1 \) \text{MeV} \). The neutronization value they estimated at the explosion epoch is roughly similar to that by Piro & Bildsten (2008), because both of the two working groups adopt similar assumptions, and in addition, the positive \( \varepsilon_{\text{URCA}} \) from the URCA pairs \(^{23}\text{(Na,Ne)}\) considered by Chamulak et al. (2008) represent a \( \sim 6\% \) correction to the effective energy delivered by the combustion of \(^{6}\text{Li}\) isotopes.

According to these considerations, the results by Chamulak et al. (2008) suffer the same limitation as the previous ones: the assumption that URCA pairs provide a heating of the convective core cannot be proved, and in addition, the role of convective URCA processes is ignored, as they cannot be included in one-zone models. In this regard, the authors observed that the large abundance of the \(^{23}\text{(Na,Ne)}\) URCA pairs could have a “dramatic effect on the properties of the convection zone.”

MR2016 computed full evolutionary models of CO WD accreting matter, by considering different accretion rates, different initial thermal contents of the accreting WD (i.e., different initial central temperatures), and different initial metallicity of the main-sequence WD progenitors. They adopted a nuclear network including 48 isotopes (up to sulfur) constructed by taking into account the results on the detailed nucleosynthesis computed in Chamulak et al. (2008). In their computation, the equations describing the physical structure and the evolution of chemical species are solved simultaneously, accounting for the effects of convective mixing by means of a diffusion equation. The energy contribution related to URCA processes are computed according to Equation (5), by setting \( \varepsilon_{W} = 0 \). As they use a full evolutionary model, MR2016 can resolve the whole temperature and density profile during the accretion and the simmering phase, so that they can evaluate the contribution of URCA processes to the local and total energy budgets of accreting WDs. Moreover, as convective mixing is directly included in the computation of models, they can also evaluate the effects of convective URCA processes on the neutronization at the explosion epoch. They show that, during the accretion, the center of the star and, then, the innermost zones cool down due to the activation of URCA shell involving \(^{23}\text{Mg}\) and \(^{23}\text{Na}\). They found, also, that the weak processes involving \(^{23}\text{Na}\) heat up the innermost zones of the accreting WD from the epoch of C-ignition and up to when the C-burning driven convection attains the \(^{23}\text{(Na,Ne)}\) URCA shell. Moreover, as also detailed in S2017, they found that the mixing of processed matter through this URCA shell becomes an efficient mechanism to refurbish the innermost burning zone with fresh \(^{23}\text{Na}\) and that the consequent \( e \)-captures on this isotope represent a source of energy as important as carbon fusion reactions. The net result is that the amount of carbon consumed to trigger the explosion is lower than what was estimated in one-zone models above, and hence, the neutronization at the dynamical break-out results is lower.

As already remarked, MR2016 assumed that the energy subtracted from the surroundings to carry down \( e \)-rich material from the URCA shell is negligible, an assumption that seems unrealistic, at least according to S2017, who computed a posteriori the total energy associated to such a process and demonstrated that it is as large as the energy losses via neutrino emission in weak processes. Moreover, MR2016 did not mention what criterion they used to establish if a layer is unstable for convection, even if it can be safely assumed that they use the Swartzschild criterion (Martinez Rodriguez 2019). As a consequence, they did not include the effects of the possible molecular weight gradient on the properties of the convective zone.

P2017 performed an analysis similar to that of MR2016, even if they used a different evolutionary code and a different nuclear network. They considered only one WD mass and a single value of the accretion rate and performed computations for three different initial metallicities of the main-sequence progenitor of the WDs. As in MR2016, the physical structure equations and those describing the evolution of chemical species are solved simultaneously, by treating the convective mixing as an advective process. In computing the evolution of accreting WDs, they set \( \varepsilon_{K} = \varepsilon_{W} = 0 \). The adopted assumptions determine the same temperature and density profiles as in MR2016 during the accretion phase. In fact, as demonstrated above, for preexisting URCA isobars, at the URCA shell, the mass and kinetic energy contributions from mother and daughter nuclei have the same absolute value and cancel each other, the neutrino/antineutrino sink remaining the only energy contribution. On the other hand, when carbon starts to be burnt in the innermost zones, P2017 estimated a negative energy contribution from the \(^{23}\text{(Na,Ne)}\) URCA process in the burning region, while it should be positive (see Figure 1); so they overestimated the amount of \(^{12}\text{C}\) consumed and, hence,
the final neutronization. At the end, during the simmering phase, their assumption about the energy contribution from the $^{32}$Na,Ne,Ne URCA process is equivalent to assume that the energy subtracted from the surroundings to carry downward e-rich material is equal to the electron kinetic energy released in e-captures. Such an assumption could be valid on average, inside the convective core, but it is not locally valid. Moreover, such an assumption assumes that the net results of URCA processes is a cooling of the accreting WD, whose effects are mimicked by arbitrarily reducing the amount of energy delivered in the burning region. As a result, they find definitely larger neutronization values at the explosion epoch and, in addition, that the amount of $^{12}$C consumed during the simmering depends on the initial metallicity of the WDs progenitors. These results are the direct consequence of the assumption concerning the energy contribution of URCA processes. In fact, the larger the initial abundance of URCA isobars, the larger the cooling effects during the accretion phase and, therefore, the larger the density value at which C-burning occurs. Hence, due to the fact that $\varepsilon_n$ does depend mainly on density, the larger the initial $Z$, the more efficiently the neutrino cooling and, hence, the larger C-consumption and final neutronization.

4. Preliminary Analysis

According to previous considerations, it comes out that all the models in the extant literature for the simmering phase of SNe Ia are not accurate in reproducing the physical and chemical processes at work in real accreting WDs. In fact, all of them are based on oversimplified assumptions that cannot be proved a priori and that, in some cases, are incorrect. The previous analysis has remarked once again the evidence that the neutronization level at the explosion epoch depends mainly on the energy contribution of URCA isotopes during the simmering phase that depends not only on local properties but also on the outward and inward circulation of $n$-rich and e-rich material, respectively; the efficiency of the latter process depends critically on the possibility that the convective zone could encompass the $^{32}$Na,Ne URCA shell.

In the following, we present models including all the energy term in Equation (3), by paying particular attention to the role played by the formation of a strong $\mu$-gradient at the URCA shell and to its effects on the physical and chemical properties of the zone unstable for convection. In our simulation, we make use of the FuNS code as in P2017 (Straniero et al. 2006; Cristallo et al. 2009). The nuclear network we adopt is slightly different from that in P2017, and it includes 200 nuclear processes linking 50 isotopes, namely $p$, $\alpha$, $n$, C(12–14), N(13–15), O(16–20), F(18–22), Ne(20–24), Na(22–25), Mg(23–28), Al(25–28), Si(27–32), P(30–32), $^{56}$Mn, $^{57}$Fe, and $^{58}$Fe. With respect to P2017, we neglect isotopes above phosphorus, because according to the extant literature (e.g., see Chamulak et al. 2008; Piersanti et al. 2017), they are scarcely produced during the simmering phase. In doing so, we also neglect the contribution by the $^{32}$Si,Si,P, $^{37}$S,Cl, and $^{39}$Ar,K URCA processes, because either their abundances are very low or they occur at densities so low that they provide a negligible contribution to both the total neutronization and the energetic of the accreting WD (see Table 1 in Piersanti et al. 2017). The only exception is the URCA triplet $^{32}$Si,Si,P whose contribution to the neutronization along the whole accreting WD is $\Delta\eta_C \sim 2 \times 10^{-4}$. On the other hand, we include the $^{28}$Si,Al,Mg and the $^{24}$Mg,Na,Ne URCA triplets that play an important role in determining both the ignition conditions and the neutronization of the accreting WDs. Reaction rates for these two URCA triplets are derived from Suzuki et al. (2016), while those for all the other URCA processes considered in the nuclear network are included as detailed in Paper I (see Section 2 and Table 1). Reaction rates for charged particle interactions are derived from the JINA REACLIB database (Cyburt et al. 2010).

To illustrate how the $^{28}$Si,Al,Mg and the $^{24}$Mg,Na,Ne URCA triplets affect the evolution during the accretion and the simmering phases, we compute a set of test models by adopting the same network as in Piro & Bildsten (2008), but including also the $^{23}$Mg,Na) and $^{56}$Fe,Mn,Cr) URCA processes. The energy contribution related to URCA processes is implemented according to Equation (3) during both the accretion and the simmering phase, and the convective mixing is modeled as a diffusive process, so that the equation describing the physical structure and the evolution of chemical species can be solved simultaneously. The initial WD structure is the same ZSUN model in P2017 (i.e., a CO WD with $M = 0.817 M_{\odot}$, whose progenitor star had solar metallicity $Z_{\odot} = 1.38 \times 10^{-2}$ - see P2017). CO-rich matter is accreted directly at $\dot{M} = 10^{-7} M_{\odot}$ yr$^{-1}$. The results relative to this reference model (Model K00—solid black line) are displayed in Figure 2, where we plot the evolution of the center of the accreting WD in the $\rho-T$ plane (upper panel) and the increase of neutronization at the center $\Delta\eta_C$ as a function of the central density (lower panel). This model is similar to those presented in MR2016 ($Z = Z_{\odot}$ and $M = 10^{-7} M_{\odot}$ yr$^{-1}$) and in S2017, and has a quite similar evolution (for instance compare the lower panel in Figure 2 and Figure 2 in S2017), even if the increase of the central neutronization at the epoch of dynamical breakout is larger (see also Table 1). This is due partly to the inclusion in Model K00 of the $^{56}$Fe,Mn,Cr) URCA triplets that produce a variation of the neutronization of $\Delta\eta_C = 2X(^{56}\text{Fe})/A(^{56}\text{Fe}) \simeq 3.75 \times 10^{-5}$ where we use $X(^{56}\text{Fe}) = 1.05 \times 10^{-3}$ (see Table 1 in P2017). The remaining discrepancy ($\Delta\eta_C \simeq 1.56 \times 10^{-4}$) has to be ascribed to the different nuclear network adopted, in particular to the fact that

![Figure 2](image-url)
the very short network used here largely underestimates the energy contribution from the $p$- and $\alpha$-capture and, hence, a larger amount of $^{12}$C has to be burnt in order to attain the explosion.

In computing Model K28, we include in the nuclear network also the $^{28}$(Si,Al,Mg) URCA triplets. We recall that the $e$-capture on $^{28}$Al has a threshold density of $\rho_{th} \approx 0.89 \times 10^9$ g cm$^{-3}$, while for $^{28}$Si it is $\rho_{th} \approx 4.24 \times 10^9$ g cm$^{-3}$. This means that when the physical conditions suitable for $e$-capture on $^{28}$Si are attained in the daughter nucleus $^{28}$Al immediately at the capture of an electron, forming $^{28}$Mg. As a matter of fact, we find that, when the central density of the accreting WD exceeds $4.17 \times 10^9$ g cm$^{-3}$, already 10% of the initial local $^{28}$Si have been converted into $^{28}$Mg. At that epoch, the chemical potential of the electrons, or the kinetic energy of electrons, involved in $e$-capture is equal to $\varepsilon_K \sim 6.011$ MeV while the mass excess energy term for the $^{28}$Al($e^-, \nu$)$^{28}$Mg and the energy of the emitted neutrino are $\varepsilon_M = -1.832$ MeV and $\varepsilon_K = 1.981$ MeV, respectively. This means that the activation of $^{28}$(Si,Al,Mg) URCA process determines a rapid heating of the innermost zone of the accreting WD, as it can be clearly seen in Figure 2. Due to the larger central temperature, $C$-burning proceeds at a higher rate, so that the innermost zones of the accreting WDs become unstable for convection at a lower density (see Table 1). After the onset of convection, the $^{28}$(Si,Al,Mg) URCA triplet does not contribute anymore to the energy budget during the simmering phase, because the density threshold to activate the $\beta$-decay of $^{28}$Mg is too low. The conversion of the initial $^{28}$Si into $^{28}$Mg produces an increase of the local neutronization at the center of $\delta_n = 2X(\varepsilon_{28}Si)/AX(\varepsilon_{28}Si) \approx 5.21 \times 10^{-5}$. This explains the larger $\Delta\eta_C$, with respect to Model K00, obtained at the epoch when convection attains the $^{23}$(Na,Ne) URCA shell. However, since the accreting WD is globally less dense, a smaller amount of carbon has to be nuclearly processed in order to trigger the explosion. As a consequence, the value of central neutronization at the epoch of dynamical breakout is larger by only $\sim$4% than that in the K00 model.

The situation is quite different for the K24 model, which includes the $^{24}$(Mg,Na,Ne) URCA triplet. In this case, the density threshold for $e$-capture on $^{24}$Na ($\rho_{th} = 5.07 \times 10^9$ g cm$^{-3}$) is slightly larger than that for the $e$-capture on $^{24}$Mg ($\rho_{th} \approx 4.16 \times 10^9$ g cm$^{-3}$). As a consequence, the activation of the $^{24}$(Mg,Na) processes does not produce any sizable effects on the physical conditions at the onset of convection, but the whole $^{24}$(Mg,Na,Ne) URCA triplet behaves as a heating process of the innermost zone of the accreting WD at the beginning of the simmering phase, due to the circulation of this isotopes forced by convective mixing. As a whole, Model K24 results less dense for a fixed temperature value, while the contribution of the considered URCA triplets to the central neutronization is $\delta_n = 2X(\varepsilon_{24}Mg)/AX(\varepsilon_{24}Mg) \approx 0.39 \times 10^{-5}$. Also in this case, as discussed above for the $^{28}$(Si,Al,Mg) URCA triplet, a smaller amount of $^{12}$C has to be burnt to produce the explosion as the structure is globally less dense. As a consequence, the central neutronization at the breakout epoch is practically the same as in Model K00 (only $\sim$0.7% lower).

The other important difference with respect to P2017 is the treatment of convective mixing as a diffusive process, whereas in that paper it was modeled according to an advective scheme. S2017 partially analyze the effects of different scheme on the neutronization level at the epoch of the explosion. To this aim, they assume that the "advective scheme" in P2017 corresponds to the mixing algorithm used in the FuNS code as described in Straniero et al. (2006). Such an algorithm was originally introduced by Sparks & Endal (1980), successively corrected by Chiefi et al. (1998), and linearized in the form currently available in the FuNS code by Straniero et al. (2006). Hereinafter we refer to this mixing algorithm as Sparks-Endal-Linearized (SEL). Such an algorithm evaluates the local variation of chemical composition via convective mixing as due to the effective coupling of a given layer with all the others in the convective-unstable zone. As a consequence, the SEL mixing scheme cannot be implemented and solved together with the equations describing the physical structure and nuclear burning along the star, and for this reason, the operator splitting technique has to be employed. S2017 found that the central neutronization at the epoch of dynamical breakout is a factor $\sim$2 larger in the model based on the SEL scheme and explained such a discrepancy as a consequence of the different mixing efficiency before the occurrence of the mixing freeze-out (Nonaka et al. 2012).

We remark that the mixing algorithm used in P2017 is not the SEL scheme assumed by S2017, but it is based on the assumption that the transport of matter by both the forward and the downward convective fluxes are regulated by an advective equation, namely the following:

$$\frac{\partial X}{\partial t} + 4\pi r^2 \rho v \frac{\partial X}{\partial m} = 0 \quad (7)$$

where $m$ and $t$ are the mass coordinate and time, while $\rho(m,t)$, $v(m,t)$, and $X_j(m,t)$ are the density, average convective velocity, and mass fraction of the isotope $j$, respectively. As in the case of diffusion, also the advective scheme can be arranged in order to solve simultaneously all the relevant equations describing the physical and chemical properties of the accreting WD. Nevertheless, advection and diffusion describe different physical processes. In fact, diffusion often refers to microscopic transport of particles (or heat), while advection describes a bulk transport of mass (or heat). In general, convection combines diffusion and advection, but the vertical flow, as driven by the thermal gradient, is mainly due to advection.

| Model     | Model K00 | Model K28 | Model K24 |
|-----------|-----------|-----------|-----------|
| $\rho_{th}$ ($10^9$ g cm$^{-3}$) | 4.105 | 4.077 | 4.070 |
| $T_{th}$ (10$^8$ K) | 1.732 | 1.728 | 1.727 |
| $\rho_{th}$ ($10^9$ g cm$^{-3}$) | 4.495 | 4.187 | 4.459 |
| $T_{th}$ (10$^8$ K) | 1.819 | 1.751 | 1.818 |
| $\Delta M(12C)$ (10$^{-2}$ M$_\odot$) | 1.324 | 1.313 | 1.233 |
| $M_{\text{max}}(\text{M}_\odot)$ | 1.213 | 1.213 | 1.213 |
| $\rho_{th}$ ($10^9$ g cm$^{-3}$) | 3.704 | 3.560 | 3.667 |
| $\Delta\eta_C(10^{-3})$ | 0.544 | 0.563 | 0.540 |

Note. $\rho_{th}$ and $T_{th}$ central density and temperature at C-ignition; $\rho_{th}$ and $T_{th}$ central density and temperature at the onset of convection; $\Delta M(12C)$ amount of $^{12}$C consumed via nuclear burning from the onset of mass accretion up to the explosion; $M_{\text{max}}(\text{M}_\odot)$ maximum extension of the convective zone; $\Delta\eta_C$ variation of the neutronization level at the center from the beginning of the accretion process up to the explosion.
Currently, a self-consistent description of turbulent convection is still missing. The choice of the scheme used to describe this process is a critical issue of the extant models of SNe Ia progenitors. Indeed, different mixing algorithms produce different mixing efficiencies, and during the simmering phase, the energy contribution from the URCA processes is taken into account according to Equation (5), whose value may be positive or negative, depending on the circulation efficiency of the URCA isobars.

To investigate in detail such an issue, we compute a second set of test models, by assuming the same short network as in Model K00 above, but varying the mixing algorithm. In particular, we compute models by using in the modelization of convective, mixing the SEL, advective, and diffusive schemes (models SEL, ADd, and DFD, respectively), but in all three cases, we apply the operator splitting technique and compute the effect of mixing after each time-step integration. At the end, we also compute a model by using the advective scheme and solving simultaneously all the equations describing the evolving WD (Model ADV). The results are displayed in Figure 3, and some relevant quantities of the computed models are listed in Table 2. Our computations confirm previous results from S2017 that the SEL mixing scheme predicts, at the explosion epoch, values for $\eta_C$ definitively larger with respect to Model K00; at the same time, we also found that Model DFD has a slightly larger ($\sim$7%) value of $\eta_C$. Moreover, Figure 3 suggests that the evolution of all decoupled models (SEL, DFD, and ADd) after convection has attained the $^{23}$(Na,Ne) URCA shell, which is quite different from that of Model K00. This clearly indicates that the operator splitting technique used to compute the effects of convective mixing underestimated the effects of heating related to the circulation of URCA isobars, because their energetic feedback is not consistently taken into account and, hence, automatically predicts larger neutronization values. The differences existing in the final value of $\eta_C$ among these models with different mixing schemes have to be ascribed to the different mixing efficiencies, the diffusive approach having the highest one, and the advective approach having the lowest one, respectively.

The evolution of Model ADV, i.e., with convective mixing modeled according to Equation (7) that is solved contemporarily to all the other equations describing the physical structure and nuclear burning, is completely different. In fact, after central convection has attained the $^{23}$(Na,Ne) URCA shell, due to the low mixing efficiency, the dredged-down $^{23}$Na undergoes e-capture during the inward motion well before it could attain the center of the accreting WD. This produces a local heating at the mass coordinate $M_{sp} \approx 0.295 M_\odot$, and very soon, the convective core splits into two convective regions: an innermost one, extending from the center to $M_{sp}$ and an outermost one up to the $^{23}$(Na,Ne) URCA shell. In the outer convective shell, the circulation of URCA isobars continues to produce an efficient heating, while the C-burning rate progressively increases close to $M_{sp}$ as the local temperature increases, so that the neutronization in this zone remains practically constant. As temperature increases at $M_{sp}$, the convective shell encompasses the $^{23}$(Na,Ne) URCA shell and progressively engulfs larger and larger portions of the accreting WD. On the other hand, in the convective core, the circulation of URCA isobars does not produce any energy contribution, because the outer border of the convective core is well confined inside the corresponding URCA shell. As a consequence, $\eta_C$ increases very slowly, as due to C-burning and subsequent production of $^{23}$Na. When the timescale for e-capture on $^{23}$Na becomes longer than the local heating timescale at $M_{sp}$, neutronization in the convective shell starts to increase due to e-capture on $^{14}$N. Therefore, at the explosion epoch, both the temperature and neutronization profiles for models exhibit a peak at $M_{sp}$, as displayed in Figure 4. Such a result is a consequence only of the mixing efficiency related to the advective mixing scheme. As a test, we arbitrarily increase by a factor of 50 the convective velocity inside the convective- unstable zone, and we find that the original convective core splits at the mass coordinate $M_{sp}^{22} \approx 0.21 M_\odot$, thus confirming the effects of the adopted mixing scheme (i.e., the adopted mixing efficiency) on the physical and chemical properties of the simmering phase.

5. Next Generation Models

5.1. Inward Motion of E-rich Material

We continue our analysis of the simmering phase of SNe Ia by using, as a case study, the model that ZSUN introduced in a previous section, the model accreting CO-rich matter at $M = 10^{-7} M_\odot$ yr$^{-1}$. We adopt the diffusive scheme for convective mixing, and we use the full nuclear network (including 52 isotopes linked by 200 nuclear processes), except the weak processes associated with the URCA $^{21}$(F,Ne) isobars. We compute first a reference model (label “REF”) by including the energy contribution from URCA isotopes according to Equation (3). Then we compute a second model by including the cooling effects due to the inward motion of e-rich material in zones unstable for convection according to Equation (5). As already recalled, the importance of the contribution $\xi_W$ was first discussed by Iben (1978) who suggested that it should be proportional to the effective inward flux of electrons and to the variation of the electronic chemical potential $\mu_e$, i.e., the Fermi energy, along the inward motion. The formula is as follows
Table 2

| Model   | Model K00 | Model SEL | Model DFd | Model ADd | Model ADV |
|---------|-----------|-----------|-----------|-----------|-----------|
| \(M_{\text{conv}}^{\text{max}}\) (M\(_e\)) | 1.213     | 1.241     | 1.211     | 1.209     | 1.258     |
| \(\Delta \eta_{\text{c}}(10^{-3})\) | 0.544     | 1.702     | 0.584     | 2.081     | 0.211     |
| \(\pi_{\text{exp}}(10^{-3})\) | 0.322     | 0.403     | 0.355     | 0.417     | 0.287     |
| \(\rho_{\text{exp}}(10^9\text{g cm}^{-3})\) | 3.704     | 3.649     | 3.729     | 3.810     | 1.942     |
| \(\Delta M^{(12}\text{C})(10^{-2} M_e)\) | 1.324     | 1.550     | 1.478     | 1.596     | 1.112     |

Note. \(M_{\text{conv}}^{\text{max}}\): maximum extension of the convective zone; \(\Delta \eta_{\text{c}}\): variation of the neutronization at the center from the beginning of the accretion process up to the explosion; \(\pi_{\text{exp}}\): neutronization at the explosion averaged over the convective zone \(M_{\text{conv}}^{\text{max}}\); \(\rho_{\text{exp}}\): density at the explosion epoch; \(\Delta M^{(12}\text{C})\): amount of \(^{12}\text{C}\) consumed via nuclear burning from the onset of mass accretion up to the explosion.

Figure 4. Temperature (upper panel) and neutronization (lower panel) profile for Model ADV at the explosion epoch. For comparison the same quantities are also plotted for Model K00 (dashed red line).

(see also Equation (8) in Iben 1978):

\[
\dot{\varepsilon}_W = -\frac{\partial \rho_e \Phi_e(M)}{\partial M}
\]  

where \(\Phi_e(M)\) is the inward flux of electrons through a sphere of mass \(M\), and it is defined as the following:

\[
\Phi_e(M) = -N_A \int_0^M \left( \frac{dY_e}{dt} - \frac{\delta Y_e}{\delta t} \right) dm
\]

(9)

where \(N_A\) is the Avogadro’s number, \(\frac{dY_e}{dt}\) is the local variation of \(Y_e\) as determined by weak processes, and \(\frac{\delta Y_e}{\delta t}\) is the mass-averaged (and therefore actual local) rate of change of the electron abundance in the core as determined by both convection and nuclear burning.

As we directly couple the solution of all the equations describing the physical and chemical structure of the accreting WD, we can release the assumption that the chemical composition in the convective core is homogeneous, so that, in implementing Equation (9), we can include directly the local variation of the free electrons number:

\[
\Phi_e(M) = -N_A \int_0^M \left( \frac{dY_e}{dt} - \frac{\delta Y_e}{\delta t} \right) dm
\]

(10)

where \(\delta t\) is the actual timestep on which the new structure is integrated. S2017 correctly observed that, when the effects of convective mixing are included in the iteration procedure, the argument of the integral, in Equation (9), reduces to the mixing term only; so by using the diffusion scheme to model convective mixing, it becomes the following:

\[
\Phi_e(M) = N_A (4\pi r^2 (M) \rho (M))^2 D(M) \frac{\partial Y_e}{\partial \rho} \bigg|_M
\]

(11)

where \(D(M)\) is the diffusion coefficient at \(M\).

We verified that both the evaluations for \(\Phi_e(M)\) in Equation (10) and (11) produce the same results when the \(\varepsilon_W\) energy term is evaluated a posteriori after the convergence of the model at each timestep. However, when we try to incorporate Equation (11) in the model computation, we found that this formulation largely overestimated the inward electron flux, so that the corresponding energy contribution \(\varepsilon_W\) becomes larger and larger, making the computation impossible. On the other hand, when we use Equation (10), we are able to perform the integration of the models. The reason for such an occurrence is not related to inconsistent assumptions in deriving the above formulation but only to how Equation (11) has been implemented. In fact, in our code, the border of convective-unstable zones and the mixing efficiency there (i.e., the diffusion coefficients) are determined at the beginning of the iteration procedure and then are maintained constant. This procedure is commonly adopted in the computation of stellar models to avoid numerical fluctuations during the integration procedure (see, for instance, Section 6.2 in Paxton et al. 2011). In the computation of \(\Phi_e(M)\), however, such a procedure introduces an inconsistency, as the mixing term at the beginning of a new iteration step is overestimated with respect to the previous converged solution. This occurrence is particularly relevant at the external border of the convective core where the new zones radiative at a previous time become now unstable for convection. As a consequence, the local inward electron flux is very large (in the previous integrated model the mixing efficiency there was zero), and this drives the model very far from the convergence domain.

In his seminal work, Iben (1978) found that when the border of the convective core approaches the \(^{23}\text{Na,Ne}\) URCA shell,
the accreting WD undergoes thermal oscillations. This is the consequence of the rapid increase of the neutrino/antineutrino emission from the $^{23}$Na,Ne URCA pairs due to both the increased abundance of these isotopes (they are produced by C-burning) and the increased density at the center that determines a larger cross section for these weak processes. When the cooling effects related to URCA processes become the dominant energy losses in the convective core, thermal oscillations start and proceed through a sequence of alternate cooling and heating phases. In the former, the URCA neutrino/antineutrino cooling is larger than the heating via nuclear burning and release of the kinetic energy of electrons from e-capture; as a consequence, the temperature in the whole convective core decreases, the mass of the convective-unstable zone reduces, and the structure undergoes a compression. In the latter phase, due to the decrease of URCA neutrino/antineutrino cooling, heating becomes the dominant process, so that the temperature in the core rises again, and the convective-unstable region grows in mass again.

Our results are displayed in Figure 5, where we report, as a function of the central temperature, the evolution of the density (upper panel) and neutronization (lower panel) at the center. The model implementing Equation (5) is labeled “WRK” (solid black line); for comparison we also plot data for the model REF (long-dashed heavy green line). As it can be seen in the upper panel of Figure 5, thermal oscillations do not occur; or better, their amplitude is very small, even if the general trend reported by Iben (1978; slow increase of central temperature and rapid compression of the whole core) is confirmed. Such an occurrence is a direct consequence of the very high resolution in mass adopted in our computation (meshes in the adopted mass grid have mass extension $\Delta M$ lower than $5 \times 10^{-4} M_e$), and this is confirmed by comparing Model WRK with the W00 model (blue line in Figure 5), which was computed by adopting a very coarse mass grid ($\Delta M \gtrsim 10^{-2} M_e$). As an additional test, we computed Model W06 (red line in Figure 5) by adopting a very fine mass grid in the innermost 0.6 $M_e$ zone ($\Delta M \lesssim 5 \times 10^{-4} M_e$) of the accreting WD and a coarse one outside ($\Delta M \gtrsim 10^{-2} M_e$). As it can be seen, when the mass of the convective core exceeds 0.6 $M_e$, a huge thermal oscillation occurs. Hence, we conclude that the characteristics of thermal oscillations depend on the adopted mass grid for the computation not only close to the $^{23}$Na,Ne URCA shell but in the whole convective core.

Iben (1982) investigated also the behavior of CO WD accreting H-rich matter; as in his previous work devoted to analyzing the effects of URCA processes in the evolution of CO core undergoing C-burning, he stopped the computation well before the final outcome could be assessed. Notwithstanding, he suggested that thermonuclear runaway triggered by C-burning at the center cannot be limited by the cooling effects at the URCA shell, so that the net results, of including the cooling effects produced by inward convective mixing of e-rich matter, are only a delay of the dynamic explosive event. Our simulation (Model WRK in Figure 5) confirms such a guess, as we obtain that, during the phase of slowly increasing temperature, the total luminosity above the $^{23}$Na,Ne URCA shell progressively increases from negative values up to zero. In this condition, convective-unstable core can encompass the URCA shell, and the heating at the center rapidly accelerates up to the explosion. However, we also found that during the thermal oscillation phase, due to the reduced efficiency of heating via circulation of e-rich material, the evolution is mainly driven by nuclear burning of $^{12}$C; this implies a large production of $^{23}$Na and, hence, a large increase of neutronization due to the $^{23}$Na(e, $\nu$)$^{23}$Ne weak process. This is clearly displayed in the lower panel of Figure 5, where we report the variation of the neutronization at the center. As it can be seen, during the thermal oscillation phase, $\eta$-rapidly increases, while later on, it remains almost flat, and then it decreases due to the fact that convective core encompass the URCA shell, and the timescale for e-capture on $^{23}$Na becomes shorter that the local heating timescale. As a matter of fact, we find that, at the dynamical breakout, Model WRK has $\rho_{\text{exp}} = 5.47 \times 10^9$ g cm$^{-3}$, $M_{\text{max conv}} = 1.065 M_e$, $\Delta M^{12}C = 4.83 \times 10^{-2} M_e$, $\Delta M_{\text{c}} = 42.86 \times 10^{-3}$, a factor 8 larger with respect to Model REF.

5.2. Criterion for Convection

As it is well known, stellar matter undergoing nuclear reactions evolves progressively from light isotopes to heavy ones, determining a continuous increase of the molecular weight $\mu$. If nuclear burning occurs in a convective zone, as in the convective core of accreting WDs during the simmering phase, at the borders of this zone, a gradient of molecular weight $\nabla_{\mu}$ occurs, and its local value can be assessed. Notwithstanding, a density gradient and the position of the borders of the layers unstable for convection should be evaluated not on the basis of the Schwarzschild criterion but according to the Ledoux criterion in Equation (6). As the ratio $\phi/\delta$ is positive, the existence of a $\mu$-gradient acts as a limiting factor in the definition of the extension of the convective zone. If the layers close to the border of the convective zone do not fulfill the Schwarzschild criterion $\nabla_{\mu} \lesssim \nabla_{\text{ad}}$, the borders of a convective zone defined according to the Ledoux criterion are secularly unstable, as a slow partial mixing through them produces the reduction of the local $\nabla_{\mu}$ value, thus allowing convection to encompass the borders themselves.

However, at the $^{23}$Na,Ne URCA shell, the situation becomes more complex for two reasons: (i) the $n$-rich isotope...
implementing the Ledoux criterion the convective core stops at the $^{23}$(Na,Ne) URCA shell. This determines a change in the slope $d \log T / d \log P$, causing a more rapid increase of the central temperature as compared to Model REF. During further evolution, the mass of the convective core remains constant ($M_{CC} \simeq 0.449 M_\odot$); this limits the circulation of e-rich matter downward and, hence, the $\varepsilon_K$ energy contribution due to e-capture on $^{23}$Na. As a consequence, a larger amount of carbon is consumed at the center to drive the accreting WD at the dynamical breakout. This determines a large increase of neutronization at the center and in the whole convective core. The further evolution does not present any particular difference, up to when the heating timescale at the center becomes shorter than that for e-capture on $^{23}$Na at the center; hence, the increase of neutronization is mainly produced by the conversion of $^{13}$N into $^{13}$C (see the change in the slope of the curves in both panels in Figure 6 at $T_C \approx 6.4 \times 10^8$ K). We find that, at the dynamical breakout, Model LED has $\rho_{\text{env}} = 4.32 \times 10^9$ g·cm$^{-3}$, $M_{\text{max}}^{\text{env}} = 0.449 M_\odot$, $\Delta M(12C) = 3.63 \times 10^{-3}$, $\Delta M_C = 1.097 \times 10^{-3}$, a factor 4.4 larger with respect to Model REF.

We repeat the computation of Model LED by changing the value of the $\alpha_{\text{OS}}$. We find that the obtained results do not change for values in the range $10^{-7} \leq \alpha_{\text{OS}} \leq 1$, clearly indicating that the local cooling at the $^{23}$(Na,Ne) URCA shell determined by neutrino/antineutrino emission prevents any upward motion of the convective eddies, thus stopping the growth in mass of the convective core. We compute additional models by applying an overshooting at the border of the convective core. We consider an exponential decreasing profile of either the convective velocity or the Eulerian diffusion coefficient, and we vary the length-scale adopted in the computation. In all the considered cases, we found that the convective core is unable to encompass the URCA shell.

The analysis performed above clearly suggested that both the inclusion of the cooling of the surroundings produced by e-rich matter carried downward by convection and the use of the Ledoux criterion to define the border of zones unstable for convection. In the layers close to the convective zone, where it results in $\nabla_L < \nabla_T < \nabla_{\text{ad}}$ (semiconvective layers), we assume that the local convective velocity is equal to

$$v_{\text{OS}} = \alpha_{\text{OS}} \cdot v_c$$

where $v_c$ is the value of the convective velocity computed via the mixing length theory according to the Schwarzschild criterion, and $\alpha_{\text{OS}}$ is a parameter whose value has been varied to study the dependence of the obtained results on the mixing efficiency in the semiconvective zones. Once $v_{\text{OS}}$ has been determined, we recompute consistently the value of $\nabla_T$. We want to remark that, on a computational point of view, the determination a priori of the real extension of the semiconvective region requires an iterative procedure. In fact, usually a molecular weight gradient occurs only at the border of the convective-unstable region. As a consequence, partial convective mixing in progressively larger zones has to be tested to individuate the real extension of a semiconvective zone. This was first pointed out by Castellani et al. (1971), and successively remarked also by Langer et al. (1983).

Our results, obtained by assuming $\alpha_{\text{OS}} = 10^{-5}$, are displayed in Figure 6 where we report the same quantities as in Figure 5, but for Model LED as compared to Model REF. The evolution of the two models is identical up to when the convective core approaches the $^{23}$(Na,Ne) URCA shell. At that epoch, due to the contemporary action of weak processes determining the local cooling and, hence, the decrease of the local luminosity, and the local decrease of molecular weight due to the conversion of $^{23}$Ne into $^{23}$Na, we find that in the model LED
factor depending mainly on the mass of the progenitor and the initial metallicity. Later on, during the simmering phase, $^{21}$Ne is produced via $n$-capture on $^{20}$Ne.

In order to illustrate the effects of the $^{21}$(F,Ne) URCA shell on the evolution of accreting WDs, we consider once again our test case, i.e., the ZSUN WD model accreting CO-rich matter at $10^{-7} M_\odot$ yr$^{-1}$, having an initial $^{21}$Ne abundance equal to 3.74 x $10^{-5}$ by mass fraction. We include all the energy contributions related to weak processes as detailed in Equation (5), and we define the borders of the zone that is unstable for convection according to the Ledoux criterion. We adopt the full nuclear network detailed above, and we compute two models, one including (Model FNe) and another not including (Model noFNe) weak processes involving $^{21}$F and $^{21}$Ne. The results are displayed in Figure 7, where we report the run of central density (upper panel), of neutronization variation at the center (middle panel) and of the mass extension of the convective core (lower panel) as a function of central temperature for these two models.

As the central density approaches the threshold value for $e$-capture on $^{21}$Ne, ($\rho_{th} = 3.78 \times 10^9$ g cm$^{-3}$), in Model FNe, the local temperature decreases due to emission of neutrino/antineutrino, the mass and kinetic energy terms in Equation (5) canceling out each other. During this phase and up to when the central density does not exceed $\sim 5 \times 10^9$ g cm$^{-3}$, the $^{21}$Ne nucleosynthesis is very scarce due to the fact that the equilibrium abundance of neutrons and of $^{17}$O abundance is very low, so that neither $\alpha$- nor $n$-channel production are efficient. As a consequence, the evolution of Model FNe is quite similar to that of Model noFNe, even if it occurs at a slightly lower temperature: central convection sets in, and when the central density reaches the corresponding $\rho_{th}$ values, URCA triplets $^{21}$(Si,Al,Mg) and $^{24}$(Mg,Na,Ne) become active. However, when the central density exceeds $4.42 \times 10^9$ g cm$^{-3}$, the border of the convective core attains the location of the $^{21}$(F,Ne) URCA shell. As a consequence, the innermost zone of the core experiences a thermal oscillation, and the convective core mass slightly reduces. The same occurs also later on when the central density exceeds $\sim 4.7 \times 10^9$ g cm$^{-3}$. Such a behavior is caused by the interplay between the URCA triplets $^{29}$(Si,Al,Mg) and $^{24}$(Mg,Na,Ne) with the $^{21}$(F,Ne) URCA shell. As a net result, it comes out that Model FNe is not heated efficiently by the circulation of isobars involved in URCA processes, as it occurs in Model noFNe. This produces the contraction of the innermost zones so that the model attains a definitively larger central density.

When the central density exceeds $\sim 5.4 \times 10^9$ g cm$^{-3}$, Model FNe experiences a huge thermal oscillation. At the beginning, the circulation of $e$-rich material from the $^{21}$(F,Ne) URCA shell inward produces the very efficient cooling of the center, while the border of the convective core remains close to the URCA shell, located at the mass coordinate $\sim 0.154 M_\odot$. Later on, due to the decrease of the temperature in the convective core, the energy production via nuclear burning decreases, so that the convective core reduces. Hence, due to the contraction of the innermost zones, the temperature starts to increase again, and the convective core grows in mass up to the $^{21}$(F,Ne) URCA shell, which is now located at the mass coordinate $\sim 0.2 M_\odot$. The occurrence of the large thermal oscillation is unavoidable. We tried to increase the spatial and temporal resolution, but the only result was to delay or anticipate its occurrence. We remark that the time and mass grid resolution adopted in the computation of this models represents the best choice to avoid spurious thermal oscillations whose only effect is to artificially increase the neutronization inside the convective core. In this regard, please note that the thermal oscillation described above and displayed in Figure 7 has practically no effects on the neutronization at the center that is a tracer of the average neutronization in the convective core (see the middle panel).

As a result, the inclusion of the $^{21}$(F,Ne) URCA pairs determines a larger central density, a smaller convective core, and a larger neutronization in the innermost zones. We want to remark that the ultimate consequence of the inclusion of the URCA pairs produces a lower total neutronization during the simmering phase in Model FNe as compared to the noFNe one. In fact, in the latter, the total variation of the neutronization along the whole CO WD $\Delta y_{bud}$ is $3.96 \times 10^{-4}$ while in the former it is $2.40 \times 10^{-4}$. Notwithstanding, at the explosion epoch, the average neutronization level in the innermost zones of the accreting WD is larger in Model FNe because the maximum mass extension of the convective core is lower (see Figure 8).

A further inspection of Figures 7 and 8 reveals that the use of the Ledoux criterion in defining the zones unstable for...
convection has an important effect in determining the thermal and chemical stratification in the zones close to the center. In particular, as soon as the nuclear timescale at the center becomes comparable to that for energy transport via convection, the mixing efficiency of nuclearily processed matter from the center outward decreases. This produces a chemical composition gradient that acts as a stabilizing mechanism in the Ledoux criterion. As a consequence, the center of the accreting WD becomes radiative. In Model FNe, this determines also a decrease of the energy flux in the layers above, so that the convective core becomes a convective shell, having its inner border at $M \sim 0.007 M_e$ and extension $\sim 0.025 M_e$. In Model noFNe, the inner border of the convective shell is located at $\sim 0.08 M_e$, even if the outer border is practically coincident with the maximum extension of the previous convective core. Such an occurrence determines the rapid increase of the temperature at the center, as clearly visible in both the upper panel of Figure 7 and lower panel of Figure 8, respectively.

7. Accreting WDs with Different Initial Z

In order to investigate the effects of different initial chemical compositions of the WD progenitors on the simmering phase, we adopt five different starting WD models, with total metallicity of the progenitor equal to $Z = (0.0245, 0.6, 1.38, 2$ and $4) \times 10^{-2}$ and labeled in the following as “Z12,” “Z63,” “Z14,” “Z22,” and “Z42,” respectively. Please note that models Z14, Z12, and Z42 are the same as ZLOW, ZSUN, and ZHIGH in P2017, respectively. We accrete CO-rich matter according to the same procedure described in P2017 at a rate of $M = 10^{-7} M_e$ yr$^{-1}$. In the computation, we adopt the same nuclear network as for Model FNe above; meanwhile, as in P2017, we adopt the advective scheme for the convection mixing. We define zones unstable for convection according to the Ledoux criterion, and we include in the energy conservation equation all the contributions related to weak processes according to Equation (5). Our results are summarized in Table 3 where we report several physical and chemical properties characterizing the simmering phase of the considered models. In Figure 9, we report the evolution of the center in the $\rho - T$ (upper panel) and the value at the center of neutronization as a function of the central density. Figure 9 displays the onset of various URCA shell and their effect on the thermal evolution of the accreting WD. In particular, as it is well known, during the accretion phase, e-captures on $^{24}$Mg, $^{23}$Na, and $^{21}$Ne determine a cooling of the innermost zones of the accreting WDs and, at the same time, an increase of the local value of neutronization. The effect of the $^{21}$(Ne,F) URCA shell is completely negligible in Model Z14, as the $^{21}$Ne abundance in the initial WD is very low (see Table 3). In this model, the central temperature continues to increase due to the homologous compression of the whole WD as determined by the continuous mass deposition. Moreover, Figure 9 also discloses that the activation of e-capture on $^{23}$Mg determines a rapid heating of the center, so that convection sets in (see the values of $\rho_{\text{lim}}$ in Table 3). Also in this case, the evolution of Model Z14 is different, as the onset of convection occurs at a lower density.

At this phase of the evolution, C-burning is fully active, so that $^{23}$Na and $^{21}$Ne$^{10}$ are produced. When central convection

\footnotetext{[10] We recall that $^{21}$Ne is produced via $\alpha$-capture on $^{20}$Ne, the neutrons being produced by the reaction chain $^{12}$C(p,$\gamma$)$^{13}$N($p^3$)$_j^3$C(\alpha,$\gamma$)$^{16}$O.}
of the mass of the convective core, while the continuous mass deposition determines the compressional heating of the center and the increase of the whole density profile. As a matter of fact, the mass location of the $^{21}\text{(Ne,F)}$ URCA shell moves outward. In this phase, nuclear burning occurs at a reduced rate, so that neutronization at the center is almost constant. Later on, when nuclear burning fully resumes, the convective core starts to increase once again, diluting the $n$-rich material and, hence, limiting the increase of $\eta_C$. When convection attains the $^{21}\text{(Ne,F)}$ URCA shell, the local $\mu$ gradient stops the transport of thermal energy and prevents any further mixing of neutronized material outward. From this moment up to the explosion, the external border of the convective-unstable zone does not change anymore.

Models with different initial metallicity experience different cooling during the presupernova phase, and this determines a different location of the $^{21}\text{(Ne,F)}$ URCA shell (the lower the initial $Z$, the more internal this URCA shell). As a consequence, the maximum extension of the convective core depends also on the initial metallicity (see Table 3). As the convective core is limited by the URCA shell, the circulation of neutron poor material is reduced so that such a heating mechanism has a very low efficiency. As a consequence, the evolution up to the explosion is driven by C-burning, which, in turn, determines a large increase of the neutronization level in the innermost region (for instance, compare the run of $\eta_C$ in Figure 9 with those in Figure 2). On the other hand, the fact that the convective core small determines a rapid increase of the temperature in the burning region.

In Figure 10, we report the run of neutronization as a function of the mass coordinate for all the computed models at the epoch of explosion.

### 8. Effects of Different $\dot{M}$, $M_{\text{WD}}$, and Cooling Age

As it is well known, for CO WDs attaining $M_{\text{Ch}}$ via mass accretion, the physical conditions at the onset of C-burning and, hence, at the explosion depend on the relative ratio of the compressional heating and of the thermal diffusion timescales. The former depends on the value of the mass transfer rate, while the latter is fixed by the thermal content of the accreting WDs, as determined by the cooling age before the onset of mass transfer, and on the WD initial mass. In order to investigate how the properties during the simmering phase depend on $\dot{M}$, $M_{\text{WD}}$, and on the cooling age, we compute three additional sets of models. In Set 1, we adopt as the initial model the $0.8 M_\odot$ CO WD with $Z = 1.38 \times 10^{-2}$, and we compute the evolutionary sequence by adopting $\dot{M} = 0.5$, 2, 6 and $9 \times 10^{-7} M_\odot$ yr$^{-1}$ (models R5m8, R2m7, R6m7, and R9m7, respectively). In Set 2, we let the same CO WD model of Set 1 cool down for additional $t_{\text{cool}} \simeq 1$ Gyr, up to when its central temperature attains $T_C = 7.9 \times 10^7$ K; then we compute two evolutionary sequences by adopting $\dot{M} = 1$ and $9 \times 10^{-7} M_\odot$ yr$^{-1}$ (models Clm7 and C9m7, respectively). In Set 3, we accrete matter at $\dot{M} = 10^{-7} M_\odot$ yr$^{-1}$ on the same initial CO WD of Set 1. When the total mass of the accreting WD is equal to $1 M_{\text{Ch}}$, we stop the accretion and let the model cool down up to when its central temperature attains the value $T_C = 1.13 \times 10^7$ K. Hence, we compute two models with the same accretion rates as in Set 2 (models M1m7 and M9m7, respectively). In Table 4, we report selected physical and chemical quantities characterizing the accretion and the simmering phases for models in Set 1, 2, and 3.

Concerning the effects of different $\dot{M}$ (models in Set 1), the behavior as well as the physical and chemical properties at the explosion of models with $\dot{M} \leq 2 \times 10^{-7} M_\odot$ yr$^{-1}$ are easily explained when considering that, decreasing the accretion rate, the amount of mass to be accreted to trigger the explosion is larger. As a consequence, the lower the accretion rate, the higher the density level along the whole accreting structure and, hence, the more external the maximum mass coordinate attained by the $^{21}\text{(F,Ne)}$ URCA shell. This implies that for lower accretion rates the mass of the convective core is larger and, hence, a greater amount of $^{12}\text{C}$ has to be burnt to produce the explosion. However, even if in the considered range of $\dot{M}$ the convective core almost doubles, and the consumed carbon increases by $\sim 35\%$, the average neutronization in the convective core decreases by only $\sim 7\%$ while the neutronization value at the center is practically unaffected (see also Figure 11). The situation for Model R6m7 with $M = 6 \times 10^{-7} M_\odot$ yr$^{-1}$ is quite different as C-burning is ignited close to the epoch of the $e$-capture on $^{21}\text{Ne}$ activation at the center. This implies that C-ignition occurs off center, due to the interplay among the compressional heating, of the inward diffusion of thermal energy from the accreted layers, and of the local cooling due to electron capture at the center. As a consequence, at the onset of the simmering phase, a convective shell forms at the mass coordinate $M_{\text{sim}}^\text{e} \simeq 0.015 M_\odot$, while the central region is cooled down by the conversion of $^{21}\text{Ne}$ into $^{21}\text{F}$. However, as the convective shell grows in mass outward, the innermost regions start to expand so that the density at center decreases and the energy contribution from the $e$-capture on $^{21}\text{Ne}$ decreases. Contemporary thermal energy flows from the convective shell inward, heating up rapidly the center so that the thermonuclear runaway occurs there. As the $^{21}\text{(F,Ne)}$ URCA shell is ineffective in this model, the resulting convective core can grow in mass up to the $^{21}\text{(Ne,F)}$ URCA shell, so that the amount of carbon to be consumed to trigger the explosion results definitively larger than that for Model R2m7 (see Table 4). Model R9m7 has the same behavior even if the compressional heating timescale is definitively shorter than the inward thermal diffusion timescale, thus determining the occurrence of C-burning very far from center. As in Model R6m7, at the onset of the simmering phase, a convective shell forms, but now its inner border is more external, and it is located at $\sim 0.024 M_\odot$. This prevents any efficient heating of the central region of the accreting WD so that the thermonuclear runaway occurs off center. The other important difference between models R6m7 and R9m7 is that the latter has a larger thermal content at the onset of the simmering phase, so that a smaller amount of $^{12}\text{C}$ has to be burnt to trigger
When considering models in Set 3, it comes out once again that for low values of the accretion rate the physical and chemical properties at the C-ignition epoch and during the simmering phase up to the explosion are largely independent on the initial mass of the accreting CO WDs. On the other hand, the thermal diffusion timescale in Model M9m7 is definitively larger than the compressional heating one, so that the evolution of the innermost \( \approx 0.5 M_0 \) is driven by the homologous compression only. As a consequence, C-ignition occurs at the center for very large values of density, and as a whole, its evolution resembles quite closely that of the models with lower CO WD initial mass and very low mass accretion rates.

9. Final Considerations

In this work, we analyze URCA processes and their interplay with convection during the simmering phase of SNe Ia progenitors. By including all the relevant physical processes, we determine accurately the physical and chemical profile of an accreting WD as well as the neutronization level at the explosion epoch. Our results suggest the following:

1. A complete nuclear network, at least up to \(^{32}\)S, should be used. In fact, by neglecting \( n \)-capture, \( \alpha \)-capture, and \( p \)-capture, the amount of \(^{12}\)C consumed during the simmering is overestimated, and consequently, the final neutronization level results are larger.

2. All the relevant equations describing the physical and chemical structure of an accreting WD as well as its time evolution should be solved as contemporaries. In fact, by neglecting \( n \)-capture, \( \alpha \)-capture, and \( p \)-capture, the amount of \(^{12}\)C consumed during the simmering is overestimated, and consequently, the final neutronization level results are larger.

3. The Ledoux criterion should be used in order to account for the existence of a \( \mu \)-gradient at a given URCA shell. Our results show that according to this criterion the convective core stops at the innermost URCA shell (i.e., the \(^{21}\)(Ne,F) one), thus reducing the inward circulation of

---

**Table 4**

| Model       | R5m8 | Z12  | R2m7 | R6m7 | R9m7 | C1m7 | C9m7 | M1m7 | M9m7 |
|-------------|------|------|------|------|------|------|------|------|------|
| \( M(10^{-3}M_0) \), yr\(^{-1} \) | 5.0  | 10.0 | 20.0 | 60.0 | 90.0 | 10.0 | 90.0 | 10.0 | 90.0 |
| \( M_{\text{acc}}(M_0) \) | 0.568| 0.567| 0.566| 0.564| 0.562| 0.567| 0.566| 0.384| 0.388 |
| \( t_{\text{acc}}(10^6 \text{ yr}) \) | 11.35| 5.665| 2.828| 0.935| 0.625| 5.668| 0.629| 3.839| 0.431 |
| \( t_{\text{bann}}(10^4 \text{ yr}) \) | 14.71| 5.853| 3.171| 0.221| 0.038| 6.097| 0.027| 6.187| 0.963 |
| \( \rho_{\text{sim}}(10^9 \text{ g cm}^{-3}) \) | 4.298| 4.250| 3.625| 1.500| 1.236| 4.239| 1.236| 4.272| 4.442 |
| \( T_{\text{sim}}(10^8 \text{ K}) \) | 1.463| 1.693| 1.901| 2.059| 2.153| 1.687| 1.933| 1.678| 1.378 |
| \( M_{\text{sim}}(10^{-2}M_0) \) | 0.000| 0.000| 0.000| 0.153| 2.365| 0.000| 14.55| 0.000| 0.000 |
| \( \rho_{\text{simmax}}(10^9 \text{ g cm}^{-3}) \) | 4.270| 4.194| 1.872| 3.743| 3.519| 4.193| 3.471| 4.200| 4.287 |
| \( T_{\text{simmax}}(10^8 \text{ K}) \) | 1.454| 1.693| 1.526| 2.394| 2.551| 1.668| 2.628| 1.652| 1.310 |
| \( M(10^{-3}C_{12}) \) | 0.313| 0.313| 0.314| 0.311| 0.312| 0.314| 0.310| 0.313| 0.312 |
| \( \Delta M(10^{-3}C_{12}) \) | 2.512| 2.089| 1.700| 4.583| 3.553| 1.811| 5.104| 2.488| 3.067 |
| \( M_{\text{ext}}(M_0) \) | 0.202| 0.152| 0.109| 0.323| 0.347| 0.162| 0.446| 0.167| 0.235 |
| \( \rho_{\text{exp}}(10^9 \text{ g cm}^{-3}) \) | 6.007| 5.447| 5.029| 3.668| 3.683| 5.564| 3.479| 5.532| 6.315 |
| \( \eta_{\text{exp}}(10^{-3}) \) | 2.726| 2.738| 2.767| 2.754| 2.320| 2.753| 2.352| 2.820| 3.182 |
| \( \tau_{\text{exp}}(10^{-3}) \) | 2.355| 2.430| 2.526| 2.188| 1.952| 2.240| 2.118| 2.503| 2.326 |
| \( M_{\text{exp}}(10^{-3}M_0) \) | 0.000| 0.000| 0.000| 0.000| 9.078| 0.000| 91.20| 0.000| 0.000 |

**Note.** \( M_{\text{sim}} \) represents the mass coordinate of the inner border of the convective zone at the onset of the simmering phase; \( M_{\text{exp}} \) is the mass coordinate where the thermonuclear runaway occurs.
e-rich isobars and the corresponding contribution to the total energy budget.

4. The inward convective motion of e-rich isobar from an URCA shell determines an effective cooling and, in addition, prevents convection encompassing the URCA shell itself. Such an occurrence determines a larger C-consumption and, hence, a larger neutronization level and a higher density at the explosion epoch.

5. The chemical and physical structure at the explosion epoch depends on the assumed mixing efficiency, as determined by different numerical algorithms used to model convective mixing.

An inspection to Table 3 and of Figures 9 and 10 clearly suggests that the properties of the accreting WDs during the simmering phase and at the explosion epoch depend on the initial metal content of the progenitor. This can be easily understood, because with a larger initial metallicity, the following conditions are true:

1. a larger abundance of URCA mother nuclei in the accreting WD and, hence, stronger cooling effects during the accretion phase;
2. a larger amount of mass to be accreted to trigger the thermonuclear runaway;
3. a more external position of the $^{21}(\text{F,Ne})$ URCA shell and, hence, larger zone unstable for convection during the simmering;
4. a larger amount of nuclear energy to be injected to trigger the explosion;
5. a larger carbon consumption and, hence, larger neutronization level.

We want to remark once again that such a result is a direct consequence of the fact that the innermost URCA shell represents a barrier for convection so that circulation of URCA isobars across the shell is inhibited, and their energy contribution to the heating up of the innermost zone is drastically reduced.

For this reason, analogously to P2017, we define a final-to-initial-neutron-excess relation by interpolating the values of $\eta_{\exp}$ as a function of $\eta_{c,0}$, thus obtaining

$$\eta_{\exp} = (1.080 \pm 0.132) \times 10^{-3} + (0.995 \pm 0.068) \times \eta_{c,0}$$

$$R^2 = 0.9930.$$  

On the other hand, the value of $\eta_{c,0}$ depends on the initial metallicity according to the following relation:

$$\eta_{c,0} = 1.213 \times 10^{-5} + 9.224 \times 10^{-2} \times Z$$

$$R = 1.000$$

so that Equation (14) becomes

$$\eta_{\exp} = (1.094 \pm 0.143) \times 10^{-3} + (9.168 \pm 0.677) \times 10^{-2} \times Z$$

$$R^2 = 0.9920.$$  

Such a relation can be used to infer the metallicity of star progenitors of galactic SNe Ia, once a reliable estimate of the neutronization level at the explosion epoch is available. Badenes et al. (2008) and Park et al. (2013) derived the neutronization level of the Tycho and Kepler Supernova based on the observed actual abundance of Mn and Cr, $\eta_{\exp,\text{Tycho}} = 4.36 \times 10^{-3}$ and $\eta_{\exp,\text{Kepler}} = 4.55 \times 10^{-3}$. By assuming that these values correspond to the average neutronization of the two supernovae at the explosion epoch, one can derive the progenitor’s metallicity to be as follows: $Z_{\text{Tycho}} = 0.0353 = 2.56Z_\odot$ and $Z_{\text{Kepler}} = 0.0373 = 2.71Z_\odot$.

However, the estimation of the remnants metallicity in Tycho and Kepler assumed that the measured Mn and Cr were synthesized in the incomplete Si burning regime, which is expected to occur above the mass affected by convection in the present models. As can be seen in Figure 10, the neutron excess above the $^{21}(\text{F,Ne})$ URCA shell is the same as in the progenitor WD before accretion begins. As a consequence, we recover the large values of Z derived by Badenes et al. (2008), Park et al. (2013) for the progenitors of these two historical SNe Ia.

Our results are in agreement with the findings by P2017, even if the physical and chemical properties of models at the explosion epoch are completely different. In fact, in P2017 not all the relevant energy contributions related to URCA processes are included consistently, and the effects of $\mu$-gradient are not taken into account in defining the zones unstable for convection. As a consequence, in that work, the amount of carbon consumed to trigger the explosion is about an order of magnitude larger than what estimated in the present work (compare the values of $\Delta M_{^{12}\text{C}}$ reported in Table 3 and those in Table 2 in P2017). As a consequence, in P2017 the total neutronization produced during the evolution is also larger, even if the average value in the convective core is almost equal to the one derived in the present work: this is a consequence of the fact that in P2017 almost the whole accreting WD is unstable for convection, meanwhile, in our models, the convective core extends only on the innermost zones.

We have found that a large accretion rate is needed to affect significantly the evolution of the WD pre-explosion. Accretion rates as large as $(6-9) \times 10^{-7} M_\odot \, \text{yr}^{-1}$ imply a high rate of compressional heating, which shortens the time needed for carbon to achieve simmering conditions first, and explosive conditions later. The result is that the $^{21}(\text{F,Ne})$ URCA pair influence is reduced, convection sets in a shell close to the center, the convective mass is larger (but far from the values close to $\sim 1 M_\odot$ found in previous works), and the explosion density is smaller, on the order of $3.6 \times 10^{9} \, \text{g} \, \text{cm}^{-3}$.

This publication is part of the project I+D+I PGC2018-095317-B-C21 funded by MICIN/AEI/1013039/501100011033 and FEDER “A way of doing Europe” (E.B. and I.D.); L.P. and O.S. acknowledge financial support from the INAF-mainstream project “Type Ia Supernovae and their Parent Galaxies: Expected Results from LSST,” O.S. and L.P. acknowledge their participation to the V:ANS project (Vanvitielli program on standard candles in astrophysics: Atomic and Nuclear physics in SNIa) supported by the Vanvitielli University.

**Appendix**

**Summary of Computed Models**

In Table 5, we list all the models presented in the text and summarize the setup adopted in their computations. The meaning of the various columns is as follows:

1. Model: label adopted in the text to address a given model;
Table 5
Computed Models and Adopted Computational Setup

| Model | $Z$ | $M_{\text{WD}}(M_\odot)$ | $M(10^{-7} M_\odot \text{yr}^{-1})$ | Nuc.Net. | Conv.Crit. | Mix.Sch. | Coupled | $\varepsilon_{\text{urca}}$ |
|-------|-----|--------------------------|-------------------------------|----------|------------|----------|---------|------------------|
| K00   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | SHORT | SCHWAR | DIF | YES | Equation (3) |
| K28   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | SHOR1 | SCHWAR | DIF | YES | Equation (3) |
| K24   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | SHORT | SCHWAR | DIF | NO | Equation (3) |
| SEL   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | SHORT | SCHWAR | DIF | NO | Equation (3) |
| ADd   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | SHORT | SCHWAR | ADV | NO | Equation (3) |
| DFd   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | SHORT | SCHWAR | DIF | NO | Equation (3) |
| ADV   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | SHORT | SCHWAR | ADV | YES | Equation (3) |
| REF   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | LONG | SCHWAR | DIF | YES | Equation (3) |
| W00   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | LONG | SCHWAR | DIF | YES | Equation (5) |
| W06   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | LONG | SCHWAR | DIF | YES | Equation (5) |
| WRK   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | LONG | SCHWAR | DIF | YES | Equation (5) |
| LED   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | LONG | LEDOUX | DIF | YES | Equation (3) |
| noFNe | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | FULL | LEDOUX | DIF | YES | Equation (5) |
| FNe   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | FULL | LEDOUX | DIF | YES | Equation (5) |
| Z14   | 2.45 $\times$ 10$^{-4}$ | 0.8 | 1.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| Z63   | 6.00 $\times$ 10$^{-4}$ | 0.8 | 1.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| Z12   | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| Z22   | 2.00 $\times$ 10$^{-2}$ | 0.8 | 1.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| Z42   | 4.00 $\times$ 10$^{-2}$ | 0.8 | 1.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| R5m8  | 1.38 $\times$ 10$^{-2}$ | 0.8 | 0.5 | FULL | LEDOUX | ADV | YES | Equation (5) |
| R2m7  | 1.38 $\times$ 10$^{-2}$ | 0.8 | 2.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| R6m7  | 1.38 $\times$ 10$^{-2}$ | 0.8 | 6.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| R9m7  | 1.38 $\times$ 10$^{-2}$ | 0.8 | 9.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| C1m7  | 1.38 $\times$ 10$^{-2}$ | 0.8 | 1.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| C9m7  | 1.38 $\times$ 10$^{-2}$ | 0.8 | 9.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| M1m7  | 1.38 $\times$ 10$^{-2}$ | 1.0 | 1.0 | FULL | LEDOUX | ADV | YES | Equation (5) |
| M9m7  | 1.38 $\times$ 10$^{-2}$ | 1.0 | 9.0 | FULL | LEDOUX | ADV | YES | Equation (5) |

2. $M_{\text{WD}}$: initial mass of the CO WD at the onset of the accretion phase;
3. $M$: adopted accretion rate;
4. Nuc.Net.: nuclear network adopted in the computation. The different possibilities are as follows:
   SHORT: same one as in Piro & Bildsten (2008), but including also the $^{25}$(Mg,Na) and $^{56}$(Fe,Cr,Mn) URCA processes;
   SHOR1: same as SHORT, but including also the $^{28}$(Si,Al,Mg) URCA triplet (see Section 4);
   SHOR2: same as SHOR1, but including also the $^{24}$(Mg,Na,Ne) URCA triplet (see Section 4);
   LONG: nuclear network including 52 isotopes of all the strong and weak nuclear processes, except for weak processes on $^{21}$F and $^{21}$Ne (see Section 6);
   FULL: same as LONG, but including also the weak processes on $^{21}$F and $^{21}$Ne (see Section 6);
5. Conv.Crit.: adopted criterion to defined zones unstable for convection (see Section 2):
   SCHWAR: Schwarzschild criterion;
   LEDOUX: Ledoux criterion;
6. Mix.Sch.: numerical scheme adopted to model convective mixing (see Section 4):
   DIF: diffusive scheme;
   ADV: advective scheme;
   SEL: Sparks-Endal-Linearized scheme
7. Coupled: YES: all the equations describing the physical and chemical evolution of accreting WD are solved simultaneously; NO: the operator splitting technique is applied;
8. $\varepsilon_{\text{urca}}$: energy contribution due to URCA processes and, more in general to weak processes, included in the equation for the conservation of energy.

ORCID iDs
Luciano Piersanti @ https://orcid.org/0000-0002-8758-244X
Eduardo Bravo @ https://orcid.org/0000-0003-0894-6450
Oscar Straniero @ https://orcid.org/0000-0002-5514-6125
Sergio Cristallo @ https://orcid.org/0000-0001-9683-9406
Inmaculada Domínguez @ https://orcid.org/0000-0002-3827-4731

References
Badenes, C., Bravo, E., & Hughes, J. P. 2008, ApJL, 680, L33
Betoule, M., Kessler, R., Guy, J., et al. 2014, A&A, 568, A22
Branch, D., & Wheeler, J. C. 2017, Supernova Explosions (Heidelberg: Springer)
Bravo, E., Isern, J., & Canal, R. 1993, A&A, 270, 288
Brown, P. J., Kuin, P., Scalzo, R., et al. 2014, ApJ, 787, 29
Castellani, V., Giannone, P., & Renzini, A. 1971, Ap&SS, 10, 355
Chamulak, D. A., Brown, E. F., Timmes, F. X., & Dupczak, K. 2008, ApJ, 677, 160
Chiefi, A., Limongi, M., & Straniero, O. 1998, ApJ, 502, 737
Christl, S., Straniero, O., Gallino, R. et al. 2009, ApJ, 692, 979
Cyburt, R. H., Amthor, A. M., Ferguson, R., et al. 2010, ApJS, 189, 240
Dominguez, I., Höflich, P., & Straniero, O. 2001, ApJ, 557, 279
Förts, A., Spyromilio, J., Taubenberger, S., et al. 2020, MNRAS, 491, 2902
Gallagher, J. S., Garnavich, P. M., Berlind, P., et al. 2005, ApJ, 634, 210
Hakobyan, A. A., Karapetyan, A. G., Barkhudaryan, L. V., Gevorgyan, M. H., & Adibekyan, V. 2021, MNRAS, 505, L52
Hamuy, M., Carter, R., Contreras, C., & Suntzeff, N. B. 2021, MNRAS, 500, 1095
Hamuy, M., Phillips, M. M., Suntzeff, N. B., et al. 1996, AJ, 112, 2398
