

THE IMPRINT OF NOVA NUCLEOSYNTHESIS IN PRESOLAR GRAINS

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ABSTRACT

Infrared and ultraviolet observations of nova light curves have confirmed grain formation in their expanding shells that are ejected into the interstellar medium by a thermonuclear runaway. In this paper we present isotopic ratios of intermediate-mass elements up to silicon for the ejecta of CO and ONe novae, based on 20 hydrodynamic models of nova explosions. These theoretical estimates will help to properly identify nova grains in primitive meteorites. In addition, equilibrium condensation calculations are used to predict the types of grains that can be expected in the nova ejecta, providing some hints on the puzzling formation of C-rich dust in $O > C$ environments. These results show that SiC grains can condense in ONe novae, in concert with an inferred (ONe) nova origin for several presolar SiC grains.

Subject headings: dust, extinction — novae, cataclysmic variables —
nuclear reactions, nucleosynthesis, abundances

Online material: color figures

1. INTRODUCTION

Classical novae are powered by thermonuclear runaways (TNRs) that occur on the white dwarf (WD) component of close binary systems (see Starrfield 1989; Kovetz & Prialnik 1997; José & Hernanz 1998, hereafter JH98; Starrfield et al. 1998 and references therein). During such violent stellar events, whose energy release is only exceeded by gamma-ray bursts and supernova explosions, about 10^{-4} to $10^{-5} M_{\odot}$ are ejected into the interstellar medium. Because of the high peak temperatures attained during the outburst, $T_{\text{peak}} \sim (2-3) \times 10^8$ K, the ejecta are enriched in nuclear-processed material relative to the solar abundances, containing significant amounts of ^{13}C , ^{15}N , and ^{17}O and traces of other isotopes, such as ^7Li , ^{20}Ne , ^{26}Al , or ^{28}Si (depending on the nova type, CO or ONe, the mass of the underlying WD, and other properties). Indeed, theoretical models of the explosion reveal an isotopic pattern that does not correspond to equilibrium CNO burning (Starrfield et al. 1972).

In order to constrain the models, several studies have focused on a direct comparison of the atomic abundances, inferred from observations of the ejecta, with theoretical nucleosynthetic predictions (JH98; Starrfield et al. 1998). Despite problems

associated with the modeling of the explosion (Starrfield 2002), such as the unknown mechanism responsible for the mixing of the accreted envelope and the outermost shells of the underlying WD, or the difficulty of ejecting as much material as inferred from observations (see also Shore 2002), there is good agreement between theory and observations with regard to nucleosynthesis. This agreement includes atomic abundance determinations (H, He, C, N, O, Ne, Na, Mg, Si, etc.) and a plausible endpoint for nova nucleosynthesis (around Ca). For some well-observed novae, such as PW Vul 1984 or V1688 Cyg 1978, the agreement between observations and theoretical predictions (see Table 5 in JH98 for details) is quite amazing. The reader is referred to Gehrz et al. (1998) for an extended list of abundance determinations in nova ejecta.

Moreover, since the nucleosynthesis path is very sensitive to details of the explosion (i.e., chemical composition, extent of convective mixing, thermal history of the envelope, etc.), the agreement between the inferred abundances and the theoretical yields not only validates the thermonuclear runaway model but also imposes limits on the nature of the mechanism responsible for the mixing. For instance, if one assumes that the mixing settles very late in the course of the explosion,

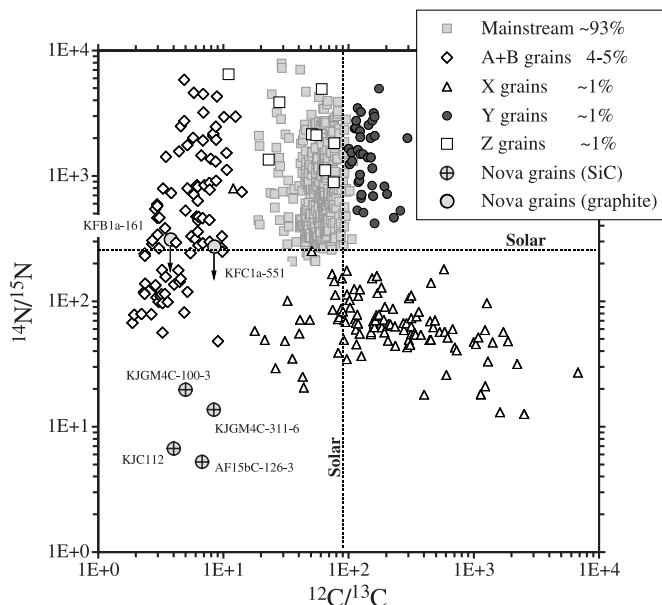


FIG. 1.—Carbon and nitrogen isotopic ratios of nova candidate grains compared with those of SiC grains of different populations (see legend for details). Silicon carbide grains have been classified into several populations based on their C, N, and Si isotopic ratios. Error bars are smaller than the symbols. [See the electronic edition of the *Journal* for a color version of this figure.]

pile-up of larger amounts of matter in the envelope would be favored since the injection of significant amounts of ^{12}C , which triggers the onset of the TNR through proton capture reactions, would be delayed. Hence, the explosion would take place in a somewhat more massive envelope, characterized by a higher ignition density (and pressure), giving rise to a more violent outburst with T_{peak} exceeding in some cases 4×10^8 K (S. Starrfield 2000, unpublished; J. José & M. Hernanz 2001, unpublished). Therefore, one would expect a significant enrichment in heavier species, beyond calcium, in the ejecta accompanying such violent outbursts. However, such an abundance pattern has never been seen in nature.

Nevertheless, a direct comparison with the elemental abundance pattern inferred from observations relies only on atomic abundances and does not pose very strict limits on nova models. In contrast, a much more precise set of constraints could be obtained if information on specific isotopic abundances were available. One good example is silicon, with three stable isotopes (i.e., ^{28}Si , ^{29}Si , and ^{30}Si) in the region of interest for nova nucleosynthesis: whereas ^{28}Si is strongly connected to the nature of the WD core (either a CO or an ONe WD¹), ^{29}Si and ^{30}Si are good indicators of the peak temperatures achieved in the explosion and of the dominant nuclear paths followed in the course of the TNR, which have a clear imprint on the overall composition of the ejecta.

Such detailed information can be (partially) obtained through the laboratory analysis of presolar grains, which yields isotopic abundance ratios. Presolar grains, found in primitive meteorites, are characterized by huge isotopic anomalies that can only be explained in terms of nucleosynthetic processes

¹ The initial mass of the progenitor star determines the number of evolutionary stages that it will undergo. Hence, stars within $2.3 \leq (M/M_{\odot}) \leq 8$ evolve through hydrogen and helium burning, leaving a CO-rich WD remnant. Stars in the mass interval $8 \leq (M/M_{\odot}) \leq 10-12$ additionally undergo carbon burning, leaving an ONe-rich remnant instead.

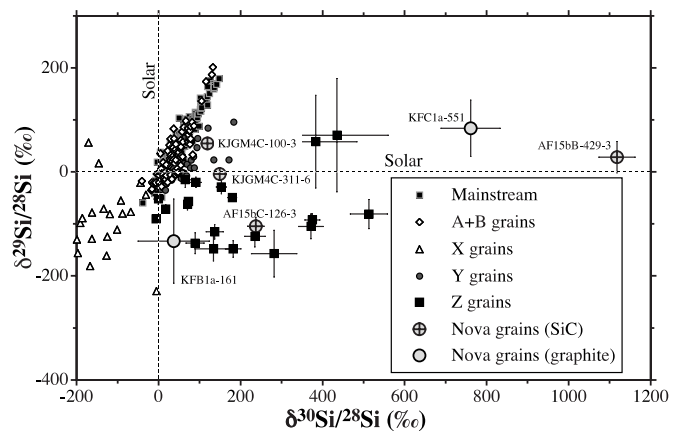


FIG. 2.—Silicon isotopic ratios of five nova candidate grains and other SiC grains. Ratios are expressed as delta values, deviations from the solar Si isotopic ratios in permil (see § 2.4 for definition). [See the electronic edition of the *Journal* for a color version of this figure.]

that took place in their stellar sources. In fact, detailed studies of these grains have opened up a new and promising field of astronomy (see Zinner 1998). So far, silicon carbide (SiC), graphite (C), diamond (C), silicon nitride (Si_3N_4), and oxides (such as corundum and spinel) have been identified as presolar grains. Ion microprobe analyses of single presolar grains have revealed a variety of isotopic signatures that allow the identification of parent stellar sources, such as asymptotic giant branch (AGB) stars and supernovae (Zinner 1998). Up to now, SiC grains have been most extensively studied and can be classified into different populations on the basis of their C, N, and Si isotopic ratios (see Figs. 1 and 2).

Infrared (Evans 1990; Gehrz et al. 1998; Gehrz 1999) and ultraviolet observations (Shore et al. 1994) of the evolution of nova light curves suggest that novae form grains in their expanding ejected shells. Both nova types, CO and ONe, behave in a similar way in the infrared immediately after the explosion, but as the envelope expands and becomes optically thin, differences in their infrared emission appear: whereas in a CO nova this phase is typically followed by dust formation, accompanied by a decline in visual brightness and a simultaneous rise in infrared emission (see Rawlings & Evans 2002; Gehrz 1999, 2002), ONe novae (erupting on more massive WDs than CO novae) are not such prolific dust producers. The reason for this is that the latter have lower-mass, high-velocity ejecta, where the typical local densities may be too low to allow the condensation of appreciable amounts of dust. Observations of the condensation of dust containing different species, such as silicates, SiC, carbon, and hydrocarbons, have been reported for a number of novae (Gehrz et al. 1998). The presence of SiC (or C-rich) dust in nova ejecta is established from spectroscopic measurements (see Table 1 in Starrfield et al. 1998 and Table 2 in Gehrz et al. 1998). It is generally believed that $\text{C} > \text{O}$ is needed for the formation of SiC and/or graphite grains. If oxygen is more abundant than carbon, essentially all C is locked up in the very stable CO molecule, and the excess O leads to formation of oxides and silicates as condensates. On the other hand, if carbon is more abundant than oxygen, essentially all O is tied up in CO and the excess C can form reduced condensates such as SiC or graphite. Since theoretical models of nova outbursts yield, on average, $\text{O} > \text{C}$, one would expect only oxidized condensates using the carbon and oxygen abundances as a sole criterion. However,

TABLE 1
PRESOLAR GRAINS WITH AN INFERRED NOVA ORIGIN

Grain	Composition	$^{12}\text{C}/^{13}\text{C}$	$^{14}\text{N}/^{15}\text{N}$	$\delta(^{29}\text{Si}/^{28}\text{Si})$	$\delta(^{30}\text{Si}/^{28}\text{Si})$	$^{26}\text{Al}/^{27}\text{Al}$	$^{20}\text{Ne}/^{22}\text{Ne}$
AF15bB-429-3 (A01)	SiC	9.4 ± 0.2	...	28 ± 30	1118 ± 44
AF15bC-126-3 (A01)	SiC	6.8 ± 0.2	5.22 ± 0.11	-105 ± 17	237 ± 20
KJGM4C-100-3 (A01)	SiC	5.1 ± 0.1	19.7 ± 0.3	55 ± 5	119 ± 6	0.0114	...
KJGM4C-311-6 (A01)	SiC	8.4 ± 0.1	13.7 ± 0.1	-4 ± 5	149 ± 6	>0.08	...
KJC112 (H95)	SiC	4.0 ± 0.2	6.7 ± 0.3
KFC1a-551 (A01)	C	8.5 ± 0.1	273 ± 8	84 ± 54	761 ± 72
KFB1a-161 (N04)	C	3.8 ± 0.1	312 ± 43	-133 ± 81	37 ± 87	...	<0.01
Solar	89	272	14
Nova models	0.3–1.8	0.3–1400	–900 to 10	–1000 to 47000	0.01–0.6	0.1–250

NOTES.—The solar N ratio in the table is that of the air. Grains AF. . . are from the Acfer 094 meteorite, whereas grains KJ. . . and KF. . . are from the Murchison meteorite. Delta values measure deviations from the solar Si isotopic ratios in permil (see § 2.4 for definition). Errors are 1σ .

this is at odds with the observation of C-rich dust detected around some novae (Gehrz et al. 1993; Starrfield et al. 1997; Gehrz 1999).

While previously the identification of presolar nova grains from meteorites relied only on low $^{20}\text{Ne}/^{22}\text{Ne}$ ratios (with ^{22}Ne being attributed to ^{22}Na decay; Amari et al. 1995; Nichols et al. 2004, hereafter N04), recently five SiC and two graphite grains that exhibit other isotopic signatures characteristic of nova nucleosynthesis have been identified (for details see Hoppe et al. 1995, hereafter H95; Amari et al. 2001, hereafter A01; Amari 2002). This discovery provides a very valuable set of constraints for nova nucleosynthesis. Table 1 summarizes the mineralogy and isotopic composition of these grains, reported in A01 and Amari (2002). The SiC grains have very low $^{12}\text{C}/^{13}\text{C}$ and $^{14}\text{N}/^{15}\text{N}$ ratios, while the graphite grains have low $^{12}\text{C}/^{13}\text{C}$ but normal $^{14}\text{N}/^{15}\text{N}$ ratios. However, the original $^{14}\text{N}/^{15}\text{N}$ ratios of these two graphite grains could have been much lower because there is evidence that indigenous N in presolar graphites has been isotopically equilibrated with terrestrial nitrogen. For example, most presolar graphite grains show a huge range in C isotopic ratios but essentially normal (terrestrial) N isotopic composition (H95). Recent isotopic imaging of C and O inside of slices of graphite spherules showed gradients from highly anomalous ratios in the center to more normal ratios close to the surface, also indicating isotopic equilibration (Stadermann et al. 2004). The $^{26}\text{Al}/^{27}\text{Al}$ ratios have been determined only for two SiC grains (KJGM4C-100-3 and KJGM4C-311-6) and are very high ($>10^{-2}$; see Fig. 3). We note that the $^{20}\text{Ne}/^{22}\text{Ne}$ ratio is only available for the graphite grain KFB1a-161 (<0.01 ; $^{22}\text{Na}/\text{C} = 9 \times 10^{-6}$; see N04), being considerably lower than the ratios predicted by nova models (see § 2). Usually, neon is incorporated in grains via implantation, since noble gases do not condense as stable compounds into grains (Amari 2002). However, the low $^{20}\text{Ne}/^{22}\text{Ne}$ ratio measured in this grain suggests that Ne has not been implanted in the ejecta, but ^{22}Ne most likely originated from in situ decay of ^{22}Na (with a mean lifetime $\tau = 3.75$ yr).

Silicon isotopic ratios of the five SiC grains are characterized by ^{30}Si excesses and close to or slightly lower than solar $^{29}\text{Si}/^{28}\text{Si}$ ratios. Whereas CO nova models (Kovetz & Prialnik 1997; Starrfield et al. 1997; JH98; Hernanz et al. 1999, hereafter H99; and unpublished data) predict close to solar $^{30}\text{Si}/^{28}\text{Si}$ and close to or lower than solar $^{29}\text{Si}/^{28}\text{Si}$, huge enrichments of ^{30}Si and close to or lower than solar $^{29}\text{Si}/^{28}\text{Si}$ ratios are obtained for ONe novae (JH98; H99; José et al. 1999, 2001, hereafter JCH99, JCH01; Starrfield et al. 1998). We have also included unpublished data on Si isotopic ratios

for grain KFB1a-161, in both Table 1 and Figure 2. Unfortunately, trace element concentrations in KFB1a grains are low and hence measurements are characterized by large errors.

The isotopic signatures of these grains qualitatively agree with current predictions from hydrodynamic models of nova outbursts. In fact, a comparison between grain data and nova models suggests that these grains formed in ONe novae with a WD mass of at least $1.25 M_{\odot}$ (A01). However, two main problems, related with the likely nova paternity of these grains, remain to be solved: (1) the challenging connection with ONe novae, which, as stated before, are not as prolific dust producers as CO novae; and (2) in order to quantitatively match the grain data, one has to assume a mixing process between material newly synthesized in the nova outburst and more than 10 times as much unprocessed, isotopically close to solar, material before grain formation.

In this paper we provide theoretical predictions for the expected isotopic composition of the nova ejecta and explore which type of condensates may form. The structure of the paper is as follows: In § 2 we summarize the main nucleosynthesis results from one-dimensional, hydrodynamic computations of nova outbursts. We report the theoretically expected isotopic ratios in the ejected nova shells, which

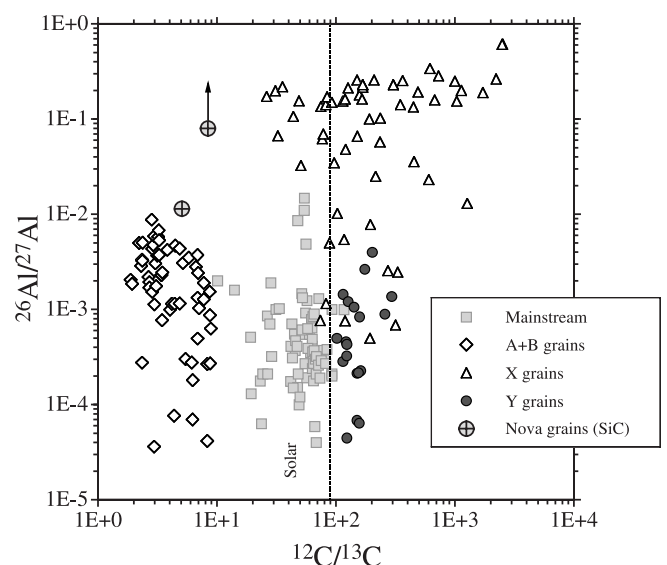


FIG. 3.—Same as Fig. 1, but for aluminum vs. carbon isotopic ratios. [See the electronic edition of the Journal for a color version of this figure.]

should also be representative of the isotopic composition of grains condensed in the ejecta. In § 3 we describe the results of chemical equilibrium condensation calculations for different types of nova ejecta. In § 4 we explore the isotopic patterns of individual ejected shells and compare them with those resulting from mean mass-averaged envelopes. Furthermore, we analyze the effect of relevant nuclear physics uncertainties on the results presented here and compare them with results obtained by other groups. A summary of the main conclusions of this paper is given in § 5.

2. THEORETICAL ISOTOPIC RATIOS IN NOVA EJECTA: MEAN MASS-AVERAGED VALUES

We have adopted two different approaches in our search for trends in the isotopic composition that may characterize the nova ejecta: (1) an analysis based on mean mass-averaged isotopic ratios for a number of species, resulting from hydrodynamic calculations of classical nova outbursts; and (2) a detailed analysis of the chemical abundance gradients found when individual ejected shells are taken into account. It is important to stress that mean mass-averaged ratios provide a global view of the nucleosynthetic imprints of the explosion. The envelopes ejected in our numerical models of the nova outburst consist of a large number of shells of different masses (which decrease outward from the envelope's base). The innermost shells are probably the most relevant ones, the reason being twofold: first, these are the shells that undergo the largest changes in chemical composition through nuclear processing (i.e., the shells that will exhibit the strongest imprints of a nova outburst), and second, because of their larger masses, material from these shells has a larger probability of condensing and forming dust and grains. Both aspects are partially taken into account by the mass-averaging process² that assigns different weights to individual shells. In contrast, a quantitative analysis based on individual shells, although in principle more detailed, can be potentially misleading: it may generate a biased view of the nucleosynthetic history since, a priori, all possible ratios found throughout the envelope (see Tables 2 and 3 and variation bars in Figs. 4–9) seem, at first glance, equally likely. It is therefore important to point out that the largest deviations from the mean are often obtained in individual shells located near the surface, in low-mass shells with a lower probability of forming grains and with isotopic features that reflect to a much lower extent the imprint of a nova outburst. In fact, the differences found in those surface layers are connected with details of the retreat of convection from the surface rather than caused by nuclear processes. It is therefore our aim first to focus this analysis on mean mass-averaged ratios and then to address the question of how robust our conclusions are when an analysis based on individual shells is performed (§ 4.1).

A full list of mean mass-averaged values, together with maximum and minimum isotopic ratios throughout the ejecta, for a sample of 20 hydrodynamic nova models is given in Tables 2 and 3. Details of the numerical code, developed to follow the course of nova outbursts, from the onset of accretion to the expansion and ejection stages, have been discussed by JH98 and are summarized in the accompanying Appendix, together with a brief description of the initial isotopic ratios adopted in the models presented here.

2.1. Nitrogen and Carbon Isotopic Ratios

The final $^{14}\text{N}/^{15}\text{N}$ ratios found in the ejected shells of nova outbursts show a wide range of variation (see Fig. 4). Explosions involving ONe WDs yield low ratios, ranging from ~ 0.3 to 4 (solar ratio = 272). In contrast, CO nova models are characterized by higher ratios, typically between ~ 3 and 100 (see Fig. 4), but as high as ~ 1400 for the extreme $0.6 M_{\odot}$ CO case. As we stress throughout this section, the nuclear activity in this low-mass CO model is so tiny that the final isotopic ratios are, for many species, close to the initial ratios of the envelope at the onset of the TNR. These differences in the final N ratios between CO and ONe models reflect differences in the main nuclear paths followed in the course of the explosions.

The synthesis of ^{15}N depends critically on the amount of ^{14}N available (both the initial one, present in the accreted material, and the amount synthesized through the CNO cycle, starting from ^{12}C). Since both CO and ONe models begin with the same initial ^{14}N (see Table 4), differences in the ejecta reflect different thermal histories during the explosion (and in particular, differences in T_{peak}): the higher peak temperatures achieved in ONe models favor proton capture reactions on ^{14}N , leading to $^{14}\text{N}(p, \gamma)^{15}\text{O}(\beta^+)^{15}\text{N}$, and are thereby responsible for the higher ^{15}N content in the ejecta. This explains also why the $^{14}\text{N}/^{15}\text{N}$ ratio decreases as the WD mass increases, for both CO and ONe models, a direct consequence of the higher temperatures achieved for more massive WDs. In summary, the $^{14}\text{N}/^{15}\text{N}$ ratio provides a means for distinguishing between CO and ONe novae: large ratios, of the order of 100–1000, are only achieved in explosions involving low-mass CO novae, according to the models discussed.

In contrast to the N isotopic ratios, both CO and ONe models yield very low $^{12}\text{C}/^{13}\text{C}$ ratios (see also Fig. 4), in the range ~ 0.3 –2 (solar ratio = 89). The dramatic reduction in the final $^{12}\text{C}/^{13}\text{C}$ ratios as compared to the initial ones is due to the very efficient synthesis of ^{13}C through $^{12}\text{C}(p, \gamma)^{13}\text{N}(\beta^+)^{13}\text{C}$, which, in turn, decreases the final amount of ^{12}C .

The effect of the WD mass on the $^{12}\text{C}/^{13}\text{C}$ ratios follows also a certain pattern, but unlike the case of N isotopic ratios, it depends on the nova type: for low-mass CO WDs, the amount of ^{13}C synthesized from proton capture reactions on ^{12}C is strongly limited by the moderate range of temperatures achieved in the explosion. However, as the mass of the WD (and hence the temperature at the envelope's base) increases, more ^{13}C is produced, leading to lower $^{12}\text{C}/^{13}\text{C}$ ratios, up to a point where the temperatures achieved in the envelope are high enough to enable significant proton captures on ^{13}C , increasing again the $^{12}\text{C}/^{13}\text{C}$ ratio. In ONe novae, the temperatures achieved during the explosions are always high enough for significant proton capture reactions to proceed on ^{13}C (which, in turn, increase the final amount of ^{14}N), leading to $^{12}\text{C}/^{13}\text{C}$ ratios that monotonically increase with the WD mass.

We stress that C ratios are highly diagnostic for identifying potential nova grain candidates since, as can be seen in Fig. 4, independently of the nova type and of the adopted WD mass, all models are characterized by an extremely narrow range of low $^{12}\text{C}/^{13}\text{C}$ values (in contrast with the wide dispersion obtained for the N ratios), definitely a characteristic signature of a classical nova explosion.

2.2. Oxygen and Neon Isotopic Ratios

Oxygen isotopic ratios depend on the nova type (i.e., CO or ONe) and the WD mass. As can be seen in Figure 5, CO models are in general characterized by moderate to large

² In fact, the analysis available in Starrfield et al. (1997), the only work that addresses a similar search of nova nucleosynthesis trends and its connection with meteorites, is exclusively based on mean mass-averaged isotopic ratios.

TABLE 2
ISOTOPIC RATIOS FOR C, N, O, AND Ne, OBTAINED FROM DIFFERENT MODELS, AS DISPLAYED IN FIGURES 4–6

Model ^a	Mass (M_{\odot})	Initial Composition	$^{12}\text{C}/^{13}\text{C}$	$^{14}\text{N}/^{15}\text{N}$	$^{16}\text{O}/^{17}\text{O}$	$^{16}\text{O}/^{18}\text{O}$	$^{20}\text{Ne}/^{21}\text{Ne}$	$^{20}\text{Ne}/^{22}\text{Ne}$
ONe1.....	1.00 (JH98)	50% ONe	0.83 [0.59–1.4]	3.6 [0.59–27.8]	10 [7.1–17.7]	23.3 [15.3–73.6]	9950 [4320–27400]	97 [89.9–103]
ONe2.....	1.15 (JH98)	25% ONe	0.85 [0.67–1.9]	1.9 [0.55–3.6]	2.3 [1.5–4.2]	11.3 [8.9–23.6]	7270 [3990–9170]	159 [121–169]
ONe3.....	1.15 (JH98)	50% ONe	0.89 [0.23–1.6]	1 [0.39–2.4]	4.6 [3.3–18.5]	22.5 [10.5–63.8]	6300 [3920–8290]	113 [100–120]
ONe4.....	1.15 (H99)	50% ONe	0.76 [0.63–2.0]	1.2 [0.36–1.9]	3.8 [2.5–5.4]	388 [309–663]	2200 [966–3540]	109 [72.9–119]
ONe5.....	1.15 (JH98)	75% ONe	0.88 [0.65–1.8]	1.2 [0.35–3.5]	6 [4.3–10.6]	35.5 [22.2–183]	6830 [4600–11900]	108 [78.1–118]
ONe6.....	1.25 (JH98)	50% ONe	0.95 [0.73–2.2]	0.82 [0.30–1.5]	1.9 [1.5–3.3]	19 [13.9–49.3]	5400 [3510–7080]	181 [113–190]
ONe7.....	1.25 (H99)	50% ONe	0.79 [0.60–2.4]	0.97 [0.28–1.6]	1.9 [1.4–2.9]	220 [188–394]	1620 [804–2770]	153 [77.9–185]
ONe8.....	1.35 (JH98)	50% ONe	1.5 [0.50–3.2]	0.41 [0.14–0.61]	1.4 [0.63–2.6]	25.3 [22.7–76.2]	3090 [2460–3340]	220 [56.9–459]
ONe9.....	1.35 (JCH01)	50% ONe	1.5 [0.54–3.1]	0.40 [0.14–0.58]	1.4 [0.68–2.9]	283 [176–529]	557 [426–641]	125 [37.2–352]
ONe10.....	1.35 (JH98)	75% ONe	1.1 [0.28–2.7]	0.25 [0.12–0.52]	1.9 [1.7–3.7]	56.9 [46.6–265]	3000 [2480–4210]	247 [79.9–1430]
ONeMg1.....	1.25	50% ONeMg	0.73 [0.52–2.4]	1.0 [0.34–1.6]	2.6 [2–8.9]	22 [11.2–211]	5060 [4190–5740]	1550 [818–2890]
CO1.....	0.6	50% CO	1.8 [1.7–1.8]	1370 [742–1960]	232 [230–236]	38,800 [33,800–43,000]	26,750 [20,300–33,400]	0.18 [0.18–0.18]
CO2.....	0.8 (H99)	50% CO	0.51 [0.42–1.8]	60 [21.4–177]	63 [56.1–178]	3960 [3400–20,600]	17,670 [3120–46,200]	0.18 [0.18–0.18]
CO3.....	0.8 (JH98)	25% CO	0.45 [0.39–0.63]	103 [21.3–245]	41.9 [38.8–66]	174 [154–436]	43,500 [12,000–89,700]	0.51 [0.51–0.53]
CO4.....	0.8 (JH98)	50% CO	0.52 [0.49–1.8]	127 [20.8–142]	60.4 [57.8–178]	502 [474–3790]	21,500 [3080–26,800]	0.18 [0.18–0.18]
CO5.....	1.0 (JH98)	50% CO	0.30 [0.29–0.70]	19 [6–267]	31.9 [30.9–46.8]	123 [95.2–423]	15,900 [5480–147000]	0.19 [0.18–0.19]
CO6.....	1.15 (JH98)	25% CO	0.71 [0.50–0.78]	3.3 [1.6–8.5]	7.6 [5.8–9.7]	22 [16.1–120]	7740 [3640–14200]	0.70 [0.61–0.72]
CO7.....	1.15 (JH98)	50% CO	0.54 [0.50–1]	3.0 [2.4–5.3]	10.6 [10.4–16.7]	62.2 [35.7–443]	5990 [2740–15500]	0.22 [0.2–0.23]
CO8.....	1.15 (H99)	50% CO	0.54 [0.50–0.93]	2.8 [2.4–13.5]	10.9 [10.3–15.7]	854 [750–3730]	2640 [1240–10800]	0.22 [0.2–0.23]
CO9.....	1.15 (JH98)	75% CO	0.39 [0.32–0.88]	5.2 [3.2–12]	19 [18.6–26.5]	103 [51.3–548]	7580 [2690–22700]	0.078 [0.07–0.082]
Solar.....	89	272	2622	498	412	14

^a All models have been computed with the nuclear reaction network described in JH98, except models ONe4, ONe7, CO2, and CO8 (H99) and models CO1 and ONe9 (JCH01). See Appendix for details.

TABLE 3
 SAME AS TABLE 2, FOR Mg, Al, AND Si ISOTOPES, AS DISPLAYED IN FIGURES 7–9

Model	Mass (M_{\odot})	Initial Composition	$^{24}\text{Mg}/^{25}\text{Mg}$	$^{26}\text{Mg}/^{25}\text{Mg}$	$^{26}\text{Al}/^{27}\text{Al}$	$\delta(^{29}\text{Si}/^{28}\text{Si})$	$\delta(^{30}\text{Si}/^{28}\text{Si})$
ONe1	1.00 (JH98)	50% ONe	0.026 [0.005–0.09]	0.090 [0.082–0.11]	0.23 [0.19–0.32]	–951 [–951 to –949]	–965 [–965 to –964]
ONe2	1.15 (JH98)	25% ONe	0.021 [0.013–0.037]	0.096 [0.076–0.13]	0.25 [0.23–0.3]	–803 [–808 to –784]	296 [–128 to 320]
ONe3	1.15 (JH98)	50% ONe	0.036 [0.004–0.076]	0.11 [0.074–0.14]	0.22 [0.16–0.27]	–852 [–853 to –848]	–645 [–744 to –624]
ONe4	1.15 (H99)	50% ONe	0.10 [0.04–0.29]	0.16 [0.14–0.18]	0.24 [0.22–0.27]	–788 [–795 to –756]	–105 [–407 to –73]
ONe5	1.15 (JH98)	75% ONe	0.21 [0.043–0.65]	0.13 [0.082–0.21]	0.22 [0.21–0.24]	–796 [–808 to –756]	–354 [–482 to –325]
ONe6	1.25 (JH98)	50% ONe	0.087 [0.032–0.19]	0.11 [0.083–0.15]	0.28 [0.25–0.30]	–701 [–734 to –585]	1380 [722–1470]
ONe7	1.25 (H99)	50% ONe	0.13 [0.05–0.33]	0.15 [0.13–0.17]	0.28 [0.26–0.32]	–655 [–698 to –447]	2020 [1230–2110]
ONe8	1.35 (JH98)	50% ONe	0.089 [0.023–0.19]	0.14 [0.12–0.16]	0.42 [0.26–0.46]	–74 [–677 to 1760]	7730 [7260–9700]
ONe9	1.35 (JCH01)	50% ONe	0.12 [0.026–0.27]	0.15 [0.14–0.17]	0.42 [0.27–0.46]	–54 [–666 to 1830]	8150 [7760–9580]
ONe10	1.35 (JH98)	75% ONe	0.19 [0.034–0.38]	0.15 [0.056–0.19]	0.35 [0.19–0.39]	–113 [–742 to 1210]	7140 [7000–7770]
ONeMg1	1.25	50% ONeMg	0.47 [0.14–1.8]	0.043 [0.018–0.053]	0.44 [0.34–0.88]	–546 [–652 to 87.8]	3780 [3500–3830]
CO1	0.6	50% CO	4.3 [4.3–4.3]	0.65 [0.65–0.65]	0.006 [0.006–0.006]	–3.3 [–3.3 to –3.3]	1.9 [1.9–1.9]
CO2	0.8 (H99)	50% CO	0.31 [0.22–1.8]	0.16 [0.15–0.35]	0.095 [0.021–0.11]	–6.3 [–6.3 to –3.3]	–1.2 [–1.20 to 1.9]
CO3	0.8 (JH98)	25% CO	0.16 [0.12–0.7]	0.14 [0.13–0.21]	0.18 [0.074–0.2]	–6.4 [–6.4 to –2.4]	–3.2 [–3.2 to 0.85]
CO4	0.8 (JH98)	50% CO	0.27 [0.25–1.8]	0.15 [0.15–0.34]	0.14 [0.023–0.14]	–6.3 [–6.3 to –3.3]	–1.2 [–1.2 to 1.9]
CO5	1.0 (JH98)	50% CO	0.060 [0.025–0.31]	0.10 [0.099–0.14]	0.39 [0.23–0.42]	–33 [–35.6 to –18.1]	4.9 [–1.9 to 6.5]
CO6	1.15 (JH98)	25% CO	0.042 [0.017–0.15]	0.096 [0.08–0.12]	0.25 [0.23–0.34]	–677 [–697 to –497]	–47.1 [–53.6 to –21.2]
CO7	1.15 (JH98)	50% CO	0.12 [0.024–0.69]	0.094 [0.073–0.18]	0.38 [0.36–0.43]	–449 [–480 to –236]	29.6 [15.8–45.5]
CO8	1.15 (H99)	50% CO	0.11 [0.028–0.61]	0.14 [0.12–0.2]	0.39 [0.35–0.43]	–435 [–478 to –243]	32.6 [19.8–52.1]
CO9	1.15 (JH98)	75% CO	0.080 [0.01–0.46]	0.070 [0.06–0.12]	0.58 [0.53–0.59]	–267 [–296 to –159]	90.1 [71.3–97.4]
Solar	7.9	1.1	0	0	0

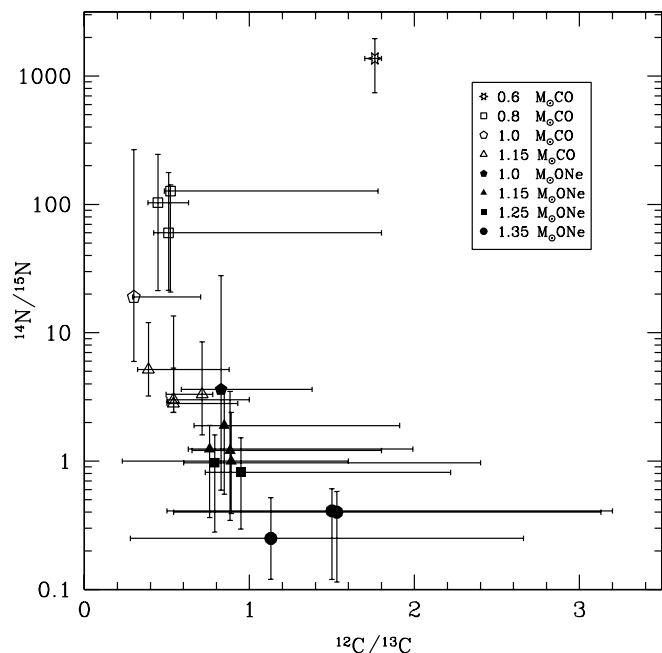


FIG. 4.—Nitrogen vs. carbon isotopic ratios, predicted by hydrodynamic models for both CO and ONe novae (see Tables 2 and 3 for details). Symbols represent mean mass-averaged ratios. Deviation bars, taking into account the gradient of composition in the ejected shells, are also shown for all models. See text for details. [See the electronic edition of the Journal for a color version of this figure.]

$^{16}\text{O}/^{18}\text{O}$ ratios, ranging from 20 to 39,000 (solar ratio = 498), and moderate $^{16}\text{O}/^{17}\text{O}$ ratios, from 8 to 230 (solar ratio = 2622). In contrast, lower ratios are, in general, found for ONe models: whereas $^{16}\text{O}/^{18}\text{O}$ ranges from 10 to 400, $^{16}\text{O}/^{17}\text{O}$ ranges from 1 to 10. It is important to stress that a recent revision of the $^{18}\text{F}(p, \alpha)$ reaction rate (see H99; Coc et al. 2000) yields, in general, higher $^{16}\text{O}/^{18}\text{O}$ ratios than those obtained with previous estimates (compare, for instance, the O ratios obtained in models ONe3 and ONe4). At the time the calculations presented in this paper were made, the $^{18}\text{F}(p, \alpha)$ rate was affected by a large uncertainty (a factor of ~ 300). Two recent experiments, at Oak Ridge and Louvain-la-Neuve, have reduced this uncertainty by a factor of ~ 5 , but clearly, a significant uncertainty still remains. However, it is important to stress that, while the uncertainty in the rate has been reduced, there is no clear indication of a significant deviation from the nominal rate that was used in our calculations; thus, our conclusions concerning the $^{16}\text{O}/^{18}\text{O}$ ratios remain essentially unaffected.

The decrease from the huge initial ratios (see Table 4) down to the values predicted for the ejecta is a measure of the nuclear processes that transform ^{16}O to $^{17,18}\text{O}$, beginning with proton captures onto ^{16}O , which require high enough temperatures to overcome the large Coulomb potential barrier. Indeed, at the typical temperatures attained in nova outbursts, the abundance of ^{16}O always decreases, since $^{16}\text{O}(p, \gamma)$ dominates over $^{15}\text{N}(p, \gamma)^{16}\text{O}$, $^{19}\text{F}(p, \alpha)^{16}\text{O}$, and $^{17}\text{F}(\gamma, p)^{16}\text{O}$. The ^{17}O is synthesized by $^{16}\text{O}(p, \gamma)^{17}\text{F}(\beta^+)^{17}\text{O}$ and can be destroyed by either $^{17}\text{O}(p, \gamma)^{18}\text{F}$ (which decays into ^{18}O) or $^{17}\text{O}(p, \alpha)^{14}\text{N}$. The dominant destruction reaction for ^{18}O is $^{18}\text{O}(p, \alpha)$. Since ONe models, which reach higher peak temperatures than CO models, synthesize larger amounts of both $^{17,18}\text{O}$, they are characterized by lower $^{16}\text{O}/^{17}\text{O}$ ratios and similar or lower $^{16}\text{O}/^{18}\text{O}$ ratios than models of CO novae.

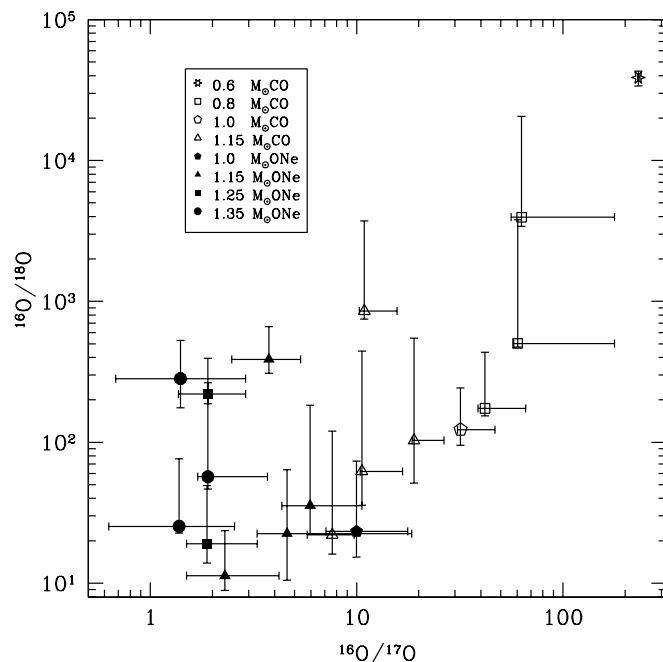


FIG. 5.—Same as Fig. 4, but for $^{16}\text{O}/^{18}\text{O}$ and $^{16}\text{O}/^{17}\text{O}$ ratios. [See the electronic edition of the Journal for a color version of this figure.]

A similar trend is found when looking at the role of the WD mass (Fig. 5): in CO models, as the mass of the WD increases (accompanied by increasing temperatures in the envelope), the $^{16}\text{O}/^{17}\text{O}$ and $^{16}\text{O}/^{18}\text{O}$ ratios both decrease. No trend is clearly seen for ONe novae: the synthesis of ^{17}O has a maximum at a temperature around 2×10^8 K (i.e., $1.25 M_{\odot}$ ONe models); however, because of the decrease in ^{16}O , the $^{16}\text{O}/^{17}\text{O}$ ratio decreases in general as the mass of the WD increases. Most ONe models show similar $^{16}\text{O}/^{18}\text{O}$ ratios, around ~ 200 – 400 with the new $^{18}\text{F}(p, \alpha)$ rates (see H99), with no clear dependence on the WD mass.

Another interesting source of information are the neon isotopic ratios. They are useful for distinguishing between CO and ONe novae: the higher initial ^{20}Ne content in ONe novae is the main reason for the much higher $^{20}\text{Ne}/^{22}\text{Ne}$ ratios found in those models (Fig. 6), ranging typically from ~ 100 to 250 (solar ratio = 14). In contrast, CO models yield $^{20}\text{Ne}/^{22}\text{Ne}$ ratios ranging only from ~ 0.1 to 0.7.

Differences between CO and ONe novae are not so extreme with regard to the $^{20}\text{Ne}/^{21}\text{Ne}$ ratio (see also Fig. 6), and in fact, ratios for the two nova types overlap at values of ~ 2500 – $10,000$. The increase in the $^{20}\text{Ne}/^{21}\text{Ne}$ ratio with respect to the initial value (see Table 4 and Appendix) reflects the fact that ^{20}Ne is scarcely modified in most nova models, since its destruction by proton capture reactions requires rather high temperatures. On the other hand, ^{21}Ne , a fragile isotope, is almost completely destroyed: first, by proton capture reactions and then, as the temperature rises, the synthesis path through $^{20}\text{Ne}(p, \gamma)^{21}\text{Na}(\beta^+)^{21}\text{Ne}$ is halted as soon as proton captures on ^{21}Na become faster than its β^+ -decay. This accounts for the large $^{20}\text{Ne}/^{21}\text{Ne}$ ratios found in both nova types.

2.3. Aluminum and Magnesium Isotopic Ratios

Similar $^{26}\text{Al}/^{27}\text{Al}$ ratios (typically, ~ 0.01 – 0.6 ; see Fig. 7) are obtained for both CO and ONe nova models. Although ^{26}Al is efficiently synthesized only in ONe novae, the larger initial amount of ^{27}Al in such novae, more than 2 orders of magnitude higher than in CO novae (and not strongly modified during the

TABLE 4
INITIAL ISOTOPIC RATIOS FOR BOTH CO AND ONe MODELS, AS A FUNCTION OF THE DEGREE OF MIXING

INITIAL RATIO ^a	CO MODELS			ONe MODELS		
	25%	50%	75%	25%	50%	75%
¹² C/ ¹³ C	4987	14782	44165	181	362	906
¹⁴ N/ ¹⁵ N	271	271	271	271	271	271
¹⁶ O/ ¹⁷ O	47,724	137,929	408,542	49,182	142,302	421,663
¹⁶ O/ ¹⁸ O	9064	26,196	77,592	9340	27,026	80,084
²⁰ Ne/ ²¹ Ne	412	412	412	56	55	55
²⁰ Ne/ ²² Ne	0.51	0.18	0.059	74	78	79
²⁴ Mg/ ²⁵ Mg	7.9	7.9	7.9	3.7	3.6	3.6
²⁶ Mg/ ²⁵ Mg	1.1	1.1	1.1	0.61	0.60	0.60

^a For all CO and ONe Models, $^{26}\text{Al}/^{27}\text{Al} = \delta(^{29,30}\text{Si}/^{28}\text{Si}) = 0$.

explosion), results in similar ratios in the two nova types. Therefore, the $^{26}\text{Al}/^{27}\text{Al}$ ratio is not a diagnostic for distinguishing between CO and ONe novae. Nucleosynthesis of ^{26}Al is complicated by the presence of a short-lived $^{26}\text{Al}^m$ ($\tau = 9.15$ s) spin isomer. The only way to synthesize the long-lived $^{26}\text{Al}^g$ isotope ($\tau = 1.04 \times 10^6$ yr) in nova explosions is through proton capture reactions on ^{25}Mg , which can yield both the ^{26}Al ground and isomeric states. Hence, the ^{25}Mg abundance is critical for the synthesis of ^{26}Al . Eventually, other isotopes, such as ^{24}Mg , ^{23}Na , and, to some extent, ^{20}Ne , can also contribute to the final ^{26}Al yield (see JCH99). The synthesis of ^{27}Al is also complicated: whereas it is mainly destroyed by $^{27}\text{Al}(p, \gamma)$, several mechanisms compete in its synthesis: one is $^{26}\text{Mg}(p, \gamma)$, with ^{26}Mg coming from its initial abundance, as well as from $^{26}\text{Al}^m$ decay [synthesized by $^{25}\text{Mg}(p, \gamma)$ or through two proton captures on ^{24}Mg , leading to the β^+ -unstable ^{26}Si]; another possibility is $^{27}\text{Si}(\beta^+)^{27}\text{Al}$, with ^{27}Si coming from both $^{26}\text{Al}^{g,m}(p, \gamma)$. There is some tendency to obtain higher $^{26}\text{Al}/^{27}\text{Al}$ ratios for more massive WDs.

Both CO and ONe nova models yield, in general, low $^{24}\text{Mg}/^{25}\text{Mg}$ (~ 0.02 – 0.3) and $^{26}\text{Mg}/^{25}\text{Mg}$ (~ 0.07 – 0.2) ratios

(see Fig. 8), except for the extreme $0.6 M_{\odot}$ CO case, with $^{24}\text{Mg}/^{25}\text{Mg} = 4.3$ and $^{26}\text{Mg}/^{25}\text{Mg} = 0.7$. The CO nova models show a rather complicated pattern because Mg synthesis is very sensitive to the maximum temperature (and hence to the adopted WD mass) attained in the explosion. Since proton captures on ^{26}Mg require high enough temperatures to overcome its large Coulomb barrier, the final ^{26}Mg abundances are, in general, very close to the initial ones (with only a small decrease for the $1.15 M_{\odot}$ CO models). Again, the $0.6 M_{\odot}$ CO model shows no imprint of nuclear activity involving ^{24}Mg , but already the $0.8 M_{\odot}$ CO model begins to show a decrease in the final ^{24}Mg yield, since at the moderate temperatures reached in this model $^{24}\text{Mg}(p, \gamma)^{25}\text{Al}$ dominates $^{23}\text{Na}(p, \gamma)^{24}\text{Mg}$. This in turn explains the increase in ^{25}Mg powered by $^{25}\text{Al}(\beta^+)^{25}\text{Mg}$. However, when the temperature reaches $\sim 2 \times 10^8$ K (as for the $1.15 M_{\odot}$ CO models), the rates for $^{24}\text{Mg}(p, \gamma)^{25}\text{Al}$ and $^{23}\text{Na}(p, \gamma)^{24}\text{Mg}$ become comparable, and hence the decrease in the final ^{24}Mg yield is halted. At the same time, $^{25}\text{Mg}(p, \gamma)$ dominates over $^{24}\text{Mg}(p, \gamma)^{25}\text{Al}(\beta^+)^{25}\text{Mg}$, which accounts for some decrease in the ^{25}Mg yield.

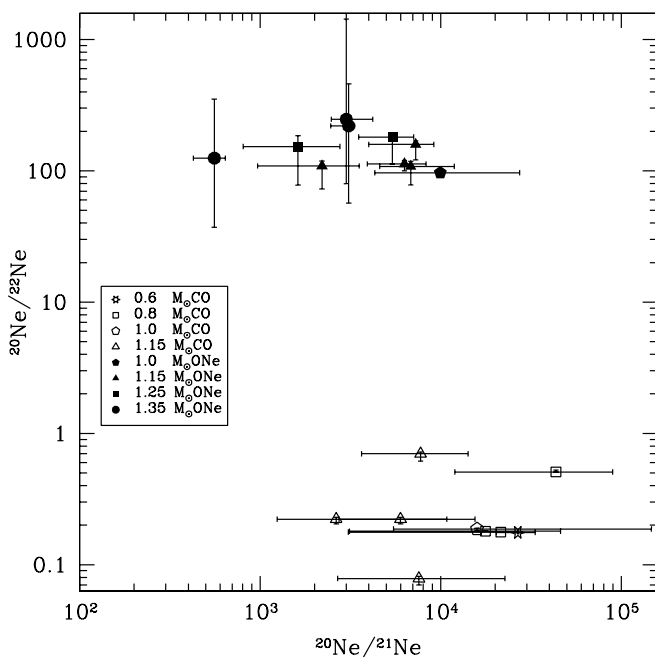


FIG. 6.—Same as Fig. 4, but for $^{20}\text{Ne}/^{21}\text{Ne}$ and $^{20}\text{Ne}/^{22}\text{Ne}$ ratios. [See the electronic edition of the Journal for a color version of this figure.]

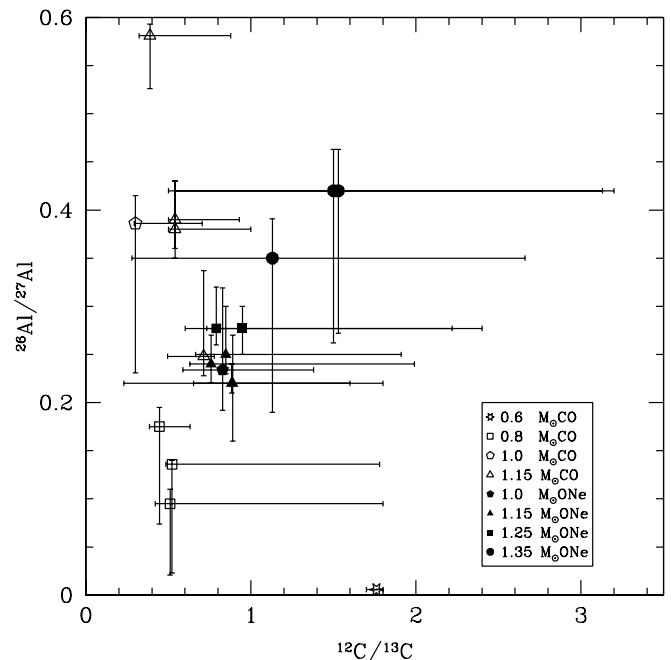


FIG. 7.—Same as Fig. 4, but for $^{26}\text{Al}/^{27}\text{Al}$ and $^{12}\text{C}/^{13}\text{C}$ ratios. [See the electronic edition of the Journal for a color version of this figure.]

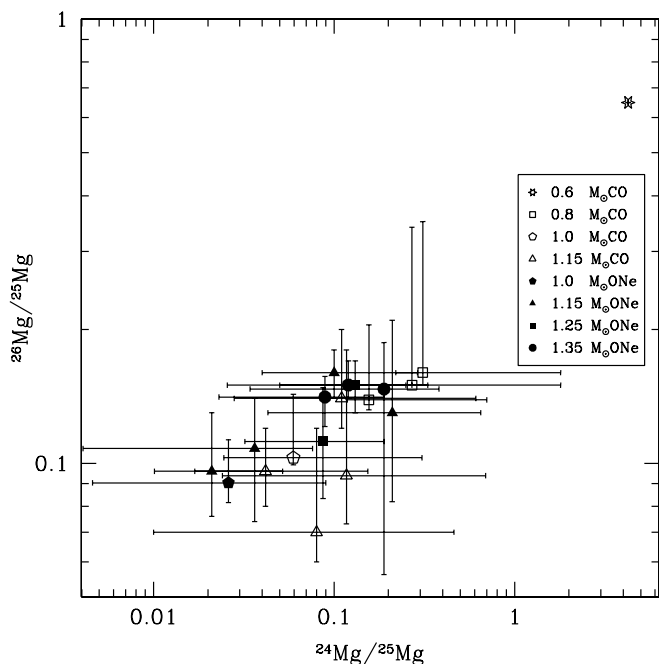


FIG. 8.—Same as Fig. 4, but for $^{24}\text{Mg}/^{25}\text{Mg}$ and $^{26}\text{Mg}/^{25}\text{Mg}$ ratios. [See the electronic edition of the Journal for a color version of this figure.]

In contrast, for the ONe models the final Mg yields do not depend much on the adopted WD mass: in all models, both the final ^{24}Mg and ^{26}Mg abundances are significantly lower than the initial values (by 2 and 1 orders of magnitude, respectively), whereas ^{25}Mg decreases by only a factor of ~ 2 . Most of the destruction of Mg isotopes takes place at temperatures around 2×10^8 K (see JCH99). The differences with respect to the results for CO models are essentially due to significant differences in the initial chemical composition (the ONe models are, for instance, much richer in ^{23}Na and $^{25,26}\text{Mg}$), which affects not only the dominant nuclear path but also the characteristic timescales of the explosion and hence the exposure time to potential proton capture reactions.

2.4. Silicon Isotopic Ratios

CO novae show, in general, a very limited nuclear activity beyond the CNO mass region because of the moderate peak temperatures attained during the explosion and also because of the lack of significant amounts of “seed” nuclei above this mass range. Therefore, hydrodynamic models of CO novae yield close to solar Si isotopic ratios in the ejecta. Only the most massive CO models (i.e., $1.15 M_{\odot}$) show marginal activity in the Si region, powered by a moderate leakage from the MgAl region through $^{26}\text{Al}(p, \gamma)^{27}\text{Si}$, followed by either $^{27}\text{Si}(p, \gamma)^{28}\text{P}(\beta^+)^{28}\text{Si}$ or $^{27}\text{Si}(\beta^+)^{27}\text{Al}(p, \gamma)^{28}\text{Si}$, which compete favorably with $^{28}\text{Si}(p, \gamma)$ and hence tend to increase the amount of ^{28}Si . In contrast, the mass fraction of ^{29}Si decreases, since destruction through $^{29}\text{Si}(p, \gamma)$ dominates synthesis by $^{28}\text{Si}(p, \gamma)^{29}\text{P}(\beta^+)^{29}\text{Si}$. The β^+ -decay of the residual ^{30}P nuclei is responsible for some marginal overproduction of ^{30}Si .

Silicon isotopic ratios are usually expressed as $\delta^{29,30}\text{Si}/^{28}\text{Si} = [(^{29,30}\text{Si}/^{28}\text{Si}) / (^{29,30}\text{Si}/^{28}\text{Si})_{\odot} - 1] \times 1000$, which represent deviations from solar abundances in permil. As shown in Figure 9, all CO models are characterized by close to or lower than solar $\delta^{29}\text{Si}/^{28}\text{Si}$ and close to solar $\delta^{30}\text{Si}/^{28}\text{Si}$. A quite different pattern is found for the ONe models, partially because of the higher peak temperatures achieved during the explosion, but also

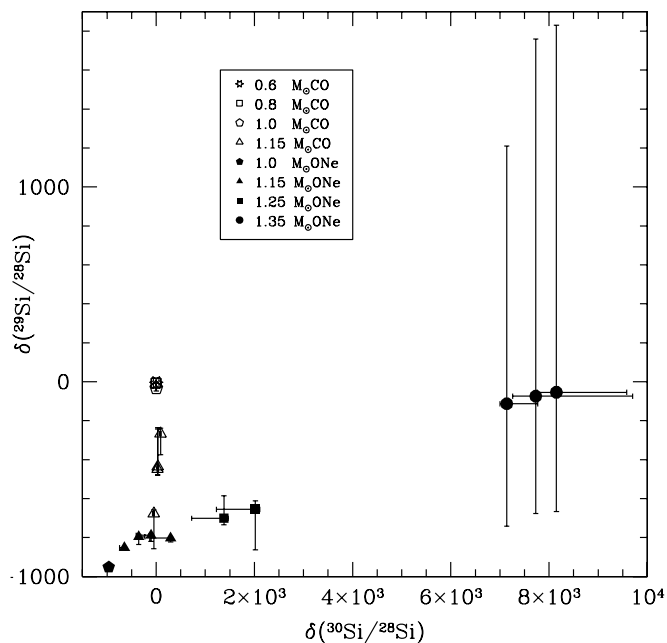


FIG. 9.—Same as Fig. 4, but for silicon isotopic ratios, expressed as delta values (deviations from the solar Si isotopic ratios in permil). [See the electronic edition of the Journal for a color version of this figure.]

because of the higher initial ^{27}Al abundance. The abundance of ^{28}Si increases from 1 to $1.25 M_{\odot}$ ONe models and then decreases a bit for $1.35 M_{\odot}$ models. This results from the fact that around $T = 10^8$ K, $^{27}\text{Al}(p, \gamma)^{28}\text{Si}$ dominates $^{28}\text{Si}(p, \gamma)$. When the temperature rises, $^{26}\text{Al}^{m,g}(p, \gamma)^{27}\text{Si}(p, \gamma)^{28}\text{P}(\beta^+)^{28}\text{Si}$ also contributes to ^{28}Si synthesis, but as the temperature reaches $\sim 3 \times 10^8$ K, destruction through proton capture reactions dominates all reactions leading to ^{28}Si synthesis. Hence, there is a maximum in the ^{28}Si production around $1.25 M_{\odot}$ ONe models (which attain $T_{\text{peak}} < 3 \times 10^8$ K).

In contrast, both ^{29}Si and ^{30}Si increase monotonically with the WD mass. They are powered by $^{29,30}\text{P}(\beta^+)^{29,30}\text{Si}$, which dominate destruction through proton capture reactions. Figure 9 shows an increase in $\delta^{30}\text{Si}/^{28}\text{Si}$ with increasing WD mass: whereas $1.0 M_{\odot}$ ONe models show a noticeable destruction of ^{30}Si , $1.15 M_{\odot}$ ONe models yield close to solar $\delta^{30}\text{Si}/^{28}\text{Si}$ values. Excesses appear for $M_{\text{wd}} \geq 1.25 M_{\odot}$, as a result of the higher temperatures attained in the envelope. On the other hand, $\delta^{29}\text{Si}/^{28}\text{Si}$ ratios are below solar and only approach close to solar values when the WD mass reaches $1.35 M_{\odot}$.

2.5. Isotopic Ratios of Elements beyond Silicon

For classical novae, nuclear activity above silicon is limited to events involving a very massive ONe WD, close to the Chandrasekhar limit, since nucleosynthesis in the Si-Ca mass region requires temperatures above 3×10^8 K to overcome the large Coulomb barriers of those elements (Politano et al. 1995; JH98; Starrfield et al. 1998). Indeed, observations of some novae reveal the presence of nuclei in this mass range in their spectra, including sulfur (Nova Aql 1982; Snijders et al. 1987; Andreã et al. 1994), chlorine (Nova GQ Mus 1983; Morisset & Pequignot 1996), and argon and calcium (Nova GQ Mus 1983, Morisset & Pequignot 1996; Nova V2214 Oph 1988, Nova V977 Sco 1989, and Nova V443 Sct 1989; Andreã et al. 1994). Models of explosions on $1.35 M_{\odot}$ ONe WDs yield large overproduction factors (i.e., $f = X_i/X_{i,\odot}$) for a number of isotopes, including ^{31}P ($f \sim 1100$), ^{32}S ($f \sim 110$), ^{33}S

($f \sim 150$), and ^{35}Cl ($f \sim 80$) (JCH01). However, the chances to measure such excesses in presolar grains are scarce. Although the predicted ^{33}S excess may provide a remarkable signature of a classical nova event, no sulfur isotopic measurements have been made so far on presolar SiC grains. Nevertheless, equilibrium condensation calculations predict that sulfides might be incorporated into SiC grains (Lodders & Fegley 1995). Analyses of graphite grains (unpublished data) yielded solar S isotopic ratios, and it is likely that the measured S is dominated by contamination or that any indigenous S has been isotopically equilibrated during the chemical separation procedure. Since ^{31}P is the only stable phosphorus isotope, no P isotopic ratios can be obtained. One chance might be to measure the ratio of two isotopes from different elements (such as P and S), but this would require information on the condensation behavior of these elements, which usually cannot be obtained. Two stable isotopes are available for chlorine, but there is not much of a chance to measure excesses in ^{35}Cl because Cl is not expected to condense into SiC. Furthermore, the standard separation procedure for SiC uses HCl, introducing a strong Cl contamination.

3. FORMATION OF GRAINS

We explored grain formation by calculating thermodynamic equilibrium condensation sequences of the ejected layers for three representative nova models, involving $1.15 M_{\odot}$ CO, $1.15 M_{\odot}$ ONe, and $1.35 M_{\odot}$ ONe WDs. In particular, we adopted the chemical composition of the first, innermost ejected layer in each case, since the largest changes in chemical composition from solar are expected to be found precisely in the innermost shells of the envelope. We note that similar results are found if nearby shells (i.e., 5 or 10 shells above the innermost ejected one) are used instead, but toward the outermost shells chemical variations are less extreme and different condensates are likely to appear. The isotopic chemical pattern of those envelope shells, calculated self-consistently by means of the hydrodynamic code (for nuclei ranging between H and Ca), was augmented by assuming solar element/Ca abundance ratios for elements heavier than Ca. Calculations were performed with the CONDOR code, and the computational procedure is similar to that described in Lodders & Fegley (1995, 1997) and Lodders (2003). The calculations use the temperature and pressure profile computed for the whole envelope with the hydrodynamic code. At temperatures where condensation occurs, total pressures are in the range of 10^{-6} to 10^{-7} bars. It should be stressed that the results of these computations only apply to the innermost ejected shell of the expanding nova ejecta and the underlying assumption is that no mixing occurs between this and overlying shells. Therefore, the calculations only describe the condensates for an extreme endmember composition of the overall ejecta from a given nova model, and additional condensates of different mineralogy that may be produced in the outer shells are not considered here. The investigation of the condensates that can form in the different ejected layers and in overall homogenized ejecta will be described elsewhere (K. Lodders et al. 2004, in preparation).

3.1. Condensates for $1.15 M_{\odot}$ CO Nova Ejecta

The atomic C/O ratio of the innermost shell of this CO nova model is ~ 0.8 , and we expected oxides and silicates as condensates, similar to those that condense from a solar composition gas. This expectation was met and the condensation

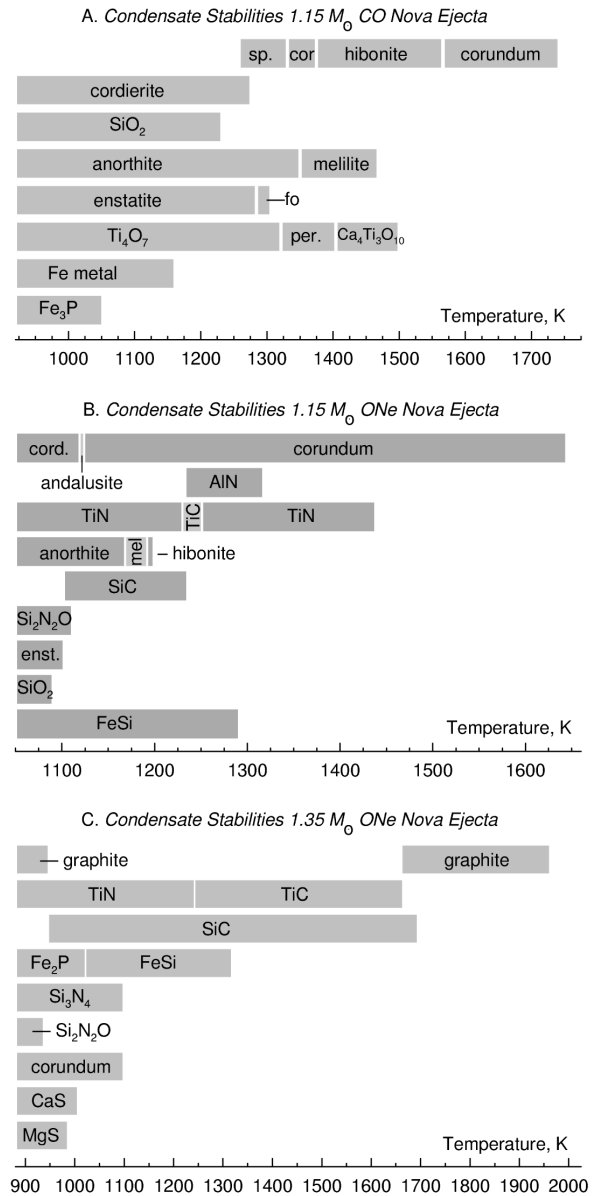


FIG. 10.—Equilibrium condensation sequences showing the different types of grains expected to form in the ejecta of three different classical nova outbursts: a $1.15 M_{\odot}$ CO WD (*top panel*), a $1.15 M_{\odot}$ ONe WD (*middle panel*), and a $1.35 M_{\odot}$ ONe WD (*bottom panel*).

sequence is shown in Figure 10 (*top panel*). The first condensate is corundum (Al_2O_3) at 1743 K, followed by hibonite ($\text{CaAl}_{12}\text{O}_{19}$) at 1567 K. Gehlenite ($\text{Ca}_2\text{Al}_2\text{SiO}_7$), the Al-rich endmember of the gehlenite-akermanite solid solution called melilite, appears in addition at 1469 K. This phase consumes Ca, which is much less abundant than Al. This limits the stability of hibonite, and corundum is stable again after melilite appears. With further decrease in temperature, melilite transforms into anorthite ($\text{CaAl}_2\text{Si}_2\text{O}_8$), and corundum into spinel (MgAl_2O_4). Cordierite ($\text{Mg}_2\text{Al}_4\text{Si}_5\text{O}_{18}$) appears together with anorthite and spinel below ~ 1280 K. In fact, cordierite is not a stable condensate in a solar composition gas but appears here because of the larger relative abundances of Mg, Si, and Al. Cordierite eventually consumes more Al so that the spinel stability is terminated. Substantial removal of Si and Mg from the gas starts when forsterite (Mg_2SiO_4) and

then enstatite (MgSiO_3) condense near 1300 K and, in addition to enstatite, SiO_2 appears as a separate phase. The first Ti-bearing condensate is a calcium titanate ($\text{Ca}_4\text{Ti}_3\text{O}_{10}$) at 1500 K, which converts to perovskite (CaTiO_3) at lower temperatures. Anorthite is the major sink for Ca, and perovskite transforms into Ti_4O_7 shortly after anorthite becomes stable. Metallic iron condenses at 1166 K and phosphorus condenses as Fe_3P at 1053 K. Iron sulfide (FeS) only forms at low temperatures of ~ 720 K. In this condensation sequence, corundum, spinel, and enstatite are the only minerals that are also found among the major presolar grain types. Reduced condensates such as SiC or graphite do not appear, which suggests that models similar to this, involving a $1.15 M_\odot$ CO WD, will not contribute to the presolar SiC or graphite grains.

3.2. Condensates for $1.15 M_\odot$ ONe Nova Ejecta

The C/O ratio of the first, innermost ejected shell is now ~ 0.7 , below unity and only slightly lower than that of the CO nova model of the same mass. If the C/O ratio were the sole criterion, oxidized condensates would be expected as well. However, by comparison to solar, the abundances of Al, Ca, Mg, and Si are fairly high in ONe ejecta, which means that the C and O chemistries will be affected by the abundances of the rock-forming elements. Figure 10 (*middle panel*) shows the calculated condensate stabilities as a function of temperature. The first condensate is corundum at 1644 K, followed by AlN and TiN. The AlN only coexists with corundum for a short temperature interval, and oxidized Al-bearing compounds (i.e., hibonite, melilite, anorthite) coexist with corundum instead at lower temperatures. Near 1120 K, corundum turns into andalusite (Al_2SiO_5) and cordierite when more of the abundant silicon is removed from the gas. The TiN stability range is interrupted for a brief temperature step from ~ 1230 to 1250 K, when TiC is more stable, but the TiC stability is limited by the appearance of SiC at 1235 K. The first Si-bearing condensate is FeSi, and SiC is the next stable one. Sinoite ($\text{Si}_2\text{N}_2\text{O}$) enters the suite of condensates at 1110 K, followed by enstatite around 1100 K and SiO_2 at ~ 1090 K. With the appearance of sinoite and enstatite the SiC stability ends. The occurrence of SiC, corundum, and enstatite in this ejected shell suggests that such intermediate-mass ONe novae could contribute to the known presolar SiC, corundum, and enstatite grain populations. However, unlike for the more massive case discussed below, Si_3N_4 , a rare presolar grain type, is not found among the condensates in this $1.15 M_\odot$ nova model.

3.3. Condensates for $1.35 M_\odot$ ONe Nova Ejecta

The condensate stabilities in the $1.35 M_\odot$ ONe nova model are shown in Figure 10 (*bottom panel*). Of the three cases investigated here, this is the only one with a C/O ratio above unity, in the specific shells considered. Condensation of graphite starts at a relatively high temperature of 1960 K. The carbides of silicon and titanium follow at 1690 and 1660 K, respectively. When TiC starts forming in addition to SiC, graphite is no longer stable because the Si and C abundances are approximately the same and SiC consumes carbon. However, graphite appears again at low temperatures (~ 950 K) when SiC is no longer stable. In addition to SiC, silicon condenses as iron silicide, silicon nitride, and silicon oxynitride. The latter two compounds are responsible for the termination of SiC stability and these compounds form mainly because the

nitrogen abundance is about 5 times that of silicon (or carbon). For similar reasons, TiC is replaced by TiN near 1240 K. Moreover, aluminum nitride begins to condense at 1080 K but corundum becomes more stable at 980 K. Ca and Mg both form sulfides below ~ 1000 K and no other calcium and magnesium compounds appear because S is more abundant than both Ca and Mg combined. The P abundance in this ejected shell is nearly 20 times larger than that of Fe and therefore all iron from iron silicide enters Fe_2P at 1020 K. These results suggest that condensates from massive ONe novae could be present among the known major presolar grain types graphite, SiC, Si_3N_4 , and corundum.

4. DISCUSSION

4.1. Mean Mass-averaged Values versus Individual Shells

In this section we summarize the trends found in our analysis of mean mass-averaged isotopic ratios (i.e., § 2) and address how they compare with a more detailed approach based on individual shell variations.

4.1.1. N and C Isotopic Ratios

The most remarkable trends found in our analysis of $^{14}\text{N}/^{15}\text{N}$ and $^{12}\text{C}/^{13}\text{C}$ ratios can be summarized as follows:

1. Large dispersion in N ratios.
2. In general, larger N ratios for CO models, with huge ratios of about ~ 100 – 1000 for low-mass models.
3. The N ratio decreases when the adopted WD mass increases, for both CO and ONe models.
4. Similar (low) C ratios for all models.
5. CO models yield, in general, lower C ratios than ONe models.

As can be seen in Figure 4, most of these trends, obtained in the framework of mean mass-averaged ratios, still hold when considering individual shells. Despite the large overlap between models, we still can argue that the $^{14}\text{N}/^{15}\text{N}$ ratios are diagnostic for distinguishing between CO and ONe novae, especially when taking into account that, according to detailed stellar evolution calculations, it is likely that WDs with masses lower than $1.05 M_\odot$ are of the CO type, whereas more massive objects would be made of ONe. This fact would eliminate the overlap between the two groups in Figure 4 with regard to N ratios. Moreover, the present plot shows no overlap at all around $^{14}\text{N}/^{15}\text{N} \sim 50$ – 1000 , which reinforces our claim that such large N ratios are characteristic of CO novae. It is hard to assess if the dependence of the N ratio on the WD mass still holds: despite the trend seen in Figure 4 for mass-averaged ratios, the big overlap makes this claim questionable if one gives the same relative importance to all individual shells (but see discussion in § 2). Concerning C ratios, it is clear from Figure 4 that all models are characterized by small ratios, regardless of the nova type, the degree of mixing, and/or the mass of the compact star. This, indeed, provides a remarkable nova signature, especially when combined with a simultaneous low N ratio (see Fig. 1). Moreover, a recent estimate of the $^{12}\text{C}/^{13}\text{C}$ ratio, ranging from 0.88 to 1.89 (Rudy et al. 2003), inferred from near-infrared spectrophotometry of the non-neon nova V2274 Cygni 2001 No. 1, seems to be fully compatible with the range of values shown in Figure 4 for some CO novae, when individual shells are taken into account. Furthermore, and for similar reasons as those mentioned above, it is hard to derive any correlation between the C ratios and the nova type when individual layers are taken into account, although

Figure 4 suggests that larger values can be reached in ONe novae. It is worth noting that the largest dispersions in the $^{12}\text{C}/^{13}\text{C}$ ratios are found in $1.35 M_{\odot}$ ONe models.

4.1.2. O and Ne Ratios

The most remarkable trends found for $^{16}\text{O}/^{17,18}\text{O}$ and $^{20}\text{Ne}/^{21,22}\text{Ne}$ are as follows:

1. Moderate to large $^{16}\text{O}/^{18}\text{O}$ ratios and moderate $^{16}\text{O}/^{17}\text{O}$ ratios for CO models.
2. Lower O ratios found, in general, for ONe models.
3. In CO models, the O ratios decrease with increasing adopted WD masses.
4. Similar $^{20}\text{Ne}/^{21}\text{Ne}$ ratios in CO and ONe models, with an overlapping region at $\sim 2500\text{--}10,000$.
5. Much larger $^{20}\text{Ne}/^{22}\text{Ne}$ ratios in ONe models than in CO models.

As shown in Figure 5, there is only a small overlap in the O ratios between CO and ONe models even when variation bars are considered. In fact, CO models are somewhat concentrated toward the upper right part of the diagram, whereas ONe models tend to cluster around the lower left corner. We can thus claim that CO models are characterized by larger $^{16}\text{O}/^{17}\text{O}$ ratios (with only a small overlap with ONe models around $\sim 10\text{--}20$) whereas a much larger overlapping region, around $\sim 30\text{--}700$, is found for $^{16}\text{O}/^{18}\text{O}$ between the two nova types. However, the largest and smallest $^{16}\text{O}/^{18}\text{O}$ ratios are still achieved in CO and ONe models, respectively. The correlation found between the O ratios and the WD mass for CO models still holds, in general, for $^{16}\text{O}/^{17}\text{O}$. However, a correlation is less pronounced for $^{16}\text{O}/^{18}\text{O}$ although models with 0.6 , 0.8 , and $1.0 M_{\odot}$ show some trend. It is worth noting that, in general, larger variations are found for $^{16}\text{O}/^{18}\text{O}$ than for $^{16}\text{O}/^{17}\text{O}$ ratios.

Similar conclusions can be made for Ne ratios when considering individual shells (Fig. 6), in particular on the remarkable differences in $^{20}\text{Ne}/^{22}\text{Ne}$ between CO and ONe novae (CO models do not show significant variations relative to the mean mass-averaged values, whereas some variations are present for $^{20}\text{Ne}/^{21}\text{Ne}$) but a larger overlap for $^{20}\text{Ne}/^{21}\text{Ne}$.

4.1.3. Al and Mg Ratios

With respect to the $^{26}\text{Al}/^{27}\text{Al}$, $^{24}\text{Mg}/^{25}\text{Mg}$, and $^{26}\text{Mg}/^{25}\text{Mg}$ ratios, the main trends can be summarized as follows:

1. High Al ratios that overlap completely for both CO and ONe novae.
2. A tendency for larger $^{26}\text{Al}/^{27}\text{Al}$ ratios to be found in nova explosions hosting more massive WDs.
3. Low Mg ratios, in general, for both CO and ONe novae.
4. The Mg ratios are nearly independent of the WD mass for ONe models, whereas they show a complicated dependence pattern in CO novae.

In general, moderate dispersions are obtained for both Mg and Al isotopic ratios. Complete overlap characterizes the Al ratio plot (see Fig. 7). While high Al ratios are found for both CO and ONe novae (providing another characteristic signature of nova outbursts), no clear dependence on the WD mass is found. However, the maximum $^{26}\text{Al}/^{27}\text{Al}$ ratios are obtained for the maximum WD masses adopted for both CO and ONe populations. The extraordinary overlap shown in Figure 8 does not allow us to discern any trend in the data for both CO and ONe models.

4.1.4. Si Ratios

Our final analysis involves the Si isotopic ratios; they are characterized by the following trends in our models of classical novae:

1. Close to or lower than solar $^{29}\text{Si}/^{28}\text{Si}$ and close to solar $^{30}\text{Si}/^{28}\text{Si}$ ratios for all CO models.
2. Close to or lower than solar $^{29}\text{Si}/^{28}\text{Si}$ ratios in all ONe models.
3. Close to or lower than solar $\delta^{30}\text{Si}/^{28}\text{Si}$ ratios for ONe models with $M_{\text{wd}} \leq 1.15 M_{\odot}$, but large ^{30}Si excesses for $M_{\text{wd}} \geq 1.25 M_{\odot}$.

In general, all models are characterized by very small dispersions among individual shells (a remarkable exception being the ^{30}Si excesses in the $1.35 M_{\odot}$ ONe models; see Fig. 9). This fact reinforces most of our conclusions based on mean mass-averaged ratios: for instance, CO models with $M_{\text{wd}} \leq 1.0 M_{\odot}$ are essentially characterized by close to solar Si isotopic ratios (i.e., $\delta^{29,30}\text{Si}/^{28}\text{Si} \sim 0$), whereas CO models of $1.15 M_{\odot}$ exhibit a noticeable lower than solar $^{29}\text{Si}/^{28}\text{Si}$ ratio. Figure 9 suggests also that the dependence of the ^{30}Si excesses on the WD mass for ONe models holds for individual shells: as the mass of the WD is increased, we move from lower than or close to solar $^{30}\text{Si}/^{28}\text{Si}$ ratios (i.e., $M_{\text{wd}} \leq 1.15 M_{\odot}$) to a region characterized by moderate to huge ^{30}Si excesses. Notice that the huge excess found in all $1.35 M_{\odot}$ models provides a valuable and characteristic isotopic signature of a classical nova outburst on a massive ONe WD. A final, interesting aspect concerns the $^{29}\text{Si}/^{28}\text{Si}$ ratios found in ONe models. As shown in Figure 9, this ratio seems to increase as the mass of the WD increases. This trend, while evident when considering only mean mass-averaged quantities, seems less robust for individual shells because of a tiny overlap between models. A remarkable situation is found for $1.35 M_{\odot}$ ONe models, which exhibit an extraordinary dispersion in $\delta^{29}\text{Si}/^{28}\text{Si}$: the analysis reveals that whereas a large number of ejected shells are characterized by lower than and close to solar ratios, a few shells show huge ^{29}Si excesses. In principle, this may open up the possibility to form grains with excesses in both ^{29}Si and ^{30}Si in specific shells.

4.2. The Effect of Nuclear Uncertainties

Nuclear uncertainties associated with specific reaction rates important for nova nucleosynthesis may affect, to some extent, the predicted isotopic ratios for a number of elements. In many cases, estimates of the impact of such nuclear uncertainties are obtained from postprocessing calculations with temperature and density profiles that, in the best cases, are taken from hydrodynamic models. Usually, such an approach has to be taken with caution, since the lack of convective mixing in these parameterized calculations tends to overestimate the influence of a given nuclear uncertainty.

According to a recent analysis of the effect of uncertainties in nuclear reaction rates for nova nucleosynthesis (Iliadis et al. 2002), present reaction rate estimates give reliable predictions for both $^{12}\text{C}/^{13}\text{C}$ and $^{14}\text{N}/^{15}\text{N}$ isotopic abundance ratios, in agreement with several tests performed with hydrodynamic models of nova outbursts. In contrast, uncertainties in several reactions can introduce large variations in the final yields for a number of species between Ne and Si. The reader is referred to Iliadis et al. (2002) for a complete list of reactions whose uncertainties may affect predictions for a number of isotopes in the framework of nova nucleosynthesis. However, because

of the parametric approach adopted in that paper, the impact of each individual reaction has to be tested properly with a full hydrodynamic calculation.

Reactions whose impact has been confirmed through a series of hydrodynamic tests include $^{17}\text{O}(p, \alpha)^{14}\text{N}$, $^{17}\text{O}(p, \gamma)^{18}\text{F}$, and $^{18}\text{F}(p, \alpha)^{15}\text{O}$, which may significantly affect the $^{17,18}\text{O}$ yields, $^{21}\text{Na}(p, \gamma)^{22}\text{Mg}$, $^{22}\text{Na}(p, \gamma)^{23}\text{Mg}$, and to some extent $^{22}\text{Ne}(p, \gamma)^{23}\text{Na}$, which may affect $^{21,22}\text{Ne}(^{22}\text{Na})$, and $^{30}\text{P}(p, \gamma)^{31}\text{S}$. Recent experiments focused on $^{18}\text{F}(p, \alpha)^{15}\text{O}$ (Bardayan et al. 2002; de Sérville et al. 2003), $^{21}\text{Na}(p, \gamma)^{22}\text{Mg}$ (Bishop et al. 2003; Davids et al. 2003), $^{22}\text{Na}(p, \gamma)^{23}\text{Mg}$ (Jenkins et al. 2004), and $^{30}\text{P}(p, \gamma)^{31}\text{S}$ (E. Rehm & C. J. Lister 2003, private communication) have substantially improved this issue. Indeed, a dramatic example is provided by $^{30}\text{P}(p, \gamma)$: the $\delta^{29}\text{Si}/^{28}\text{Si}$ values are substantially reduced when the upper limit for the $^{30}\text{P}(p, \gamma)$ rate, instead of the nominal one, is adopted (see JCH01 for details). Moreover, larger differences are found for $\delta^{30}\text{Si}/^{28}\text{Si}$: the ^{30}Si excesses obtained with the nominal rate increase by up to a factor of ~ 6 if the lower limit is adopted or even turn into deficits with the upper limit. Clearly, a better determination of this critical rate is needed in order to provide more robust predictions for the Si isotopic ratios.

4.3. Comparison with Other Calculations

We compared our theoretical predictions with results obtained from similar hydrodynamic models of nova outbursts by Kovetz & Prialnik (1997) and Starrfield et al. (1997) for CO novae and by Starrfield et al. (1998) for ONe novae. It is worth mentioning that only mean mass-averaged ratios have been considered in the above-mentioned papers; consequently, our comparison will be restricted to this particular approach. In general, there is good agreement with the calculations reported by Kovetz & Prialnik (1997) and Starrfield et al. (1998) for novae hosting CO WD cores, in particular for $^{12}\text{C}/^{13}\text{C}$ and $^{16}\text{O}/^{17}\text{O}$ ratios. One difference involves the range of $^{14}\text{N}/^{15}\text{N}$ ratios predicted for nova outbursts. The very high $^{14}\text{N}/^{15}\text{N}$ ratios reported by Kovetz & Prialnik (1997) and Starrfield et al. (1997) are obtained in explosions that achieve low peak temperatures (i.e., involve low-mass WDs), for which $^{14}\text{N}(p, \gamma)^{15}\text{O}(\beta^+)^{15}\text{N}$ is not very efficient, thus reducing the ^{15}N content and increasing the final $^{14}\text{N}/^{15}\text{N}$ ratio. This interpretation is fully consistent with the results presented in this paper for the $0.6 M_{\odot}$ CO WD model, which achieves the highest N ratio. Other differences may result from the specific reaction rate libraries adopted, from details of the treatment of convective transport, or from additional input physics.

Concerning ONe models, there is also an excellent agreement with the calculations reported by Starrfield et al. (1998) for many isotopic ratios, including $^{12}\text{C}/^{13}\text{C}$, $^{26}\text{Al}/^{27}\text{Al}$, and $\delta^{29,30}\text{Si}/^{28}\text{Si}$. We stress that, besides the expected differences attributable to the specific choice of input physics, as mentioned above, the main source of differences is probably the specific prescription adopted for the initial amounts of O, Ne, and Mg in the outer shells of the WD, where mixing with the solar-like accreted material takes place. Whereas calculations by Starrfield et al. (1998) assume a core composition based on hydrostatic models of carbon-burning nucleosynthesis by Arnett & Truran (1969), rather enriched in ^{24}Mg (with ratios $^{16}\text{O}:^{20}\text{Ne}:^{24}\text{Mg} \sim 1.5:2.5:1$), we use a more recent prescription, taken from stellar evolution calculations of intermediate-mass stars (Ritossa et al. 1996), for which the ^{24}Mg content is much lower ($^{16}\text{O}:^{20}\text{Ne}:^{24}\text{Mg} = 10:6:1$). It is worth mentioning that calculations based on the Arnett & Truran (1969) abundances yield an unrealistically high con-

tribution of novae to the Galactic ^{26}Al content, in contradiction with the results derived from the COMPTEL map of the 1809 keV ^{26}Al emission in the Galaxy (see Diehl et al. 1995), which points toward young progenitors (Type II supernovae and Wolf-Rayet stars). For the purpose of comparison, we list in Tables 2 and 3 model ONeMg1, for which we assumed a $1.25 M_{\odot}$ ONeMg WD, with the chemical abundances given by Arnett & Truran (1969). As expected, this model agrees much better with the chemical patterns of the ejecta in the series of models of nova outbursts reported by Starrfield et al. (1998).

4.4. The Formation of C-rich Dust in CO Novae

The equilibrium condensation sequences reported in § 3 predict, for the first time, the types of grains that can be expected to form in the ejecta of both CO and ONe novae. This includes some contribution to the major presolar grain types, namely, corundum (CO and ONe novae), silicon carbide (ONe novae), and silicon nitride (only in massive ONe novae). These results confirm that SiC grains are likely to condense in ONe novae, giving support to the inferred ONe nova origin to presolar SiC and graphite grains recently discovered in the Murchison and Acfer 094 meteorites (A01; Amari 2002). Indeed, silicon carbide and/or carbon dust formation has been inferred through infrared measurements in a number of ONe novae, such as Nova Aql 1982 or Nova Her 1991 (see details in Gehrz et al. 1998). Nevertheless, it is important to point out that we may be facing a problem of limited statistics so conclusions exclusively based on the experimental determinations for only seven grains can induce a clear bias in our global picture of classical nova outbursts. It is our hope that the recent implementation of new devices, such as the NanoSIMS (Stadermann et al. 1999a, 1999b; Hoppe 2002), will soon improve the statistics and will help us to extract conclusions on a firmer basis, providing in turn a tool to constrain theoretical nova models. We note that three additional nova candidate grains have recently been located (Nittler & Hoppe 2004).

A puzzling preliminary result obtained in our analysis of equilibrium condensation sequences is that reduced condensates such as SiC or graphite do not form in CO novae (at least for the selected $1.15 M_{\odot}$ CO case) and hence they will not contribute to presolar SiC or graphite grains. Whereas a much deeper analysis of ejecta from a wider sample of CO nova models is required to confirm this result, it remains to be understood which mechanism is responsible for the formation of C-rich dust seen in infrared analyses of CO novae, a feature that seems to be common in many explosions of this type (see Gehrz et al. 1998). Possible explanations include a mechanism capable of dissociating the CO molecule (see Clayton et al. 1999 for a radiation-based mechanism to dissociate the CO molecule in a supernova environment), which would drive the condensation sequence out of equilibrium conditions (however, some aspects of the chemistry where the CO molecule is absent have been investigated by Ebel & Grossman 2001, showing that SiC formation is still unlikely). In this respect, the recent spectrophotometric studies of CO emission in nova V2274 Cygni 2001 No. 1 (Rudy et al. 2003) at two different epochs suggest that, whereas emission from the first overtone of carbon monoxide is seen about 18 days after outburst, the absence of such CO emission at 370 days is an indication of partial destruction of the CO molecules. Among the mechanisms proposed are photodissociation and photoionization (see Shore & Gehrz 2004), charge transfer reactions (Rawlings 1988; Liu et al. 1992), and dissociation by He^+ ions (Lepp et al. 1990). In addition, Scott (2000) has suggested that

rotation-driven latitudinal abundance gradients may affect dust formation. Other alternatives involve possible contamination of the outer layers of the main-sequence companion (during the previous evolution of the WD progenitor), which in some cases may lead to C enrichment in those shells, or scenarios leading to nova explosions with significant C-enriched envelopes, which may lead to C > O ejecta. In this respect, we have performed a hydrodynamic simulation of a $0.6 M_{\odot}$ CO WD, identical to the model previously discussed in this paper but with a slightly different composition for the outermost layers of the WD core (for which 60% ^{12}C and 40% ^{16}O has been adopted instead of the usual $^{12}\text{C}/^{16}\text{O} = 1$). The results of this test suggest that indeed the outermost ejected envelope is C-rich, allowing for the formation of C-rich dust. Finally, the recent update of the solar abundances (see Lodders 2003), which reduce the C and O content in the solar mixture by about $\sim 50\%$, may help to condense C-rich dust in CO novae thanks to the presence of Si, Mg, and Al atoms (in a similar way as described for ONe novae). Hydrodynamic tests to validate this possibility are currently under way.

5. CONCLUSIONS

In this paper we presented a detailed analysis of isotopic ratios in the ejecta of classical novae, for nuclei up to Si, based on a series of 20 hydrodynamic models of the explosion. Analyses based on both global mean mass-averaged ratios and composition gradients through individual shells were presented. From this study, we conclude that nova grains are, in general, characterized by low C ratios, high Al ratios, and close to or slightly lower than solar $^{29}\text{Si}/^{28}\text{Si}$ ratios. Other predicted isotopic ratios are specific of each nova type (CO or ONe): for instance, we expect that grains condensed in the ejecta from massive ONe novae will exhibit significant ^{30}Si excesses (with the possibility of a ^{29}Si excess not being ruled out in the outermost ejected shells), whereas those resulting from explosions in CO novae will show close to solar $^{30}\text{Si}/^{28}\text{Si}$ ratios. Indeed, our study suggests that the ejecta from ONe novae are characterized by low N ratios whereas CO novae show a large dispersion in the N ratios, with values ranging from ~ 0.1 to more than 1000. With respect to Ne, ONe novae are characterized by large $^{20}\text{Ne}/^{21}\text{Ne}$ and $^{20}\text{Ne}/^{22}\text{Ne}$ ratios, whereas CO novae show large $^{20}\text{Ne}/^{21}\text{Ne}$ but small $^{20}\text{Ne}/^{22}\text{Ne}$ ratios. However, it is worth noting that predictions of N and Ne isotopic ratios in the grains are difficult because of a likely

N isotopic equilibration and also because ^{22}Ne excesses could come from both ^{22}Ne implantation and ^{22}Na in situ decay.

In addition, we report on equilibrium condensation sequences that predict, for the first time, the types of grains that are expected to form in the ejecta of both CO and ONe novae. Our analysis shows that the ejecta of $1.15 M_{\odot}$ CO novae are likely contributors to the known presolar populations of corundum, spinel, and enstatite grains. The $1.15 M_{\odot}$ ONe novae can produce corundum and enstatite grains, as well as SiC grains. The more massive $1.35 M_{\odot}$ novae allow formation of corundum, silicon carbide, and silicon nitride grains. This analysis points out that SiC grains are likely to condense in the ejecta from ONe novae and supports the inferred ONe nova origin of the sample of presolar SiC and graphite grains isolated from the Murchison and Acfer 094 meteorites (A01; Amari 2002). Among the presolar oxide grains discovered so far, no oxide grain with a nova signature has been discovered to date, although they are likely to condense in most (if not all) nova explosions, according to this work. These grains would be clearly identified by huge ^{17}O and somewhat smaller ^{18}O excesses.

We expect that these theoretical estimates will help to correctly identify nova grains embedded in primitive meteorites. Indeed, the recent development of new instruments, such as the NanoSIMS, is expected to lead to future identification of nova grains. The improved spatial resolution and sensitivity of this instrument, together with its capability to measure simultaneously several isotopes, open new possibilities, including measurements of elemental and isotopic compositions inside the grains (see Stadermann et al. 2002). Such accurate sources of information will help to constrain nova models in a much more precise way.

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APPENDIX

NUCLEOSYNTHESIS AND NOVA MODELS

Tables 2 and 3 list the mean mass-averaged isotopic ratios in the ejected envelopes from a sample of 20 hydrodynamic models of classical nova outbursts (JH98; H99; JCH01; and unpublished data). Calculations have been carried out by means of the one-dimensional, implicit, Lagrangian, hydrodynamical code SHIVA. Minimum and maximum ratios for individual ejected shells (in square brackets) are also given for completeness.

Each model listed in Tables 2 and 3 is characterized by the mass of the underlying WD, as well as by the initial envelope composition, which distinguishes explosions taking place on WDs hosting either CO or ONe cores. As discussed in JH98, the models assume mixing between material from the outermost core and the solar-like accreted envelope (see also Starrfield et al. 1998 and references therein) in order to mimic the unknown mechanism responsible for the enhancement in metals, essentially ^{12}C , which ultimately powers the explosion. To parameterize this process, different degrees of mixing, ranging from 25% to 75%, have been considered and are also indicated in the tables. The adopted composition of the outer layers for CO WDs is $X(^{12}\text{C}) = 0.495$, $X(^{16}\text{O}) = 0.495$, and $X(^{22}\text{Ne}) = 0.01$ (Salaris et al. 1997). For ONe WDs we used $X(^{16}\text{O}) = 0.511$, $X(^{20}\text{Ne}) = 0.313$, $X(^{12}\text{C}) = 9.16 \times 10^{-3}$, $X(^{23}\text{Na}) = 6.44 \times 10^{-2}$, $X(^{24}\text{Mg}) = 5.48 \times 10^{-2}$, $X(^{25}\text{Mg}) = 1.58 \times 10^{-2}$, $X(^{27}\text{Al}) = 1.08 \times 10^{-2}$, $X(^{26}\text{Mg}) = 9.89 \times 10^{-3}$, $X(^{21}\text{Ne}) = 5.98 \times 10^{-3}$, and $X(^{22}\text{Ne}) = 4.31 \times 10^{-3}$. These values correspond to the composition of the remnant of a $10 M_{\odot}$ Population I star, evolved from the H-burning main-sequence phase up to the thermally pulsing super-AGB stage (Ritossa et al.

1996). Solar abundances were taken from Anders & Grevesse (1989). For comparison, we include also results from model ONeMg1, where the adopted chemical composition of the WD core is taken from carbon-burning nucleosynthesis calculations by Arnett & Truran (1969) (see § 4.3 for details). We note that this rather old prescription for the WD is the one assumed in all calculations of ONe(Mg) novae published by Starrfield's group up to now.

Table 4 summarizes the different initial isotopic ratios in the sample of models presented in this paper. Although the initial ^{13}C abundance is the same in both CO and ONe novae (i.e., a fraction of the solar content, depending on the adopted degree of mixing), the much higher initial ^{12}C content in CO models results in initial $^{12}\text{C}/^{13}\text{C}$ ratios different from those in ONe models: they range from ~ 5000 to $44,000$ (25%–75% mixing) for CO and from ~ 180 to 900 (25%–75% mixing) for ONe models. The initial $^{14}\text{N}/^{15}\text{N}$ ratio for all models is solar. Because oxygen in the WD (for both CO and ONe models) is almost pure ^{16}O , the initial oxygen ratios are extremely high. Values depend on the degree of mixing, which strongly modifies the ^{16}O content in the envelope: for CO models, the initial $^{16}\text{O}/^{18}\text{O}$ ratios range from 9100 (25% mixing) to $78,000$ (75% mixing), whereas $^{16}\text{O}/^{17}\text{O}$ ratios range from $48,000$ (25%) to $410,000$ (75%). Similar values are found for ONe models: $^{16}\text{O}/^{18}\text{O}$ ratios range from 9500 (25%) to $80,000$ (75%), whereas $^{16}\text{O}/^{17}\text{O}$ ratios range from $49,000$ (25%) to $420,000$ (75%). The $^{20}\text{Ne}/^{22}\text{Ne}$ initial ratios are larger in ONe novae (i.e., 74–78, depending on the degree of mixing) than in CO novae (0.06 for 75% mixing to 0.5 for 25% mixing). The initial $^{20}\text{Ne}/^{21}\text{Ne}$ ratio is 412 (the solar value) for the CO models, whereas a value around 55 corresponds to the ONe models. CO models have initially solar Mg ratios (i.e., $^{26}\text{Mg}/^{25}\text{Mg} = 1.1$; $^{24}\text{Mg}/^{25}\text{Mg} = 7.9$), since Mg is only present in the accreted material. In contrast, ONe models are characterized by $^{26}\text{Mg}/^{25}\text{Mg} = 0.6$ and $^{24}\text{Mg}/^{25}\text{Mg} = 3.6$ (where Mg from both core material and accreted envelope is taken into account). In both cases, the initial isotopic ratios are nearly independent of the adopted degree of mixing. Finally, all silicon initially present in the envelope comes from the WD companion in solar proportions (i.e., $\delta^{29,30}\text{Si}/^{28}\text{Si} = 0$), regardless of the nova type and the degree of mixing.

Most CO and ONe models listed in Tables 2 and 3 have been computed with the same nuclear reaction network, consisting of ~ 100 isotopes, ranging from ^1H to ^{40}Ca and linked through a network containing 370 nuclear reactions (details can be found in JH98). Exceptions are models ONe4, ONe7, CO2, and CO8, for which updated $^{18}\text{F}+p$ rates have been used (see H99 for details), and models CO1 and ONe9, for which both $^{18}\text{F}+p$ and S-Ca updated rates have been taken into account (see JCH01 for details).

REFERENCES

- Amari, S. 2002, *NewA Rev.*, 46, 519
 Amari, S., Gao, X., Nittler, L. R., Zinner, E., José, J., Hernanz, M., & Lewis, R. S. 2001, *ApJ*, 551, 1065 (A01)
 Amari, S., Lewis, R. S., & Anders, E. 1995, *Geochim. Cosmochim. Acta*, 59, 1411
 Anders, E., & Grevesse, N. 1989, *Geochim. Cosmochim. Acta*, 53, 197
 Andrea, J., Drechsel, H., & Starrfield, S. 1994, *A&A*, 291, 869
 Arnett, W. D., & Truran, J. W. 1969, *ApJ*, 157, 339
 Bardayan, D., et al. 2002, *Phys. Rev. Lett.*, 89, 262501
 Bishop, S., et al. 2003, *Phys. Rev. Lett.*, 90, 162501
 Clayton, D. D., Liu, W., & Dalgarno, A. 1999, *Science*, 283, 1290
 Coc, A., Hernanz, M., José, J., & Thibaud, J. P. 2000, *A&A*, 357, 561
 Davids, B., et al. 2003, *Phys. Rev. C*, 68, 055805
 de Séréville, N., et al. 2003, *Phys. Rev. C*, 67, 052801
 Diehl, R., et al. 1995, *A&A*, 298, 445
 Ebel, D. S., & Grossman, L. 2001, *Geochim. Cosmochim. Acta*, 65, 469
 Evans, A. 1990, in *The Physics of Classical Novae*, ed. A. Cassatella & R. Viotti (Berlin: Springer), 253
 Gehrz, R. D. 1999, *Phys. Rep.*, 311, 405
 ———. 2002, in *Classical Nova Explosions*, ed. M. Hernanz & J. José (New York: AIP), 198
 Gehrz, R. D., Truran, J. W., & Williams, R. E. 1993, in *Protostars and Planets III*, ed. E. H. Levy & J. I. Lunine (Tucson: Univ. Arizona Press), 75
 Gehrz, R. D., Truran, J. W., Williams, R. E., & Starrfield, S. 1998, *PASP*, 110, 3
 Hernanz, M., José, J., Coc, A., Gómez-Gomar, J., & Isern, J. 1999, *ApJ*, 526, L97 (H99)
 Hoppe, P. 2002, *NewA Rev.*, 46, 589
 Hoppe, P., Amari, S., Zinner, E., & Lewis, R. S. 1995, *Geochim. Cosmochim. Acta*, 59, 4029 (H95)
 Iliadis, C., Champagne, A., José, J., Starrfield, S., & Tupper, P. 2002, *ApJS*, 142, 105
 Jenkins, D., et al. 2004, *Phys. Rev. Lett.*, 92, 031101
 José, J., Coc, A., & Hernanz, M. 1999, *ApJ*, 520, 347 (JCH99)
 ———. 2001, *ApJ*, 560, 897 (JCH01)
 José, J., & Hernanz, M. 1998, *ApJ*, 494, 680 (JH98)
 Kovetz, A., & Prialnik, D. 1997, *ApJ*, 477, 356
 Lepp, S., Dalgarno, A., & McCray, R. 1990, *ApJ*, 358, 262
 Liu, W., Dalgarno, A., & Lepp, S. 1992, *ApJ*, 396, 679
 Lodders, K. 2003, *ApJ*, 591, 1220
 Lodders, K., & Fegley, B., Jr. 1995, *Meteoritics Planet. Sci.*, 30, 661
 ———. 1997, in *Astrophysical Implications of the Laboratory Study of Presolar Materials*, ed. T. J. Bernatowicz & E. Zinner (New York: AIP), 391
 Morisset, C., & Pequignot, D. 1996, *A&A*, 312, 135
 Nichols, R. J., Jr., Kehm, K., Hohenberg, C. M., Amari, S., & Lewis, R. S. 2004, *Geochim. Cosmochim. Acta*, submitted (N04)
 Nittler, L. R., & Hoppe, P. 2004, *Lunar Planet. Sci. Conf.*, 35, 1598
 Politano, M., Starrfield, S., Truran, J. W., Weiss, A., & Sparks, W. M. 1995, *ApJ*, 448, 807
 Rawlings, J. M. C. 1988, *MNRAS*, 232, 507
 Rawlings, J. M. C., & Evans, A. 2002, in *Classical Nova Explosions*, ed. M. Hernanz & J. José (New York: AIP), 270
 Ritossa, C., García-Berro, E., & Iben, I. 1996, *ApJ*, 460, 489
 Rudy, R. J., Dimpfl, W. L., Lynch, D. K., Mzuk, S., Venturini, C. C., Wilson, J. C., Puetter, R. C., & Perry, R. B. 2003, *ApJ*, 596, 1229
 Salaris, M., Domínguez, I., García-Berro, E., Hernanz, M., Isern, J., & Mochkovitch, R. 1997, *ApJ*, 486, 413
 Scott, A. D. 2000, *MNRAS*, 313, 775
 Shore, S. N. 2002, in *Classical Nova Explosions*, ed. M. Hernanz & J. José (New York: AIP), 175
 Shore, S. N., & Gehrz, R. D. 2004, *A&A*, 417, 695
 Shore, S. N., Starrfield, S., González-Riestra, R., Hauschildt, P. H., & Sonneborn, G. 1994, *Nature*, 369, 539
 Snijders, M. A. J., Batt, T. J., Roche, P. F., Seaton, M. J., Spoelstra, T. A. T., & Blades, J. V. C. 1987, *MNRAS*, 228, 329
 Stadermann, F. J., Bernatowicz, T., Croat, T. K., Zinner, E., Messenger, S., & Amari, S. 2002, *Lunar Planet. Sci. Conf.*, 33, 1796
 Stadermann, F. J., Croat, T. K., Bernatowicz, T. J., Amari, S., Messenger, S., Walker, R. M., & Zinner, E. 2004, *Geochim. Cosmochim. Acta*, in press
 Stadermann, F. J., Walker, R. M., & Zinner, E. 1999a, *Lunar Planet. Sci. Conf.*, 30, 1407
 ———. 1999b, *Meteoritics Planet. Sci.*, 34, A111
 Starrfield, S. 1989, in *Classical Novae*, ed. A. Evans & M. Bode (New York: Wiley), 123
 ———. 2002, in *Classical Nova Explosions*, ed. M. Hernanz & J. José (New York: AIP), 89
 Starrfield, S., Gehrz, R., & Truran, J. W. 1997, in *Astrophysical Implications of the Laboratory Study of Presolar Materials*, ed. T. J. Bernatowicz & E. Zinner (New York: AIP), 203
 Starrfield, S., Truran, J. W., Sparks, W. M., & Kutter, G. S. 1972, *ApJ*, 176, 169
 Starrfield, S., Truran, J. W., Wiescher, M. C., & Sparks, W. M. 1998, *MNRAS*, 296, 502
 Zinner, E. 1998, *Annu. Rev. Earth Planet. Sci.*, 26, 147