The i-process yields of rapidly accreting white dwarfs from multicycle He-shell flash stellar evolution models with mixing parametrizations from 3D hydrodynamics simulations

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ABSTRACT
We have modelled the multicycle evolution of rapidly accreting CO white dwarfs (RAWDs) with stable H burning intermittent with strong He-shell flashes on their surfaces for \(0.7 \leq M_{\text{RAWD}}/M_\odot \leq 0.75\) and \([\text{Fe/H}]\) ranging from 0 to \(-2.6\). We have also computed the i-process nucleosynthesis yields for these models. The i process occurs when convection driven by the He-shell flash ingests protons from the accreted H-rich surface layer, which results in maximum neutron densities \(N_n, \text{max} \approx 10^{13}–10^{15} \text{cm}^{-3}\). The H-ingestion rate and the convective boundary mixing (CBM) parameter \(f_{\text{top}}\) adopted in the one-dimensional nucleosynthesis and stellar evolution models are constrained through three-dimensional (3D) hydrodynamic simulations. The mass ingestion rate and, for the first time, the scaling laws for the CBM parameter \(f_{\text{top}}\) have been determined from 3D hydrodynamic simulations. We confirm our previous result that the high-metallicity RAWDs have a low mass retention efficiency (\(\eta \sim 10\) per cent). A new result is that RAWDs with \([\text{Fe/H}] \lesssim -2\) have \(\eta \gtrsim 20\) per cent; therefore, their masses may reach the Chandrasekhar limit and they may eventually explode as SNeIa. This result and the good fits of the i-process yields from the metal-poor RAWDs to the observed chemical composition of the CEMP-r/s stars suggest that some of the present-day CEMP-r/s stars could be former distant members of triple systems, orbiting close binary systems with RAWDs that may have later exploded as SNeIa.

Key words: Sun: abundances – Galaxy: abundances – binaries: close – stars: interiors – stars: white dwarfs – nuclear reactions, nucleosynthesis, abundances.

1 INTRODUCTION
The type Ia supernovae (SNeIa) are thermonuclear explosions of carbon–oxygen (CO) white dwarfs (WDs) (e.g. Hoyle & Fowler 1960; Hillebrandt & Niemeyer 2000; Hillebrandt et al. 2013; Churazov et al. 2014; Livio & Mazzali 2018). In the single degenerate (SD) channel of SNeIa progenitors, which was originally proposed by Schatzman (1963) and Whelan & Iben (1973), it is assumed the CO WD is a primary star of a close binary system with a main-sequence, sub-giant, or red giant branch companion. The secondary star fills its Roche lobe and donates matter from an H-rich envelope to the WD, and, as a result, the WD will explode when its growing mass \(M_{\text{WD}}\) approaches the Chandrasekhar limit \(M_{\text{Ch}} \approx 1.38M_\odot\). Initially, the primary star of such a binary system was an intermediate-mass star with \(M \approx 2.5–7M_\odot\), the upper boundary of this mass interval depending on the amount of convective boundary mixing and C-burning rate (Chen et al. 2014). It left a core – the CO WD – after having lost the rest of its mass during a common-envelope event, when it arrived at the asymptotic giant branch (AGB) and filled its Roche lobe. Given that the CO cores of AGB stars can grow in mass only up to \(\sim 1M_\odot\) (Chen et al. 2014), the SD channel can work only if the accreted H-rich matter is first
processed into He and then into C and O, while being efficiently retained on the WD.

Fig. 9 of Nomoto (1982) summarizes the results of the previous investigations of H accretion onto CO WDs at different rates $\dot{M}_{\text{acc}}$ (an update of this figure can be found in Nomoto et al. 2007). It shows that stable burning of accreted H occurs in a very narrow interval of $\dot{M}_{\text{acc}}$ around a value of $10^{-9}$ M$_\odot$ yr$^{-1}$ that linearly increases with $M_{\text{WD}}$. At the lower rates, H is processed into He via thermal flashes that become stronger when $\dot{M}_{\text{acc}}$ decreases, eventually leading to thermonuclear runaways typical for the classical nova. At the higher rates, the non-processed H accumulates in an expanding envelope, so that such a rapidly accreting WD would be resembling a red giant.

However, an important aspect of the evolution of rapidly accreting WDs is the fact that even if accreted H is burning stably the He shell will ignite in a thermonuclear runaway when enough He has been accumulated as H-shell burning ash, just as in a thermal-pulse AGB star (Cassisi, Iben & Tornambè 1998). A few consecutive He-shell flashes at the end of H accretion were computed by Idan, Shaviv & star (Cassisi, Iben & Tornambè 1998). A few consecutive He-shell flashes at the end of H accretion were computed by Idan, Shaviv & Shaviv (2013). Following Nomoto (1982), various outcomes of pure He accretion onto CO WDs at different values of $\dot{M}_{\text{acc}}$ have been studied in a number of papers (e.g. Piersanti, Tornambè & Yungelson 2014; Wang et al. 2015). The data obtained in such simulations are used in models of binary population synthesis to estimate a theoretical SN Ia rate for the SD channel, simply assuming that the He accretion rate matches that of H in the regime of stable H burning (e.g. Han & Podsiadlowski 2004).

Recently, Denissenkov et al. (2017a, hereafter Paper I) have presented the results of the first stellar evolution computations in which rapid accretion and stable burning of H on CO WDs were repeatedly interrupted by strong He-shell flashes that were followed by mass loss caused either by the super-Eddington luminosity wind or by the common-envelope event resulting from the expansion of the WD envelope overflowing its Roche lobe, after which the H accretion resumed. These simulations show a low efficiency of He retention ($\eta_{\text{He}} \lesssim 10$ per cent) and, because all the processes that accompany the rapid H accretion were taken into account, they provide estimates of the low retention efficiency of the total accreted mass ($\eta \approx \eta_{\text{He}}$) by the rapidly accreting WDs (RAWDs).\footnote{We define an RAWD as a WD that accretes H rapidly enough for its stable burning to be maintained on the WD surface. This definition is slightly different from the one used by Lepo & van Kerkwijk (2013) whose RAWDs can have the higher accretion rates and lose mass via the optically thick wind.} In one of the models even a negative value of $\eta_{\text{He}}$ was found meaning that $M_{\text{WD}}$ was decreasing with time. Given that the binary population synthesis models predict an immediate split of the He-shell flash convection zone which would prevent any mixing of protons and the burning products of p capture on $^{12}$C into the hottest bottom region of the pulse-driven convection zone, Herwig et al. (2014) have shown that in three-dimensional (3D) hydrodynamic simulations the He ingestion triggers a Global Oscillation of Shell H-ingestion (GOSH) which drastically rearranges the structure of the He-shell flash convection zone. So far it has not been possible to follow the long-term evolution of this event past the first GOSH. The one-dimensional (1D), spherically symmetric nucleosynthesis simulations of Herwig et al. (2011) adopted the approach that mixing between the upper layer in which protons and $^{12}$C react and the bottom layer in the pulse-driven convection zone, where neutrons can be released via the $^{13}$C($\alpha$,n)$^{16}$O reaction, continues until the observed chemical composition of Sakurai’s object has been reproduced as well as possible. This delayed-split scenario is not yet fully supported by the initial 3D hydrodynamics simulation results by Herwig et al. (2014). The initial GOSH in those simulations happens at a time when the amount of protons consumed is still insufficient to explain the neutron exposure required to explain observations, and the evolution past the initial GOSH in three dimensions remains unclear.

With an ongoing supply of protons, most of which are transformed into neutrons in the above reactions, the neutron number density at the bottom of the He zone can reach a value of $N_n \sim 10^{11}$ cm$^{-3}$ intermediate between the values characteristic of the $s$ ($N_n \lesssim 10^{10}$ cm$^{-3}$) and $r$ ($N_n \gtrsim 10^{20}$ cm$^{-3}$) process. Very-late TP objects such as the post-AGB star Sakurai’s object will only experience one H-ingestion event. Their nucleosynthesis production is very unique, but the impact on a galactic chemical evolution scale is negligible. RAWDs on the other hand continuously accrete from a close binary companion, and can potentially experience dozens of He-shell flashes followed by mass-loss episodes before the concomitant changes in the binary system parameters terminate the rapid H accretion. Along with their low mass retention efficiency, this makes the RAWDs a potentially important galactic sources of heavy elements, with distinct elemental and isotopic abundance signatures different from those produced in the $s$ and $r$ process. For example, i process from RAWDs can make a significant contribution to the first n-capture peak of the Solar system abundance distribution, as demonstrated by combining RAWD i-process yields with stellar population synthesis and galactic chemical evolution models (Côté et al. 2018).

\footnote{A segment of a WD cooling track where the evolution with nearly constant luminosity and increasing effective temperature changes to the evolution with both of them decreasing.}
RAWDs are not the only possible sites of i-process nucleosynthesis triggered by H ingestion into a He convective zone. The other sites can be low-metallicity and low-mass thermally pulsing AGB stars (Iwamoto et al. 2004; Lugaro et al. 2012), He-core flash in low-metallicity RGB stars (Campbell, Lugaro & Karakas 2010), super-AGB stars (Jones et al. 2016), and He- and He-burning shell merger in Population-III massive stars (Banerjee, Qian & Heger 2018; Clarkson, Herwig & Pignatari 2018).

In Paper I, we have considered the rapid \((M_{\text{acc}} = 1-2 \times 10^{-7} \text{M}_\odot \text{yr}^{-1})\) accretion of only solar-composition matter on CO WDs with the masses \(0.65 \text{M}_\odot, 0.73 \text{M}_\odot, \) and \(1 \text{M}_\odot.\) The present work extends the set of our RAWD models to sub-solar metallicities, while keeping their masses close to \(M_{\text{WD}} \approx 0.73 \text{M}_\odot.\) The main goals of this paper are to describe the methods that we use to simulate the multicycle evolution of RAWDs (Section 2.1) and the i-process nucleosynthesis in their He-flash convection zones during H ingestion (Section 2.2), and to present the results of our new computations of the RAWD evolution and i-process yields for a range of metallicity (Section 4) that have been used in Côté et al. (2018). Section 3 describes the 3D hydrodynamic simulations that support our 1D estimates of H mass ingestion rates for the RAWD models. Section 5 concludes the paper.

2 METHODS USED FOR 1D STELLAR EVOLUTION AND NUCLEOSYNTHESESIMULATIONS

2.1 The RAWD evolution

The new RAWD models are computed with the revision 7624 of the MESA stellar evolution code (Paxton et al. 2011, 2013). We use the reaction rates from the JINA Reaclib database (Cyburt et al. 2010) and the MESA default equation of state. The nuclear network includes 31 species, from neutron to \(^{28}\text{Si},\) that can participate in 60 reactions of the pp chains, four CNO cycles, NeNa and MgAl cycles, as well as the He (\(\alpha\))\(^{12}\text{C}(\alpha, \gamma)^{16}\text{O}, \alpha^{16}\text{O}(\alpha, \gamma)^{20}\text{Ne}, \alpha^{20}\text{Ne}(\alpha, \gamma)^{24}\text{Mg}, \alpha^{24}\text{Mg}, \alpha^{13}\text{C}(\alpha, n)^{16}\text{O}, \) and other \((\alpha, \gamma), (\alpha, n)\) and \((\alpha, p)\) reactions with the included isotopes) and C burning. The initial mixtures of elements and isotopes are prepared using the Solar system chemical composition of Asplund et al. (2009) that is scaled to specified values of \([\text{Fe/H}].\) We assume that for \([\text{Fe/H}] \leq -0.7\) the initial mixtures are \(\alpha\)-element enhanced with \([\alpha/\text{Fe}] = +0.4.\) The appropriate Type 1 and Type 2 (with enhanced C and O abundances) OPAL and low-temperature molecular opacities have been prepared for such mixtures (Denissenkov et al. 2017b) and used in our computations.

The CO WD models are made with the inlists from the MESA test suite example make_CO_wd. With these inlists the MESA code first computes the evolution of an intermediate-mass star from the pre-main sequence to the completion of the first He-shell TP on the AGB (blue curves in Fig. 1, initial masses given in each panel), then the Blöcker AGB wind parameter (Blöcker 1995) is increased from 0.1 to 5 to mimic the enhanced mass loss caused by a common-envelope interaction in a close binary system in which the AGB star overflows its Roche lobe. As a result, the model star leaves the AGB and evolves towards and down the WD cooling track (green curves in the same Figure).

\[^{3}[A/B] = \log_{10}(N(A)/N(B)) - \log_{10}(N(C)/N(C))\), where \(N(A)\) and \(N(B)\) are the abundances (number densities or mass fractions) of the nuclides A and B.

Our computations include convective boundary mixing (CBM) adopting the exponentially decaying diffusion boundary (Herwig et al. 1997):

\[D(r) = D_0 \exp \left(-\frac{2|r-r_0|}{H_P}\right),\]

where \(D_0\) is a value of the convective diffusion coefficient provided by the mixing-length theory (MLT) and \(H_P\) is the pressure scale height, both evaluated at \(r = r_0\) in the vicinity of the respective convective boundary. For the boundaries of the H and He convective cores we have adopted the value of \(f = 0.014\) that is close to the one constrained by the position of the terminal-age main sequence (TAMS) in a large number of stellar clusters (Herwig 2000). For the top and bottom boundaries of the He-flash convective zone we use the values of \(f_{\text{top}} = 0.1\) and \(f_{\text{bot}} = 0.008\) that are equal or close to those obtained in the multidimensional hydrodynamic simulations of the He-shell flash convection by Herwig et al. (2007). The value for \(f_{\text{bot}}\) is consistent with a number of abundance observables of \(\alpha\)-process elements and H-deficient stars (Herwig 2005; Werner & Herwig 2006; Battino et al. 2016). Our stellar models evolve through the phases of convective H and He core burning and on the RAWD phase they experience convective He-shell burning, therefore the above constraints for the values of \(f\) can be applied to them. We provide further support from new hydrodynamic simulations for our choice of \(f_{\text{top}}\) in Section 3.

When the CO WD model cools down to \(\log_{10} L/L_\odot = -2,\) we initiate a slow accretion of H-rich matter on it with \(M_{\text{acc}} = 10^{-8} \text{M}_\odot \text{yr}^{-1}\) using the MESA mass-change control parameter in a new inlist with the same as before input physics. A higher value of \(M_{\text{acc}}\) at this stage would result in MESA iterations having not converged. We assume that the accreted matter has the initial chemical composition of the binary. Because the accretion rate is now lower than the one required for stable H burning, our model exhibits mild H-shell flashes, each of them being followed by the expansion of its envelope. To stop the model from becoming a red giant, we enforce a mass loss by implementing the MESA super-Eddington wind prescription with an artificially reduced value of \(L_{\text{max}}\) so that the model returns to the accretion phase and continues to make nova-like loops on the Hertzsprung–Russel diagram (a grey curve in the top-left-hand panel of Fig. 1; it is shown only for the progenitor of the solar-metallicity RAWD model A). During the stable H burning at a higher accretion rate a RAWD remains near the high-\(L_{\text{eff}}\) and high-\(L\) ‘knee’ of a nova loop (e.g. Wolf et al. 2013), therefore we switch the accretion rate to a higher value in a model that is located near the knee. We adjust a value for \(M_{\text{acc}}\) that would guarantee stable H burning and allow relatively large time-steps between consecutive evolutionary models.

The RAWDs spend most of their time stably burning the accreted H at the knees of nova loops, during which they should be seen as super-soft X-ray sources, unless being obscured by the ejected circum-binary matter (van den Heuvel et al. 1992; Lepò & van Kerkwijk 2013; Woods & Gilfanov 2016). Just as in thermal-pulse AGB stars, when a critical mass of He is accreted from the H-burning shell, a He-shell flash occurs causing an expansion of the accreted envelope (Figs 2 a and f). Whereas the mass loss via the radiation-driven super-Eddington wind becomes less efficient at sub-solar metallicities, a fast and efficient mass loss can still be assumed for a star in a close binary system when it expands and overflows its Roche lobe. Therefore, a slightly modified MESA scheme for the Roche-lobe wind has been implemented in the present work to model the mass loss by our RAWD models during their expansion driven by the He-shell flashes. While the MESA code approximates
the Roche-lobe mass-loss rate as

\[ M_{RL} = M_{RL,0} \exp \left( \frac{R - R_{RL}}{H_{RL}} \right), \]

where \( M_{RL,0} \) is a base value for which we use \( 10^{-3} M_\odot \text{ yr}^{-1} \), \( R_{RL} \) is the Roche-lobe radius of the mass-losing star, and \( H_{RL} \) is the Roche-lobe wind scale height, we prefer to use the following prescription:

\[ M_{RL} = M_{RL,0} \left( \frac{R}{R_{RL}} \right)^6 \]

for \( R > 0.2 R_{RL} \). The implementation of our prescription for the Roche-lobe mass loss, that is equivalent to the MESA approximation for \( |R - R_{RL}| \ll R_{RL} \) if one uses \( H_{RL} = R_{RL}/6 \), results in a smoother evolution of the RAWD models. After having lost a certain amount of mass with the Roche-lobe wind, which determines the mass retention efficiency, our RAWD model returns to the knee, and the H mass accretion resumes.

### 2.2 The RAWD i-process nucleosynthesis

The i process in an RAWD commences when the top of the He convective zone reaches the bottom of its H-rich surface layer, soon after the He-shell flash peak luminosity. At this moment, the He-shell convection begins to ingest protons (the left dotted line in Fig. 2(d) marks the moment when the H luminosity suddenly increases by more than two orders of magnitude as a result of the beginning H ingestion). It should be noted that our MESA computations find such H ingestion in all of our RAWD models even when \( f_{\text{top}} = 0 \) (Paper I), i.e. even when convective boundary mixing at the top boundary of the He convective zone is not included (although the H ingestion rate \( M_{\text{H}} \) does positively correlate with \( f_{\text{top}} \)).

The i process in our RAWD models is simulated in post-processing computations similar to those carried out to model the i-process nucleosynthesis in Sakurai’s object (Herwig et al. 2011; Denissenkov et al. 2018). These computations use the NuGrid...
Figure 2. The evolution of luminosities, total mass, H-ingestion rate and temperature profile for the RAWD model C from the middle-left-hand panel of Fig. 1. Panel a shows the changes of the total luminosity with the radius during eleven consecutive He-shell flashes (the evolution goes counterclockwise in the cycles) for which the corresponding changes of the H and He luminosities are shown in panel b and the changes of the total mass in panel c. The expansion of the RAWD is bounded by its assumed $2 M_\odot$ Roche-lobe radius (equation 2). Panels d and e show how the luminosities and H-ingestion rate change with time during the 6th He-shell flash (the black line segment in panel a). The wiggles on the $L_H$ and $\dot{M}_H$ curves are caused by the fact that the CBM prescription (1) is non-local and therefore its implementation in MESA is decoupled from the solution of stellar structure equations. Panel f: the evolution of the temperature profile in the RAWD during the 6th He-shell flash. The black dashed curve was used in the post-processing computations of nucleosynthesis in the He convective zone.

multizone post-processing nucleosynthesis parallel code mppnp (Pignatari et al. 2016) customized for the H-ingestion He-shell problem. The input data for this code include a static or time-dependent structure of the He convective zone (this work uses the first option), i.e. the radius $r$, temperature $T$, density $\rho$, and convective diffusion coefficient $D_{conv}$ at each point of its mass mesh, the chemical compositions of the He zone and of the ingested matter, the mass ingestion rate $\dot{M}_{ing}$ and its duration $t_{ing}$.

Whereas the ingested matter has the initial chemical composition of the binary, the composition of the He convective zone at the beginning of H ingestion is obtained by processing the initial mixture through complete H burning followed by its processing via partial He burning, until the increasing $C$ abundance matches its value from the corresponding MESA RAWD model, for which we use the NuGrid single-zone code ppm (cf. Denissenkov et al. 2018).

Our i-process nucleosynthesis simulations include $\sim$1000 isotopes and $\sim$15000 reactions. The reaction rates for these simulations are taken from the same list of references as in Denissenkov et al. (2018). We adopt an equally spaced 100-zone mass grid for the He-shell region by interpolating the stellar structure variables to the new mesh. At each time-step $\Delta t$, we add $X_k^0 \dot{M}_{ing} \Delta t$ mass of the $k$th isotope from the envelope to the top $\Delta M = 1 - 4 \times 10^{-4} M_\odot$ of the He shell that occupies $\sim$10 mass zones, as described in Appendix A.

2.3 The mass ingestion rate and duration

The mass ingestion rate $\dot{M}_{ing}$ is determined from a combination of constraints from 3D simulations (see Section 3) and the MESA RAWD stellar evolution simulations. Given the modelling choices described in the previous section the 1D stellar evolution simulations predict the ingestion of H-rich envelope material into the He-shell flash convection zone as it expands outward in Lagrangian
coordinate. The protons ingested into the convection zone lead to a H-burning luminosity \( L_H \) which reflects the mass ingestion rate through the relation \( M_H = X_{\text{surf}} M_{\text{ing}} \), where \( X_{\text{surf}} \) is the H mass fraction at the RAWD surface, and

\[
M_H \approx \frac{L_H}{\varepsilon_H},
\]

where \( \varepsilon_H \) is the energy released per one gram of burned H.

Because only the first two reactions of the CNO cycle are fast enough to occur in the He-shell convective zone during a convective overturn time, we assume that \( \varepsilon_H = 0.667 \epsilon_{\text{CNO}} \), where \( \epsilon_{\text{CNO}} \approx 6.3 \times 10^{18} \text{erg g}^{-1} \) is the energy released per one gram of H transformed into He in the full CNO cycle, and the factor 0.667 is the fraction of this energy produced per one gram of consumed H in the reactions \( ^{12}\text{C}(p, \gamma)^{13}\text{N} \) and \( ^{13}\text{N}(\nu, p)^{12}\text{C} \).

For the duration of mass ingestion \( t_{\text{ing}} \), we adopt the values estimated from our MESA RAWD models. We use \( M_H(t) \) curves, like the one shown in Fig. 2(e), to estimate a time interval \( t_{\text{ing}} \) during which \( M_H \) remains close to its maximum value wiggling around some nearly constant mean value, the latter giving us an estimate of \( M_H \). We check that the estimated values of \( t_{\text{ing}} \) do not exceed their upper limits constrained by the condition \( t_{\text{ing}} < M_{\text{env}}/M_{\text{ing}} \). The adopted values of \( M_{\text{ing}} \) and \( t_{\text{ing}} \) used in the different simulations are summarized in Table 1. Although this is a somewhat subjective method of choosing the \( M_{\text{ing}} \) and \( t_{\text{ing}} \) input data for the post-processing nucleosynthesis simulations, we think its accuracy is consistent with our using a static temperature profile, like the black dashed curve in Fig. 2(f), in these simulations.

### 3 Ingestion Rates and Convective Boundary Mixing Parameters from 3D Hydrodynamic Simulations

The ingestion of material from the stable layer into the convection zone is the result of complex mixing processes at the convective boundary that may involve global, large-scale flow modes revealed in full 4π 3D hydrodynamic simulations (Woodward, Herwig & Lin 2015). As mentioned in Section 2.1 we adopt in 3D simulations the exponentially decaying CBM model with an efficiency parameter \( f_{\text{loop}} \) to describe this mixing.

Both the mass ingestion rate and the convective boundary parameter \( f_{\text{loop}} \) can be determined from hydrodynamic simulations, as demonstrated by Jones et al. (2017). For this purpose we have performed 3D hydrodynamic simulations of the He-shell flash convection zone and H ingestion in a RAWD using the PPMstar code (Woodward et al. 2015). It is an explicit Cartesian-grid-based code for 3D hydrodynamics built around the Piecewise-Parabolic Method (PPM; Woodward & Colella 1981, 1984; Colella & Woodward 1984; Woodward 1986, 2007). The code advects the fractional volume of the lighter fluid in a two-fluid scheme using the Piecewise-Parabolic Boltzmann method (PPB; Woodward 1986; Woodward et al. 2015). Thanks to PPB’s use of sub-cell information, it needs two to three times fewer grid cells along all three axes than PPM to reach the same level of fidelity in the advection of a quantity, like the multifluid mixing fraction, whose value is conserved along stream lines. The code was designed with strong emphasis on parallel efficiency and it has performed past simulations of shell convection on up to 440000 CPU cores on the NCSA Blue Waters computer (Herwig et al. 2014; Woodward et al. 2015).

The radial stratification of simulations is based on model A from Paper I. Like model A in this paper, model A in Paper I had the solar initial chemical composition, but it had a bit lower mass of 0.65 M⊙. We consider the point in time 7.44hr after the beginning of the second He-shell flash in that model, when its He luminosity has dropped to \( 4.10 \times 10^8 \text{L}_\odot \) from its maximum value of \( 7.4 \times 10^{10} \text{L}_\odot \), and H ingestion into the He shell has just started.

The initial stratification of the 3D simulations follows the same approach as in Woodward et al. (2015) and approximates the 1D model with three polytropes: a lower stable layer (radial range 6 Mm < \( r < 33.5 \) Mm), and an upper stable layer (33.5 Mm < \( r < 50.0 \) Mm).

We neglect radiation pressure, which contributes less than 25 per cent to the total pressure in the 1D model, and we use the equation of state for a monatomic ideal gas. To obtain a similar overall stratification with the slightly different equation of state, we use a small mean molecular weight \( \mu_0 = 0.3 \) for the fluid \( F_1 \) initially filling the upper stable layer. The rest of the simulation domain contains fluid \( F_2 \) with \( \mu_2 = 1.4 \). The two fluids are allowed to react with each other via the \( ^{12}\text{C}(p, \gamma)^{13}\text{N} \) reaction, assuming that \( F_1 \) contains 88.6 per cent of protons and \( F_2 \) contains 20.4 per cent of \( ^{12}\text{C} \) by number. The subsequent decay of \( ^{13}\text{N} \) is not considered. Convection in the He shell is driven by volume heating applied between the radii 7.9 Mm and 8.9 Mm.

The 3D simulations are done in 4π geometry on a Cartesian grid. We measure the luminosity dependence of the mass ingestion rate using runs E8, E13, and E15 (see Table 2), which cover a range of 1.4 dex in the driving luminosity \( L_{\text{He}} \) at the grid resolution of 7683.

### Table 1. Summary of the one-dimensional RAWD simulation parameters (\( t_{\text{He}}^{\text{max}} \) is the He luminosity at the beginning of H ingestion).

<table>
<thead>
<tr>
<th>model</th>
<th>[Fe/H]</th>
<th>( M_{\text{WD}} (M_\odot) )</th>
<th>( M_{\text{acc}} (M_\odot \text{ yr}^{-1}) )</th>
<th>( \log_{10}(L_{\text{He}}^{\text{max}}/L_\odot) )</th>
<th>( \log_{10}(L_{\text{He}}/L_\odot) )</th>
<th>( M_{\text{ing}} (M_\odot \text{ yr}^{-1}) )</th>
<th>( t_{\text{ing}} (\text{yr}) )</th>
<th>( \eta (%) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0.0</td>
<td>0.70</td>
<td>( 2.6 \times 10^{-7} )</td>
<td>10.9</td>
<td>9.1</td>
<td>( 2.3(35) \times 10^{-12} )</td>
<td>0.17(0.024)</td>
<td>–</td>
</tr>
<tr>
<td>B</td>
<td>−0.7</td>
<td>0.71</td>
<td>( 1.7 \times 10^{-7} )</td>
<td>9.5</td>
<td>8.5</td>
<td>( 2.0 \times 10^{-12} )</td>
<td>0.054</td>
<td>4.9</td>
</tr>
<tr>
<td>C</td>
<td>−1.1</td>
<td>0.71</td>
<td>( 1.5 \times 10^{-7} )</td>
<td>9.3</td>
<td>8.4</td>
<td>( 4.0 \times 10^{-12} )</td>
<td>0.042</td>
<td>4.9</td>
</tr>
<tr>
<td>D</td>
<td>−1.55</td>
<td>0.71</td>
<td>( 1.5 \times 10^{-7} )</td>
<td>9.3</td>
<td>8.5</td>
<td>( 4.2 \times 10^{-12} )</td>
<td>0.083</td>
<td>9.6</td>
</tr>
<tr>
<td>E</td>
<td>−2.0</td>
<td>0.74</td>
<td>( 1.7 \times 10^{-7} )</td>
<td>8.7</td>
<td>8.1</td>
<td>( 3.5 \times 10^{-12} )</td>
<td>0.060</td>
<td>27</td>
</tr>
<tr>
<td>F</td>
<td>−2.3</td>
<td>0.75</td>
<td>( 1.5 \times 10^{-7} )</td>
<td>9.2</td>
<td>8.6</td>
<td>( 2.4 \times 10^{-11} )</td>
<td>0.058</td>
<td>19</td>
</tr>
<tr>
<td>G</td>
<td>−2.6</td>
<td>0.75</td>
<td>( 1.5 \times 10^{-7} )</td>
<td>8.5</td>
<td>8.0</td>
<td>( 6.7 \times 10^{-12} )</td>
<td>0.087</td>
<td>29</td>
</tr>
</tbody>
</table>

### Table 2. Summary of our 3D PPMstar simulations.

<table>
<thead>
<tr>
<th>run</th>
<th>grid</th>
<th>( \log_{10}(L_{\text{He}}/L_\odot) )</th>
<th>( L_{\text{He}}/L_\odot )</th>
<th>( M_{\text{ing}} (M_\odot \text{ yr}^{-1}) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>E8</td>
<td>7683</td>
<td>9.95</td>
<td>0.025</td>
<td>( 2.32 \times 10^{-10} )</td>
</tr>
<tr>
<td>E10</td>
<td>1536</td>
<td>9.95</td>
<td>0.034</td>
<td>( 3.26 \times 10^{-10} )</td>
</tr>
<tr>
<td>E13</td>
<td>7683</td>
<td>10.65</td>
<td>0.019</td>
<td>( 9.45 \times 10^{-10} )</td>
</tr>
<tr>
<td>E15</td>
<td>7683</td>
<td>11.35</td>
<td>0.020</td>
<td>( 5.04 \times 10^{-9} )</td>
</tr>
</tbody>
</table>
and one high-resolution run E10 (1536³) with the same driving luminosity as its 768³ equivalent E8.

Hydrogen ingestion starts as soon as the first upwelling plumes reach the upper convective boundary. After a few convective overturns a balance between hydrogen ingestion and burning is reached and the convective-reactive flow becomes quasi-stationary. We do not find a GOSH in these H-ingestion simulations which differs from the results found in the case of Sakurai’s object (Herwig et al. 2014). This corresponds to the result from 1D stellar evolution reported in Paper I and also found in most simulations here that the H-ingestion does not cause a split of the convection zone, with the exception of model A where a split happens in very late phases (see Section 4).

Fig. 3 shows that the ingestion process is dominated by large scales. The average hydrogen luminosity $L_H$ reaches 2–3 per cent of the driving luminosity $L_{He}$ (Table 2). Ingestion events localized in time and space can have a much stronger influence on the flow than these small values suggest, but they are not strong enough to launch a global ingestion instability such as the GOSH phenomenon observed by Herwig et al. (2014) in their 3D simulations of Sakurai’s object.

The amount of mass $M_{ing}(t)$ ingested into the convection zone by a time $t$ is the sum of mass $M_p(t)$ present in the convection zone at this time and mass $M_b(t)$ burnt in the convection zone by this time. The radius $r_{ub}$ of the upper boundary of the convection zone increases in time as a result of both mass ingestion and thermal expansion. We define $r_{ub}$ to be the radius at which the radial gradient of the root-mean-square horizontal velocity $v_h(r)$ reaches a local maximum and we integrate the density of the ingested fluid up to the radius $r_{top} = r_{ub} - H_{v_{h u b}}$, where the velocity scale height $H_{v_{h u b}} = (\partial v_h/\partial r)^{-1}$ is evaluated at $r_{ub}$. The subtraction of $H_{v_{h u b}}$ mitigates issues related to the large contrast in the concentration of the ingested fluid between the boundary region and the bulk of the convection zone (for details, see Jones et al. 2017). To find the burnt mass $M_b(t)$, we compute the mass burning rate from spherically averaged profiles of temperature, density, and fractional volume of ingested fluid at regularly spaced points in time; $M_b(t)$ is then obtained by time integration. The resulting time dependence of $M_p, M_b,$ and $M_{ing}$ in run E10 is shown in Fig. 4. We obtain the ingestion rate $\dot{M}_{ing}$ by fitting a straight line to $M_{ing}(t)$.

Fig. 5 shows that the ingestion rate scales in proportion to the driving luminosity $L_{He}$, in agreement with the results of Jones et al.
(2017) on the O-shell convection in massive stars. This is likely to be caused by the fact that the ingestion rate is limited by the amount of work needed to be done to overcome the buoyancy of the ingested material, as argued by Spruit (2015). The ingestion rate in run E8 (7680) is only 26 per cent lower than in run E10 (15360) of the same luminosity, which provides an idea of the resolution dependence of these entrainment rate results.

The ingestion rates measured in 1D models B–F are close to the scaling relation established by the 3D hydrodynamic simulations, which may not have been expected given the numerous differences between the 1D and 3D models. For the RAWD models in Fig. 5, we have used the He-shell luminosity at the moment when the top of the He convective zone reaches the bottom of the H-rich envelope and, as a result of this, the H-burning luminosity quickly increases. These luminosities are somewhat lower than their corresponding peak He luminosities (e.g. panel d in Fig. 2). In any case, the ingestion rates adopted for our nucleosynthesis simulations are consistent with the results obtained with the 3D hydrodynamic simulation in the sense that they are close to the scaling relation between the driving luminosity of the convection and the ingestion rate established by three 768-grid hydrodynamic simulations.

We can also obtain information on the convective boundary mixing efficiency at the top of the He-shell flash convection zone. As in Jones et al. (2017) we determine the $f$ parameter from the evolution of the spherically averaged radial profiles of the abundance of the H-rich fluid that is entrained into the He-shell flash convection zone (Fig. 6). This is done by solving the inverted Lagrangian diffusion equation which gives the diffusion coefficient profile at a time $t$ by means of a 1D diffusion process. The time difference $\Delta t$ is usually taken to be one or a few convective overturning time-scales. The details of this procedure have been improved somewhat over the approach in Jones et al. (2017) and will be described elsewhere in detail. For the high-resolution E10 simulation we determine $f_{\text{top}} = 0.434$.

This value is larger than the one appropriate for our 1D simulations because the E10 hydrodynamic simulation has been performed at a driving luminosity that is 141 times higher than, for example, the He-burning luminosity in the stellar evolution run C (log $L_{\text{He}} = 7.8$). In order to scale the $f$ value obtained in our higher luminosity hydro simulation to the actual lower luminosity of the stellar evolution RAWD model we use the scaling relationship

$$f_{\text{top}} \propto L_{\text{drive}}^{1/3}$$

(4)

between the driving luminosity of a shell convection and the convective boundary mixing parameter $f_{\text{top}}$ at the top of the convection zone. This relationship has been derived from a series of new 1536-grid 3D hydrodynamic simulations of O-shell convection as in Jones et al. (2017), but with an improved version of the PPMSTAR code (Fig. 7). These simulations, the $f$ determination and the resulting scaling law will be described in detail elsewhere. However, this relationship can be motivated within a simplistic picture in which the CBM $f$ parameter is a measure of how much convective plumes can deform the convective boundary and penetrate into it before being decelerated by their negative buoyancy upon entering the stably stratified regions. It would be the momentum of the convective plume that determines the level of penetration and boundary deformation. The convective velocity scales with one-third power with the luminosity (Biermann 1932; Porter & Woodward 2000; Müller & Janka 2015; Jones et al. 2017). The scaling relation equation (4) reflects this reasoning: $f \propto L^{1/3}$.

Applying relation equation (4) to scale $f_{\text{top}} = 0.434$ from the E10 driving luminosity of log $L_{\text{He}} = 9.95$ to the driving luminosity of the stellar evolution model C at the time of H ingestion (log $L_{\text{He}} = 7.8$) we obtain $f_{\text{top}} = 0.08$. This provides strong support for the value $f_{\text{top}} = 0.10$ that we have adopted in the RAWD stellar evolution simulation which are all at a similar He-burning luminosity at the time of ingestion.

4 RESULTS

4.1 The RAWD multicycle evolution

Using the methods described in Section 2.1, we have computed the evolution of seven RAWD models with the metallicities and WD masses listed in Table 1 along with other model parameters. The evolutionary tracks of six of these models are shown in Fig. 1, where their initial masses are also indicated. Except the solar-metallicity model A, we have simulated the RAWD evolution for many cycles, typically more than five, (orange curves in Fig. 1). The multicycle evolution of the solar-metallicity RAWD models was discussed in Paper I. Black curve segments in Fig. 1 highlight the relatively short-living evolutionary phase of the He-shell TP whose serial number is specified for each model and that has been chosen for the i-process post-processing nucleosynthesis computations in Section 4.2.
Figure 8. The changes of the total mass of our RAWD models caused by the accretion and Roche-lobe mass loss.

All of the RAWD models have nearly the same initial central temperature with $\log_{10} T_c \approx 7.2$ and use the same WD Roche-lobe radius $R_{\text{RL,WD}} = 2 R_\odot$ in equation (2) that corresponds to the orbital period $P \approx 1.2$ d for a secondary mass of $\sim 2 M_\odot$.

The new computations of the RAWD multicycle evolution confirm the conclusion about their low mass retention efficiency (the last column of Table 1 and Fig. 8) made in Paper I, at least for $\text{[Fe/H]} \gtrsim -2$. A new result is that $\eta$ increases when $\text{[Fe/H]}$ decreases below $-2$ to fractions of $\approx 20$ to 30 per cent in models E, F, and G. The 0.75 $M_\odot$ RAWD model with $\text{[Fe/H]} = -2.6$ (model G) has the highest value of $\eta = 29$ per cent and a lower He peak luminosity of $\log_{10} L_{\text{He}}^{\text{max}} = 8.5$ (the last row in Table 1) compared to models E and F. According to these models, the SD channel of SNIa progenitors may still work at very low metallicities.

The revealed trend of $\eta$ with $\text{[Fe/H]}$ is probably caused by the lower opacity of the metal-poor accreted matter. It allows the accumulated He layer to cool down and therefore be compressed by the gravity more efficiently. As a result, the He flash starts at a lower mass of the He shell and achieves a lower peak luminosity, which is reflected in different amplitudes of the RAWD mass changes in the models with $\text{[Fe/H]} < -1.55$ (E and F) compared to the models with $\text{[Fe/H]} \geq -1.55$ (B, C, and D) in Fig. 8 and in their different values of $\log_{10} L_{\text{He}}^{\text{max}} / L_\odot$ (Table 1). Besides, during the expansion of the RAWD following the He-shell flash, its envelope cools down faster, because of the lower opacity, in the low-metallicity models, and the RAWD returns to the accretion phase after having lost a smaller fraction of the accreted matter.

As for the RAWD models with $\text{[Fe/H]} \geq -1.55$, they stubbornly want to expand to red giant dimensions and it is only because of our implementation of the Roche-lobe mass loss that they can expand only up to $R \approx R_{\text{RL,WD}}$ (Fig. 2a) and retain this radius until a significant fraction of the accreted matter is gone with the wind. To test if this behaviour is affected by our choice of the MESA mass-loss algorithm, we have switched to the super-Eddington wind prescription in model C, enforcing it to work only when $R \approx R_{\text{RL,WD}}$. With this modification, we have reproduced the results obtained for model C with the Roche-lobe mass-loss prescription.

We have also addressed the frequently raised concern that a much larger number of He-shell flashes than we have simulated in our RAWD models could lead to significant changes in the RAWD multicycle evolution. To test this, we have extended the computed number of the He-shell flashes in model C up to 42 from the initial 10. The results of this long-run simulations are plotted in Fig. 9. Its top panel demonstrates that the WD central temperature has increased only by 6 per cent. Our analysis of the RAWD mass variations presented in the middle panel shows that the mass retention efficiency has varied between 5.8 per cent and 8.8 per cent in this run, still remaining below 10 per cent, as it was in the initial 10 cycles. Finally, the bottom panel reveals that the He peak luminosity has not changed at all.

4.2 The RAWD i-process nucleosynthesis yields

Before presenting the yields, we will summarize how we choose the input data for the RAWD i-process nucleosynthesis computations. For this purpose, we use model C (Fig. 2), as an example. We begin with selecting a representative RAWD He-shell flash that does not stand out; therefore, this cannot be the first He-shell flash that usually is stronger than the others. The only exception to this rule in this paper is the solar-metallicity model A. We find it to be much more difficult to compute multiple cycles with stable H burning interrupted by strong He-shell flashes at a high metallicity, probably because of the high opacity of the envelope matter. Because the multicycle evolution of the solar-metallicity RAWD models has already been discussed in Paper I, and the i-process yields calculated for the first He-shell flash in the solar-metallicity RAWD model in this work are similar to those presented in Paper I, we have...
followed only one He-shell flash in the new model A. As for model C, we have chosen the 6th flash (panels a, b, and c in Fig. 2). We have used its corresponding H-burning luminosity (panel d) to estimate the H-ingestion rate with equation (3) (the purple curve in panel e). The dashed line in panel e is our estimate of an average $\dot{M}_H$ value for this model that was used to calculate the parameter $M_{\text{ing}} = (M_H/X_{\text{start}})$ listed in Table 1. The vertical dotted lines in panel e constrain $t_{\text{ing}}$. The left line is chosen close to the beginning of H ingestion. The position of the right line is less certain. It marks the beginning of the fast decline of $M_H$. The H-ingestion time in the RAWD models is usually less than 1 month ($t_{\text{ing}} \lesssim 0.08$ yr), except the first phase of H-ingestion in model A. Like in the solar-metallicity model A from Paper I, the updated model A has two phases of H ingestion, the longer lasting slow-ingestion phase with $M_{\text{ing}} = 2.2 \times 10^{-12} \, M_\odot \, s^{-1}$ and $t_{\text{ing}} = 0.17$ yr, followed by the shorter fast-ingestion phase with $M_{\text{ing}} = 3.5 \times 10^{-11} \, M_\odot \, s^{-1}$ and $t_{\text{ing}} = 0.024$ yr, both included in our nucleosynthesis computations.

It usually takes about hundred time-steps for the MESA code to evolve a RAWD model through the entire H-ingestion phase, meaning that the time interval between two consecutive models on this evolutionary phase is hundreds of minutes, corresponding to tens of convective turnover times of the He-shell flash convection. This is too long for the i-process nucleosynthesis simulations that usually require a time-step of the order of minutes (Herwig et al. 2011). We cannot reduce the MESA time-steps, because the MLT adopted in MESA to describe convection is formulated in terms of time and spatial averages. Time steps smaller than about 10 times the convective turnover time would violate this assumption. Therefore, we simply take the temperature, density, radius, and MLT convective diffusion coefficient profiles from a MESA model in the middle of the H-ingestion phase, when the accreted envelope is already expanding (e.g. the dashed $T$ profile in panel f), and use them to set up our post-processing nucleosynthesis simulations. Note that Herwig et al. (2011) were able to reproduce the surface abundances of heavy elements measured in Sakurai’s object by Asplund et al. (1999) using a 1D model similar to this one.

The post-processing simulations of the i-process nucleosynthesis were carried out using the methods described in Section 2.2. The nucleosynthesis simulations provide the abundance distributions in the He convective zone and the surface abundances at the top of the He shell for the selected (∼1000) isotopes for $0 \leq t \leq t_{\text{ing}}$. At the end, we allow the surface abundances to decay for 1 Gyr. Fig. 10 shows the evolution of the maximum neutron number density in the He shell for our RAWD models. The peak value of the $N_{n, \text{max}}(t)$ curves increases with a decrease of $[\text{Fe}/H]$ because the total mass fraction of the isotopes that capture neutrons decreases with the metallicity, while the production of neutrons, which is controlled by the $^{12}\text{C}$ abundance in the He shell and the total amount of ingested H, remains approximately the same. In other words, the neutron source is primary, while the i-process seeds are secondary. The steep increase of $N_{n, \text{max}}$ at the end of its evolution in model A marks the beginning of the short fast-ingestion phase that appears to be common for solar-metallicity RAWDs (Paper I). It demonstrates that $N_{n, \text{max}}$ increases with $M_{\text{ing}}$ for models with the same metallicity.

Fig. 11 shows the final surface elemental abundances divided by the solar abundances from Asplund et al. (2009) of the RAWD models. As in the $s$ process, the global heavy-element distribution shifts to higher mass elements at lower metallicity (Clayton 1968; Busso et al. 2001). We therefore, generally speaking, expect to see local elemental i-process signatures in higher mass second- and third-peak species at lower metallicity, whereas the i-process signature may be most prominently detected in lower mass, first-peak elements at higher, solar-like metallicities. Accordingly, it had been proposed in Paper I that the solar-metallicity RAWDs could be contributors of first-peak elements to the Solar system abundance distribution. Côté et al. (2018) have used our RAWD i-process yields in a framework that included Galactic chemical evolution and binary-star population synthesis models to confirm this hypothesis.

An example for solar-like metallicity i-process abundances includes those of Sakurai’s object that indeed shows large enhancements in the first peak n-capture elements (around $Z = 40$) as shown in Fig. 11. The details of the abundance patterns of Sakurai’s object are, however, better reproduced by the very-late TP post-AGB star models of Herwig et al. (2011) that feature peak neutron...
densities and H ingestion rates that are both more than two orders of magnitude higher compared to the RAWD models.

4.3 A possible relation of the metal-poor RAWDs to the CEMP-r/s stars

An example for i-process abundance patterns in the second-peak elements expected at lower metallicity may be found in the subclass of carbon-enhanced metal-poor (CEMP) stars with abundance patterns that appear to be enhanced with both r- and s-process elements (e.g. Beers & Christlieb 2005; Masseron et al. 2010; Bisterzo et al. 2012; Lugaro et al. 2012). Dardelet et al. (2015) and Hampel et al. (2016) have done one-zone nucleosynthesis simulations of i-process conditions to demonstrate that the abundances of heavy elements observed in these CEMP-r/s stars can be reproduced by an n-capture process with neutron densities $n = 10^{12} – 10^{15}$ cm$^{-3}$.

As an example, Fig. 12 shows the best $\chi^2$ fit of the i-process elemental yields from our RAWD models G and F to the surface chemical composition of the CEMP-r/s star CS31062-050 that has the metallicity [Fe/H] = $-2.42$ (Johnson & Bolte 2004) intermediate between those assumed for our models F and G. The only exception to the otherwise excellent agreement is the discrepant Ba abundance that requires further investigation. For other CEMP-r/s stars, a good agreement with the RAWD i-process yields can be found as well, including their lower Ba abundances. The low-metallicity RAWD models are at this point the first and only models in which nucleosynthesis calculation directly post-processing complete stellar evolution models can reproduce the complete abundance patterns observed in CEMP-r/s stars. We therefore propose that CEMP-r/s stars that have been well reproduced with i-process models should be referred to as CEMP-i stars.

Our findings suggest a new scenario for the formation of CEMP-r/s, or in this case CEMP-i, stars, that takes into account our finding that the RAWDs with [Fe/H] $\lesssim -2$ may reach the Chandrasekhar mass and explode as SNeIa. It is based on the fact that the mass retention efficiency of our RAWD models significantly increases when [Fe/H] decreases below $-2$ (the last column of Table 1), and it is supported by our calculations of the evolution of binary-star parameters. For these calculations, we have used the isotropic re-emission model of mass transfer in which a fraction $\beta$ of matter accreted by the primary star (RAWD) is lost from the binary system. In our case, $\beta = 1 - \eta$, where $\eta$ is the mass retention efficiency. This model and its equations are described by Postnov & Yungelson (2014) in their section 3.3.3. We have solved these equations to model the evolution of the semimajor axis $a$, the mass ratio $q = M/M_2$, and the RAWD mass $M_1$ for two sets of the binary initial parameters. In both cases, we start with $M_1 = 2.5 M_\odot$ and use $M_2 = -1.5 \times 10^{-3} M_\odot$ yr$^{-1}$. The first case assumes that $\eta = 10$ per cent, which includes the RAWD models with [Fe/H] $\gtrsim -2$, while the second case has $\eta = 30$ per cent and demonstrates a possible evolution of the binary-system parameters for RAWDs with [Fe/H] $\lesssim -2$. In both cases, we have stopped the calculations at $q = 2$.

In the first case, $M_1$ grows from 0.75 $M_\odot$ to only $\sim 0.9 M_\odot$, while in the second case $M_1$ starts growing from 0.85 $M_\odot$ and it does reach the Chandrasekhar limit, and the RAWD ends its life as a SNII. If this is true, then some of the present-day CEMP-i stars could be former tertiary members of hierarchical triple systems in which they had been orbiting a close binary system with a RAWD. A series of dozens He-shell flashes on the RAWD, each being followed by the RAWD expansion and mass loss, could enrich the tertiary star with the products of i-process nucleosynthesis. This enrichment scenario is similar to the one proposed to explain abundance anomalies in CEMP-s stars by accretion of material lost by their AGB star binary companions (e.g. Abate, Pols & Stancliffe 2018, and references therein). A difference is that tertiary stars in hierarchical triple systems are farther away from their polluting primary components. This should result in a stronger dilution of the accreted material. But, on the other hand, heavy elements produced in the i process in AGB stars are already diluted in their relatively massive envelopes before they are ejected, while the dilution in a thin envelope of a RAWD is negligible. If thermohaline mixing in envelopes of accreting stars leads to an even stronger dilution depends on the efficiency (time-scale) of this mixing, which still remains uncertain (Stancliffe et al. 2007; Denissenkov & Pinsoneault 2008). For example, thermohaline mixing can be suppressed by strong horizontal turbulent diffusion (Denissenkov 2010). Finally, when the RAWD exploded as SNII, the tertiary star, that became a CEMP-i star by that moment, would leave the triple system because of a decreased gravitational pull to the center of mass that has suddenly lost $\sim 1.4 M_\odot$. Those CEMP-i stars would not be binaries anymore. If the RAWD does not explode as SNII, the CEMP-i star would be possibly in a wider orbit around compact binary and would show a long binary period superimposed with the very short period of the compact RAWD binary. Thus, CEMP-i stars can be both single stars and binaries in this scenario. Our scenario is supported by the recent finding that the fractions of binary and triple star systems in stellar populations significantly increase with a decreasing metallicity (Fuhrmann et al. 2017; Badenes et al. 2018; Moe, Kratter & Badenes 2019).

This scenario can potentially explain the observational bias against finding CEMP-r/s stars in globular clusters that may be caused by the destruction of wide triple systems through close star encounters in dense cores of globular clusters. Our scenario differs from the triple-system scenario for the formation of CEMP-r/s stars discussed by Abate, Stancliffe & Liu (2016), in which a primary massive star was assumed to produce the r-process elemental abundances during its SN explosion and the presence of a secondary AGB star was required to make s-process elements. The plausibility of
our scenario could be checked with triple-star population synthesis simulations, but this is out of the scope of this paper.

5 SUMMARY

We have used the MESA stellar evolution code (Paxton et al. 2011, 2013) to compute the models of CO WDs rapidly accreting H-rich matter, assuming that this matter is donated by normal (main-sequence, sub-giant, or red giant branch) components of the WDs in close binary systems. Such stellar configurations can result from common envelope events, after the primary AGB star components fill their Roche lobes, loose almost entire envelopes above the WD cores in unstable mass transfers, and the binary systems become tighter via the transformation of their orbital energies into the kinetic energy of the ejecta. When the secondary components of these post-common-envelope systems expand and fill their own Roche lobes by or after the end of the main-sequence evolution, they begin to donate H-rich matter to their, by this time cooled-down, WD primary components. We assume that the mass accretion rate is rapid enough, $M_{\text{acc}} \sim 10^{-7}$ $M_\odot$ yr$^{-1}$, for the accreted H to be stably burning on the WD surface, resulting in the accumulation of a He shell. When its mass reaches a critical value, the He shell will experience a thermal flash in which some fraction of He will be transformed into C. If the sequence of the stable H burning intermittent with the He-shell flashes is not accompanied by a significant mass-loss by the WD, then its mass may eventually reach the Chandrasekhar limit. This convective–reactive process transforms almost every proton ingested at the top of the He shell into a neutron at its bottom; the WD, then its mass may eventually reach the Chandrasekhar limit. If the sequence of the stable H burning intermittent with the He-shell flashes is not accompanied by a significant mass-loss by the WD, then its mass may eventually reach the Chandrasekhar limit.

$\sim 10^{12}$ – $10^{15}$ cm$^{-3}$. These values are intermediate between those characteristic of the $s$ and $r$ processes; therefore, the ensuing n-capture process is called the i-process (Cowan & Rose 1977).

The important input parameters – the ingestion rate and convective boundary mixing efficiency – have been determined and constrained through a series of 3D hydrodynamic simulations of the RAWD He-shell flash convection. The estimates of the mass ingestion rate obtained from 1D RAWD models are consistent with the 3D hydrodynamic simulations. The convective boundary mixing efficiency parameter adopted in our stellar evolution simulations at the top of the He-shell flash convection boundary is in agreement with the luminosity scaling law for the CBF $f_{\text{bg}}$ parameter presented here.

It is interesting that the i-process nucleosynthesis yields predicted by our metal-poor ([Fe/H] $\lesssim -2$) RAWD models almost perfectly fit the abundances of heavy elements measured in some CEMP-n/s stars (Fig. 12). These are the CEMP-i stars. Given that the same RAWD models have the higher mass retention efficiencies and can potentially become SNeIa, we propose that CEMP-i stars used to be RAWDs. When the RAWD exploded as a SNIa, the tertiary star, polluted by the products of i-process nucleosynthesis that had taken place on the RAWD, got loose from the system and is now seen as a single star. If the RAWD has not exploded yet, the CEMP-i star can still be a member of a triple system and would then show signs of binality.

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APPENDIX A: THE IMPLEMENTATION OF MASS INGESTION IN RAW MODELS

Let us denote $X_i^k$ and $X_o^k$ the mass fractions of the $k$th isotope, respectively, inside and outside the convective He shell in the vicinity of its outer boundary. We assume that the envelope matter ingested into the He shell gets immediately distributed within an ingestion zone with the mass $\Delta M \ll M_{\text{tot}} - M_{\text{bot}}$, where $M_{\text{bot}}$ and $M_{\text{tot}}$ are the mass coordinates of the bottom and the top of the He shell. The input parameters here are $X_i^k, \Delta M$, and the mass ingestion rate $\dot{M}_{\text{ing}}$. Because of a small size of the ingestion zone, we assume that all the isotopes are linearly distributed in it, i.e.

$$X_i^k(M) = X_i^k(M_{\text{max}}) - \left(\frac{M_{\text{max}} - M}{\Delta M}\right) \left[X_i^k(M_{\text{max}}) - X_i^k(M_{\text{min}})\right],$$

where $M_{\text{min}} \leq M \leq M_{\text{max}}$, $M_{\text{max}} = M_{\text{top}}$, and $M_{\text{min}} = M_{\text{bot}} - \Delta M$.

The total mass of the $k$th isotope in this distribution is

$$\Delta M_i^k = \int_{M_{\text{min}}}^{M_{\text{max}}} X_i^k(M) dM = \frac{1}{2} \left[X_i^k(M_{\text{min}}) + X_i^k(M_{\text{max}})\right] \Delta M.$$

After $\Delta M_{\text{ing}} = \dot{M}_{\text{ing}} \Delta t$ of the envelope matter is ingested, the mass of the $k$th isotope in the ingestion zone becomes

$$\Delta M_i^k = \Delta M_i^k + X_i^k \Delta M_{\text{ing}} - \delta M_i^k,$$

where

$$\delta M_i^k = \int_{M_{\text{max}} - \Delta M_{\text{ing}}}^{M_{\text{max}}} X_i^k(M) dM = \frac{1}{2} \left[X_i^k(M_{\text{min}}) + X_i^k(M_{\text{max}})\right] \Delta M_{\text{ing}}$$

is the mass of the $k$th isotope in the mass $\Delta M_{\text{ing}}$ that replaces the ingested mass outside the He shell (we assume that the mixing between the He shell and the envelope does not change the chemical composition, $X_i^k$, of the latter.)

The change of the average mass fraction of the $k$th isotope in the ingestion zone is therefore

$$\langle X_i^k(M) \rangle = \frac{\Delta M_i^k}{\Delta M} = \frac{X_i^k(M_{\text{max}}) - X_i^k(M_{\text{min}})}{\Delta M},$$

or as a ramp increase

$$\langle X_i^k(M) \rangle = X_i^k(M) + 2 \frac{(M - M_{\text{min}})}{\Delta M} \langle X_i^k \rangle$$

for $M_{\text{min}} \leq M \leq M_{\text{max}}$. In this work, we have chosen the latter option.

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