EVOLUTIONARY CALCULATIONS OF PHASE SEPARATION IN CRYSTALLIZING WHITE DWARF STARS

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ABSTRACT

We present an exploration of the significance of carbon/oxygen phase separation in white dwarf stars in the context of self-consistent evolutionary calculations. Because phase separation can potentially increase the calculated ages of the oldest white dwarfs, it can affect the age of the Galactic disk as derived from the downturn in the white dwarf luminosity function. We find that the largest possible increase in ages due to phase separation is \( \sim 1.5 \) Gyr, with a most likely value of approximately 0.6 Gyr, depending on the parameters of our white dwarf models. The most important factors influencing the size of this delay are the total stellar mass, the initial composition profile, and the phase diagram assumed for crystallization. We find a maximum age delay in models with masses of \( \sim 0.6 \ M_\odot \), which is near the peak in the observed white dwarf mass distribution. In addition, we note that the prescription that we have adopted for the mixing during crystallization provides an upper bound for the efficiency of this process, and hence a maximum for the age delays. More realistic treatments of the mixing process may reduce the size of this effect. We find that varying the opacities (via the metallicity) has little effect on the calculated age delays. In the context of Galactic evolution, age estimates for the oldest Galactic globular clusters range from 11.5 to 16 Gyr and depend on a variety of parameters. In addition, a 4–6 Gyr delay is expected between the formation of the globular clusters and the formation of the Galactic thin disk, while the observed white dwarf luminosity function gives an age estimate for the thin disk of \( 9.5^{\pm 1.1} \) Gyr, without including the effect of phase separation. Using the above numbers, we see that phase separation could add between 0 and 3 Gyr to the white dwarf ages and still be consistent with the overall picture of Galaxy formation. Our calculated maximum value of \( \lessapprox 1.5 \) Gyr fits within these bounds, as does our best-guess value of \( \sim 0.6 \) Gyr.

Subject headings: dense matter — equation of state — stars: evolution — white dwarfs

1. ASTROPHYSICAL CONTEXT

The phenomenon of phase separation and crystallization exists within the larger context of white dwarf cooling. Since the time of Mestel’s original treatment (Mestel 1952), much work has been done both to improve the input physics of the models and to make more complete observations of the white dwarf luminosity function (WDLF). In 1987, Winget et al. (1987) showed that the observed downturn in the WDLF could be understood in terms of a finite age for the Galactic disk, and that the WDLF could therefore in principle be used to determine an age for the local Galactic disk. Using the preliminary results from Liebert, Dahn, & Monet (1988, hereafter LDM) for the observed WDLF, they obtained an age for the local Galactic disk in the range 7–10 Gyr. Since then, Wood has made more detailed calculations using improved input physics, Galactic evolution models, and WD parameters to constrain this age even further (Wood 1990, 1992, 1995). Historically, these developments were foreshadowed by Schwarzschild (1958), Schmidt (1959), and D’Antona & Mazzitelli (1978), all of whom considered white dwarf evolution in a Galactic context.

Two observational surveys within the last 10 yr stand out in their importance to the field. First, Liebert et al. (1988) produced a WDLF containing 43 cool field WDs, which was the largest such sample size up to that point. More recently, Oswalt et al. (1996) produced a WDLF of 50 cool WDs in wide binaries. Using the models of Wood (1995), the LDM sample yields an age for the Galactic disk of \( \sim 7.5 \pm 1 \) Gyr, while the Oswalt et al. (1996) sample gives an age of \( 9.5^{\pm 1.1} \) Gyr. Taking the error estimates at face value, these results differ by 2 \( \sigma \). Wood & Oswalt (1998) conducted Monte Carlo simulations and found that it is unlikely that both samples are consistent with the same parent population. Further investigations will be needed to resolve the cause of this discrepancy.

In addition to the uncertainties in the observed WDLF, the way we treat various physical processes in white dwarf interiors greatly affects the ages that we derive for them. After the prediction in the early 1960s that white dwarfs should undergo a phase transition and crystallize as they cool (Abrikosov 1960; Kirzhnits 1960; Salpeter 1961), Mestel & Ruderman (1967) and Van Horn (1968) estimated that the associated release of latent heat during this process would be large enough to delay the cooling of white dwarfs significantly. Lamb & Van Horn (1975) included this energy release as part of their evolutionary calculations of a \( 1 \ M_\odot \) pure carbon white dwarf.

Stevenson (1977) was the first to propose a phase-separation model that might affect white dwarf cooling times by providing an additional source of energy analogous to the release of latent heat. This model had a carbon core with trace amounts of iron. In a later model, Stevenson (1980) suggested that a uniform mixture of carbon and oxygen would become chemically differentiated as a result of the crystallization process. Because such a redistribution of elements could lower the binding (nonthermal) energy of the star, the change in energy would be added to the thermal energy, and hence to the luminosity, of the star.
This would increase the time for a white dwarf to cool to a given luminosity, and would extend the apparent age of the Galactic disk as derived from the WDLF.

Estimates of the amount by which the age of the local Galactic disk might be extended have ranged from 0.5 to 6 Gyr (Mochkovitch 1983; Barrat, Hansen, & Mochkovitch 1988; Garcia-Berro et al. 1988; Chabrier et al. 1993; Segretain & Chabrier 1993; Hernanz et al. 1994; Segretain et al. 1994; Isern et al. 1997; Salaris et al. 1997), although recent estimates have been on the smaller end of this range; e.g., Salaris et al. (1997) calculate a delay of ~1.0 Gyr. Most of this spread in calculated age delays comes from differences in the assumed phase diagram, although the assumed C/O profile also has a large effect.

In the context of Galactic evolution, age estimates for the oldest Galactic globular clusters range from 13–16 Gyr (Pont et al. 1998) to 11.5 ± 1.3 Gyr (Chaboyer et al. 1998) and depend on a variety of parameters. In addition, a 4–6 Gyr delay is expected between the formation of the globular clusters and the formation of the Galactic thin disk (e.g., Burkert, Truran, & Hensler 1992; Chiappini, Matteucci, & Gratton 1997), while the observed white dwarf luminosity function gives an age estimate for the thin disk of 9.5 ± 1.1 Gyr (Oswalt et al. 1996), without including the effect of phase separation. Using the above numbers, we see that phase separation could add anywhere from 0 to 3 Gyr to the white dwarf ages and still be consistent with the overall picture of Galaxy formation.

In this paper, we examine the sensitivity of this calculated age delay to the various physical assumptions by varying the initial C/O profile of the white dwarf models, their total mass, and their H and He surface-layer masses. In addition, we examine the effect of using two different published phase diagrams for the phase separation process, that of Segretain & Chabrier (1993) and that of Ichimaru, Iyetomi, & Ogata (1988).

Our work improves upon previous calculations of the age delay in that we use self-consistent evolutionary models. In particular, our models use the modern OPAL opacities (Iglesias & Rogers 1993) instead of the older Cox-Stewart opacities, and we are able to treat self-consistently the age delay as a function of total stellar mass, instead of using a relation scaled by mass for the connection between the core temperature and the surface luminosity. Finally, we are able to examine surface-layer masses suggested by more recent asteroseismological investigations (Clemens 1993).

2. THE PHYSICS OF PHASE SEPARATION

2.1. Chemical Redistribution

Our present physical picture for the phenomenon of phase separation in white dwarf stars is as follows. As a white dwarf cools, it eventually reaches a temperature at which its central regions begin to crystallize. This occurs when the thermal energy of the ions becomes much smaller than the energy of the Coulomb interactions between neighboring ions. As a result, the ions settle into lattice sites and lose the ability to move freely in three dimensions.

If the white dwarf interior is initially a mixture of C and O, then recent calculations indicate that the solid that crystallizes will have a higher O content than the fluid from which it formed (Ichimaru et al. 1988; Segretain & Chabrier 1993). Thus, the crystallizing region of the white dwarf becomes O-enhanced and the fluid layer overlying this region becomes C-enhanced. Since the C is slightly less dense than the O at a given pressure, this C-enhanced fluid layer is mixed via a Rayleigh-Taylor instability (Mochkovitch 1983; Isern et al. 1997) with the layers above, and C is transported outward from the center. As the white dwarf continues to crystallize, the O-enhanced crystalline core also continues to grow, with the net result that O is transported inward in the white dwarf and C is transported outward. Thus, the chemical composition profile after significant crystallization has occurred is different from the profile before crystallization.

Just how different this profile is depends on the particular phase diagram adopted for the process. In a “spindle” diagram, the solid that forms always has an enhanced concentration of the higher charge element (in this case oxygen), and the temperature of crystallization of the mixture lies between those of the individual elements. An “azeotropic” diagram differs in that there is a range of concentrations for which crystallization takes place below the temperature of crystallization of either of the pure elements. This is somewhat analogous to the phenomenon of “supercooling.” Finally, a “eutectic” phase diagram is one in which there is a near total separation of the higher and lower charged ions upon crystallization, resulting in a segregation of the two chemical species.

Stevenson’s original phase diagram (Stevenson 1980) was a eutectic phase diagram, with C and O being immiscible in the solid phase, with the result that a pure-O core would be formed in the models during crystallization. Using a density-functional approach, Barrat et al. (1988) calculated a phase diagram of spindle type. In this case, the solid that forms is a C/O alloy, but with the O content of the solid enhanced relative to that of the fluid out of which it formed.

This problem was revisited by Ichimaru et al. (1988). They found that Stevenson’s initial prediction of a eutectic phase diagram was an artifact of his use of the random-alloy mixing (RAM) model for the internal energies in the solid phase. By comparison with Monte Carlo simulations, they found that the linear mixing formula is more accurate for the solid phase. They then used density-functional theory to derive a phase diagram of azeotropic type, which is shown by the dashed line in Figure 1. This diagram is similar to the spindle diagram, with the exception that there is a range of compositions for which the crystallization temperature is less than the crystallization temperature for either of the pure compositions.

Most recently, Segretain & Chabrier (1993) used a density-functional approach to derive phase diagrams for arbitrary binary-ionic mixtures as a function of $Z_1/Z_2$, where $Z_1$ and $Z_2$ are the nuclear charges of the two chemical species. For C and O ($Z_1/Z_2 = 0.75$), they obtain a phase diagram of spindle type, which is shown by the solid line in Figure 1.

As shown in Figure 1, these diagrams of Ichimaru et al. (1988) and Segretain & Chabrier (1993) differ slightly in the composition changes during crystallization, as well as in the temperatures at which crystallization takes place. As a result, they produce different chemical composition profiles after crystallization and different age delays.

We mention that the validity of the azeotropic phase diagram of Ichimaru et al. (1988) has recently been called into question by DeWitt, Slattery, & Chabrier (1996). DeWitt et al. find that the azeotropic diagram of Ichimaru et al. results from their use of the fitting function of Ogata &
increases. Thus, \( E_{\text{grav}} \) becomes more negative, as does \( E_{\text{coul}} \). \( E_{\text{deg}} \), however, becomes more positive, since the Fermi energy of the electrons increases with increasing density. Summing these contributions, we find that there is a net decrease in \( E_{\text{bind}} \) for the models considered here. Because of conservation of energy, this energy must be used to increase the thermal energy of the ions, which are the only significant repository of thermal energy in the cores of our white dwarf models. This energy, then, is available to be radiated away and acts as an additional luminosity source.

The various contributions to the photon luminosity \( L \) and the neutrino luminosity \( L_\nu \) of the white dwarf may be formally written as (e.g., Isern et al. 1997; Chabrier 1998)

\[
L + L_\nu = \int_0^{M_{\text{ WD}}} \left[ C_e \frac{dT}{dt} + T \left( \frac{\partial P}{\partial T} \right)_{V,X_0} \frac{dV}{dt} + \left( \frac{\partial u}{\partial X_0} \right)_{T,V} \frac{dX_0}{dt} \right] \, dm
\]

where \( V = 1/\rho \) is the specific volume, \( X_0 \) is the mass fraction of the heavier of the two chemical species (in this case oxygen), and \( u \) is the internal energy per unit mass, which contains thermal, electron degeneracy, and Coulomb contributions. The first term in the integrand on the right-hand side of equation (2) is due to the heat capacity of the core, which includes the release of the latent heat of crystallization, while the second term gives the contribution to the luminosity due to volume changes, and is usually small in white dwarfs, since the pressure \( P \) is only a weak function of the temperature. The final term gives the luminosity due to the changing chemical composition profiles within the white dwarf. This is the term we will study in our numerical calculations.

As a check on the direct evolutionary calculations, we can estimate the age delay produced by a given energy release. If we denote by \( dE \) a small amount of energy that is released during the process of phase separation, and if we assume that this energy is quickly radiated, then we can calculate an estimated age delay, \( t_d \),

\[
t_d = \int \frac{dE}{L}.
\]

In the context of a sequence of evolutionary models, this integral is operationally a sum, since a given model is computed at discrete points in time, luminosity, etc. Furthermore, since the energy \( \Delta E_i \) is released between luminosities \( L_{i-1} \) and \( L_i \), say, the average luminosity at which the energy is released is approximately \( (L_{i-1} + L_i)/2 \), so the discrete version of equation (3) becomes

\[
t_d = \sum_i \frac{\Delta E_i}{(L_{i-1} + L_i)/2}.
\]

We have used equation (4) as an alternate prescription to calculate age delays. For the larger energy releases, \( t_d \) computed in this way agrees with the delay calculated from the self-consistent evolutionary calculations, and for small energy releases it provides a better estimate, since the small energies can become masked in the numerical noise of the evolutionary calculations.
3. NUMERICS

The basis for these calculations is WDEC, the white dwarf evolutionary code, as described in Lamb & Van Horn (1975) and Wood (1990). Our current version uses the updated OPAL opacity tables (Iglesias & Rogers 1993; Wood 1993). We use the additive volume technique to treat the equation of state of the C/O mixture in the cores of our models.

3.1. The Melting Curve

Our criterion for crystallization is given by the phase diagram that we adopt, with the following caveat. Our equation of state (EOS) is based on the Lamb EOS code (Lamb & Van Horn 1975), which has $\Gamma \approx 160$ at crystallization. Here, $\Gamma \equiv Z^2 e^2 / \langle r \rangle k_B T$ is the ratio of Coulomb energy between neighboring ions to each ion’s kinetic energy. More recent calculations indicate that $\Gamma \approx 180$ (Ogata & Ichimaru 1987). As a result, our values for the crystallization temperature of C, $T_{C, \text{crystal}}$, are too high by a factor of $\sim 180/160 = 1.125$.

To remedy this situation, we could simply adjust $T_{C, \text{crystal}}$ downward accordingly, and we have done this for a few runs. This is inconvenient, however, because it places us at the edge of our EOS tables, which were calculated with $\Gamma \approx 160$. Instead, we apply a correction factor to our calculated age delays that takes into account the fact that crystallization/phase separation occurs at lower central temperatures, and therefore lower luminosities, than is calculated directly in our models. This correction to the calculated age delays is typically of the order of 25%. For example, an age delay due to phase separation computed with $\Gamma \approx 160$ might be $\sim 1$ Gyr, but with the more physical value of $\Gamma \approx 180$ would be $\sim 1.25$ Gyr. We find this procedure of computing the age delays based on the corrected luminosity during crystallization to be accurate to within 1%–2%.

3.2. Implementation in WDEC

The calculation of the evolutionary sequences is “quasi-static” in the sense that we compute a sequence of static models separated by finite steps in time. Each static model represents the cooling white dwarf at a different age and luminosity. We include the physics of phase separation using the same approximation; we assume that the timescale for any mixing that occurs is short compared to the individual evolutionary time steps (see §3.3 of this paper; Mochkovitch 1983; Isern et al. 1997), and we assume that the binding energy released by this process can be modeled by some suitably chosen local energy generation rate, $\epsilon_{\text{ps}}$ (e.g., Isern et al. 1997).

The phase-separation calculation can therefore be broken into three sections. The first part involves obtaining the changing composition profile as a function of the crystallized mass fraction, while the second part is the calculation of the cumulative energy released, also as a function of the crystallized mass fraction. The final part is the calculation of the value of $\epsilon_{\text{ps}}$, which is the energy locally deposited per unit mass per unit time. Our implementation of the complete problem is self-consistent in that we let $\epsilon_{\text{ps}}$ vary as the compositional profile changes due to crystallization, as WDEC iterates to a converged model.

The first part of the overall problem relates to the composition of the crystallizing layers. Using the phase diagram of Segretain & Chabrier (1993) or Ichimaru et al. (1988), we compute the final composition profile of the model given the initial profile, before doing a full evolutionary calculation. This is possible because the composition of the crystals that are forming is determined solely by the mass fractions of C and O present in the fluid phase, and not by the temperature and density of the medium (the temperature and density of course determine when the fluid crystallizes, but given that it is crystallizing, the chemical composition of the solid is determined solely by the composition of the fluid). We are therefore required to compute only once, at the onset of crystallization, the composition profile as a function of the crystallized mass fraction. At subsequent evolutionary times, we use this relation and the current crystallized mass fraction to interpolate onto the composition grid, which is a computationally convenient procedure.

We take this same approach for calculating the energy released. At the onset of crystallization, we calculate the total amount of energy released as a function of $M_{\text{crystal}}/M_*$. Using the relation

$$\delta E = \int_0^{M_{WD}} \left( \frac{\partial u}{\partial X_C} \right)_{T,V} \delta X_C dm,$$

where $\delta E$ is the binding energy released by the composition change $\delta X_C$. Since these changes in composition are with respect to the precrystallization state, we are in effect holding both the temperature and density profiles constant for all subsequent phases of crystallization. Holding the temperature profile constant is a quite reasonable approximation, since the vast majority of the mass in the white dwarf model is strongly degenerate for the temperature range of interest. Similarly, we expect the changes in the density profile to be small ($\delta \rho / \rho \simeq 1\%$) even in the presence of composition changes. This is a consequence of the fact that the equations of state for carbon and oxygen are very similar in the strongly degenerate regime, i.e., $\mu_e$, the atomic mass per electron, is 2.0 for both elements. This suggests that this approach would not necessarily be as accurate for carbon and iron, since $\mu_e = 2.15$ for iron.

Figure 2 provides the final justification for our assumptions. The filled circles represent the energy released as computed self-consistently at each evolutionary time step, and the solid line shows the calculated energy released assuming a static density and temperature profile as described in the above paragraph. The best agreement is for smaller amounts of crystallization, since these models differ the least from the initial static model. Even near complete crystallization, however, the difference between the two values is less than 0.5%, justifying our assumptions. Computationally, it is very convenient to compute the energy release just once at the outset and then interpolate using the present value of the crystallized mass fraction. This allows WDEC to avoid doing a calculation of the energy release for each iteration of each model, which would significantly affect the speed of the calculations.

Because all our calculations are done on evolutionary timescales, we do not have any information about the actual dissipative processes that are responsible for depositing the energy of phase separation locally. Indeed, without an accurate hydrodynamic model of the mixing process, this is not possible. Fortunately, it is more important to know the total energy released rather than exactly how this
energy is deposited within the white dwarf model. This is because the core has a very high thermal conductivity, which tends to smooth out the temperature distribution. Thus, wherever the energy is initially deposited, it will soon be shared throughout the core; indeed, an isothermal core was an assumption of the original theory of Mestel (1952), and it is still a very accurate description of the physics in the interiors of white dwarfs (e.g., García-Berro et al. 1996; Segretain et al. 1994). We therefore choose $\epsilon_{ps}$ such that the local temperature is increased by the same fractional amount throughout the core, i.e., $\delta T/T = \text{const.}$, while we simultaneously require that the total energy deposited in this way is equal to the energy released as a result of phase separation in a given time step. This is somewhat analogous to the analytical approach outlined by Isern et al. (1997), although we developed our approach for ease of numerical implementation.

There is one final adjustment that we make to the value of $\epsilon_{ps}$ as calculated above. It is due to the fact that WDEC calculates models quasi-statically, so that $\epsilon_{ps}$ is assumed to have been constant during the last time step taken, when in fact it may have changed by a substantial amount. Put another way, the value of $\epsilon_{ps}$ that WDEC calculates should be associated with the average luminosity of the present and previous time steps, not just the current luminosity. Thus, WDEC is implicitly calculating a delay based on

$$t_d = \sum \frac{\Delta E_i}{L_i},$$

(6)

instead of the expression in equation (4). We can remedy this situation by an appropriate rescaling of $\epsilon_{ps}$. If we rescale $\Delta E_i$, and hence $\epsilon_{ps}$, by $2L_i/(L_i + L_{i-1})$, then equation (6) is transformed into equation (4), and we recover the correct age delay due to crystallization when implemented in the evolution code. In the limit in which our time steps are very small, the above prescription is not necessary, but such small time steps would be computationally inconvenient with regard to both cpu time and numerical convergence.

3.3. Consistency Checks

We use three different initial C/O profiles in our analysis. In Figure 3 we show the oxygen composition in the core both before (curve a) and after (curves b and c) crystallization has taken place. We have taken a homogeneous 50:50 C/O initial distribution and assumed complete mixing of the overlying fluid layers as crystallization takes place. This should place an upper limit on the extent that phase separation can have on any particular model. We note that the composition profile after crystallization reached assuming the Segretain & Chabrier (1993) phase diagram agrees well with that given in Chabrier et al. (1993).

Figure 4 shows a different initial oxygen profile computed by Salaris et al. (1997) for a 0.61 $M_\odot$ white dwarf model. This profile was obtained by considering nuclear reaction processes in the white dwarf progenitor. Here we use a modified algorithm for mixing, which reduces to the “complete mixing” algorithm when applied to an initially flat distribution. When a shell crystallizes, we check to see whether the enhanced carbon content of the innermost fluid shell now has more carbon than the shell overlying it. If it does, then we mix the two shells and perform the same comparison with the next shell farther out, mixing all three shells if necessary. In this way we move outward through the fluid until further mixing no longer decreases the carbon content of the fluid between this point and the crys-
F. 4.—Same as Fig. 3, except that the initial C/O profile (solid line, a) is that computed by Salaris et al. (1997) for a 0.61 $M_\odot$ white dwarf model. Dotted and dashed curves (b and c) are the final profiles assuming the Segretain & Chabrier (1993) and Ichimaru et al. (1988) phase diagrams, respectively. Note that the oxygen mass fraction at the very center increases by only about 15% during crystallization in this case, compared with a 40% increase for the central value in Fig. 3. Thus, less energy is liberated.

tallization boundary. This is physically reasonable, since carbon is, in this sense, “lighter” than oxygen, so these layers should be mixed by a convective instability.

For completeness, we use a third profile taken from Wood (1990, 1995). It is designed to be representative of C/O profiles calculated in Mazzitelli & D'Antona (1986) and D'Antona & Mazzitelli (1989), who also consider nuclear reaction rates. Algebraically, it is given by

$$X_O = \begin{cases} 
0.75, & 0.0 \leq q \leq 0.5, \\
0.75 - 1.875(q - 0.5), & 0.5 < q \leq 0.9, \\
0.0, & 0.9 < q \leq 1.0, 
\end{cases}$$

(7)

where $q = M_i/M_* \text{ and } X_C = 1 - X_O$.

Our treatment of the mixing process provides an upper bound for the efficiency of this process. If we were to perform a more self-consistent calculation, we would compute the Brunt–Väisälä frequency for a given chemical composition profile in the model and mix those layers that were convectively unstable and whose computed timescales for mixing were shorter than the individual time steps in our evolutionary calculations. An analytical approach to this more detailed problem is given in Isern et al. (1997) and Mochkovitch (1983). Here we merely note that a typical value of $N^2$ for a Rayleigh-Taylor unstable region in the cores of our models is $\sim 10^{-2}$, yielding a timescale for the mixing instability of $1/|N| \sim 10^5$ s, which is clearly shorter than the relevant timescales for evolution.

4. A SIMPLE TEST PROBLEM

As a check of the standard approach to treating phase separation, we performed a simplified treatment of that given in Xu & Van Horn (1992), in which they calculate the change in binding energy of a zero-temperature C/Fe white dwarf that undergoes phase separation. In order to do this, we have written a separate code that implements the equations for a zero-temperature degenerate electron gas. Our approach is simpler in that we do not include Coulomb effects in our EOS calculations, so our approach is essentially pure Chandrasekhar theory (Chandrasekhar 1939). We do, however, include relativistic effects, which Xu & Van Horn are unable to treat. In testing this approach to phase separation, we compute the energy released as a result of phase separation in two different ways. First, we directly compute the global change in the binding energy. Second, we use the expression for the local energy release and integrate this over the mass of the model, as given in equation (5). To further simplify things, we have taken the initial state to be one in which the distribution of Fe and C is uniform throughout the model, and we have taken the final state to be a pure Fe core surrounded by a pure C mantle.

Figure 5 shows the results for differing initial fractions of C and Fe. For instance, for an initial 50:50 C/Fe distribution, we calculate an energy release of about $1.9 \times 10^{48}$ ergs, with less than a 5% relative error between the two methods. Even this small amount of error decreases as we approach a pure Fe or C initial state. This is because the density and composition changes before and after phase separation are now smaller, which makes the local calculation more accurate. For instance, if the model is 99% C uniformly distributed initially, then after phase separation most of it (in fact, 99%) is pure C. The contrast between 99% and 100% is small enough that the local density and composition changes are also small ($\delta X_C \lesssim 1\%$, $\delta \rho/\rho \lesssim$...
which means that the approximation involved in making the infinitesimal variations in equation (5) finite is more accurate. We note that it is possible to perform such a simplified treatment for a C/Fe white dwarf model and still obtain meaningful results, while for a C/O model it would not be possible. This is because \( \mu_c = 2.0 \) for both C and O, while \( \mu_c = 2.15 \) for Fe. Thus, ignoring Coulomb effects, C and O have identical equations of state, while C and Fe are still nontrivially different in this approximation.

The results of this test problem (Fig. 5) convince us that by applying equation (5) we are correctly calculating the change in the binding energy of the configuration, and thus the amount of thermal energy that has been liberated from the amount of change in the binding energy of the configuration, and thus the amount of thermal energy that has been liberated from the amount of thermal energy that has been liberated from structural sources. This shows us that the overall approach to the problem that we use here, and that has been used in the past, is sound and accurately describes the physics of phase separation.

5. RESULTS

5.1. 0.6 \( M_\odot \) White Dwarf Models

In Table 1 we give an evolutionary list of our fiducial sequence (other sequences are available from the author upon request). This sequence is more than just a convenient reference model for the rest of our calculations. Given the observed peak of the masses of isolated white dwarfs in the vicinity of 0.6 \( M_\odot \) (Weidemann & Koester 1983; Weidemann & Yuan 1989; Bergeron et al. 1995; Lamontagne et al. 1997), this model will be the most useful in our comparisons with the white dwarf population as a whole. For the surface-layer masses, we have taken and computed this effect in different surface-layer masses later in this section.

We now wish to consider the effect of phase separation on actual evolutionary sequences. We compute this effect in two different ways. Using the first method, we compare sequences in which the physics of phase separation is included with those in which it is not. Taking two such sequences, we first perform a spline fit for each sequence's age over a fixed luminosity grid, and then we calculate the difference in ages at each luminosity. The results are shown by the solid line in Figure 6, and indicate an age delay at complete crystallization of about 1.5 Gyr; this is for an initially homogeneous 50:50 C/O profile with an assumed metallicity of \( Z = 0.000 \) in the opacities. We note that all the age delays computed here are for complete crystallization of our models, and hence represent upper limits to the possible age delays. Operationally, our models are almost completely crystallized near the observed luminosity turnndown at \( \log L/L_\odot \sim -4.5 \), so that there is at most a 1% change in our calculated age delays if we consider only models that have not yet cooled beyond this point.

The other method involves applying equation (4) to a sequence undergoing phase separation, which is shown by the dotted line in Figure 6. This yields an asymptotic value for the age delay of 1.38 Gyr, which is within 5% of the age difference computed with the first method. This result indicates that the basic physics that is operating is well described by equation (4), i.e., the energy being released by phase separation is mostly being radiated in a given time step. For the remainder of the results quoted here, the age delays have been calculated using this second method (eq. [4]), since this proves to be more accurate for cases involving smaller energy releases and age delays.

We now study the effect of the initial composition profile on the age delays. We use three different profiles: one that is a homogeneous 50:50 mix (Fig. 3), one calculated by Salaris et al. (1997) (Fig. 4), and one given by equation (7). Our results are summarized in Table 2, where the columns labeled SC and I/O indicate that we have used the phase separation approximation.
operating is well described by eq. (3). Because less matter is being redistributed in the initially stratified case, we would expect less energy to be released as a result. Using the phase diagram of Segretain & Chabrier (1993) applied to a 0.6 M_⊙ white dwarf model, we find that in the homogeneous case 2.38 \times 10^{46} \text{ ergs} are released, whereas in the initially stratified case shown in Figure 4, only 1.03 \times 10^{46} \text{ ergs} are released. These energies result in age delays of 1.38 Gyr and 0.62 Gyr, respectively. Thus, the initial composition profile has a large effect on the calculated age delays. In addition, the Ichimaru et al. (1988) phase diagram produces smaller composition changes and hence smaller values, reducing the Segretain & Chabrier age delays by approximately one-third.

We now consider the effect of a nonzero metallicity in the opacity tables. The effect of varying the metallicity from Z = 0.000 to Z = 0.001 results in a change of less than 0.016% in the energies released, and is barely detectable numerically. The main effect of changing the metallicity is to affect the luminosity range at which the phase-separation energy is released, which in turn affects the age delay, \( t_{\text{delay}} \). For both the homogeneous and stratified case, the average luminosity during crystallization changes by less than 3% as Z is varied from 0.000 to 0.001, and hence \( t_{\text{delay}} \) also changes by less than 3%. Thus, the age delay is essentially insensitive to the metallicity assumed for the opacities.

Finally, we summarize the effect of different surface-layer masses in Table 3. For \( M_{\text{He}}/M_\odot = 10^{-3} \) and \( M_{\text{He}}/M_\odot = 10^{-5} \) (composition 1), we find maximum age delays of 1.45 Gyr, and for \( M_{\text{He}}/M_\odot = 10^{-4} \) and \( M_{\text{He}}/M_\odot = 10^{-6} \) (composition 2), our maximum calculated age delay is 1.56 Gyr. These values represent increases of 5% and 13%, respectively, over the age delays calculated in our fiducial model. For clarity, we note that these calculations are for the age differences introduced by phase separation alone at these new surface-layer masses; the white dwarf ages themselves change significantly with He layer mass, which produces a decrease in the calculated ages (without including phase separation) of ~0.75 Gyr for each order of magnitude increase in \( M_{\text{He}} \). Again, we find that varying the metallicity in the opacities has a small effect on these numbers, at only the 1% level.

### 5.2. The Mass Dependence

The mass of the white dwarf model affects the process of phase separation in two main ways, as is illustrated in Figure 7. First, a more massive white dwarf has a higher gravity, so that more energy is released by the subsequent rearrangement of matter. Second, the luminosity at which

| INITIAL PROFILE | SC (Gyr) | IIO (Gyr) |
|-----------------|---------|----------|
| 50:50 homogeneous | 1.38    | 0.99     |
| Stratified (Salaris et al. 1997) | 0.62    | 0.39     |
| Stratified (Wood 1995) | 0.30    | 0.20     |

**TABLE 2**

**Age Delays for 0.6 M_⊙ Models**

| INITIAL PROFILE | SC (Gyr) | IIO (Gyr) |
|-----------------|---------|----------|
| 50:50 homogeneous | 1.45    | 1.04     |
| Stratified (Salaris et al. 1997) | 0.66    | 0.42     |
| Stratified (Wood 1995) | 0.32    | 0.21     |

**TABLE 3**

**Age Delays for 0.6 M_⊙ Models with Different Surface-Layer Masses**

**FIG. 6.—Solid line:** Self-consistent calculation of the age difference between two 0.6 M_⊙ white dwarf evolutionary sequences with Z = 0.0, one of which is undergoing phase separation. **Dotted line:** Result of applying eq. (4) to the evolutionary sequence undergoing phase separation, which yields an asymptotic value for the age delay of ~1.4 Gyr. At complete crystallization (log \( L/L_\odot \sim -4.6 \)), the value given by the direct evolutionary calculation is within 5% of this, indicating that the basic physics that is operating is well described by eq. (3).

Diagrams of Segretain & Chabrier (1993) and Ichimaru et al. (1988), respectively. Near the centers of these models, we found that the initial/final oxygen mass fraction changed by only about 15% in the initially stratified case in Figure 4.

**TABLE 2**

**AGE DELAYS FOR 0.6 M_⊙ MODELS**

| INITIAL PROFILE | SC (Gyr) | IIO (Gyr) |
|-----------------|---------|----------|
| 50:50 homogeneous | 1.38    | 0.99     |
| Stratified (Salaris et al. 1997) | 0.62    | 0.39     |
| Stratified (Wood 1995) | 0.30    | 0.20     |

**Note:** SC and IIO indicate phase diagrams of Segretain & Chabrier 1993 and Ichimaru et al. 1988, respectively.

**TABLE 3**

**AGE DELAYS FOR 0.6 M_⊙ MODELS WITH DIFFERENT SURFACE-LAYER MASSES**

| INITIAL PROFILE | Composition 1 | Composition 2 |
|-----------------|---------------|---------------|
|                 | SC (Gyr)     | IIO (Gyr)     |
| 50:50 homogeneous | 1.45    | 1.04     |
| Stratified (Salaris et al. 1997) | 0.66    | 0.42     |
| Stratified (Wood 1995) | 0.32    | 0.21     |

**Note:** SC and IIO indicate phase diagrams of Segretain & Chabrier 1993 and Ichimaru et al. 1988, respectively.
crystallization occurs is higher for a more massive white dwarf, which tends to lessen the age delay for a given energy release. For example, even though the total energy released in a 1.2 $M_\odot$ model increases by a factor of $\approx 10$, the average luminosity increases by a factor of $\approx 30$, and hence there is a net decrease in the time delay relative to the 0.6 $M_\odot$ sequence.

The competition of these two effects suggests that there may be a mass for which there is a maximum age delay for a fixed composition profile. This is indeed the case, as is demonstrated in Figure 8. We find that the 0.6 $M_\odot$ white dwarf models have maximum age delays for a given composition profile (this was also found by Segretain et al. 1994). The calculated age delay is only weakly dependent upon the metallicity, as can be seen from the small difference between the solid and dashed curves. It is strongly dependent upon the initial profile, however, which can decrease the energy release, and hence the age delays, by a factor of 3 or more, as shown in Figure 8.

From the preceding calculations, we find that the age delay we calculate is near the maximum possible with respect to this parameter. In terms of the initial C/O profile, however, the situation is reversed. For a 0.61 $M_\odot$ white dwarf model, the profile calculated by Salaris et al. (1997) reduces the age delay by a factor of $\approx 2$ from the 50:50 homogeneous case. Using the profile of Wood (1995), which is based on results from Mazzitelli & D'Antona (1986) and D'Antona & Mazzitelli (1989), the reduction factor is $\approx 5$.

If we take as our best guess the initial profile of Salaris et al. (1997), assume a 0.6 $M_\odot$ white dwarf model with $M_{\text{He}}/M_*=10^{-2}$ and $M_{\text{H}}/M_*=10^{-4}$, and use the Segretain & Chabrier (1993) phase diagram, then we obtain an age delay of $\approx 0.6$ Gyr.

6. CONCLUSIONS

We find a maximum age delay of $\approx 1.5$ Gyr due to phase separation for our fiducial white dwarf model ($M_*=0.6$ $M_\odot$), and a best-guess age delay of $\approx 0.6$ Gyr. Salaris et al. (1997) have recently calculated a value of $\approx 1$ Gyr using the evolutionary models of Wood & Winget (1989). If we scale their value to our present models (assuming an average luminosity during crystallization for their models of log $L/L_\odot \approx -4.1$), then we obtain 0.75 Gyr, which is in basic agreement with our estimate of 0.62 Gyr. The differ-
ences in these models are mainly due to the different surface-layer masses adopted; more recent asteroseismological analyses of the class of DAs suggests that the appropriate surface-layer masses are $M_{\text{H}}/M_* \sim 10^{-2}$ and $M_{\text{H}}/M_* \sim 10^{-4}$ (Clemens 1993, 1995), and these are the values that we have assumed.

The most important factors influencing the size of the calculated age delay are the total stellar mass and the initial composition profile. We find that the largest age delays occur in models with masses of $\sim 0.6 M_\odot$ near the peak in the observed white dwarf mass distribution. The best current theoretical initial C/O profile produces models with smaller age delays, of $\sim 0.6$ Gyr. In addition, if we use the phase diagram of Ichimaru et al. (1988) instead of the Segretain & Chabrier (1993) phase diagram, then our age delays are reduced by about one-third. We note that the prescription we have adopted for the mixing during crystallization provides an upper bound for the efficiency of this process, and hence a maximum for the age delay. More realistic treatments of the mixing process may reduce the age delay. We find that varying the opacities (via the metallicity) and varying the surface-layer masses has only a small effect ($\lesssim 10\%$) on the calculated age delays.

Our calculations do not take into account the possible age delays introduced by the phase separation of heavier trace-element species, such as $^{22}$Ne, which may produce significant age delays of 2–3 Gyr (Segretain et al. 1994; Hernanz et al. 1994). These species would arise from the initial abundance of metals in the main-sequence stars that later evolved into white dwarfs. This effect may only be important for Population I stars, however, and would not therefore affect the calculated ages of the cool white dwarfs that populate the turndown in the WDLF, since these white dwarfs were formed very early in the history of the Galaxy (Hernanz et al. 1994).

In the context of Galactic evolution, age estimates for the oldest Galactic globular clusters range from 13–16 Gyr (Pont et al. 1998) to $11.5 \pm 1.3$ Gyr (Chaboyer et al. 1998), and depend on a variety of parameters. In addition, a 4–6 Gyr delay is expected between the formation of the globular clusters and the formation of the Galactic thin disk (e.g., Burket et al. 1992; Chiappini et al. 1997), while the observed white dwarf luminosity function gives an age estimate for the thin disk of $9.5^{+1.6}_{-1.4}$ Gyr (Oswalt et al. 1996), without including the effect of phase separation. Using the above numbers, we see that phase separation could add anywhere from 0 to 3 Gyr to the white dwarf ages and still be consistent with the overall picture of Galaxy formation. Our calculated maximum value of $\lesssim 1.5$ Gyr fits within these bounds, as does our best-guess value of $\lesssim 0.6$ Gyr.

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