On the Impact of $^{22}\text{Ne}$ on the Pulsation Periods of Carbon–Oxygen White Dwarfs with Helium-dominated Atmospheres

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Abstract

We explore changes in the adiabatic low-order g-mode pulsation periods of 0.526, 0.560, and 0.729 $M_\odot$, carbon–oxygen white dwarf models with helium-dominated envelopes due to the presence, absence, and enhancement of $^{22}\text{Ne}$ in the interior. The observed g-mode pulsation periods of such white dwarfs are typically given to 6–7 significant figures of precision. Usually white dwarf models without $^{22}\text{Ne}$ are fit to the observed periods and other properties. The rms residuals to the $\Delta P/P < 0.3\%$. We find average relative period shifts of $\Delta P/P \approx \pm 0.5\%$ for the low-order dipole and quadrupole g-mode pulsations within the observed effective temperature window, with the range of $\Delta P/P$ depending on the specific g-mode, abundance of $^{22}\text{Ne}$, effective temperature, and the mass of the white dwarf model. This finding also in concert with stellar models, can probe the elemental composition profiles of a white dwarf model, our study on the impact of $^{22}\text{Ne}$ can provide new inferences on the derived interior mass fraction profiles. We encourage routinely including $^{22}\text{Ne}$ mass fraction profiles, informed by stellar evolution models, to future generations of white dwarf model-fitting processes.

Unified Astronomy Thesaurus concepts: White dwarf stars (1799); Stellar pulsations (1625); Stellar evolution (1599); Non-radial pulsations (1117); Stellar physics (1621)

1. Introduction

Photons emitted from stellar surfaces and neutrinos released from stellar interiors may not directly reveal all that we want to know about the internal constitution of the stars. For example, a direct view of the chemical stratification from the core to the surface is hidden. These interior abundance profiles matter: they impact a star’s opacity, thermodynamics, nuclear energy generation, and pulsation properties. The stellar models, in turn, are used to interpret the integrated light of stellar clusters and galaxies (e.g., Alston et al. 2020), decipher the origin of the elements (e.g., Arcones et al. 2017; Placco et al. 2020), predict the frequency of merging neutron stars and black holes (Giacobbo & Mapelli 2018; Abbott et al. 2020; Farmer et al. 2020; Marchant & Moriya 2020), and decipher the population(s) of exploding white dwarfs (WDs) that underlay Type Ia supernova cosmology (e.g., Miles et al. 2016; Rose et al. 2020).

Neutrino astronomy, in concert with stellar models, can probe the isotopic composition profiles in energy producing regions of the Sun (Borexino Collaboration et al. 2018, 2020) and nearby ($d \lesssim 1$ kpc) pre-supernova massive stars up to tens of hours before core collapse (e.g., Patton et al. 2017; Simpson et al. 2019; Mukhopadhyay et al. 2020). Steller seismology, also in concert with stellar models, can probe the elemental composition profiles in pulsating stars from the upper main sequence (e.g., Simón-Díaz et al. 2018; Pedersen et al. 2019; Balona & Ozyur 2020) through the red giant branch (e.g., Hekker & Christensen-Dalsgaard 2017; Hon et al. 2018) to WDs (e.g., Hermes et al. 2017; Giammichele et al. 2018; Córsico et al. 2019; Bell et al. 2019; Bischoff-Kim et al. 2019; Althaus et al. 2021).

Most of a main-sequence star’s initial metallicity $Z$ comes from the carbon-nitrogen-oxygen (CNO) and $^{56}\text{Fe}$ nuclei inherited from its ambient interstellar medium. All of the CNO piles up at $^{14}\text{N}$ when H-burning on the main sequence is completed because the $^{14}\text{N}(p,\gamma)^{15}\text{O}$ reaction rate is the slowest step in the H-burning CNO cycle. During the ensuing He-burning phase, all of the $^{14}\text{N}$ is converted to $^{22}\text{Ne}$ by the reaction sequence $^{14}\text{N}(\alpha,\gamma)^{18}\text{F}(e^{-}\nu_{e})^{19}\text{O}(\alpha,\gamma)^{22}\text{Ne}$. The abundance of $^{22}\text{Ne}$ when He-burning is completed is thus proportional to the initial CNO abundance of the progenitor main-sequence star. The weak reaction in this sequence powers the neutrino luminosity during He-burning (e.g., Serenelli & Fukugita 2005; Farag et al. 2020) and marks the first time in a star’s life that the core becomes neutron rich. For zero-age main sequence (ZAMS) masses between $\approx 0.5 M_\odot$ (Demarque & Mengel 1971; Prada Moroni & Straniero 2009; Gautschy 2012) and $\approx 7 M_\odot$ (Becker & Iben 1979, 1980; García-Berro et al. 1997), depending on the treatment of convective boundary mixing (Weidemann 2000; Denissenkov et al. 2013; Jones et al. 2013; Farmer et al. 2015;
De Geronimo et al. 2019; Bischoff-Kim et al. 2019; Giammichele et al. 2017, 2018; Charpinet et al. 2019; Lecoanet et al. 2016; Constantino et al. 2015, 2016, 2017; Giammichele et al. 2018; Timmes et al. 2018; Charpinet et al. 2019; De Gerőnimo et al. 2019. Additional details of the input physics are given in Appendix B, and the MESA r12115 files to reproduce our work are available at doi:10.5281/zenodo.4338180.

2.2. Mass Fraction Profiles

Figure 1 shows the mass fraction X(^12C/Z) profiles of the resulting 0.56 M_☉ DB WD model, where A is the number of nucleons and Z is the number of protons. Brown boxes divide the mass fraction profiles into three regions according to their origins and uncertainties. The X(^12C) and X(^16O) profiles in the innermost ≈90% of mass region are determined during core and shell He-burning. The main uncertainties in this region are the ^12C(α,γ)^16O reaction rate (e.g., deBoer et al., 2017), and the treatment of convective mixing boundaries during core H- and He-burning (e.g., Constantino et al. 2015, 2016, 2017).

The CO and X(^4He) profiles between ≈1% and ≈10% of the exterior WD mass originate from shell He-burning during the thermal pulse phase of evolution on the AGB. Most of the total He mass is held in this region. The primary uncertainties in this region are the number of thermal pulses and convective boundary layer mixing. The number of thermal pulses a model undergoes is sensitive to the mass resolution, time resolution, mass loss rate, and the treatment of convective boundaries (Iben & Renzini 1983; Herwig 2005; Karakas & Lattanzio 2014). The sharp change in all the mass fractions at ≈1% of the exterior WD mass marks the extent reached by the convective boundary during the last thermal pulse.

CO profiles in this region may also be subject to other mixing processes. For example, the magnitude of the X(^12C) “bump” is subject to the strength and duration of the thermohaline instability, which occurs when material is stable to convection according to the Ledoux criterion, but has an inverted molecular weight gradient (Baines & Gill 1969; Brown et al. 2013; Garaud 2018; Bauer & Bildsten 2018).

The X(^4He) profile of the outer ≈0.1%–1% of the WD mass is determined by shell H-burning. All of the initial CNO mass fractions have been converted to ^14N. The main uncertainties in this region are the number of thermal pulses during the AGB phase of evolution, late or very late thermal pulses (Bloecker 1995a, 1995b; Bloecker 2001), and mechanisms to remove the residual high entropy, H-rich layer to produce a DB WD from single and binary evolution (e.g., D’Antona & Mazzitelli 1990; Althaus & Benvenuto 1997; Parsons et al. 2016).

The X(^22Ne) profile is essentially flat and spans the inner ≈99% by mass. As discussed in Section 1, X(^22Ne) is created from ^14N during H-burning.

2.3. Constructing Ab Initio WD Models

Starting from a set of pre-main-sequence (pre-MS) initial conditions, accurate predictions for the properties of the resulting WD model, especially the mass fraction profiles, do not exist due to the nonlinear system of equations being approximated. In addition, evolving a stellar model from the pre-MS to a WD can be resource intensive. It can thus be useful for systematic studies to build ab initio WD models (e.g.,

2. A Baseline WD Model

2.1. Input Physics

We use MESA version r12115 (Paxton et al. 2011, 2013, 2015, 2018, 2019) to evolve a 2.1 M_☉, Z = 0.02 metallicity model from the ZAMS through core H-burning and core He-burning. After winds during the thermal pulses on the AGB have reduced the H-rich envelope mass to 0.01 M_☉, the remaining hydrogen is stripped from the surface to form a young, 0.56 M_☉ DB WD. This model is tuned to match the observed and inferred properties of KIC 08626021 (Bischoff-Kim et al., 2014; Giammichele et al. 2018; Timmes et al. 2018; Charpinet et al. 2019; De Gerőnimo et al. 2019). Additional details of the input physics are given in Appendix B, and the MESA r12115 files to reproduce our work are available at doi:10.5281/zenodo.4338180.

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WDEC, Bischoff-Kim & Montgomery 2018). By ab initio we mean calculations that begin with a WD model, as opposed to a WD model that is the result of a stellar evolution calculation from the pre-MS. A potential disadvantage (or advantage) of ab initio WD models is the imposed initial mass fraction profiles may not be attainable by a stellar model evolved from the pre-MS. Throughout the remainder of this article we use a new capability, \texttt{wd\_builder}, to construct ab initio WD models in \texttt{MESA} of a given mass and chemical stratification.

The initial structure of an ab initio WD model is approximated as an isothermal core and a radiative envelope in hydrostatic equilibrium. Here, we specify an initial WD mass of 0.56 \(M_\odot\), the same WD mass as produced by the stellar evolution calculation. The imposed \(X(\text{He}), X(12\text{C}), X(14\text{N}), X(16\text{O}),\) and \(X(22\text{Ne})\) profiles are taken from the stellar evolution derived mass fraction profiles of Figure 1 and normalized to sum to unity in each cell. Henceforth we refer to this ab initio model as the “baseline model.”

For ab initio WD models we use He-dominated, \(\log(H/\text{He}) = -5.0\), model atmosphere tables spanning 5000 K \(\leq T_{\text{eff}} \leq 40,000\) K that were provided by O. Toloza (2019, private communication) using the Koester WD atmosphere software instrument (Koester 2010). These tabulated atmospheres for DB WDs are publicly available as a standard atmosphere option as of \texttt{MESA r12115}. In addition, we use five element classes for the diffusion classes \(^4\text{He}, ^{12}\text{C}, ^{14}\text{N}, ^{16}\text{O},\) and \(^{22}\text{Ne}\.\) Otherwise, all of the physics implementations and modeling choices are as described in Section 2.1.

The initial baseline model is then evolved with \texttt{MESA}. As the model is not in thermal equilibrium, there is an initial transient phase lasting a few thermal timescales that is disregarded. The thermal timescale is \(\tau_{\text{th}} \approx E_{\text{th}}/L_{\text{tot}} \approx 0.67\) Myr, where \(E_{\text{th}}\) is the thermal energy of the WD and \(L_{\text{tot}}\) is the photon plus neutrino luminosity. Specifically, we set the zero-point to be 1.5 thermal timescales (\(\approx 1\) Myr) after the transient reaches its peak luminosity. The evolution terminates when \(L_{\text{tot}}\) falls below \(\log(L/L_\odot) = -2.5\).

Figure 2 shows the cooling properties of the baseline model. Plasmon neutrino emission dominates the energy loss budget at \(T_{\text{eff}} \geq 25,000\) K (e.g., Vila 1966; Kutter & Savedoff 1969; Winget et al. 2004; Bischoff-Kim & Montgomery 2018). Photons leaving the WD surface begin to dominate the cooling as the electrons transition to a strongly degenerate plasma (van Horn 1971). The luminosity becomes proportional to the enclosed mass, \(L_v \propto M_v\), in this model only when \(T_{\text{eff}} \lesssim 20,000\) K (Timmes et al. 2018). Energy transport in the interior is dominated by conduction, driven primarily by electron-ion scattering. Energy transport in the outer layers is dominated by radiation or convection associated with the partial ionization of He at \(T_{\text{eff}} \approx 30,000\) K.

Figure 2 also shows the diffusion of the initial mass fractions as the baseline WD model cools to \(T_{\text{eff}} = 30,000, 15,000,\) and \(12,138\) K (corresponding to the termination at \(\log(L/L_\odot) = -2.5\)). Element diffusion of \(^{22}\text{Ne}\) is modest for the baseline \(0.56\) \(M_\odot\) DB WD model. Depletion of the \(^{22}\text{Ne}\) mass fraction at \(\log(1 - M_r/M) \approx -1.9\) has occurred by the time the model has cooled to \(T_{\text{eff}} \approx 30,000\) K. As the model cools further, the surface regions in the tail of the He-dominated layer further deplete and a small \(^{22}\text{Ne}\) bump forms and propagates inwards toward the center. The timescale for \(^{22}\text{Ne}\) to travel from near the surface to the center of this WD model is \(\tau_{\text{df}} \approx 2 \Gamma \bar{Z}/\rho_0^{1.25}\) Gyr \(\approx 30\) Gyr (Isern et al. 1991; Bravo et al. 1992; Bildsten & Hall 2001; Deloye & Bildsten 2002; Camisassa et al. 2016), where \(\bar{Z}\) is the mean charge of the material, \(\Gamma\) is the electrostatic to thermal energy ratio, and \(\rho_0\) is the baryon mass density in units of \(10^6\) g cm\(^{-3}\). Thus, the \(X(^{22}\text{Ne})\) profile does not significantly change as the 0.56 \(M_\odot\) baseline model evolves to \(\log(L/L_\odot) = -2.5\) in \(\approx 350\) Myr. More massive WDs show larger amounts of \(^{22}\text{Ne}\) sedimentation over the same time period (Camisassa et al. 2016). WD cooling data suggests a significant enhancement due to \(^{22}\text{Ne}\) diffusion (Cheng et al. 2019; Bauer et al. 2020), but does not effect the

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**Figure 1.** Mass fraction profiles of the 0.56 \(M_\odot\) DB WD resulting from the evolution of the 2.1 \(M_\odot\) ZAMS model.
Figure 2. Evolution of baseline model’s photon luminosity $L$ and neutrino luminosity $L_{\nu}$ (left top), effective temperature $T_{\text{eff}}$ and radius $R$ (left middle), and central temperature $T_c$ and central density $\rho_c$ (left bottom). Time begins a few thermal timescales after the ab initio WD is released. Gray bands show the luminosity and $T_{\text{eff}}$ range of currently observed DBV WD (Montreal White Dwarf Database, Dufour et al. 2017). Mass fraction profiles are shown at $T_{\text{eff}} = 30,039$ K (right top), 15,012 K (right middle), and 12,092 K (right bottom) and at the end of the evolution. Initial mass fraction profiles are shown as solid curves and the diffusing mass fraction profiles are shown as dotted curves.
baseline model until it cools to effective temperatures lower than considered here \( T_{\text{eff}} \lesssim 10,000 \text{ K} \).

### 2.4. Pulsation Periods of the Baseline Model

Having established the structural and composition profiles of a cooling baseline WD model, we now consider the g-mode pulsation periods. Some of the material is classic (e.g., Unno et al. 1989; Fontaine & Brassard 2008), but we also derive and verify the accuracy of an approximation formula for the Brunt–Väisälä frequency in WDs that allows physical insights into why the low-order g-mode pulsation periods change due to variations in the mass fraction of \(^{22}\text{Ne}\). This material is essential for establishing that the baseline model, before introducing any modifications to the chemical profiles, produces pulsation periods that are commensurate with the observations of DBV WDs.

Figure 3 shows a diagram of the propagation of the dipole \( \ell = 1 \) (top) and quadrupole \( \ell = 2 \) (bottom) g-modes at \( L = 0.01 \, L_\odot \), for the baseline WD model. The Lamb frequency \( (S_p, \text{ orange}), \) Brunt–Väisälä frequency \( (N, \text{ blue}), \) radial order \( n = 1, 2, 3, \) and 10 eigenfrequencies (dotted black lines), nodes in the radial eigenfunction (filled circles), and the g-mode period of each radial order are labeled.

- **Figure 3.** Diagram of the propagation of the dipole \( \ell = 1 \) (top) and quadrupole \( \ell = 2 \) (bottom) g-modes at \( L = 0.01 \, L_\odot \) for the baseline WD model. The Lamb frequency \((S_p, \text{ orange})\), Brunt–Väisälä frequency \((N, \text{ blue})\), radial order \( n = 1, 2, 3, \) and 10 eigenfrequencies (dotted black lines), nodes in the radial eigenfunction (filled circles), and the g-mode period of each radial order are labeled.

Radial overtone number, the \( \ell = 1 \) periods are \( \sim \sqrt{3} \) longer than the \( \ell = 2 \) periods, due to the local dispersion relation for low-frequency g-modes \( \sigma_n \) scaling as

\[
\sigma_n^2 \approx \ell (\ell + 1) N^2/(k_r^2 r^2),
\]

where \( k_r \) is the radial wavenumber. The Brunt–Väisälä frequency \( N \) is

\[
N^2 = \frac{g^2 \rho X_T}{P \chi_\rho} (\nabla_{\text{ad}} - \nabla_T + B),
\]

where \( g \) is the gravitational acceleration, \( \rho \) is the mass density, \( P \) is the pressure, \( T \) is the temperature, \( \chi_T \) is the temperature exponent \( \partial (\ln P)/\partial (\ln \rho)|_{\Gamma,\mu_f} \), \( \chi_\rho \) is the density exponent \( \partial (\ln P)/\partial (\ln T)|_{\Gamma,\mu_f}, \) \( \nabla_{\text{ad}} \) is the adiabatic temperature gradient, \( \nabla_T \) is the actual temperature gradient, and \( B \) accounts for composition gradients (e.g., Hansen & Kawaler 1994; Fontaine & Brassard 2008). Bumps in the \( N \) profile of Figure 3 correspond to transitions in the \( X(^{16}\text{O}), X(^{12}\text{C}), \) and \( X(^{4}\text{He}) \) profiles. The implementation of Equation (2) in MESA is described in Section 3 of Paxton et al. (2013).

An approximation for \( N^2 \) in the interiors of WDs that yields physical insights begins by assuming \( \nabla_{\text{ad}} \) is much larger than \( \nabla_T \) and \( B \). Then

\[
N^2 = \frac{g^2 \rho X_T}{P \chi_\rho} \nabla_{\text{ad}}.
\]

In the interior of a WD the ions are ideal and dominate the temperature derivatives of an electron degenerate plasma. Substituting the pressure scale height \( H = P/(\rho g) \) and Equation 3.110 of Hansen & Kawaler (1994)

\[
\chi_T = \frac{\rho}{P} \frac{k_B T}{\mu_I m_p}
\]

into Equation (3) gives

\[
N^2 = \frac{1}{H^2 \chi_\rho} \frac{k_B T}{\mu_I m_p} \nabla_{\text{ad}},
\]

where \( k_B \) is the Boltzmann constant, \( \mu_I = 1/(\sum X_i/A_i) \) is the ion mean molecular weight, and \( m_p \) is the mass of the proton. Equation (3.90) of Hansen & Kawaler (1994) shows \( \nabla_{\text{ad}} = (\Gamma_3 - 1)/\Gamma_1 \), where \( \Gamma_1 \) is the first adiabatic index and \( \Gamma_3 = k_B/(\mu_I m_p c_v) \) is the third adiabatic index, where in the gas phase the ideal specific heat capacity is \( c_v = 3k_B/(2\mu_I m_p) \). The sentence beneath Equation 3.112 of Hansen & Kawaler (1994) thus notes that \( \Gamma_3 - 1 = 2/3 \) for the ions in the gas phase \( (\Gamma_3 - 1 = 1/3 \) in the liquid phase). Combining these expressions, yields the approximation

\[
N^2 = \frac{2}{3 \Gamma_1 \chi_\rho} \frac{k_B T}{H^2 \mu_I m_p}.
\]

Figure 4 compares the approximation in Equation (6) with the full \( N^2 \) calculation from MESA. The difference at \( r/R \approx 0.5 \) corresponds to the \( X(^{16}\text{O}) \rightarrow X(^{12}\text{C}) \) transition, at \( r/R \approx 0.8 \) to the \( ^{12}\text{C} \) bump and at \( r/R \approx 0.9 \) to the transition to a He-dominated atmosphere. Except for the outermost layers and regions where the composition gradients are significant, the agreement is sufficient to use Equation (6) as a scaling relation.
Figure 4. Comparison of the approximation for $N^2$ (blue curve) in Equation (6) and the full calculation of $N^2$ from MESA (green curve).

for building physical insights. We always use, however, the full $N^2$ calculation from MESA for any quantitative analysis.

It is useful to reference features of the baseline model with respect to mass or radius. Figure 5 thus shows the mass–radius relation of the baseline model at $\log(L/L_\odot) = -2.5$ with key transitions labeled.

Figure 6 shows the low-order g-mode pulsation periods as the baseline WD model cools. The periods increase due to $N^2$ decreasing as the cooling progresses, per Equation (6). Higher radial orders have steeper slopes due to the periods scaling with $k$, in Equation (1). The increase in the number of MESA models at $T_{\text{eff}} \approx 30,000$ K is due to the partial ionization of He, which leads to envelope convection in relatively hot DBV WDs. The change in slope at $T_{\text{eff}} \approx 20,000$ K is due to the luminosity becoming proportional to the enclosed mass, $L_{\text{r}} \propto M_{\text{r}}$, as the plasmon neutrino emission becomes insignificant.

In Appendix A we show that the low-order g-mode pulsation periods of the baseline model calculated with GYRE are only weakly dependent on the mass and temporal resolution of the MESA calculations.

3. The Impact of $^{22}\text{Ne}$

Having established the cooling properties and g-mode pulsation periods of a baseline model whose mass fraction profiles are from a stellar evolution model, we now explore changes in the g-mode pulsation periods due to changes in the $^{22}\text{Ne}$ mass fraction profile shown in Figure 1. We consider three modifications: replacing $^{22}\text{Ne}$ with $^{14}\text{N}$, a metal-free model, and a super-solar metallicity model.

3.1. Putting $^{22}\text{Ne}$ into $^{14}\text{N}$

Replacing $X(\text{Ne})$ with $X(\text{N})$ is a model for the reaction sequence $^{14}\text{N}(\alpha,\gamma)^{18}\text{F}(e^-\nu\gamma)^{18}\text{O}(\alpha,\gamma)^{22}\text{Ne}$ either physcally not occurring or being ignored. Figure 7 shows the relative differences in the low-order g-mode pulsation periods from this composition change. All of the relative differences are negative, implying the pulsation periods in models that exclude $^{22}\text{Ne}$ are longer than the corresponding pulsation periods in models that include $^{22}\text{Ne}$. The magnitude of the relative period differences span $\approx 0.25\%–1\%$ over the range of currently observed DBV WDs, with the $g_{1,1}$ and $g_{1,2}$ modes showing the largest differences at cooler $T_{\text{eff}}$. The change in the slopes at $T_{\text{eff}} \approx 20,000$ K is due to plasmon neutrino emission becoming insignificant, and thus the luminosity becoming proportional to the enclosed mass, $L_{\text{r}} \propto M_{\text{r}}$.

What drives these g-mode period changes? Replacing an isotope that has a larger mass number with an isotope that has a smaller mass number decreases $\mu_t$. This replacement also increases $H$ through the mechanical structure and equation of state (EOS) of the CO WD. Figure 8 shows the relative differences in the $H^2$, $\mu$, $\Gamma_1$, $\chi_{\rho}$, and $T$ contributions to $N^2$ in Equation (6). These changes collectively determine the magnitude and sign of the period change relative to the baseline model. For this $X(\text{Ne}) \rightarrow X(\text{N})$ model, the overall positive changes in $\mu_t$ and $T$ are countered by the negative changes from $H^2$, $\Gamma_1$, and $\chi_{\rho}$. The magnitude of the relative difference in $H^2$ drives the net result of a smaller $N^2$ and thus longer g-mode periods. The nearly uniform negative change in $H^2$ implies a change in the radius of the WD model. We find $(R_B - R_{14\text{N}})/R_B \approx -0.4\%$, meaning that the $X(\text{Ne}) \rightarrow X(\text{N})$
model has a larger radius than the model with $^{22}\text{Ne}$. This is expected given differences in the electron fraction of a WD.

Figure 9 compares the weight functions of the baseline model with $^{22}\text{Ne}$ and the model where the $^{22}\text{Ne}$ has been replaced with $^{14}\text{N}$. Following Kawaler et al. (1985), the weight function is

$$
\frac{d\zeta}{dr} = \frac{\int \left[ C(y, r) + N(y, r) + G(y, r)\rho^2 \right] d\rho}{\int_0^R T(y, r) \rho^2 d\rho},
$$

where $C(y, r)$ varies with the Lamb frequency, $N(y, r)$ contains the Brunt–Väisälä frequency, $G(y, r)$ involves the gravitational eigenfunctions, $T(y, r)$ is proportional to the kinetic energy density, and $y = (y_1, y_2, y_3, y_4)$ are the Dziembowski (1971) variables. The frequency of an adiabatic mode is then

$$
\nu^2 = \zeta = \int_{r=0}^{r=R} \frac{d\zeta}{dr} \cdot dr.
$$

The weight function for the two models is dominated by the $N(y, r)$ term except for the surface layers. Figure 9 shows that the net effect of the $^{22}\text{Ne} \rightarrow ^{14}\text{N}$ composition change is a shift in $\zeta$, the area under the weight function curves, toward smaller frequencies of the low-order $g$-modes.

Figure 9 illustrates the relative percentage differences between the weight function curves. Most of the changes in $\zeta$ occur at the CO transition region ($r/R \approx 0.45$, see Figure 5), $^{12}\text{C}$ bump ($r/R \approx 0.8$), and at the transition to a He-dominated atmosphere ($r/R \approx 0.9$). The changes in these regions get as large as $\sim 10\%$. We identify the dipole $g$-mode of radial order $n = 2$ as being more sensitive to the location and gradient of $\mu_I$ at the CO transition ($r/R \approx 0.5$) than other low-order $g$-modes.

### 3.2. Zero Metallicity and Super-solar Metallicity

Replacing $X(^{14}\text{N})$ with $X(^{4}\text{He})$ and $X(^{22}\text{Ne})$ with $X(^{12}\text{C})$ is a model for ignoring the birth metallicity of the ZAMS star, CNO burning on the main sequence, and the $^{14}\text{N}(\alpha, \gamma)^{18}\text{F}(e^+\nu_e)^{18}$
O(\alpha, \gamma) \; ^{22}\text{Ne} \; \text{reaction sequence during He-burning. Most}
\text{studies of the pulsation periods of observed WDs use zero-metallicity DBV WDs when}
deriving the interior mass fraction profiles, although see Camisassa et al. (2016) for a counter-
example. Alternatively, doubling X(^{14}\text{N}) \text{at the expense of } X(^{4}\text{He}) \text{and doubling } \chi_{^{22}\text{Ne}} \text{at the expense of } X(^{12}\text{C}) \text{is a model for a super-solar metallicity DBV WD.}

Figure 10 compares the relative change in the low-order g-mode pulsation periods of the zero
and super-solar metallicity models. The period differences are negative for the zero-
metallicity model and positive for the super-solar metallicity model. The zero-metallicity DBV WD model has longer periods
than the baseline model, which in turn has longer periods than the super-solar metallicity model. The relative period differences of the zero-metallicity models, averaged over
the $T_{\text{eff}}$ evolution, are $\Delta P(g_{1,1}) \simeq -0.57 \text{s}$, $\Delta P(g_{1,2}) \simeq -0.40 \text{s}$, $\Delta P(g_{2,1}) \simeq -0.52 \text{s}$, and $\Delta P(g_{2,2}) \simeq -0.40 \text{s}$. For the super-
solar metallicity models the averaged absolute period differences are $\Delta P(g_{1,1}) \simeq 0.66 \text{s}$, $\Delta P(g_{1,2}) \simeq 0.45 \text{s}$, $\Delta P(g_{2,1}) \simeq 0.46 \text{s}$, and $\Delta P(g_{2,2}) \simeq 0.35 \text{s}$. Over the $T_{\text{eff}}$ range of currently observed DBV WDs, the mean relative period change of the dipole modes is
0.57% and the maximum of relative period change is 0.88%. The relative period change of the quadrupole modes is smaller, with a mean of 0.33% and a maximum of 0.63%.

Figure 11 shows the relative differences in the $H^{2}$, $\mu_{r}$, $\Gamma_{1}$, $\chi_{r}$, and $T$ contributions to $N^{2}$ of Equation (6) for the zero and super-solar metallicity models. These changes collectively
determine the magnitude and sign of the period change relative to the baseline model. For the zero-metallicity models, the combined positive changes in $\mu_{r}$ and $T$ are counteracted by the collective negative changes from $H^{2}$, $\Gamma_{1}$, and $\chi_{r}$. The net change is negative, resulting in smaller $N^{2}$ and longer g-mode periods.

Figure 9. Weight functions of the low-order g-modes for baseline model with $^{22}\text{Ne}$ (black curves) and a baseline model where $^{22}\text{Ne}$ has been replaced with $^{14}\text{N}$ (green curves). The subpanels show the relative percentage differences between the two curves. The profiles shown are when the two models have cooled to $\log(L/L_{\odot}) = -2.5$. Nodes in the radial eigenfunctions are marked by filled circles.
Similar reasoning for the super-solar metallicity models leads to a net positive change, resulting in larger \( N^2 \) and smaller g-mode periods. The magnitude of the difference in \( H^2 \) drives the overall result for both metallicity cases. The nearly uniform changes in \( H^2 \) imply changes in the radii, and we find \( (R_B - R_Z)/R_B \approx \pm 0.4\% \) with zero-metallicity models having smaller radii and super-solar metallicity models having larger radii.

Interrogating further the composition dependence, the top panels of Figure 12 compare the mass fraction profiles of the \( X^{22}\text{Ne} \approx 0.02 \) baseline and zero-metallicity at 30,000, 15,000, and 12,100 K as a function of mass coordinate. Element diffusion is operating in both models. The middle panels show the relative differences in these mass fraction profiles, with the \( ^{22}\text{Ne} \) and \( ^{14}\text{N} \) offsets zeroed out. The C and O differences at \( \log(1 - M_{2}/M) \approx -0.25 \), from Figure 5, correspond to the C/O transition at \( r/R \approx 0.9 \). These relative differences are the largest at 30,000 K, reaching \( \approx 7.5\% \) for \( ^{16}\text{O} \) and \( \approx -6\% \) for \( ^{4}\text{He} \). The relative differences at 15,000 and 12,100 K have about the same magnitude, \( \approx 7.5\% \) for \( ^{16}\text{O} \) and \( \approx -1\% \) for \( ^{4}\text{He} \). The relative mass fraction differences span a larger range of \( \log(1 - M_{2}/M) \) as the models cool due to element diffusion. The bottom panels of Figure 12 show the corresponding relative difference in the \( \mu_I \) profiles. As \( \mu_I \) is calculated by dividing the mass fraction of an isotope by its atomic weight, the relative differences in the mass fraction profiles are reduced in the \( \mu_I \) profiles. The \( \mu_I \) profile for 12,100 K in terms of a mass coordinate is the same as the \( \mu_I \) profile in Figure 11 in terms of a radial coordinate.

We also computed the relative period differences between the \( X^{22}\text{Ne} \approx 0.02 \) baseline and zero-metallicity model with diffusion turned off to disentangle structural and diffusion effects. The results are shown in Figure 13. While there is a slight variation from the zero-metallicity gray curves shown in Figure 10, mostly in the higher order \( g_{10,1} \) and \( g_{10,2} \) modes, the magnitude of the relative differences remains the same. This further suggests that the period shifts are a direct consequence of the presence or absence of \( ^{22}\text{Ne} \).
4. Trends in the Period Changes with the WD Mass

Using the same physics and numerical choices as used for the 0.56\,\text{M}_\odot baseline model, we evolved a Z = 0.02, 1.1\,\text{M}_\odot ZAMS stellar model from the pre-main sequence to a 0.526\,\text{M}_\odot DB WD, and a Z = 0.02, 3.6\,\text{M}_\odot ZAMS model to a 0.729\,\text{M}_\odot DB WD. This initial to final mass mapping is similar to that in Table 1 of Camisassa et al. (2016). Relative to the 0.56\,\text{M}_\odot baseline model, the 0.526\,\text{M}_\odot WD model has a thicker He layer and a more abbreviated extent of\,X(22\,\text{Ne}). Conversely, the 0.729\,\text{M}_\odot WD model has a smaller\,12\,\text{C} bump, a thinner He layer, and a more extended\,X(22\,\text{Ne}) profile. These mass fraction profiles were imposed on the 0.526\,\text{M}_\odot and 0.729\,\text{M}_\odot ab initio WD models, respectively.

Figure 14 shows the diffusion of these initial mass fraction profiles as the ab initio WD models cool to $T_{\text{eff}} \approx 30,000$\,K, then $\approx 15,000$\,K and finally to $\approx 12,000$\,K (corresponding to the termination at $\log(L/L_\odot) \approx -2.5$). Element diffusion is more pronounced for the more massive 0.729\,\text{M}_\odot DB WD model due to its larger surface gravity. An enhancement forms in the\,X(22\,\text{Ne}) profile at $\log(1 - M_r/M) \approx -2.0$ by the time the 0.729\,\text{M}_\odot model has cooled to $T_{\text{eff}} \approx 30,000$\,K. As the model further cools, the\,X(22\,\text{Ne}) bump grows in amplitude as it propagates inwards toward the center through the He-dominated outer layers. The\,X(22\,\text{Ne}) bump generates an increase in the local $N^2$ in the regions it propagates through from a locally larger $\mu_l$ and a smaller compensating $H^2$. The regions trailing the\,X(22\,\text{Ne}) bump are depleted of\,X(22\,\text{Ne}), causing a decrease in the local $N^2$ in these regions.

We find longer low-order g-mode periods for the more massive WD, consistent with Camisassa et al. (2016). As was done for the 0.56\,\text{M}_\odot baseline model, we replace $X(14\text{N})$ with $X(\text{He})$ and $X(22\text{Ne})$ with $X(12\text{C})$ to generate a zero-metallicity ab initio DB WD model. We also double $X(14\text{N})$ at the expense of $X(\text{He})$ and double $X(22\text{Ne})$ at the expense of $X(12\text{C})$ to generate a super-solar metallicity DB WD.

Figure 15 compares the relative change in the low-order g-mode pulsation periods of the zero and super-solar metallicity 0.526\,\text{M}_\odot and 0.729\,\text{M}_\odot DB WD models. As for the 0.56\,\text{M}_\odot baseline model, the relative period differences are mostly symmetric about the reference model’s $Z = 0.02$ metallicity. For the 0.526\,\text{M}_\odot models, the relative period changes found for the 0.526\,\text{M}_\odot baseline model, we replace $X(14\text{N})$ with $X(\text{He})$ and $X(22\text{Ne})$ with $X(12\text{C})$ to generate a zero-metallicity DB WD model. We also double $X(14\text{N})$ at the expense of $X(\text{He})$ and double $X(22\text{Ne})$ at the expense of $X(12\text{C})$ to generate a super-solar metallicity DB WD.

There are a few trends in the relative period differences with respect to the WD mass. For the zero-metallicity $n = 2$ and $n = 10$ g-modes, the average relative differences in the observed $T_{\text{eff}}$ range increase with increasing mass. For example, as the WD mass is increased from 0.526 to 0.560\,\text{M}_\odot, we find the average relative period differences increase by factors of 1.74, 1.22, 2.43, and 1.46, for the $g_{2,1}$, $g_{2,2}$, $g_{10,1}$, and $g_{10,2}$ modes, respectively. As
models due to the presence, absence, and enhancement of $^{22}\text{Ne}$ as the models cool through the observed range of effective temperatures. We found mean relative period shifts of $\Delta P/P \approx \pm 0.5\%$ for the low-order dipole and quadrupole g-mode pulsations within the observed effective temperature window, with a range of $\Delta P/P$ that depends on the specific g-mode, mass fraction of $^{22}\text{Ne}$, effective temperature, and mass of the WD model. Shifts in the pulsation periods due to the presence, absence, or enhancement of $X^{(22}\text{Ne})$ mostly arise from a competition between the pressure scale height and ion mean molecular weight.

Low-mass DAV WDs, the ZZ Ceti class of stars, have pulsation periods in the 100–1500 s range (e.g., Vincent et al. 2020). Comparing low-mass DAV WDs produced from stellar evolution models with and without diffusion of $^{22}\text{Ne}$, Camisassa et al. (2016) found that the $^{22}\text{Ne}$ sedimentation induces mean period differences of $\approx 3\,\text{s}$, reaching maximum period differences of $\approx 11\,\text{s}$. For the more massive DAV WD models, where sedimentation of $^{22}\text{Ne}$ is stronger, they found mean period differences of $\approx 15\,\text{s}$ between when diffusion is on and off, and a maximum period differences of $\approx 47\,\text{s}$. Comparatively, our article focuses on DBV WD models, the evolution of the pulsation periods as the DBV WD models cool, and the impact of $^{22}\text{Ne}$ being present, absent, or enhanced in the WD interior. Nevertheless, we conducted an experiment of turning element diffusion off in our 0.56 $M_{\odot}$ baseline model. At $\log(L/L_{\odot}) = -2.5$, we found an absolute mean difference for $n = 1$ to $n = 11$ of $\approx 16\,\text{s}$, with a maximum period difference at $n = 9$ of $\approx 56\,\text{s}$. This maximum difference equates to a $\approx 7\%$ relative difference between when diffusion is on and off. Our period changes are slightly higher than those found in Camisassa et al. (2016)’s 0.833 $M_{\odot}$ model, and much larger than the differences found in their 0.576 $M_{\odot}$ model. These differences could be a result of DAV versus DBV models, as DAV models have different cooling times than DBV models. In addition, Camisassa et al. (2016) computed their period differences at $\log(L/L_{\odot}) = -2.80$ and $\log(L/L_{\odot}) = -2.93$ for their 0.576 and 0.833 $M_{\odot}$ models, respectively. These are dimmer than the $\log(L/L_{\odot}) = -2.5$ used for our calculations. Our maximum radial order is found up to 11 at this luminosity, while Camisassa et al. (2016) used more radial orders, with a maximum radial order of 50.

Giammichele et al. (2018) compared the g-mode pulsation periods of a pure oxygen core surrounded by a pure helium envelope with those from an oxygen-dominated core with $X^{(22}\text{Ne}) = 0.02$ surrounded by a pure helium envelope. They found including $^{22}\text{Ne}$ yields shorter periods, with mean period differences of $\approx 0.1\%$. We find a mean period shift that is about five times larger in our 0.56 $M_{\odot}$ baseline model. This difference may be caused by the contrast in the composition of the models, which in turn causes variances in the local mean molecular weight and pressure scale height scaling described by Equation (6).

Are 1% or less period differences important? The g-mode periods of observed DBV WDs are found from a Fourier analysis of the photometric light curves and are typically given to 6–7 significant figures of precision. Usually zero-metallicity WD models (i.e., without $^{22}\text{Ne}$) are fit to the observed g-mode periods and other properties (e.g., $T_{\text{eff}}$, log g). The rms residuals to the $\approx 150–400\,\text{s}$ low-order g-mode periods are typically in the range of $\sigma_{\text{rms}} \lesssim 0.3\,\text{s}$ (e.g., Bischoff-Kim et al. 2014), for a
fit precision of $\sigma_{\text{rms}}/P \lesssim 0.3\%$. Our finding of a mean relative period shift of $\Delta P/P \simeq \pm 0.5\%$ induced by including $^{22}\text{Ne}$ in WD models suggests a systematic offset may be present in the fitting process of specific WDs when $^{22}\text{Ne}$ is absent. As part of the fitting process involves adjusting the composition profiles of the model WD, this study on the impact of $^{22}\text{Ne}$ can inform inferences about the derived interior mass fraction profiles. We encourage routinely including $^{22}\text{Ne}$ mass fraction profiles, informed by stellar evolution models, to future generations of DBV WD model fitting processes.

The adiabatic low-order g-mode pulsation periods of our DB WD models depend upon simplifying assumptions in the stellar evolution calculations (e.g., convective boundary layer mixing, shellular rotation), uncertainties (e.g., mass loss history, stripping of the residual thin H layer, thickness of the He-dominated atmosphere), and unknown inherent systematics. We hypothesize that these model dependencies and systematics could mostly cancel when dividing one model result by another model result, such as when calculating the relative period shifts $\Delta P/P$. We anticipate exploring a larger range of models,

Figure 14. Mass fractions profiles for the 0.52 $M_\odot$ (left column) and 0.73 $M_\odot$ (right column) ab initio DB WD models at $T_{\text{eff}} \simeq 30,000$ K (top), 15,000 K (middle), and at the end of the evolution (bottom). Initial mass fraction profiles are shown as solid curves and the diffusing mass fraction profiles are shown as dotted curves.
similar in approach to Fields et al. (2016), to test this conjecture in future studies.

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Software: MESA (Paxton et al. 2011, 2013, 2015, 2018, 2019, http://mesa.sourceforge.net), MESSDK 20190830 (Townsend 2019a, 2019b), wd_builder https://github.com/jschwab/wd_builder, GYRE (Townsend & Teitler 2013; Townsend et al. 2018, https://github.com/rhtownsend/gyre), Montreal White Dwarf Database (Dufour et al. 2017, http://www.montrealwhitedwarfdatabase.org), matplotlib (Hunter 2007), and NumPy (van der Walt et al. 2011).

Appendix A

Convergence Studies

In this appendix we demonstrate that the pulsation periods of the baseline model are only weakly dependent on the details of the mass and temporal resolution of the MESA + GYRE calculations.

A MESA parameter controlling the mass resolution is \( \text{max}_\text{dq} \), the maximum fraction a model’s mass in one cell. That is, the minimum number of cells in a model is \( N_{\text{min cells}} = 1/\text{max}_\text{dq} \). We use \( N_{\text{min cells}} = 5000 \) for all the results reported. MESA can also adaptively refines its mesh based on a set of mesh functions. The maximum cell-to-cell variation of these functions is maintained around the value of the control \( \text{mesh} \_\text{delta} \_\text{coeff} \). We use \( \text{mesh} \_\text{delta} \_\text{coeff} = 1 \) for all the results reported. Primarily as a result of these two mesh parameters, the total number of cells in the baseline model is \( \approx 8000 \) cells.

A MESA parameter controlling the time resolution is the largest change in the central temperature allowed over a timestep, \( \text{delta} \_\text{lgT} \_\text{cntr} \_\text{limit} \). For all the reported results, we use \( \text{delta} \_\text{lgT} \_\text{cntr} \_\text{limit} = 0.001 \). MESA
can also adaptively adjust the timestep based on other criteria, but this setting dominates the size of every timestep as the baseline WD model cools. The total number of timesteps in the baseline model is \(1000\) and varies roughly linearly with \(\delta \log T_{\text{cntr}}\).

Figure 16 shows changes in the low-order g-mode periods for different minimum mass resolutions as the baseline WD models cool. We use the notation \(g_{n\ell}\) for a g-mode of order \(n\) and degree \(\ell\). The minimum mass resolution of 5000 cells, used for all the results reported, is shown by the black horizontal lines.

\[ N_{\text{min cells}} = 25,000 \] causes maximum relative period changes of 0.000022\% at \(g_{1,1}\) to 0.028\% at \(g_{10,1}\). These maximum relative period changes are, respectively, a factor of \(\approx 20,000\) to 20 smaller than the relative period change caused by including or excluding \(^{22}\)Ne.

\[ N_{\text{min cells}} = 5,000 \] is used for all the results reported, is shown by the black horizontal lines.

### Figure 16

Relative differences in the \(g_{1,1}\), \(g_{1,2}\), \(g_{2,1}\), \(g_{2,2}\), \(g_{10,1}\), and \(g_{10,2}\) pulsation periods for different minimum mass resolutions as the baseline WD models cool. We use the notation \(g_{n\ell}\) for a g-mode of order \(n\) and degree \(\ell\). The minimum mass resolution of 5000 cells, used for all the results reported, is shown by the black horizontal lines.

### Figure 17

Relative differences in the \(g_{1,1}\), \(g_{1,2}\), \(g_{2,1}\), \(g_{2,2}\), \(g_{10,1}\), and \(g_{10,2}\) pulsation periods for different temporal resolutions as the baseline WD models cool. The largest change in the central temperature allowed over a timestep, \(\delta \log T_{\text{cntr\_limit}} = 0.001\), used for all the results reported, is shown by the black horizontal lines.

\[ \delta \log T_{\text{cntr\_limit}} = 0.001 \] is used for all the results reported, is shown by the black horizontal lines.
that follows 49 isotopes from $^1$H to $^{34}$S, including $^{22}$Ne. This model with 5 times more time resolution, delta_lgT_cntr_limit = 0.0002, has average period changes of 0.00061 s for $g_{1,1}$, -0.00077 s for $g_{1,2}$, 0.0034 s for $g_{2,1}$, 0.0010 s for $g_{2,2}$, 0.0021 s for $g_{0,1,1}$, and 0.0014 s for $g_{1,0,2}$. The average period changes are a factor of ≈1000 smaller than the average period changes caused by including or excluding $^{22}$Ne.

Appendix B
Input Physics Details

In this appendix we briefly discuss the salient physics used in our MESA models.

B.1. Thermodynamics

The MESA r12115 EOS is a blend of the OPAL (Rogers & Nayfonov 2002), SCVH (Saumon et al. 1995), PTEH Pols et al. (1995), HELM (Timmes & Swesty 2000), and PC (Potekhin & Chabrier 2010) EOSs. The MESA EOS also covers the late stages of WD cooling where the ions in the core crystallize (e.g., Bauer et al. 2020). WD interiors lie in the PC region of the MESA EOS, which provides a semi-analytic EOS treatment for arbitrary composition. The default in MESA version 12115 is to account for each species of ion with mass fraction greater than $10^{-3}$ when calling the PC EOS. Therefore, changing the interior composition in a WD model, such as including or excluding $^{22}$Ne, self-consistently changes the thermodynamics.

B.2. Opacities

MESA r12115 divides the radiative opacity tables into two temperature regimes, low ($T \lesssim 10^4$ K) and high ($T \gtrsim 10^4$ K). For the stellar evolution calculations from the pre-MS to a WD we use the Ferguson et al. (2005) low-temperature regions, and for the high-temperature regions we use the OPAL Type I opacities (Iglesias & Rogers 1996), smoothly transitioning to the OPAL Type II opacities (Iglesias & Rogers 1996) starting at the end of core H-burning. In our WD models, the radiative opacities are provided by the OPAL Type 2 tables, which are functions of the hydrogen mass fraction $X$, metal mass fraction $Z$, and the C/O-enhancements. Thus, for the same temperature and density, our $X(\text{^{22}Ne}) \rightarrow X(\text{^{14}N})$ replacement in Section 3.1 does not change the radiative opacities. Our $X(\text{^{14}N}) \rightarrow X(\text{^{4}He})$ and $X(\text{^{22}Ne}) \rightarrow X(\text{^{12}C})$ replacements to generate zero-metallicity ab initio DB WD in Section 3.2 decreases Z in the He-dominated envelope and increases the C enhancement in the interior. Conversely, our doubling $X(\text{^{14}N})$ at the expense of $X(\text{^{4}He})$ and doubling $X(\text{^{22}Ne})$ at the expense of $X(\text{^{12}C})$ to generate a super-solar metallicity ab initio DB WD in Section 3.2 increases Z in the He-dominated envelope and decreases the C enhancement in the interior. Electron conduction opacities are from Cassisi et al. (2007), which are the relevant opacity in the WD interior. The conduction opacities are a function of the mean atomic number $\bar{Z}$, which MESA evaluates using the full composition vector in each cell.

B.3. Nuclear Reaction Networks

We use MESA’s mesa49.net, a nuclear reaction network that follows 49 isotopes from $^1$H to $^{34}$S, including $^{22}$Ne. This impact of this reaction network on the properties of CO WDs from Monte Carlo stellar models is discussed by Fields et al. (2016). All forward thermonuclear reaction rates are from the Joint Institute for Nuclear Astrophysics (JINA) reachlib version 2.2 2017 October 20 (Cyburt et al. 2010). Inverse rates are calculated directly from the forward rates (those with positive $Q$-value) using detailed balance, rather than using fitted rates. The nuclear partition functions used to calculate the inverse rates are from Rauscher & Thielemann (2000). Electron screening factors for both weak and strong thermonuclear reactions are from Chugunov et al. (2007) with plasma parameters from Itoh et al. (1979). All the weak rates are based (in order of precedence) on the tabulations of Langanke & Martínez-Pinedo (2000), Oda et al. (1994), and Fuller et al. (1985). Thermal neutrino energy losses are from Itoh et al. (1996).

B.4. Mass Loss

The implementations of mass loss in MESA r12115 are based on a number of observationally and theoretically motivated prescriptions, but uncertainties remain on line-driven and dust-driven winds (Dupree 1986; Willson 2000; Boulanger et al. 2019). We follow the mass loss settings used by the MESA Isochrones and Stellar Tracks isochrones (Dotter 2016; Choi et al. 2016), with a combination of the Reimers mass loss prescription (Reimers 1975) with $\eta = 0.1$ on the red giant branch and a Blöcker mass loss prescription (Bloeker 1995a) with $\eta = 0.5$ on the AGB.

B.5. Rotation and Magnetic Fields

MESA r12115 implements the inherently 3D process of rotation by making the 1D shellular approximation (Zahn 1992; Meynet & Maeder 1997), where the angular velocity is constant over isobars. The transport of angular momentum and material due to rotationally induced instabilities is followed using a diffusion approximation (e.g., Endal & Sofia 1978; Pinsonneault et al. 1989; Heger et al. 2000; Maeder & Meynet 2003, 2004; Suijs et al. 2008) for the dynamical shear instability, secular shear instability, Eddington–Sweet circulation, Goldreich–Schubert–Fricke instability, and Taylor–Spruit dynamo. See Heger et al. (2000) for a description of the different instabilities and diffusion coefficients.

Magnetic fields are implemented in MESA using the formalism of Heger et al. (2005), where a magnetic torque due to a dynamo (Spruit 2002) allows angular momentum to be transported inside the star. The azimuthal and radial components of the magnetic field are modeled as $B_\phi \sim r \sqrt{(4\pi \rho) \omega_A}$ and $B_r \sim B_\phi / (r k)$, respectively, where $r$ is the radial coordinate, $\omega_A$ the Alfvén frequency, and $k$ the wavenumber. These magnetic fields provide a torque of $S = B_r B_\phi / (4\pi)$, which slows down the rotation rate by decreasing the amount of differential rotation (Heger et al. 2005).

We initialize rotation by imposing a solid body rotation law, $\Omega / \Omega_{\text{crit}} = 1.9 \times 10^{-4}$, at the ZAMS. ZAMS is defined as being where the nuclear burning luminosity is 99% of the total luminosity, and the rotation rate is normalized by the surface critical rotation rate $\Omega_{\text{crit}} = \sqrt{(1 - L / L_{\text{Edd}}) c M / R^3}$, where $c$ is the speed of light, $M$ is the mass of the star, $R$ is the stellar radius, $L$ is the luminosity, and $L_{\text{Edd}}$ is the Eddington luminosity. The initial magnetic field is set to $B_r = B_\phi = 0$. Effects from rotationally induced mass loss are not included.

B.6. Element Diffusion

Element diffusion is implemented in MESA r12115 following Thoul et al. (1994), and described in Section 3 of Paxton et al. (2018). All isotopes in the reaction network are categorized
into classes according to their atomic masses, each of which has a representative member whose properties are used to calculate the diffusion velocities. Diffusion coefficients are calculated, by default, according to Stanton & Murillo (2016), whose formalism is based on binary collision integrals between each pair of species in the plasma. The diffusion equation is then solved using the total mass fraction within each class. From the ZAMS to the construction of the DB WD, we use the 10 element classes $^{1}\text{H}$, $^{3}\text{He}$, $^{4}\text{He}$, $^{12}\text{C}$, $^{14}\text{N}$, $^{16}\text{O}$, $^{20}\text{Ne}$, $^{22}\text{Ne}$, $^{24}\text{Mg}$, and $^{28}\text{Si}$.

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