Mixing in classical novae: a 2-D sensitivity study

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ABSTRACT

Context. Classical novae are explosive phenomena that take place in stellar binary systems. They are powered by mass transfer from a low-mass, main sequence star onto a white dwarf. The material piles up under degenerate conditions and a thermonuclear runaway ensues. The energy released by the suite of nuclear processes operating at the envelope heats the material up to peak temperatures of ~(1–4) × 10^8 K. During these events, about 10^{-3}–10^{-5} M⊙, enriched in CNO and other intermediate-mass elements, are ejected into the interstellar medium. To account for the gross observational properties of classical novae (in particular, a metallicity enhancement in the ejecta above solar values), numerical models assume mixing between the (solar-like) material transferred from the companion and the outermost layers (CO- or ONe-rich) of the underlying white dwarf.

Aims. The nature of the mixing mechanism that operates at the core-envelope interface has puzzled stellar modelers for about 40 years. Here we investigate the role of Kelvin-Helmholtz instabilities as a natural mechanism for self-enrichment of the accreted envelope with core material.

Methods. The feasibility of this mechanism is studied by means of the multidimensional code FLASH. Here, we present a series of 9 numerical simulations performed in two dimensions aimed at testing the possible influence of the initial perturbation (duration, strength, location, and size), the resolution adopted, or the size of the computational domain on the results.

Results. We show that results do not depend substantially on the specific choice of these parameters, demonstrating that Kelvin-Helmholtz instabilities can naturally lead to self-enrichment of the accreted envelope with core material, at levels that agree with observations.

Key words. novae, cataclysmic variables – nuclear reactions, nucleosynthesis, abundances – convection – hydrodynamics – instabilities – turbulence

1. Introduction

Classical novae are cataclysmic stellar events. Their thermonuclear origin, theorized by Schatzmann (1949, 1951) and Cameron (1959) – see also Gurevitch & Lebedinsky (1957) and references therein – has been established through multi-wavelength observations and numerical simulations pioneered by Sparks (1969), who performed the first 1-D, hydrodynamic nova simulation. These efforts helped to establish a basic picture, usually referred to as the thermonuclear runaway model (TNR), which has been successful in reproducing the gross observational properties of novae, namely the peak luminosities achieved, the abundance pattern, and the overall duration of the event; see Starrfield et al. (2008), José & Shore (2008), José & Hernanz (2007) for recent reviews.

Many details of the dynamics of nova explosions remain to be explored. In particular, there are many observed cases of nonspherical ejecta, inferred from line profiles during the early stages of the outburst and from imaging of the resolved ejecta, including multiple shells, emission knots, and chemical inhomogeneities. Although the broad phenomenology of the outburst can be captured by 1-D calculations, it is increasingly clear that the full description requires a multidimensional hydrodynamical simulation of such outbursts. To match the energetics, peak luminosities, and the abundance pattern, models of these explosions require mixing of the material accreted from the low-mass stellar companion with the outer layers of the underlying white dwarf. In fact, because of the moderate temperatures achieved during the TNR, a very limited production of elements beyond those from the CNO-cycle is expected (Starrfield et al. 1998, 2009; José & Hernanz 1998; Kozvet & Prialnik 1997; Yaron et al. 2005), and the specific chemical abundances derived from observations (with a suite of elements ranging from H to Ca) cannot be explained by thermonuclear processing of solar-like material.

Mixing at the core-envelope interface represents a likely mechanism.

The details of the mixing episodes by which the envelope is enriched in metals have challenged theoreticians for nearly 40 years. Several mechanisms have been proposed, including diffusion-induced mixing (Prialnik & Kovetz 1984; Kovetz & Prialnik 1985; Iben et al. 1991, 1992; Fujimoto & Iben 1992), shear mixing at the disk-envelope interface (Durisen 1977; Kippenhahn & Thomas 1978; MacDonald 1983; Livio & Truran 1987; Kutter & Sparks 1987; Sparks & Kutter 1987), convective overshoot-induced flame propagation (Woosley 1986), and...
mixing by gravity wave breaking on the white dwarf surface (Rosner et al. 2001; Alexakis et al. 2004). The multidimensional nature of mixing has been addressed by Glasner & Livne (1995) and Glasner et al. (1997, 2005, 2007) with 2-D simulations of CO-novae performed with VULCAN, an arbitrarily Lagrangian Eulerian (ALE) hydrocode capable of handling both explicit and implicit steps. They report an effective mixing triggered by Kelvin-Helmholtz instabilities that produced metallicity enhancements to levels in agreement with observations. Similar studies (using the same initial model as Glasner et al. 1997) were conducted by Kercek et al. (1998, 1999) in 2-D and 3-D, respectively. Their results, computed with the Eulerian code PROMETHEUS, displayed mild TNRs with lower peak temperatures and velocities than Glasner et al. (1997) and insufficient mixing. While Glasner et al. (1997) argue that substantial mixing can naturally occur close to peak temperature, when the envelope becomes fully convective and drives a powerful TNR. Kercek et al. (1998) conclude instead that mixing must take place much earlier: if it occurs around peak temperature, it leads to mild explosions or to events that do not resemble a nova.

The differences between these studies have been carefully analyzed by Glasner et al. (2005), who conclude that the early stages of the explosion, before TNR ignition when the evolution is quasi-static, are extremely sensitive to the adopted outer boundary conditions. They show that Lagrangian simulations, in which the envelope is allowed to expand and mass is conserved, lead to consistent explosions. In contrast, in Eulerian schemes with a “free outflow” outer boundary condition, the choice adopted in Kercek et al. (1998), the outburst can be artificially quenched. The scenario was revisited by Casanova et al. (2010), who show that simulations with an Eulerian scheme – the FLASH code – and a proper choice of the outer boundary conditions can produce deep-mixing of the solar-like accreted envelopes with core material. The puzzling results reported by Kercek et al. (1998) stress the need for a systematic evaluation of the effect that different choices of model parameters (e.g. the intensity and location of the initial temperature perturbation, resolution, or size of the computational domain) may have on the results. To this end, we performed a series of 9 numerical simulations in 2-D aimed at testing the influence of these parameters on the level of metal enhancement of the envelope. Here we report the results of these simulations.

Our paper is organized as follows. In Sect. 2 we explain our input physics and initial conditions. Then Sect. 3 is devoted to studying the mixing at the core-envelope interface for our fiducial model. In Sect. 4 the effects of the size of the initial perturbation are analyzed, while in Sect. 5 we discuss the effects of the size of the computational domain. In Sect. 6 we quantify the influence of the grid resolution. Finally, in Sect. 7 we discuss the significance of our results and draw our conclusions.

2. Input physics and initial conditions

The simulations reported here were performed with FLASH, a parallelized explicit Eulerian code, based on the piecewise parabolic interpolation of physical quantities for solving the hydrodynamical equations, and with adaptive mesh refinement (see Fryxell et al. 2000). As in Casanova et al. (2010), we used the same initial model as Glasner et al. (1997) and Kercek et al. (1998): a 1 $M_\odot$ CO white dwarf that accretes solar composition matter ($Z = 0.02$) at a rate of $5 \times 10^{-7} M_\odot$ yr$^{-1}$. The model was evolved spherically (1-D) and mapped onto a 2-D cartesian grid, when the temperature at the base of the envelope reached $\approx 10^8$ K. It initially comprised 112 radial layers – including the outermost part of the CO core – and 512 horizontal layers. The mass of the accreted envelope was about $2 \times 10^{-3} M_\odot$. Nuclear energy generation is handled through a network of 13 species ($^1$H, $^4$He, $^{12}$C, $^{13}$C, $^{14}$N, $^{15}$N, $^{16}$O, $^{17}$O, and $^{19}$F), and connected through 18 nuclear reactions. We adopted the conductive and radiative opacities from Timmes (2000) and an equation of state based on Timmes & Swesty (2000). Periodic boundary conditions were imposed on both sides of the computational domain with vertical hydrostatic equilibrium with an outflow constraint at the top and a reflecting constraint at the bottom on the velocity (see Zingale et al. 2002). A summary of the main characteristics of the 9 models computed in this work is given in Table 1, where $H$ is the distance from the perturbation to the initial core-envelope interface, $R_c$ and $R_i$, $\delta T$, and $\delta t$ are the size, strength, and duration of the temperature perturbation, and $Z$ the mass-averaged metallicity of the envelope at the end of the calculations.

3. 2-D simulations of mixing at the core-envelope interface

In this section, we describe the basic features of our fiducial model A, as a framework for further discussion of the effect of the parameter choices on our results. A movie, showing the development of Kelvin-Helmholtz instabilities, in terms of the $^{12}$C content, up to the time when the convective front hits the upper computational boundary, ModelA-2D.wmv, is available online or at http://www.fen.edu/users/jjose/downloads.html. The simulation was performed for the condition of model A, as summarized in Table 1.

For all sequences reported in this work, the relaxation of the initial model to guarantee hydrostatic equilibrium, together with the small amount of numerical viscosity – in contrast with the simulations performed by Glasner et al. (1997) – requires an initial perturbation close to the core-envelope interface to trigger the onset of instabilities early in the calculations. The initial perturbation is applied to the temperature using four parameters: strength, location, size and duration. Model A assumes a top-hat temperature perturbation wherever $((x - x_0)/R_c)^2 + ((y - y_0)/R_i)^2 \leq 1$, where $x$ and $y$ are the space coordinates measured from the center of the perturbation, $(x_0, y_0)$, and $R_c$ and $R_i$ indicate its spatial extent. We fixed $x_0 = 5 \times 10^7$ cm in all sequences. The strength of the perturbation is 5% in temperature in all cases but one (see Table 1). It is 1 km wide, applied only during the initial timestep (that is, the temperature is fixed only during $10^{-10}$ s), and imposed on the core-envelope interface ($y_0 = 5.51 \times 10^9$ cm). The resolution adopted in model A is $1.56 \times 1.56$ km, and the size of the computational domain is $800 \times 800$ km.

The initial perturbation drives a shear flow that triggers the formation of instabilities (Fig. 1), about 150 $s$ after the start of the simulation. As soon as material from the core is mixed into the envelope, small convective cells develop. At this early stage, the fluid has a large Reynolds number, with a characteristic eddy length of 50 km, fluid velocities ranging between $v = 10^7 - 10^8$ cm s$^{-1}$, and a dynamic viscosity $\nu$ of $2 \times 10^7$ P. The

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1 The different values adopted for $R_c$ and $R_i$ in models F and G are used to account for the assumption of a rectangular (rather than square) computational domain.

2 The dynamic viscosity evaluates the resistance to flow of a fluid under an applied force. More precisely, it is defined as the tangential force per unit area required to move one horizontal plane with respect to the other at unit velocity when maintaining a unit distance apart by the fluid.
the simulation when the core-envelope interface temperature is \( T_{\text{base}} \). At this final stage, the mean mass-averaged metallicity of the calculations because of the Eulerian nature of the FLASH code. When the matter crosses the outer boundary of the computational domain, we stop the simulation even for a point-like TNR ignition.

To quantify the influence of the initial perturbation on our results, we have performed a series of 2-D hydrodynamic tests for a set of different durations, strengths (intensities), locations and sizes of the perturbation. For simplicity, a top-hat perturbation, centered at \( x_0 = 5 \times 10^7 \) cm, has been adopted in all models reported in this work.

The effect of the duration of the perturbation was checked by means of a test case (model B), identical to model A but with a perturbation lasting for 10 s. As shown in Table 1, the characteristic timescales for model B, such as the time required for the first instabilities to show up, \( T_{\text{KH}} \), or the time needed by the convective front to hit the outer boundary, \( t_Y \), become shorter. The role played by a temperature perturbation can be understood in terms of the energy injected into the envelope: the longer the duration of the perturbation, the larger the energy injected, and thus, the shorter the characteristic timescales of the TNR. This has little effect, however, on the overall metallicity enhancement in the envelope since a final CNO mass fraction of \( 0.212 \) was found in model B, whereas \( 0.224 \) resulted in model A.

Both models A and B assumed temperature perturbations of \( \Delta T \sim 5\% \) during the initial timestep \( (\sim 10^{-10} \text{ s}) \). To test the possible influence of the strength of the perturbation, a test case with \( \Delta T \sim 0.5\% \) (model C) has also been computed. As shown in Table 1 and Fig. 4, the time evolution of models A and C is very similar, and hence, similar final mean CNO mass fractions at the end of the simulations were found (with \( Z = 0.209 \) in model C).

The effect of the location of the perturbation along the vertical axis has also been studied: whereas model A assumed a temperature perturbation of \( \sim 5\% \), applied at the innermost envelope shell \( (y_0 = 5.51 \times 10^8 \) cm), in model D, a similar perturbation was placed \( \sim 5 \) km above the core-envelope interface \( (y_0 = 5.515 \times 10^5 \) cm). Both models exhibit a very similar temporal evolution, with almost identical times for the appearance of the first instabilities and for the time required to reach the outer boundary. Similar envelope mean CNO mass fractions \( (0.224 \text{ and } 0.235, \text{ respectively}) \) were also found.

Finally, the influence of the size of the perturbation has also been analyzed. Whereas model D was evolved with an initial temperature perturbation of size \( \sim 5\% \), \( R_x = 1 \) km and \( R_y \sim 1 \) km, model E assumed \( R_x = 5 \) km and \( R_y = 5 \) km. As before, very similar characteristic timescales (see Table 1) and final mean CNO mass fractions \( (0.235 \text{ and } 0.209, \text{ respectively}) \) were found.

To summarize, the specific choice of the parameters that define the initial perturbation has a negligible effect on metallicity enhancement of the envelope.

## 5. Effect of the size of the computational domain

The choice of the computational domain represents a compromise between computational time requirements and numerical accuracy. Several considerations constrain its minimum size. On one hand, the merger of large convective eddies often found in 2-D simulations may be severely affected by the adoption of a small computational domain. On the other hand, nova outbursts eventually result in mass ejection. With an Eulerian code such as FLASH, it is not possible to track the material that flows off the grid, and hence, it is important to use domains that are as large as possible along the radial direction (while being sufficiently wide along the horizontal axis). Unfortunately, when the initial 1-D model is mapped into the 2-D grid, and relaxed to guarantee hydrostatic equilibrium, densities quickly underflow values for large heights (Zingale et al. 2002).

The specific size adopted for most of the models computed in this work, i.e. \( 800 \times 800 \) km, is a bit smaller than those used in Glasner et al. (1997) \( \sim 0.1 \pi \text{rad} \), in spherical-polar coordinates – and in Kercek et al. (1998) \( 1800 \times 1100 \) km, in a cartesian, plane-parallel geometry. In this section, we analyze possible dependences of the results on the adopted size of the computational domain. To this end, two additional simulations were performed. In the first one (model F), a wider computational domain has been adopted (i.e., \( 1600 \times 800 \) km). In the second (model G), aimed at testing the influence of the vertical (radial) length, a...
Table 1. Models computed.

<table>
<thead>
<tr>
<th>Model</th>
<th>$H$ (km)</th>
<th>$R_x R_y$ (km$^2$)</th>
<th>$\delta T$ (s)</th>
<th>$\Delta t$ (s)</th>
<th>Resolution $\delta x \times \delta y$ (km)</th>
<th>$t_{KH}$ (s)</th>
<th>$t_T$ (s)</th>
<th>$Z$</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>0</td>
<td>$1 \times 1$</td>
<td>5%</td>
<td>$10^{-10}$</td>
<td>$1.56 \times 1.56$</td>
<td>800 x 800</td>
<td>155</td>
<td>496</td>
</tr>
<tr>
<td>B</td>
<td>0</td>
<td>$1 \times 1$</td>
<td>5%</td>
<td>10</td>
<td>$1.56 \times 1.56$</td>
<td>800 x 800</td>
<td>28</td>
<td>347</td>
</tr>
<tr>
<td>C</td>
<td>0</td>
<td>$1 \times 1$</td>
<td>0.5%</td>
<td>$10^{-10}$</td>
<td>$1.56 \times 1.56$</td>
<td>800 x 800</td>
<td>155</td>
<td>493</td>
</tr>
<tr>
<td>D</td>
<td>5</td>
<td>$5 \times 5$</td>
<td>5%</td>
<td>$10^{-10}$</td>
<td>$1.56 \times 1.56$</td>
<td>800 x 800</td>
<td>154</td>
<td>496</td>
</tr>
<tr>
<td>E</td>
<td>0</td>
<td>$2 \times 1$</td>
<td>5%</td>
<td>$10^{-10}$</td>
<td>$1.56 \times 1.56$</td>
<td>1600 x 800</td>
<td>156</td>
<td>486</td>
</tr>
<tr>
<td>F</td>
<td>0</td>
<td>$1 \times 1.25$</td>
<td>5%</td>
<td>$10^{-10}$</td>
<td>$1.56 \times 1.56$</td>
<td>800 x 1000</td>
<td>156</td>
<td>526</td>
</tr>
<tr>
<td>G</td>
<td>0</td>
<td>$1 \times 1$</td>
<td>5%</td>
<td>$10^{-10}$</td>
<td>$0.39 \times 0.39$</td>
<td>800 x 800</td>
<td>162</td>
<td>584</td>
</tr>
<tr>
<td>H</td>
<td>0</td>
<td>$1 \times 1$</td>
<td>5%</td>
<td>$10^{-10}$</td>
<td>$0.39 \times 0.39$</td>
<td>800 x 800</td>
<td>268</td>
<td>893</td>
</tr>
</tbody>
</table>

Fig. 2. Mach number at two different moments of the simulation, $t = 230$ s (left panel) and 496 s (right panel), for model A.

Fig. 3. Left panel: propagation of the convective front as a function of time, for models A, H, and I. Right panel: temperature profile versus radius at two different times, $t = 0$ s (solid line; $T_{\text{base}} = 9.84 \times 10^7$ K) and $t = 496$ s (dashed line; $T_{\text{base}} = 1.64 \times 10^8$ K), for model A.

The specific length adopted along the vertical direction (see model G), while unimportant for the time of appearance of the instabilities (around 155 s after the start of the simulation, as in model A), affects the time required to reach the outer boundary, located 200 km above the value adopted for model A. Moreover, the larger extension of the computational domain along the radial (vertical) direction allows the convective eddies to pump additional metal-rich core material into the envelope compared with above a threshold value the course of the TNR is insensitive to the adopted width, in agreement with the sensitivity study performed by Glasner et al. (2007).
all the simulations reported previously in this paper. Indeed, the
mean, mass-averaged metallicity in model G achieves the largest
value of all the simulations reported, \( \sim 0.291 \). This result sug-
gests that the likely mean mass-averaged metallicity driven by
Kelvin-Helmholtz instabilities should be \( Z \approx 0.3 \). In summary,
we conclude that the size of the computational domain, above a
certain threshold value, has little influence on the physical quan-
tities that are more directly related with the mixing process at the
core-envelope interface.

6. Effect of the grid resolution

All simulations discussed so far (e.g., models A to G) were per-
formed with a resolution of \( 1.56 \times 1.56 \) km, a value similar to
the minimum resolutions adopted in Glasner et al. (2007) which
is roughly \( \sim 1.4 \times 1.4 \) km, and in Kercek et al. (1998), \( 1 \times 2 \) km.
To quantitatively assess the possible effect of the resolution, two
additional test cases were computed with exactly the same in-
put parameters as in model A but with two different resolutions:
\( 1 \times 1 \) km (model H) and \( 0.39 \times 0.39 \) km (model I)\(^3\).

As shown in Table 1, the increase in resolution produces a
delay in the time required for the first instabilities to develop,
\( t_{KH} \). This seems to be a numerical artifact. In models with a
coarser resolution, the larger size of the blocks artificially gen-
erates a larger numerical diffusion compared to models with
a finer resolution (a similar resolution dependence is clearly seen
as well in the Kercek et al. 1998, simulations). Actually, the
difference of ratios in the initial build up times (i.e., (model I-
model A)/(model H-model A)) scales approximately as the zone
size dimensions to the power of two. This is a purely numeri-
cal perturbation that forces the development of instabilities. To
test this hypothesis, we computed an additional test case (not
included in Table 1), identical to model A but without any ini-
tial perturbation. The onset of the instabilities in such an ex-
traordinarily low numerical diffusion regime is substantially delayed.
The simulations reported by Glasner et al. (1997) also show
the early appearance of instabilities in a model with substi-
tional numerical noise: \( \text{within a very short time (about 10 s), the} \)

\(^3\) For comparison, whereas a maximum number of 5300 blocks are ad-
ministered in model A, the number of blocks increases up to 83,000 in
model I. The total CPU time spent in both simulations, using 256 pro-
cessors of the MareNostrum supercomputer, has been 3 and 110 kh,
respectively.

numerical noise (round-off) seeds an intense convective flow in
the envelope without any artificial perturbations.

A similar behavior is also found for the time required for the
convective front to reach the outer boundary, \( t_{f} \), and for the his-
tory of the nuclear energy generation rate (Fig. 4). As expected,
filamentary structures and convective cells are better resolved in
the finer resolution model I, compared to those computed with
somewhat coarser grids (models A and H; see Fig. 5). These mi-
nor differences do not, however, show significant variations in
the final, mean CNO abundances achieved in the envelope: while
\( Z \sim 0.224 \) in model A, models H and I yield 0.201 and 0.205, by
mass, respectively. Similar agreement is found in the peak tem-
peratures achieved and in the overall nuclear energy generation
rates (Fig. 4).

Thus, the adopted resolution has not a critical effect for the
mixing models presented in this work. The variation in the final
mean CNO abundance in the envelope, under the range of reso-
lutions adopted, is only about 12\% (when comparing results for
models A, H, and I), a quite reasonable value.

7. Discussion and conclusions

In this paper we have reported results for a series of nine 2-D
numerical simulations that test the influence of the initial pertur-
bation (duration, strength, location, and size), the resolution of
the grid, and the size of the computational domain on the results.
We have shown that mixing at the core-envelope interface pro-
ceds almost independently of the specific choice of such initial
parameters, above threshold values.

The study confirms that the metallicity enhancement inferred
from observations of the ejecta of classical novae can be ex-
plained by Kelvin-Helmholtz instabilities, powered by an ef-
fective mesoscopic shearing resulting from the initial buoyancy.
Fresh core material is efficiently transported from the outermost
layers of the white dwarf core and mixed with the approximately
solar composition material of the accreted envelope. As soon as
\(^{12}\)C and \(^{16}\)O are dredged up, convection sets in and small con-
vective cells appear, accompanied by an increased nuclear en-
ergy generation rate. The size of these convective cells increases
in time. Eventually, these cells merge into large convective ed-
dies with a size comparable to the envelope height. The range
of mean mass-averaged envelope metallicities obtained in our
simulations at the time when the convective front hits the outer

\[ \text{Fi}g. \; \text{4.} \; \text{Left panel: time evolution of the nuclear energy generation rate (in erg s}^{-1} \text{) for the 9 models computed in this work. Right panel: final CNO mass fraction versus radius.} \]
boundary, $0.21-0.29$, matches the values obtained for classical novae hosting CO white dwarfs.

It is, however, worth noting that the convective pattern is actually produced by the adopted geometry (e.g., 2-D), forcing the fluid motion to behave very differently than 3-D convection (Shore 2007; Meakin & Arnett 2007). Nevertheless, the levels of metallicity enhancement found in our 2-D simulations will likely remain unaffected by the limitations imposed by the 2-D geometry (Arnett, private communication). Fully 3-D simulations aimed at testing this hypothesis are currently underway.

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Fig. 5. Snapshots of the $^1$H (upper panels) and $^{12}$C (middle panels) mass fractions at $t \sim 395$ s (model A; left panels), and 688 s (model I; right panels). Lower panels: the number of blocks administered, at this stage, is 3184 for model A, and 43 800 for model I. In both simulations, FLASH divides each block in 8 cells. Structures such as vortexes are better resolved in the finer resolution model I.
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