PROPERTIES OF DEFLAGRATION FRONTS AND MODELS FOR TYPE Ia SUPERNOVAE

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ABSTRACT

Detailed models of the explosion of a white dwarf that include self-consistent calculations of the light curve and spectra provide a link between observational quantities and the underlying explosion model. These calculations assume spherical geometry and are based on parameterized descriptions of the burning front. Recently, the first multidimensional calculations for nuclear burning fronts have been performed. Although a fully consistent treatment of the burning fronts is beyond the current state of the art, these calculations provide a new and better understanding of the physics. Several new descriptions for flame propagation have been proposed by Khokhlov et al. and Niemeyer et al. Using various descriptions for the propagation of a nuclear deflagration front, we have studied the influence on the results of previous analyses of Type Ia supernovae, namely, the nucleosynthesis and structure of the expanding envelope.

Our calculations are based on a set of delayed detonation models with parameters that give a good account of the optical and infrared light curves and of the spectral evolution. In this scenario, the burning front first propagates in a deflagration mode and subsequently turns into a detonation. The explosions and light curves are calculated using a one-dimensional Lagrangian radiation-hydro code including a detailed nuclear network.

We find that the results of the explosion are rather insensitive to details of the description of the deflagration front, even if its speed and the time from the transition to detonation differ almost by a factor of 2. For a given white dwarf (WD) and a fixed transition density, the total production of elements changes by less than 10%, and the distribution in the velocity space changes by less than 7%. Qualitatively, this insensitivity of the final outcome of the explosion to the details of the flame propagation during the (slow) deflagration phase can be understood as follows: for plausible variations in the speed of the turbulent deflagration, the duration of this phase is several times longer than the sound crossing time in the initial WD. Therefore, the energy produced during the early nuclear burning can be redistributed over the entire WD, causing a slow preexpansion. In this intermediate state, the WD is still bound but its binding energy is reduced by the amount of nuclear energy. The expansion ratio depends mainly on the total amount of burning during the deflagration phase. Consequently, the conditions are very similar under which nuclear burning takes place during the subsequent detonation phase. In our example, the density and temperature at the burning front changes by less than 3%, and the expansion velocity changes by less than 10%. The burning conditions are very close to previous calculations which used a constant deflagration velocity. Based on a comparison with observations, those required low deflagration speeds (∼2%–3% of the speed of sound). Exceptions to the similarity are the innermost layers of ∼0.03–0.05 M⊙. Still, nuclear burning is in nuclear statistical equilibrium, but the rate of electron capture is larger for the new descriptions of the flame propagation. Consequently, the production of very neutron-rich isotopes is increased. In our example, close to the center Yc is about 0.44, compared to 0.46 in the model with constant deflagration speed. This increases the 48Ca production by more than a factor of 100 to 3.6·6 M⊙.

Conclusions from previous analyses of light curves and spectra on the properties of the WD and the explosions will not change, and even with the new descriptions, the delayed detonation scenario is consistent with the observations. Namely, the central density results with respect to the chemical structure of the progenitor and the transition density from deflagration to detonation do not change. The reason for this similarity is the fact that the total amount of burning during the long deflagration phase determines the restructuring of the WD prior to the detonation. Therefore, we do not expect that the precise, microphysical prescription for the speed of a subsonic burning front has a significant effect on the outcome.

However, at the current level of uncertainties for the burning front, the relation between properties of the burning front and of the initial white dwarf cannot be obtained from a comparison between observational and theoretical predictions by one-dimensional models. Multidimensional calculations are needed (1) to get inside the relations between model parameters such as central density and properties of the deflagration front and its relation to the transition density between deflagration and detonation and (2) to make use of information on asphericity that is provided by polarization measurements.

These questions are essential to test, estimate, and predict some of the evolutionary effects of SNe Ia and their use as cosmological yardsticks.

Subject headings: nuclear reactions, nucleosynthesis, abundances — supernovae: general — turbulence

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1. INTRODUCTION

The standard scenario for Type Ia supernovae (SNe) consists of massive carbon-oxygen white dwarfs (WDs) with masses close to the Chandrasekhar mass which accrete through Roche lobe overflow from an evolved companion star (Whelan & Iben 1973; Nomoto & Sugimoto 1977). In these accretion models, the explosion is triggered by compressional heating. From the theoretical standpoint, the key question is how the flame ignites and propagates through the white dwarf. Several models within this general scenario have been proposed in the past, including detonations, deflagrations, and delayed detonations, where the flame starts as a deflagration and turns into a detonation later on (Khokhlov 1991; Yamaoka et al. 1992; Woosley & Weaver 1994). The latter scenario and its variation, “pulsating delayed detonation,” seem to be the most promising because, from the general properties and the individual light curves and spectra, it can account for the majority of SNe Ia events (e.g., Höflich & Khokhlov 1996, and references therein). We note that with the discovery of the super-soft X-ray sources, potential progenitors have been found (e.g., van den Heuvel et al. 1992; Rappaport et al. 1994). What we observe as a supernova event is not the explosion itself but the light emitted from a rapidly expanding envelope produced by the stellar explosion. As the photosphere recedes, deeper layers of the ejecta become visible. A detailed analysis of the light curves and spectra gives us the opportunity to determine the density, velocity, and composition structure of the ejecta and to provide a direct link to observations. A successful application of observational constraints requires both accurate early light curves and spectral observations and detailed theoretical models that are coupled tightly with the hydrodynamical calculations (e.g., Harkness 1991; Höflich, Khokhlov, & Müller 1991; Bravo et al. 1996). According to previous results, normal bright SNe Ia can be explained by delayed detonation and pulsating delayed detonation models (e.g., SN 94D, Höflich 1995). During the deflagration phase, the deflagration velocity is 3% of the sound speed. In general, a transition from deflagration to detonation is required at densities of about \((2-2.5) \times 10^7 \text{ g cm}^{-3}\). Central densities of the initial WDs are \(\approx 2.0 \times 10^7 \text{ g cm}^{-3}\). Despite their success, hydrodynamical models are limited by the parametrized description of the burning front and the ad hoc adjustment of the density at which the deflagration turns into a detonation.

Recently, significant progress has been made toward a better understanding of the propagation of nuclear burning fronts. The first multidimensional hydrodynamic calculations of the deflagration fronts have been performed (Khokhlov 1995; Niemeyer & Hillebrandt 1995). A basic, qualitative understanding of the mechanism which leads to a transition from a deflagration to a detonation phase may have been achieved (Khokhlov, Oran, & Wheeler 1997a, 1997b; Niemeyer & Woosley 1997). Qualitatively, the results agree between different hydrodynamical numerical simulations, but a full description of the deflagration in the entire white dwarf and consistent calculations of the transition require high resolution in three dimensions, which is beyond the current state of the art. Note that the cited references rely on their subgrid models and do not resolve the turbulent deflagrations on small scales and for appropriate Reynolds numbers. Moreover, the transition from a deflagration to a detonation is still not well understood.

Here the question is addressed of how our results of the explosions vary if we use descriptions for the deflagration front which use functional relations derived from three-dimensional calculations.

1.1. Hydrodynamics

The explosions are calculated using a one-dimensional radiation-hydro code, including nuclear networks (Höflich & Khokhlov 1996, and references therein). This code solves the hydrodynamical equations explicitly by the piecewise parabolic method (Collela & Woodward 1984) on \(\approx 1000\) radial zones. The zones near the burning front are subdivided to properly track its propagation. The code includes the solution of the frequency-averaged radiation transport implicitly via moment equations, expansion opacities, and a detailed equation of state. The frequency-averaged variable Eddington factors and mean opacities are calculated by solving the frequency-dependent transport equations. About 1000 frequencies (in 100 frequency groups) and about 500 depth points are used. Nuclear burning is taken into account using a network of 218 nuclei (Thielemann, Nomoto, & Hashimoto 1996; Höflich, Wheeler, & Thielemann 1998, and references therein).

1.2. Description of the Burning Front

We have considered three cases:

Case 1.—\(v_{\text{burn}} = \text{const } v_{\text{sound}}\). In our previous investigations, \(\text{const} = 0.03\) has been found to give the best fits to observations. This corresponds to the fractal dimension \(D = 2\) in the description of Woosley & Weaver (1994) which suggested \(D = 2 - 2.5\).

Cases 2 and 3.—Here we assumed that \(v_{\text{burn}} = \max(v_l, v_t)\), where \(v_l\) and \(v_t\) are the laminar and turbulent velocities, respectively. The term \(v_l\) is calculated in accord with Khokhlov et al. (1997a).

Intrinsically, turbulent combustion is a three-dimensional problem. It is driven on large scales by the buoyancy of the burning products. The turbulent cascade penetrates down to very small scales, and makes the rate of deflagration independent of the microphysics. Turbulent combustion in a uniform gravitational field and in static conditions singles out the propagation of the flame against gravity \(g\). Chemical combustion experiments have been performed in confined environments, so-called combustion chambers. These experiments can be reproduced by numerical simulations. The propagation speed can be described by

\[
v_t = C_1 \sqrt{\alpha_T g L_f};
\]

\[
C_1 = 0.5, \quad \alpha_T = (x - 1)/(x + 1),
\]

\[
\alpha = \rho^2(t_{\text{burn}})/\rho^2(t_{\text{burn}}),
\]

where \(\alpha_T\) is the Atwood number, \(L_f\) is the characteristic length scale, and \(\rho^+\) and \(\rho^-\) are the densities in front of and behind the front, respectively. However, despite the success in terrestrial experiments, the basic assumptions of both a uniform gravitational field and static conditions are violated in the rapidly expanding envelopes of SNe Ia. The main effect of expansion is the freezeout of the turbulence on scales of \(L_f\) where the turbulent velocity due to Rayleigh-Taylor instabilities is comparable to the differential expansion velocities on those scales, i.e.,

\[
v_t \approx v_{\text{exp}} = L_f/\tau_{\text{ex}}.
\]
Based on this idea, Khokhlov et al. (1997b) suggested using the average turbulent velocity (eq. [1]), using \( z \) for uniform, static conditions, and using the mean expansion timescale determined by one-dimensional simulations \( \tau_{\text{exp}} \approx \frac{dt}{d \ln R_{\text{WD}}} \). He found for the propagation speed of the turbulent burning front

\[
v_i = 0.0474 \times \sqrt{(gL_f)},
\]

As a third case for the description, we followed the recipe of Khokhlov but did some modifications by taking \( \alpha, L_f \), and \( \tau_{\text{exp}} \) directly from the hydro at the location of the burning front. Freezeout was assumed when the radius of a mass element had doubled after being burned. \( C_1 \) in equation (1) has been varied. Note that a variation in \( C_1 \) is equivalent to scaling the relative length scale for the freezeout. We varied \( C_1 \) in the range to cover a parameter space which includes both the descriptions suggested by Khokhlov et al. (1997b) and Niemeyer & Woosley (1997).

2. RESULTS

The influence of the description of the deflagration front has been studied with the example of a set of delayed detonation models based on the same C-O WD with a mass of 1.39 \( M_\odot \) and a central density \( \rho_c = 2.0 \times 10^9 \) g cm\(^{-3} \). In all cases, a transition density \( \rho_{\text{tr}} \) of 2.3 \( \times 10^7 \) g cm\(^{-3} \) has been assumed. The description of the deflagration front has been varied. The deflagration velocity is taken to be 3\% of the speed of sound \( v_{\text{sound}} \) for model m2z02y24i5, and the approximation of Khokhlov is used for m2z02y24i4. Equation (1) has been used for models m2z02y24i1–3 with \( C_1 = 0.15, 0.20, \) and 0.25; m2 indicates that the initial mass of the progenitor was 2 \( M_\odot \), with solar metallicity \( z02 \) and 24\% of helium.

In Figure 1, the velocity of the burning front is shown as a function of time. In general, the speed of the burning front is mainly determined by the turbulent speed but the very early time. As can be expected, the transition density is reached later in time for smaller \( v_d \) because of the lower energy production per time and, consequently, the slower preexpansion.

The final density, velocity, and chemical structures for the most important elements are given in Figures 2 and 3. Overall, the structures are very similar because the total energy release depends on the amount of the released energy and the initial structure of the WD. Even the chemical structure, or more precisely, the location of transition between different regimes of burning (e.g, from partial to total Si burning), varies between all models by \( \leq 7\% \) in the space of the final expansion velocity. The total production of the most abundant elements changes by only 4\% and 3\% with mean values of 0.60 and 0.16 \( M_\odot \) for \( M(^{56}\text{Ni}) \) and \( M(\text{Si}) \), respectively.

At first, one may expect a rather high sensitivity of the final burning products to details of the description of the burning front (Hillebrandt, Niemeyer, & Woosley 1997), but the tests show a low insensitivity (Figs. 2 and 3). This result can be understood by the very nature of delayed detonation models. Qualitatively, the low dependence on the details of the flame propagation during the (slow) deflagration phase can be understood as follows: the production of intermediate-mass elements depends on the expansion of the outer envelope before the burning front “arrives.” This preexpansion occurs during the deflagration phase. For plausible variations in the speed of the turbulent deflagration, the duration of this phase is several times longer than the sound crossing time in the initial WD. Therefore, the energy produced during the early nuclear burning can be redistributed over the entire WD, causing a smooth lifting/preexpansion. In this intermediate state the WD is still bound, but its binding energy is reduced by the amount of nuclear energy. The expansion ratio depends mainly on the total amount of burning during the deflagration phase. Consequently, the conditions are very similar under which nuclear burning takes place during the subsequent detonation phase. In Figure 4, the burning conditions just behind the front are given for the two extreme models with \( C_1 = 0.15 \) and \( C_1 = 0.25 \). The durations of the deflagration phase are about 1.7 and 2.9 s, respectively. Both models start with laminar deflagrations, i.e., the conditions under which nuclear burning takes place are similar. After about 3\% of a solar mass has been burned, the flame speed is determined by the turbulent speeds which differ by about 70\%, and the densities behind the burning front differ significantly. However, in all models, the temperatures are sufficiently high to burn up to nuclear statistical equilibrium (NSE), and mainly \( ^{56}\text{Ni} \) is produced. After burning about 0.25 and...
Fig. 2.—Final density and velocity as a function of mass for different models (see text)

Fig. 3.—Same as Fig. 2, but final chemical composition as a function of mass
0.32 \( M_\odot \) for the models with low and high turbulent speeds, the transition from burning as a deflagration to a detonation is triggered. Outside \( \approx 0.4 \ M_\odot \), the densities, and consequently the temperatures, are very similar. Note that the layers of partial burning are located above \( \approx 0.6 \ M_\odot \). The expansion rate, relevant for the timescale for adiabatic cooling, differs by only 7%. This similarity in the burning conditions explains the insensitivity in the final chemical profiles. The preexpansion is even more similar than can be expected from this argument because the hydrodynamical timescale for energy redistribution is \( \approx 1 \) s. Thus, the influence of the burning in the deflagration mode during the few tenths of a second before turning to a detonation is reduced with respect to the preexpansion of the outer layers. As soon as the front turns into a detonation, the remainder of the WD is burned almost “instantly” or better on nuclear timescales.

For comparison, the properties at the burning front are given for the description with \( v_{\text{burn}}/v_{\text{sound}} = \text{const} = 0.03 \), where the constant has been tuned to give good agreement between the observed and calculated light curves and spectra (Höflich & Khokhlov 1996). This description for the burning front is widely used in the literature and is consistent with constraints from the nucleosynthesis (e.g., Brachwitz et al. 1998; Khokhlov 1991; Thielemann et al. 1997; Woosley & Weaver 1994, with \( D = 2 \)).

In the outer layers, the burning conditions are virtually identical to including the expansion rate. Slightly less material is burned during the deflagration phase because the front is faster and the initial energy production is larger during the first second (see Fig. 1).

Note that the amount of burning under very high densities, and therefore, the production of neutron-rich isotopes in the central region of the WD, depends sensitively on the speed of the front which, in the more realistic descriptions, starts with laminar speed. In comparison with previous calculations, this boosts the production of very neutron-rich isotopes such as \(^{48}\text{Ca}\). In our example and close to the
center, $Y_e$ is about 0.44 compared to 0.46 in the model with constant deflagration speed. This increases the $^{48}\text{Ca}$ production from $2 \times 10^{-6} M_\odot$ to $3 \times 10^{-6} M_\odot$. This increase is the same for all models which start with the laminar deflagration speed. For a systematic study of different flame speeds, see Brachwitz (1999). Note that the production of neutron-rich isotopes depends on the central density and whether the ignition is very close to the center or off-center. In principle, this opens a new window for detailed analyses of the progenitor and the ignition process in SNe Ia.

3. CONCLUSIONS

The final structure of the expanding envelope is rather insensitive to the detailed description of the burning front during the deflagration phase. It depends mainly on global quantities, namely, the total amount of burning during the deflagration phase and the sound speed in the initial white dwarf. Therefore, the detailed, spatial structure of the burning front cannot be expected to change the result of the explosion. Findings from previous analyses of light curves and spectra on the properties of the WD and the explosions will not change, and the new description of the deflagration front is consistent with the observations. The central density, conclusions about the chemical structure of the progenitor, and the transition density from deflagration to detonation do not change. Differentially, evolutionary effects and their consequences for observations can be explored by parameterized models.

The validity of the transition density found in previous analyses also poses a question for our understanding of the transition from turbulent deflagration to detonation, which is not well understood. Currently, it is attributed to the Zeldovich mechanism, which is based on the adiabatic mixing of burned and unburned material where the entropy increase in the unburned fuel triggers the explosion (Khokhlov et al. 1997b; Niemeyer & Woosley 1997). The relative change over the entire range of parameterizations discussed here corresponds to a change in the transition density of $\approx 10\%$. This leaves us with the puzzle of why the theoretical estimates for the transition density are lower by a factor of about 0.7 (Khokhlov et al. 1997b) and 0.4 (Niemeyer & Woosley 1997). For reasons, we can only speculate. The differences between the latter values may indicate the size of the uncertainties in our understanding of this transition process. Other, not yet considered, microscopic effects may be involved. Another possibility may be that we measure two different things. To derive the model parameters from the observations, we measure the mean density at the burning front when the transition occurs, whereas the theoretical considerations provide information on the location where the transition occurs. Perhaps the fragmentation of the burning front causes the transition to occur somewhat ahead of the mean front. If this interpretation is correct, it may indicate that the detonation is started at about 10\%–20\% (in radius) ahead of the mean front. In either case, further investigation is needed and will provide new insights into the properties of nuclear burning fronts.

We note that the nucleosynthesis, optical and IR spectra, and light curves seem to require that, after an initial phase of slow burning, the front moves with velocities close to the speed of sound to keep up with the expanding outer layers and to burn carbon under low-density conditions. The latter is indicated by a strong line at 1.05 $\mu$m, which can be attributed to Mg II and indicates expansion velocities in excess of 15,000–16,000 km s$^{-1}$. In general, unburned carbon is restricted to velocities above 20,000 km s$^{-1}$ (Fisher et al. 1997; Höflich 1995; Höflich & Khokhlov 1996; Khokhlov 1991; Nugent et al. 1997; Thielemann et al. 1996; Wheeler et al. 1998). Both the high velocities in Mg and C are consistent with delayed detonation models but inconsistent with the classical deflagration model W7 (Nomoto, Thielemann, & Yokoi 1984). Extended mixing of the inner, Ni-rich layers can be excluded from the spectra (Höflich, Wheeler, & Thielemann 1998). Mixing of the outer layers may produce Si lines at high velocity (Nugent et al. 1997) but does not solve the problems with Mg and C. Although the evidence favors the delayed detonation scenario in the framework of one-dimensional models, this does not necessarily imply that a transition from detonation occurs. Alternatively, the flame may propagate as a very fast deflagration wave (W. Hillebrandt 1999, private communication). In this context, the transition density may be attributed to the region where the turbulent flame is not driven anymore by Rayleigh-Taylor instabilities.

Finally, we also have to stress the limits of one-dimensional models with a parameterized description of the deflagration front and the need for multidimensional hydrodynamical calculations and radiation transport. These limits are critical to understand details of the explosion and to use SNe Ia as distance indicators on cosmological scales (e.g., Schmidt et al. 1998; Perlmutter et al. 1999). Observations clearly show a strong relation between the maximum brightness and the decline relation (Phillips 1993). From theoretical models, the amount of radioactive nickel produced during the explosion of a massive white dwarf has been identified as the basic quantity which provides this relation. The relation is independent of details of the model; however, the amount of $^{56}\text{Ni}$ actually produced depends on a combination of free parameters in the models such as the central density and chemical composition of the WD and the propagation of the burning front. If these parameters are varied independently within the limits indicated by individual fits to observations (Höflich & Khokhlov 1996), we expect a spread around the mean maximum brightness/decline relation of $\approx 0.4$ mag, which is consistent with the spread based on the Cerro Tololo Inter-American Observatory data published by Hamuy et al. (1996). Recent redefinitions of statistical methods and new observations suggest a much tighter relation with a spread of $\approx 0.12$ mag (Schmidt et. al. 1998). This narrow spread cannot be understood in the context of the parameter range allowed by current analyses. This tight relation may hint at an underlying coupling of the progenitor, the accretion rates, and the propagation of the burning front. However, on the current level of uncertainties, the relation between the free parameters such as transition density and initial central density cannot be deduced from a comparison between observation and theoretical predictions by one-dimensional models. Multidimensional calculations are needed to test, e.g., the relation between chemical composition or central density of the white dwarf and the properties of the deflagration front and its relation to the transition density between deflagration and detonation.

In general, the chemical signature of the deflagration phase is “wiped out” by the detonation front, but a small trace may remain at the interface between complete and incomplete Si burning (Thielemann et al. 1996). Although not conclusive, recent polarization measurement may
already hint at the existence of such inhomogeneities in the chemical structure (Wang, Wheeler, & Höflich 1998).

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