DEEP MIXING IN EVOLVED STARS. I. THE EFFECT OF REACTION RATE REVISIONS FROM C TO Al

S. Palmerini1, M. La Cognata2,3, S. Cristallo4, and M. Busso1

1 Dipartimento di Fisica, Università di Perugia, and INFN, Sezione di Perugia, Italy; sara.palmerini@fisica.unipg.it
2 Dipartimento di Metodologie Fisiche e Chimiche per l’Ingegneria, Università di Catania, Catania, Italy
3 Laboratori Nazionali del Sud-INFN, Catania, Italy
4 Departamento de Física Teórica y del Cosmos, Universidad de Granada, Spain

Received 2010 August 31; accepted 2010 December 17; published 2011 February 3

ABSTRACT

We present computations of nucleosynthesis in low-mass (LM) red giant branch (RGB) and asymptotic giant branch (AGB) stars of Population I experiencing extended mixing. We adopt the updated version of the FRANEC evolutionary model, a new post-process code for non-convective mixing and the most recent revisions for solar abundances. In this framework, we discuss the effects of recent improvements in relevant reaction rates for proton captures on intermediate-mass (IM) nuclei (from carbon to aluminum). For each nucleus, we briefly discuss the new choices and their motivations. The calculations are then performed on the basis of a parameterized circulation, where the effects of the new nuclear inputs are best compared to previous works. We find that the new rates (and notably the one for the $^{14}$N($p, \gamma$)$^{15}$O reaction) imply considerable modifications in the composition of post-main-sequence stars. In particular, the slight temperature changes due to the reduced efficiency of proton captures on $^{14}$N induce abundance variations at the first dredge-up (especially for $^{17}$O, whose equilibrium ratio to $^{16}$O is very sensitive to the temperature). In this new scenario, presolar oxide grains of AGB origin turn out to be produced almost exclusively by very low mass stars ($M \lesssim 1.5$–$1.7 \, M_\odot$), never becoming C-rich. The whole population of grains with $^{18}$O/$^{16}$O below $0.0015$ (the limit permitted by first dredge-up) is now explained. Also, there is now no forbidden area for very low values of $^{17}$O/$^{16}$O (below 0.0005), contrary to previous findings. A rather shallow type of transport seems to be sufficient for the CNO changes in RGB stages. Both thermohaline diffusion and magnetic-buoyancy-induced mixing might provide a suitable physical mechanism for this. Thermohaline mixing is in any case certainly inadequate to account for the production of $^{26}$Al on the AGB. Other transport mechanisms must therefore be at play. In general, observational constraints from RGB and AGB stars, as well as from presolar grains, are well reproduced by our approach. The nitrogen isotopic ratios remain consistent in mainstream SiC grains and remain an exception. For the low values measured in them (i.e., for $^{14}$N/$^{15}$N $\lesssim 2000$), we have no explanation. Actually, for the several grains with subsolar nitrogen isotopic ratios, no known stellar process acting in LM stars can provide a clue. This might be an evidence that some form of contamination from cosmic ray spallation occurs in the interstellar medium, adding fresh $^{15}$N to the grains.

Key words: nuclear reactions, nucleosynthesis, abundances – stars: AGB and post-AGB – stars: carbon – stars: evolution – stars: low-mass

Online-only material: color figures

1. INTRODUCTION

It is now well known that the processes of astration and gas return to the interstellar medium (ISM) of galaxies are dominated by low mass (LM) and intermediate mass (IM) stars, i.e., by objects below $M = 7$–$8 \, M_\odot$ (Sedlmayr 1994). These stars share the properties of ascending, after the main sequence (MS), along the red giant branch (RGB) and then, after the occurrence of core-He burning, of terminating their evolution through the so-called asymptotic giant branch (AGB) stage (see, e.g., Iben & Renzini 1983). Here, they lose mass efficiently thanks to radial pulsations and to radiation pressure on dust grains (Herwig 2005; Guandalini et al. 2006; Guandalini & Busso 2008), powering stellar winds that are enriched with the products of the internal nucleosynthesis. Tiny solid particles that are abundant in such winds maintain the signature of those nuclear processes; samples of the most refractory among those grains can be recovered from pristine meteorites (Zinner 2005) and this fact has enormously increased the importance of AGB stars as laboratories for the study of H- and He-burning nucleosynthesis (Busso et al. 1999).

At the end of the AGB stage, the residual envelopes of LM and IM stars are lost through the fast ejection of hot, ionized materials (“superwind”); the circumstellar envelope, heated and illuminated by the now hot central star, forms a planetary nebula, while the dense, electron-degenerate core starts a blueward path, which will ultimately give birth to a white dwarf (van Winckel 2003; Prada Moroni et al. 2004). Stars crossing these evolutionary stages not only represent a fundamental component of the baryonic mass in galaxies, but also provide us with invaluable information on the physical and chemical evolution of the universe. As a consequence, understanding moderately massive stars is a necessary condition for addressing a large part of modern astrophysical problems.

Unfortunately, embarrassing gaps exist in our knowledge of their physics, and fundamental problems affecting their evolution are not properly treated in common stellar modeling. This is not surprising, as stellar models provide only a basic description, where important physical mechanisms (like convection) are oversimplified and others (e.g., rotation) are often neglected (see, e.g., Charbonnel & Lagarde 2010 for a recent outline of the importance of rotation). One big problem affecting LM-star evolutionary models is revealed by the isotopic ratios of CNO elements, as observed in stellar photospheres (Harris et al. 1985; Cottrell & Sneden 1986; Gilroy & Brown 1991; Shetrone et al. 2000).
Some chemical peculiarities concerning light elements are found very early in stellar evolution (Richard et al. 2005; Meléndez et al. 2009). Others appear slightly after the occurrence of the first dredge-up (hereafter FDU), where stars populate the so-called bump of the luminosity function (Charbonnel 2004). Then, the appearance of anomalies in the composition continues along the AGB phase (Wasserburg et al. 1995; Busso et al. 2010). The observational basis for these issues has grown considerably over the years, confirming chemical anomalies at least from Li to O, but sometimes extending up to Mg and Al, including the production of the unstable isotope $^{26}$Al (see, e.g., Pilachowsky et al. 1993; Charbonnel 2004; Gratton et al. 2000; Grundahl et al. 2002; Busso et al. 2003; Zimmer et al. 2006; Nittler et al. 2008).

The above evidence suggests that non-convective transport mechanisms occur in stars, especially during the RGB and AGB phases, linking the envelope to regions where proton captures occur (Sweigart & Mengel 1979; Boothroyd et al. 1994; Charbonnel 1995; Charbonnel & Do Nascimiento 1998; Nollett et al. 2003).

Secular mixing phenomena are made easy in RGB and AGB phases immediately following an episode of convective dredge-up, because the H-burning shell then advances in homogeneous regions, so that the natural barrier opposed to mass transport by the chemical stratification is not present. Mechanisms of this kind might actually account for several puzzling astrophysical observations: from the evolution of $^3$He in the ISM (Wasserburg et al. 1995; Glöckler & Geiss 1998; Bania et al. 2002) to the complex phenomena of production and destruction of Li in the Galaxy, including its long-term decline in MS stars (Spiegel 1972; Spiegel & Zahn 1992; Zahn 1994) and its re-production in some (rare) red giants (Charbonnel & Balachandran 2000; Guandalini et al. 2009). Further implications of slow mixing probably include the anti-correlations displayed by globular cluster stars, involving IM elements up to Mg and appearing already on the MS (Gratton et al. 2001; Thévenin et al. 2001; Ramirez & Cohen 2002). Indeed, they might have been inherited from LM or IM progenitors, hosting proton captures in mixing episodes through the AGB phases.

Among the processes involved in deep-mixing phenomena, those induced by rotation certainly play an important role (Zahn 1992; Charbonnel & Lagarde 2010), despite difficulties in explaining the observed isotopic ratios (Palacios et al. 2006). Actually, a shear layer at the contact between the differentially rotating convective envelope and the more rigid stellar core is a natural expectation (see, e.g., Kippenhahn & Weigert 1990). A rigid-body behavior in the cores of LM stars might be linked to an interplay of magnetic coupling and meridional circulation (Eggenberger et al. 2005).

Despite formal differences in the various approaches, most models considered the chemical mixing induced by rotational effects (and the associated angular momentum transport) as a diffusive process, with parameterized diffusion coefficients (see, e.g., Denissenkov et al. 1998; Denissenkov & Weiss 1996, 2000; Weiss et al. 2000). This approach has also been adopted in modeling massive, stratified stars, e.g., by Maeder & Meynet (2004). Alternatives were presented by Denissenkov & Tout (2003), who studied the effects of gravitational waves, and by Busso et al. (2007b), Nordhaus et al. (2008), and Denissenkov et al. (2009). These last authors discussed the effects of magnetic buoyancy induced by a stellar dynamo. The presence of such a dynamo in red giants finds support in some stellar variability observations (Andrews et al. 1988) and in the identification of surface magnetic fields in AGB stars (Herpin et al. 2006). Stellar magnetic fields and their instabilities have been also the object of previous detailed analysis (Spruit 1999, 2002). However, by far the most popular scenario in recent years has been that of the so-called thermohaline diffusion, induced by an inversion in the molecular weight $\mu$. In H-burning regions, this is due to the activation of the reaction $^3$He$+^3$He $\rightarrow$ $^4$He + $2p$ (Eggleton et al. 2006, 2008; Charbonnel & Zahn 2007), which locally reduces $\mu$. The phenomenon is more general than that; it was recognized a long time ago and it can occur in other astrophysical contexts (Stothers & Simon 1969; Ulrich 1972).

Diffusive mechanisms (such as thermohaline mixing) and circulation phenomena (like those induced e.g., by magnetic buoyancy) yield abundance changes that are sometimes indistinguishable from one another, sometimes remarkably different, depending on the velocity of the transport. Verifying whether observations can provide clues to discriminate among the possible physical processes generating the mixing is therefore an important issue. In this respect, interesting new elements have been offered by the recent two-dimensional and three-dimensional calculations by Denissenkov (2010) and Denissenkov & Merryfield (2011). After presenting our results, we shall comment in some detail on these elements, especially in comparison with what magnetic buoyancy can a priori provide.

Before the effects of different, uncertain mixing models can be addressed, however, one has to ascertain preliminarily that the normalizing solar abundances and the nuclear input data are very reliable. Using all the upgrades that have appeared in the recent literature on these issues, we intend to revisit the coupled processes of proton capture nucleosynthesis and of deep mixing in red giants, starting from a parameterized analysis (where nuclear and abundance effects are best compared to previous work). In Section 2, we shall present a general discussion of the set of new inputs adopted, postponing to a dedicated Appendix the details of the reaction rate choices made. In Section 3, we shall then discuss recent upgrades and novelties in stellar models, and in Section 4 we shall briefly illustrate our technique for performing parameterized, post-process deep-mixing calculations, starting from the results of the Frascati Raphson-Newton Evolutionary Code (FRANEC) evolutionary code. Then, in Sections 5 and 6 we shall outline the nucleosynthesis results thus obtained, showing some comparisons with previous work, with stellar observations and with presolar grain measurements of isotopic abundances for elements from C to Al. Comments on recent models for thermohaline mixing in two and three dimensions are then added in a dedicated section (Section 7). Finally, some general conclusions are drawn in Section 8.

2. ON RECENT REVISIONS IN SOLAR ABUNDANCES AND REACTION RATES

A previous parameterized analysis of extra mixing in red giants (Nollett et al. 2003) adopted solar abundances from Anders & Grevesse (1989), coupled with the general prescriptions for reaction rates from the compilation by Angulo et al. (1999), hereinafter simply referred to as NACRE. Since then, however, many remarkable changes have occurred in both sets of input data. For what concerns solar abundances, we shall in general adopt the compilation by Asplund et al. (2009), implying a total solar metallicity of $Z = 0.014$. We underline that not only...
elemental, but also isotopic abundances (as updated in that review) are important for our analysis; this is especially the case for CNO elements. In particular, we shall make use of the new estimate of the $^{15}\text{N}/^{14}\text{N}$ ratio, as deduced by studies of the Jovian atmosphere (Fouchet et al. 2000; Owen et al. 2001); for it we adopt the value derived by the Composite Infrared Spectrometer on board the Cassini spacecraft ($2.22 \pm 0.52 \times 10^{-3}$, a factor 1.5 smaller than the terrestrial one), which should represent the initial solar-system ratio (Fouchet et al. 2004). For oxygen isotopes, we use the values suggested by Asplund et al. (2009).

Concerning reaction rates, recent extensive reviews have been published, summarizing the main experimental changes that occurred after the NACRE compilation, which has been the reference in most papers on LM stellar nucleosynthesis over the last decade. In particular, we shall discuss here (and adopt throughout the paper) the recommendations by Adelberger et al. (2010) and Iliadis et al. (2010), hereinafter simply referred to as ADE10 and ILI10, respectively. We instead will not analyze the work by Beard et al. (2010), which pertains to stellar burning at much higher temperatures and to nucleosynthesis in high-density environments. The work by ADE10 represents the follow-up of the solar-fusion cross section review by Adelberger et al. (1998) and includes subsequent measurements for reactions of the $pp$ chain and of CNO cycles. It mainly focuses on low-energy cross sections, aiming at extracting the rates for temperatures typical of the solar core ($T_9 \sim 0.015$). Reaction rates are obtained analytically, in terms of low-energy expansion coefficients of the astrophysical S(E) factors. The work by ILI10 is instead an upgrade of the previous review by Iliadis et al. (2001). In the new work, several improvements have been introduced: (1) a wider range of atomic masses is covered, from $A = 14$ to $A = 40$, including some CNO reactions; (2) updated nuclear physics inputs are considered in the reaction-rate evaluation; and (3) a calculation procedure improved with respect to that of Iliadis et al. (2001) is adopted. In detail, a suited probability distribution is first associated with each nuclear physics quantity entering the reaction-rate calculation. Then, the reaction rate itself is computed for a set of input parameter values, which are randomly taken, each following the appropriate distribution. In this way, a probability density function is generated for the reaction rate reflecting the distribution of the input values. This method has the important consequence that the upper and lower limits are estimated with a statistically appropriate approach, so that a really meaningful value can be attributed to them. Moreover, the rate is obtained over a wide temperature range $T_9 = 0.01–10$, almost reaching down to the values found at the inner borders of the convective envelopes in LM AGB stars ($T_9 \sim 0.005$). It is worth remarking that both compilations, ADE10 and ILI10, account for the recent results obtained using indirect techniques, such as the Coulomb dissociation (Baur & Rebel 1994), the asymptotic normalization coefficient (Mukhamedzhanov et al. 1997), and the Trojan Horse method (La Cognata et al. 2010). This makes the low-temperature reaction rates more accurate, thanks to the possibility of extending the measurements down to the astrophysical range of energies.

The recommended reaction rates from ADE10 and ILI10 differ, sometimes remarkably, from NACRE results. This fact (and the search for the descending astrophysical consequences) is the main motivation for the present work.

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4 For the temperature, we adopt the notation where $T_9 = T/10^9$ K.

Proton capture reactions on nitrogen and oxygen isotopes are of special importance in our context. In particular, the reaction rate for $^{14}\text{N}(p, \gamma)^{15}\text{O}$ by ADE10, reduced by 50% with respect to NACRE, leads to a reduction in the efficiency of the whole CNO cycling, moving the H-burning shell toward inner stellar regions, where temperature and density values are higher by about 10% and 25%, respectively, as compared to previous stellar models. We shall discuss at length the important consequences of these facts on the envelope abundances. As we shall see, the new data for proton captures on $^{17}\text{O}$ also induce non-negligible effects on the surface composition. By contrast, no appreciable effect derives from the reduction in the rate for $^{15}\text{N}(p, \gamma)^{16}\text{O}$; in fact, it modifies the abundances of $^{16}\text{O}$, $^{18}\text{O}$, and $^{19}\text{F}$ only in the regions where the H-burning efficiency is maximum ($T_9 > 0.05$), which are untouched by deep mixing.

In general, we use the ADE10 compilation for $pp$-chain and CNO-cycle reactions, while we adopt the results from ILI10 for heavier nuclei. However, the two compilations partly overlap in the region of CNO isotopes. In this case, specific choices are necessary. In order to motivate them, and to illustrate some of the novelties with respect to NACRE, we present in the Appendix a discussion of the available data for some key reactions and of our choice.

3. THE NEW REFERENCE STELLAR MODELS

Although we have certainly achieved a general understanding of the conditions under which chemical elements are produced in stellar interiors, a predictive nucleosynthesis theory based on first principles is still lacking. A basic missing ingredient in developing such a theory is a full understanding of the transport properties of the turbulent, reactive plasma in which the elements are forged. Within this context, many efforts have been spent in the past years to study the properties of the final phases of stellar evolution by means of theoretical models calculated with the FRANEC (see Chieffi et al. 1998). As an example, the extra-mixing calculations by Nollett et al. (2003), which represent the starting point of the present work, were based on FRANEC stellar structures taken from Straniero et al. (2003). This last work presented parameterized theoretical formulae permitting a simple derivation of the physical characteristics of AGB stars (such as, for example, the number of thermal pulses, the mass of the H-exhausted core and of the envelope, the stellar surface temperature, and so on), by interpolating the outputs of previously computed full AGB evolutionary models (Straniero et al. 1997).

Calculations presented in this paper are instead based on new evolutionary stellar sequences. In particular, we have performed complete runs (from the pre-MS to the end of the AGB phase) for a 1.5 $M_\odot$ star and a 2.0 $M_\odot$ star, with a metallicity equal to the most recent estimate for the solar one (Cristallo et al. 2009a). The runs have been performed with an upgraded version of the FRANEC code. Computations extending up to the RGB tip have also been included for various other masses in the range 1–1.7 $M_\odot$. Our calculations do not yet include rotation, nor the effects of magnetic fields. However, as compared to older versions of the FRANEC stellar code (Straniero et al. 1997), the models here adopted represent a remarkable improvement, including several new or updated ingredients.

Among the main differences with respect to the previous scenario we recall: (1) the use of a full nuclear network coupled with the physical evolution; (2) the adoption of an updated solar chemical distribution (in particular, the code...
was revised in the previous years using the Lodders 2003 compilation).\(^5\) We remind the readers that older versions of FRANEC instead used the abundances by Anders & Grevesse (1989), which are substantially different; (3) the use of new reaction rates of which crucial is in particular the new value for the \(^{14}\text{N}(p, \gamma)^{15}\text{O}\) reaction (see the Appendix for details); in the FRANEC code this update was originally implemented when the measurements by the LUNA experiment became available (Imbriani et al. 2005; Lemut et al. 2006); (4) the adoption of new low-temperature opacities and their modifications during the evolution, when the envelope chemical admixture is changed by dredge-up and becomes carbon-rich (Cristallo et al. 2007); (5) the choice, in evolved stages, of a mass-loss law different from the one previously used; this relation is similar to the one by Vassiliadis & Wood (1993), but adopts more recent data on the period–luminosity relation of AGB stars (see Straniero et al. 2006 for details); and (6) a different treatment of the radiative/convective interfaces. In particular, instead of using a bare Schwarzschild criterion, an exponentially decreasing profile of the convective velocities was applied at the inner border of the convective envelope (Cristallo et al. 2009a).

In any case, for a discussion of the improvements outlined above as items (1) to (6) and their motivations, see especially Cristallo et al. (2009a). For the scope of this present work, we want primarily to underline that the adoption of a full nuclear network (although it increases the computational time by a factor of 15–40, depending on the case considered) has proven to be crucial. Indeed, computing a set of full models, with the evolution of all the abundances coupled to the structural and thermodynamic evolution, provides bona fide abundance templates on which to verify the outcomes of any post-process calculation done for specific purposes. These post-process studies remain necessary whenever many parameterized cases have to be considered (as is required, e.g., in the present paper). However, now their results (in terms of predicted abundances) can be validated carefully on a smaller set of complete FRANEC calculations. The inclusion of the extended network into the code was primarily motivated by other reasons. In fact, the usual approach of stellar models, to consider only the limited set of reactions controlling the energy generation, becomes completely inadequate whenever the physical properties of a structure are strongly related to the chemical composition changes. This is, e.g., the case of proton ingestion by the convective shell driven by the first fully developed thermal pulse, which is found to occur in very metal-poor AGB stars and during the post-AGB phases at larger metallicities. (Cristallo et al. 2009b; Herwig et al. 2010). More generally, the code with the full network can now be used to compute the nucleosynthesis in the He shell (including r-processing). In the context of the present work, verifying the post-process calculations with a (limited) set of checks from real, full-network models was important for those isotopes whose abundances are strongly influenced by the third dredge-up (TDU). Typical examples of this are nuclei such as \(^{22}\text{Ne}\) (produced in the thermal pulses), \(^{23}\text{Na}\) (derived from proton and neutron captures on these same \(^{22}\text{Ne}\) seeds), and \(^{26}\text{Al}\). This last nuclide is produced in the innermost H-burning zones, hence sinks into the He-rich buffer and is subsequently largely destroyed by neutron captures at high temperature \((T \gtrsim 3 \times 10^9 \text{ K})\) during the thermal pulses. The surviving part is then dredged up again to the H-rich layers and can account for \(^{26}\text{Al}/^{27}\text{Al}\) ratios in the envelope of up to \((5–6) \times 10^{-3}\), which are then enhanced by the extra production of \(^{26}\text{Al}\) in deep-mixing episodes.

As we shall discuss in the next section, the new models imply important changes in the surface isotopic composition already on the RGB, at the occurrence of the FDU. In the final evolutionary stages, then, the effects are even more substantial, especially for what concerns the chemical enrichment of the envelope along the AGB.

4. COMPUTATIONS OF EXTENDED MIXING AND NUCLEOSYNTHESIS

Starting from the models described in Section 3, and also for the sake of comparison with previous works, we have performed parametric calculations, using as free parameters the rate of mass transport (\(\dot{M}\)) and the maximum temperature sampled by the circulating material \((T_P)\), as in Nollett et al. (2003). This last parameter is best expressed by the difference \(\Delta = \log T_H - \log T_P\), where \(T_H\) is the temperature at which the energy from the H-burning shell is maximum. We shall use this definition of \(\Delta\) throughout this paper; similarly, we shall often express \(\dot{M}\) in units of \(10^{-6} M_\odot\text{yr}^{-1}\) and we shall call this parameter \(\dot{M}_c\). We present here results mainly for solar-metallicity models of 1.5 and 2.0 \(M_\odot\), although some extra models have been computed for comparisons with observational data. The post-process calculations have been run in a way similar to what was done previously, using older releases of the same FRANEC stellar code (Nollett et al. 2003; Busso et al. 2010). The post-process program is new and its results have been cross-checked with the ones obtained with the older code (Palmerini et al. 2008).

During RGB stages, we have introduced extra mixing after the “bump” of the luminosity function (BLF), i.e., after the H-burning shell has erased the chemical discontinuity left behind by the FDU (Charbonnel 1996). We shall not repeat here a general discussion of RGB physics, as it can be easily found elsewhere (Charbonnel & Balachandran 2000; Palacios et al. 2006). Our procedure in the calculations was illustrated by Palmerini et al. (2009). We have followed the abundance changes during the transport due to nucleosynthesis (partial derivatives with respect to time) and to displacements across the integration grid (partial derivatives with respect to distance, times the mixing velocity). One can write this synthetically as

\[
\frac{dY_i}{dt} = \frac{\partial Y_i}{\partial t} + \dot{M} \cdot \frac{\partial Y_i}{\partial M}.
\]

Here, \(Y_i\) is the abundance by mole of the \(i\)th isotope (for the present work, this index spans from H to \(^{33}\text{S}\)). In agreement with Nollett et al. (2003) and with Busso et al. (2010), the fractional areas \(f_u\) and \(f_d\) occupied by the upward and downward mass transport, respectively, have been set to be \(f_u = f_d = 0.5\). When this analysis is pursued in the framework of a slow-mixing process, the physical mechanism of transport and its velocity do not play any crucial role; the nucleosynthesis results are then controlled by the path integral of reaction rates over the trajectory. If needed, the mixing velocity \(v\) can then be simply derived as \(v = \frac{1}{\dot{M}} \cdot \frac{M}{\Delta t}\). As discussed elsewhere (Guandalini et al. 2009), very different scenarios may open up when the transport is operated by fast mechanisms, suitable to yield mixing timescales comparable to (or shorter than) the \(^7\text{Be}\) half-life. The complex processes associated with the nucleosynthesis

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\(^5\) In the post-process, we use the updated suggestions by Asplund et al. (2009): the small differences between this set of data and the one used in the stellar code do not introduce any consequences on the model physics.
of LM nuclei such as D, $^3$He, Li, Be, and B will however be postponed to another work.

After core-He exhaustion, our LM stellar models evolve to the AGB stages. General discussions of the physics of AGB stars can be found in Iben & Renzini (1983), Busso et al. (1999), and Straniero et al. (2006). On the AGB, deep-mixing phenomena can occur again, and there is actually clear observational evidence of their presence in those final stages (Busso et al. 2010). In fact, below the convective envelopes of these evolved red giants, conditions of chemical homogeneity similar to those found after FDU are established. During the early phase of the AGB (before thermal pulses start), this is guaranteed by the previous advancement of the H-burning shell during the first ascent to the RGB. Subsequently, when thermal instabilities from the He shell begin, the downward envelope expansions (which follow them) easily reach down to the H–He discontinuity, even before the penetration into the He-rich layers occurs, a fact that formally defines the TDU. In most of the interpulse periods of AGB phases, therefore, H burning proceeds in an initially homogeneous environment, i.e., in conditions suitable for the occurrence of extra mixing.

Various experimental constraints (and in particular the spread in the C, O, and AI isotopic ratios shown by presolar grains) suggest that the details of the transport vary from star to star (Nollett et al. 2003). One also has to mention that not all the physical mechanisms suitable to drive chemical transport in first-ascent red giants are necessarily available also during the AGB stages. This was discussed by Busso et al. (2010), who showed that the operation of deep mixing on the RGB, by whatever cause, consumes $^3$He, so that the molecular weight inversion generated by its burning occurs in AGB phases only if the amount of $^3$He remaining in the envelope is substantial (as is the case in the work by Charbonnel & Lagarde 2010). Otherwise, different physical mechanisms must be looked for. In Figure 1, we better illustrate this point reporting the variation of the molecular weight ($\mu$) across the radiative region of a 1.5 and a 2.0 $M_\odot$ star with solar metallicity. While for nitrogen the change is mainly due to the choice of the new solar value, the large shifts for $^{17}$O/$^{16}$O. In order to show the selective relevance of the model choices in producing these results, Table 2 shows the same nitrogen and oxygen ratios as obtained: (1) updating, with respect to Nollett et al. (2003), the physical model parameters, the value of the solar metallicity, and the $^{14}$N($p, \gamma$)$^{15}$O rate; and (2) also adding all the other new reaction rates by ADE10 and IL110, as discussed in Section 2, and using the new (Jovian) value for the solar system nitrogen isotopic ratio. This is done for masses 1, 1.2, 1.5, and 1.7 $M_\odot$ at the solar metallicity.

A comparison between the two tables makes it clear that, while for nitrogen the change is mainly due to the choice of the new solar value, the large shifts for $^{17}$O/$^{16}$O descend from both the new abundances and the new rate for proton captures on $^{14}$N. Indeed, both the mentioned upgrades induce small but effective changes in the stellar structure, in particular in the temperature profile and in the maximum penetration of the envelope during the FDU. These changes are obviously larger for stars where CNO cycling is more efficient: our 2 $M_\odot$ model has $^{17}$O/$^{16}$O = $5.14 \times 10^{-3}$, which is more than twice the value of $2 \times 10^{-3}$, previously used in deep-mixing models by Nollett et al. (2003), taken from Boothrotd & Sackmann (1999), for the same stellar mass are also shown.

Table 1 reveals that the estimates for the surface isotopic ratios of nitrogen and oxygen at FDU can change by large amounts with respect to previous findings. In particular, this is true for $^{15}$N/$^{14}$N and for $^{17}$O/$^{16}$O. In order to show the selective relevance of the model choices in producing these results, Table 2 shows the same nitrogen and oxygen ratios as obtained: (1) updating, with respect to Nollett et al. (2003), the physical model parameters, the value of the solar metallicity, and the $^{14}$N($p, \gamma$)$^{15}$O rate; and (2) also adding all the other new reaction rates by ADE10 and IL110, as discussed in Section 2, and using the new (Jovian) value for the solar system nitrogen isotopic ratio. This is done for masses 1, 1.2, 1.5, and 1.7 $M_\odot$ at the solar metallicity.

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Table 1

| Case        | $^{15}$N/$^{14}$N | $^{17}$O/$^{16}$O | $^{18}$O/$^{16}$O |
|-------------|-------------------|-------------------|-------------------|
| $Z = 0.01$  | $6.17 \times 10^{-4}$ | $1.44 \times 10^{-3}$ | $1.57 \times 10^{-3}$ |
| $Z = 0.014$ | $6.27 \times 10^{-4}$ | $1.21 \times 10^{-3}$ | $1.59 \times 10^{-3}$ |
| $Z = 0.02$  | $6.65 \times 10^{-4}$ | $9.99 \times 10^{-4}$ | $1.62 \times 10^{-3}$ |
| Nollett et al. (2003) | $1.81 \times 10^{-3}$ | $8.10 \times 10^{-4}$ | $1.62 \times 10^{-3}$ |

Figure 1. Relative variation of the molecular weight ($\Delta \mu/\mu$) for a 1.5 and a 2.0 $M_\odot$ star with solar metallicity in the layers hosting $^3$He burning above the H shell. The $\mu$ inversion, present on the RGB, is also preserved on the AGB only if a sufficient supply of $^3$He remains. For the AGB, different lines refer to different abundances of $^3$He resulting from the previous RGB phase, as indicated by the labels.
and Boothrotd & Sackmann (1999). In this case, the new adopted solar metallicity increases the $^{17}\text{O}/^{16}\text{O}$ by about 65%, while the new $^{14}\text{N}(p, \gamma)^{15}\text{O}$ adds a further 35%. In the upper panel of Figure 2, we report the temperature profile at central hydrogen exhaustion characterizing four different 2 $M_\odot$ models, run with different inputs:

1. standard case (we shall call it ST); it has the new $\sigma_{^{14}\text{N}}$ and $Z = Z_\odot$.
2. test case with the new $\sigma_{^{14}\text{N}}$ and $Z = 0.02$ (this is the solar metallicity derived by adopting the compilation of Anders & Grevesse (1989));
3. test case with the old $\sigma_{^{14}\text{N}}$ and $Z = Z_\odot$; and
4. test case with the old $\sigma_{^{14}\text{N}}$ and $Z = 0.02$.

The standard case shows the highest temperature profile, this fact leading to a larger $^{17}\text{O}/^{16}\text{O}$ production in the region between 0.3 and 0.8 $M_\odot$ (the inner core is not affected by the FDU, while the external layers have temperatures too low to activate H burning). In the lower panel of Figure 2, we report the abundances of hydrogen, $^{16}\text{O}$, and $^{17}\text{O}$ in our ST case. In the same panel, we also report the maximum penetration of the convective envelope during FDU. Note that, although the envelope penetrates less than in the other tests ($M_{\text{bottom}}^1 = 0.302$, $M_{\text{bottom}}^2 = 0.290$, $M_{\text{bottom}}^3 = 0.298$, $M_{\text{bottom}}^4 = 0.286$), the standard case shows the largest $^{17}\text{O}/^{16}\text{O}$ value, because of the strong dependence of this ratio on the temperature (see also Nollett et al. 2003, their Figure 8 in particular).

A look at Table 2 reveals that the other reaction rate changes of Section 2, including those directly affecting oxygen isotopes, induce smaller (although non-negligible) effects than $^{14}\text{N}(p, \gamma)^{15}\text{O}$ does, in altering the isotopic mix of oxygen at FDU.

**Figure 3** shows this in graphic form: filled squares represent the isotopic composition of oxygen at FDU in our models, while open squares show the previous situation, with the old NACRE rates and the old choice for the solar metallicity. The new results confine the stellar masses that produced presolar oxide grains within a very narrow range (1 $\leq M/M_\odot \leq 1.5$–1.7).

The abundances seen at the surface of a giant star near the tip of the RGB are then known to differ from those at FDU for the effects of deep-mixing phenomena, which change the envelope composition by linking its material to the hotter, radiative regions above the H-burning shell, where proton captures re-establish a varying abundance distribution, starting from an initially homogeneous composition. Figure 3 also shows the effects of deep mixing on the RGB (for a specific choice of parameters defined in Section 4, namely, $\Delta = 0.15$ and $M_\text{mix} = 0.3$). As the figure shows, presolar grains with a moderate $^{18}\text{O}$ destruction, mainly of group 1 (Nittler et al. 1997), can be reproduced by this rather extreme processing, provided the stellar masses are confined within the range mentioned above. This range is the same where most MS–S stars are found. We shall come back to this point in the next section. Most grain data, however, and especially those with both $^{16}\text{O}/^{16}\text{O}$ and $^{17}\text{O}/^{16}\text{O}$ ratios below 0.0012, require even more extended processing than presented in the figure and clearly indicate that deep mixing cannot occur only on the RGB (see Busso et al. 2010).

Typical abundance profiles created by nucleosynthesis in the region below the convective envelope after the luminosity bump are shown in the four panels of Figure 4 taken from our 2 $M_\odot$ model with initially solar composition. In regions below $\Delta \simeq 0.2$ (corresponding to the vertical dashed line), the $^3\text{He}+^4\text{He}$ reaction starts to dominate over the $^3\text{He}+^3\text{He}$ one (which induces the inversion of the molecular weight); this is therefore the innermost position from which thermonuclear diffusion can start. Note that some remarkable abundance changes, including the production of a critical nucleus like $^{23}\text{Na}$ and an enhanced destruction of $^{18}\text{O}$, occur immediately below this zone. If only thermonuclear mixing were to take care of transporting matter to the envelope on the RGB, then we would confirm the recent findings by Charbonnel & Lagarde (2010), according to which these nuclei are only moderately affected in RGB stages. If instead deeper mixing mechanisms were at play, this result would change drastically (as we have shown for the destruction of $^{18}\text{O}$ in Figure 3).

**Table 2**

| Mass ($M_\odot$) | $^{15}\text{N}/^{14}\text{N}$ | $^{17}\text{O}/^{16}\text{O}$ | $^{18}\text{O}/^{16}\text{O}$ |
|-----------------|----------------|----------------|----------------|
| 1.0             | $2.07 \times 10^{-3}$ | $3.81 \times 10^{-4}$ | $1.89 \times 10^{-3}$ |
| 1.2             | $1.51 \times 10^{-3}$ | $4.82 \times 10^{-4}$ | $1.73 \times 10^{-3}$ |
| 1.5             | $1.09 \times 10^{-3}$ | $1.11 \times 10^{-3}$ | $1.58 \times 10^{-3}$ |
| 1.7             | $9.16 \times 10^{-4}$ | $2.04 \times 10^{-4}$ | $1.51 \times 10^{-3}$ |
| 2.0             | $7.79 \times 10^{-4}$ | $4.32 \times 10^{-4}$ | $1.44 \times 10^{-3}$ |

(a) Updating only the $^{14}\text{N}(p, \gamma)^{15}\text{O}$

| Mass ($M_\odot$) | $^{15}\text{N}/^{14}\text{N}$ | $^{17}\text{O}/^{16}\text{O}$ | $^{18}\text{O}/^{16}\text{O}$ |
|-----------------|----------------|----------------|----------------|
| 1.0             | $1.19 \times 10^{-3}$ | $3.89 \times 10^{-4}$ | $1.90 \times 10^{-3}$ |
| 1.2             | $8.64 \times 10^{-4}$ | $4.86 \times 10^{-4}$ | $1.74 \times 10^{-3}$ |
| 1.5             | $6.27 \times 10^{-4}$ | $1.21 \times 10^{-3}$ | $1.59 \times 10^{-3}$ |
| 1.7             | $5.25 \times 10^{-4}$ | $2.33 \times 10^{-3}$ | $1.52 \times 10^{-3}$ |
| 2.0             | $4.46 \times 10^{-4}$ | $5.14 \times 10^{-4}$ | $1.45 \times 10^{-3}$ |

(b) ADE10 and ILI10 rates; new [$^{15}\text{N}(p, \gamma)^{15}\text{O}$]
Figure 3. Oxygen isotopic mix in the envelope of our solar-metallicity stars at FDU (solid line). Full squares indicate the mass of the model star. As a comparison, open symbols (see arrows for associations) show the composition at FDU (solid line). Full squares indicate the mass of the model star. As a Figure 3.

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The observations do require deep mixing at a moderate temperature ($\Delta = 0.2$) and at circulation rates $0.1 \leq \dot{M}_0 \leq 0.3$. From the value of $\Delta$, we infer that thermohaline diffusion might be adequate for this purpose. However, warnings on the mixing velocity and on the diffusion coefficient deduced from the recent literature have to be considered and we refer to Section 7 for a more detailed discussion of that.

6. RESULTS FOR THE AGB PHASES

For the late evolutionary stages, the adoption of low-temperature C-enhanced opacity tables, the choice of a new mass-loss law (limiting the number of thermal pulses), and the different treatment of the inner convective border all play important roles in controlling the final results. While, again, we direct the reader to Cristallo et al. (2009a) for details, we need to recall here that the present AGB models have now a much stronger support from observations. In particular, the new opacities and the more effective mass-loss law adopted combine in inducing lower values for the surface temperature and luminosity. In their turn, these facts affect the colors, the infrared emission, and the luminosity function of AGB stars. The new models now succeed in reproducing the photometric and spectroscopic properties of their observational counterparts; in particular, the low effective temperatures of C stars at different metallicities (Dominguez et al. 2004; Busso et al. 2007a) can now be naturally reproduced. The stronger mass-loss rates reduce the number of thermal pulses, but the introduction of an exponentially decaying profile of convective velocities at the base of the envelope increases the TDU efficiency, so that the final enhancement factor of the most significant nuclei produced by AGB stars is roughly the same as before; but now, this is obtained together with a good reproduction of the luminosity functions of AGB stars (S. Cristallo et al. 2011, in preparation).

We also recall that the adopted algorithm for the velocity profile accounts, after each TDU episode, for the formation of a tiny region enriched in protons where, subsequently, $^{13}$C can be formed. This is the radiative $^{13}$C pocket where ($\alpha$, $n$) captures on $^{13}$C provide the main neutron source for the synthesis of heavy $n$-rich nuclei in AGB stars (see, e.g., Busso et al. 1999; Lagudo et al. 2003; Straniero et al. 2006; Sneden et al. 2008), thus complementing the production of lighter $n$-rich nuclei in massive stars (see, e.g., Busso & Gallino 1985; Raiteri et al. 1993; Küppeler et al. 1994; Pignatari et al. 2010). We however postpone the analysis of this important point to a future paper.

Deep mixing in thermally pulsing AGB stages operates on the composition established above the H-burning shell and homogenized after each pulse by the penetration of the convective envelope. Typical abundances established in those layers by proton captures are shown in Figure 6. Previous works (Nollett et al. 2003; Busso et al. 2010) lead us to expect that extra mixing mainly operates down to regions where $\Delta = 0.1$ to 0.2. These zones can be identified in Figure 6 from the two vertical lines. As is clear from the plots, remarkable nucleosynthesis changes are induced there by the adoption of the new reaction rates (continuous curves). Among the differences, we find a larger destruction of $^3$He, $^{15}$N, $^{18}$O, $^{19}$F, and a lower production of $^{23}$Na. This last result is directly linked to a lower destruction of $^{21}$Ne and $^{22}$Ne. The important radioactive nucleus $^{26}$Al is more effectively produced than before, but only marginally. In general, these results can be interpreted as due to the slight increase in the temperature induced by the new (less
efficient) rate for $^{14}\text{N}(p, \gamma)^{15}\text{O}$, superimposed to the specific effects of the rate changes for heavier nuclei, which have a more local and limited effect.

As a consequence of the model changes and of the reaction rate updates, the combined operation of mixing and nucleosynthesis now leads to abundance changes in the envelope such as those summarized in Figure 7 (for the 1.5 $M_\odot$ model) and in Figure 8 (for the 2 $M_\odot$ model). For these examples we adopted, on the RGB, $\Delta = 0.2$, $M_6 = 0.3$; on the AGB, $\Delta = 0.1$, $M_6 = 1$. These choices are only for the
Deep-mixing results for CNO nuclei, as compared to observations in RGB stars of metallicities close to solar. (a) Model curves correspond to $\Delta = 0.2$; they proceed from right to left, starting from abundances at FDU. Continuous lines refer to $M_0 = 0.3$, dotted lines to $M_0 = 0.1$, dashed lines to $M_0 = 0.03$. Black stars show post-FDU stars from M67 (Brown 1987; Gilroy & Brown 1991); asterisks are data from Cottrell & Sneden (1986); open squares refer to open clusters by Mikolaitis et al. (2010). Circles refer to data from Smiljanic et al. (2009), in particular to NGC5822 (filled symbols) and NGC2360 (open symbols). Only solar-metallicity models are shown to avoid excessive crowding; for the N/O ratio vs. C/O, Models of $Z = 0.02$ and $Z = 0.01$ have been included. (c) Same comparison, for the N/O vs. C/O ratios.

Figure 5. Deep-mixing results for CNO nuclei, as compared to observations in RGB stars of metallicities close to solar. (a) Model curves correspond to $\Delta = 0.2$; they proceed from right to left, starting from abundances at FDU. Continuous lines refer to $M_0 = 0.3$, dotted lines to $M_0 = 0.1$, dashed lines to $M_0 = 0.03$. Black stars show post-FDU stars from M67 (Brown 1987; Gilroy & Brown 1991); asterisks are data from Cottrell & Sneden (1986); open squares refer to open clusters by Mikolaitis et al. (2010). Circles refer to data from Smiljanic et al. (2009), in particular to NGC5822 (filled symbols) and NGC2360 (open symbols). Only solar-metallicity models are shown to avoid excessive crowding; for the N/O ratio vs. C/O, Models of $Z = 0.02$ and $Z = 0.01$ have been included. (c) Same comparison, for the N/O vs. C/O ratios.

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purposes of illustration. A wider sample of cases for the AGB is presented in the subsequent Figures 9 and 10.6

The above figures referring to the AGB clearly show a sawtooth trend in the abundances of several nuclei. This is induced by the sudden mixing with He-rich layers at the TDU, after each pulse. This trend is therefore pronounced for those nuclei that are largely affected by He-shell nucleosynthesis, in particular $^{12}$C, $^{22}$Ne, and $^{26}$Al.

The $^{12}$C-enrichment by TDU is partly compensated by its destruction in deep-mixing processing during the interpulse stages. The operation of such mixing at even moderate efficiencies has the consequence of preventing the formation of a carbon star in our models up to 1.7 $M_\odot$. Combining this with the changes already discussed for the FDU (see Figure 3), we can say that both MS, S giants, and oxide presolar grains should be produced mainly by stars below this mass limit. Obviously, a more massive star (2 $M_\odot$ or more), before becoming C-rich, also passes through an O-rich phase in which TDU induces MS–S spectral characteristics; but the high efficiency of dredge-up we have now implies that this is a not a dominant phase. This can be inferred from the limited amount of mass lost in the O-rich stages by a 2 $M_\odot$ model star becoming C-rich. For moderate extra mixing ($M_0 = 0.1$), this amounts to 0.4 and 0.2 $M_\odot$ at solar metallicity and at $Z = 0.01$, respectively. By contrast, more than 1 $M_\odot$ is lost in the C-rich phase. On the contrary, a 1.5–1.7 $M_\odot$ star loses its whole envelope (at least 0.8 $M_\odot$) in the form of O-rich material. If one also considers that the initial mass function favors these lower masses, one can deduce that most of the observed MS and S giants should have masses smaller than those of C-stars and that among these LM AGB stars, one should find the typical progenitors of presolar oxide grains. There is obviously a possibility of finding MS–S giants among IM stars ($M \geq 4 M_\odot$), where hot bottom burning prevents the formation of a carbon star. However, for them, the expected isotopic ratios (especially for oxygen) are very peculiar (e.g., with large excesses of $^{17}$O). Iliadis et al. (2008), by comparisons of IM models with presolar grain abundances, convincingly excluded any connection of IM stars with presolar grains; presently, we do not know of even one single grain really requiring an IM progenitor.

We can see this in some more detail. As already stated in the discussion of the RGB phases and in Busso et al. (2010), oxygen isotopes in presolar oxide grains provide the most stringent constraints to stellar models with deep mixing, due to the fragility of $^{18}$O and to the sensitivity of $^{17}$O to temperature. Figure 11 (top panel) presents these constraints, showing the oxygen isotopic ratios in group 1 and group 2 grains together with model curves. These last two include typical trends for extra-mixing on the RGB at a moderate efficiency, plus the results of deep-mixing computations for the AGB, for various choices of the parameters. Note that, by including the 1 $M_\odot$ star, the entire range of $^{17}$O/$^{16}$O ratios is covered. This is another effect of the model changes and corrects previous findings, based on older models, according to which the grains at the extreme left would not be explained by deep mixing at solar metallicity (Wasserburg et al. 1995; Nollett et al. 2003). The main reason for the change lies in the more marked destruction of $^{18}$O at moderate temperatures, where $^{17}$O is still untouched (see Figure 4, second panel). This is also found in very low mass models and allows their deep-mixing curves to proceed almost

For researchers interested in comparisons with specific observations and looking for an even wider series of cases, we can provide the nucleosynthesis results upon request.
Figure 6. Distribution of abundances for some light and intermediate-mass isotopes in the radiative layers between the H-burning shell and the convective envelope. The plots are taken roughly at mid-AGB evolution. The two vertical dotted lines and the dashed one indicate the places where $\Delta = 0.2, 0.1,$ and $0.0$, respectively. This last position corresponds to the layer where the energy from the H-burning shell is maximum (and $T = T_H$).

Vertically, covering a previously unaccounted area (not only the new rate for the $^{14}$N$(p, \gamma)^{15}$O, but also those for proton captures on $^{17}$O are important in determining this result). The top panel of Figure 11 also shows that the whole set of abundances for $^{18}$O-poor grains (with $^{18}$O/$^{16}$O ratios below 0.0015) is explained by models for stars below about 1.7 $M_\odot$, those that will never become C-rich.

The bottom panel of Figure 11 then presents the $^{17}$O/$^{16}$O ratio as a function of $^{26}$Al/$^{27}$Al, for those grains that show $^{26}$Mg excesses from $^{26}$Al decay. Here, $M_6$ is assumed to be 3 (a rather
efficient case). Again, most of the experimental results can be accounted for. A small group of grains, for \(^{17}O/^{16}O\) ratios below 0.001 and \(^{26}Al/^{27}Al\) above 0.01, are however not explained. We note here that no modification to the model might cover this area. In fact (as an example), mixing at higher temperature would improve the fit to \(^{26}Al\) but would worsen the problem for \(^{17}O\).
Figure 9. Evolution in time of the abundances by mole in the envelope of a 1.5 $M_\odot$ star during the AGB stages. The plot considers three values of the mixing parameter $\dot{M}_b$ and two choices of the temperature parameter $\Delta$.

(A color version of this figure is available in the online journal.)

Figure 10. Same as in the previous figure, for a stellar model of 2 $M_\odot$.

(A color version of this figure is available in the online journal.)

(not to mention the fact that mixing at higher temperatures would affect the stellar structure). There appears to be a unique way out: an increase in the reaction rate for the $^{25}$Mg($p, \gamma$)$^{26}$Al reaction at temperatures sufficiently small to minimize the effects on $^{17}$O. In the energy region of our interest, the main contribution to that reaction rate is due to resonances (at 58 and 93 keV) for which
but not in $^{17}$O can be accounted for (e.g., using $\Delta = 0.05$ and $M_0 = 3$ in a 1.2 $M_\odot$ stellar model). We leave this suggestion for future verifications. Recent new measurements (especially from LUNA) are still under analysis, so that we do not have to wait for a long time before knowing if this prediction can be confirmed.

It is important to note that the same presence of $^{26}$Al in the grains, at a level higher than allowed by TDU (which therefore requires the operation of deep mixing), poses crucial constraints to the physical mechanisms driving the transport of processed matter. Contrary to what was found on the RGB, here, thermohaline diffusion cannot be an explanation. Not only is its effectiveness cast in doubt (or at least reduced) by the previous $^3$He consumption, but the layers where it starts are too cool to host any $^{26}$Al. This is evident in Figure 6. On the AGB, the $^3$He+$^4$He reaction prevails over the $^3$He+$^4$He for $\Delta \geq 0.15$, identified in the first panel of the figure by the kink in the $^3$He curve. As the third panel shows, the entire abundance of $^{26}$Al is produced inside that zone. As we showed in Figure 11, accounting for $^{26}$Al in oxide grains requires $\Delta \leq 0.1$. We believe this is a clear demonstration that thermohaline mixing is not the mechanism driving extra mixing in AGB stars. Even its action as a driver of magnetic-like mechanisms (Denissenkov et al. 2009) would not be useful in this case. Completely different physical processes, like, e.g., magnetic buoyancy (Busso et al. 2010) or gravitational waves (Denissenkov & Tout 2003), must necessarily be at play.

Further comparisons with observations are provided by Figure 12. Its top panel shows $C$/$O$ and $^{12}$C/$^{13}$C measurements in both O-rich (Smith & Lambert 1990) and C-rich stars (Abia et al. 2002, 2010). The curves refer to our 1.5 and 2 $M_\odot$ models; the more massive case is the one that reaches $C/O \geq 1$; its C-rich phases are marked by a continuous line, the O-rich ones by a dashed line. We adopted $\Delta = 0.2$ and 0.25 and $M_0$ values as indicated by the labels. It is clear that the entire area of the observations is well accounted for. This figure offers a rather global test from stars, complementing those derived from oxide grains; it confirms that the general scenario is correct and that MS, S, and C stars indeed have the masses we suggested and do experience extra mixing.

A much less satisfactory comparison is presented in the bottom panel of Figure 12, where our model curves are superimposed to the nitrogen and carbon isotopic ratios measured in SiC grains of AGB origin (see, e.g., Zinner 2005; Zinner et al. 2006 and references therein). The models in the figure are labeled by the values of $\Delta$ and $M_0$. They mainly cover the area of the so-called mainstream SiC grains (with carbon isotopic ratios between 20 and 100), but only for the upper part of the range in $^{14}$N/$^{15}$N. Models with rather extreme $M_0$ values can reach the area where the grains with the lowest $^{12}$C/$^{13}$C ratios (called A and B grains) are present, but again only for very high values of $^{14}$N/$^{15}$N. These inconsistencies are not new and have characterized the attempts at explaining the nitrogen isotopes in SiC grains for many years (see, e.g., Nollett et al. 2003). Note that the two filled circles, at $^{14}$N/$^{15}$N = 680 and 5300, refer to the only (very uncertain) data from red giants (Wannier et al. 1991) and confirm that the RGB ratios must indeed be much larger than solar. The low values of many SiC grains have no explanation at all in LM stars. In principle, in the so-called $^{13}$C-pocket of AGB stars a reservoir of $^{13}$N exists. This is generated through proton and neutron captures on $^{14}$N and later, at the onset of the thermal pulse, by the activation of the $^{13}$O(p, $\alpha$) reaction; see also Abia et al. (2010). However, these zones are not expected.

Figure 11. Curves show the oxygen and aluminum isotopic mix in the envelopes of our solar-metallicity stars during advanced evolutionary stages, as compared to presolar oxide grain compositions (WUSTL Presolar Database: http://presolar.wustl.edu/~pgd/). Top panel: FDU is indicated by the heavy, long-dashed curve; short-dashed lines and the shaded area show the effects of deep mixing on the RGB, for $\Delta = 0.2$ and $M_0 = 0.1$. This is a more moderate choice than in Figure 3. The symbols for data points are the same as in Figure 3. Continuous lines refer to model results for extra mixing on the AGB (all models for a specific mass are named (a)–(e) from left to right). They correspond to specific choices of the parameters, namely, (a) $\Delta = 0.1$, $M_0 = 1$; (b) $\Delta = 0.1$, $M_0 = 3$; (c) $\Delta = 0.15$, $M_0 = 1$; (d) $\Delta = 0.15$, $M_0 = 3$; (e) $\Delta = 0.2$, $M_0 = 1$. Models for $M \leq 1.7 M_\odot$ explain essentially all the data for $^{15}$O-poor grains. In particular, group 2 grains mainly derive from very low mass stars (below $1.5 M_\odot$). Bottom panel: the $^{17}$O/$^{16}$O ratio as a function of the $^{26}$Al content. Continuous lines refer to $\Delta = 0.05$; long-dashed lines to $\Delta = 0.1$; short-dashed lines to $\Delta = 0.15$; dotted lines to $\Delta = 0.20$. Model curves proceed from left to right, converging toward the equilibrium $^{17}$O/$^{16}$O at the specific temperature of each model. Most of the data points are accounted for (see the text for the parameter values).

large uncertainties are still present (Iliadis et al. 1996). We found that by artificially increasing their strengths by a factor of five with respect to present recommendations, grains rich in $^{26}$Al
ratios of mainstream grains and of some A+B grains is reproduced, but our
plotted. Again, the model parameters are indicated. The range of carbon isotopic
meteorites, as a function of their $^{12}\text{C}$

e.t al.2006; WUSTL Presolar Database:http://presolar.wustl.edu/

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Contamination of the grains with solar-system nitrogen seems
unlikely, as there is no correlation between the $^{15}\text{N}$ concentration
to be ejected and their $^{15}\text{N}$ is subsequently destroyed in the
convective thermal pulse.

7. TOWARD A PHYSICAL MODEL OF DEEP MIXING: CONSTRANTS ON THE PARAMETERS

As a concluding remark we want to make some comments on the recent numerical simulations of extra mixing presented
by Denissenkov (2010) and by Denissenkov & Merryfield (2011). The work was performed through an analogy with current research on thermohaline processes known to occur in the oceans. The models were then extrapolated to the stellar case, on the basis of the values of physical parameters valid for red giants; this treatment has obviously a much higher degree of sophistication than simple one-dimensional analysis, but still includes very uncertain parameters. Anyhow, the authors suggest that the molecular weight inversion generated by $^3\text{He}$ burning induces finger-like mixing structures whose average aspect ratio (length over diameter, $\alpha = l/a$) is too small by a factor of at least 50 as compared to what observations require; this aspect ratio, squared, is proportional to the diffusion coefficient in a diffusive treatment, or to the mixing velocity in a circulation-like transport (Nollett et al. 2003). In Denissenkov (2010), it is also mentioned that, to explain the small number of red giant stars that are very Li-rich ($\log e(\text{Li}) \gtrsim 1.5$), the aspect ratios found with thermohaline processes would be too small by a factor of several hundreds; in fact, producing $^7\text{Li}$ is known to require a kind of transport fast enough to save to the envelope the short-lived $^7\text{Be}$ ($t_{1/2} = 53$ days). These results would indicate that thermohaline diffusion alone is insufficient to drive the chemical mixing required in red giants; as a way out, in Denissenkov & Merryfield (2011) it is suggested that a coupling with magnetically induced effects might be more promising.

The above results should now be verified by other authors before final conclusions can be drawn. Recent works either attribute the isotopic mix of red giants to the effects of thermohaline mixing alone (Charbonnel & Zahn 2007) or confirm that models for this process find velocities that are too low by a large factor, thus opening the possibility that other mechanisms contribute (Cantiello & Langer 2010).

Although our simple parameterized approach cannot enter into such details, we found the treatment by Denissenkov (2010) and Denissenkov & Merryfield (2011) very instructive as a tool for inferring the required properties of suitable physical models for extra mixing. In particular, it was suggested (Busso et al. 2007b; Denissenkov et al. 2009) that magnetic buoyancy might offer an alternative to thermohaline diffusion, with the advantage that it would not require a nuclear process as a trigger, hence it would have a wider possibility of application (e.g., to the formation of the $^{13}\text{C}$-pocket). If magnetic buoyancy were to proceed through the development of $\Omega$-shaped loops, formed as instabilities in MHD waves (Parker 1994), then each such structure would look like a couple of mixing fingers. Let us make two illustrative examples to derive limits on the aspect ratios of these structures. As a zero-order approximation we can consider the model by Busso et al. (2007b), neglecting any slowing process (such as phase mixing or thermal exchanges, see, e.g., Spruit 1999, 2002; Denissenkov et al. 2009). In this case, the buoyancy velocity is close to the Alfvén velocity and the aspect ratio, for the cases discussed by Busso et al. (2007b),
turns out to be in the range of 200–300, suitable for Li production. Alternatively, a less efficient transport is obtained if the instabilities can travel only a limited distance before interfering destructively (by phase mixing) with previously detached buoyant structures. The length covered is \( l = \frac{\pi \Delta t_A}{t} \), where \( t \) is the available time for the mixing and \( \Delta t_A \) is the crossing time of Alfvén waves over the star’s dimension (Spruit 1999; Busso et al. 2004). In such a case, using the parameters of RGB phases by Busso et al. (2007b), one can easily derive that the aspect ratio becomes very small (of the order of 0.1–0.2), suitable to account for the requirements of the carbon isotopic ratios in red giants and with a correspondingly small velocity. A similarly small velocity is obtained for the magneto-thermohaline mixing by Denissenkov et al. (2009). These are only very simplified, order-of-magnitude estimates, derived just by applying to magnetic buoyancy some of the considerations by Denissenkov (2010). However, they are already sufficient to underline that this kind of mechanism now deserves to be carefully modeled, as it seems to be able to cover the whole range of mixing velocities (or aspect ratios) required by virtually all the chemical peculiarities displayed by red giant stars.

8. CONCLUSIONS

In this paper, we presented revised parametric calculations of nucleosynthesis in extra-mixing phenomena occurring in LM evolved stars, from the phase of the FDU up to the end of the AGB stage. The calculations were based on a recent revision of the FRANEC code and on a new extra-mixing code. They incorporated several physical and numerical improvements, among which new opacities for carbon-rich envelopes, a new algorithm for the determination of the convective border position, and a new mass-loss law.

With these tools we explored the coupling of nucleosynthesis and non-convective mixing in RGB and AGB phases, comparing the results obtained with the old NACRE compilation of reaction rates with those descending from the adoption of new recommendations from ADE10 and ILI10. In particular, the specific choices made for each updated reaction rate were discussed extensively in the Appendix.

It was found that, among the new reaction rates, the ones inducing the largest variations in the evolution of LM stars are those for proton captures on \(^{14}\text{N}\) and (to a lesser extent) for proton captures on \(^{17}\text{O}\). Similar conclusions were recently obtained in another context by Magic et al. (2010). We obtained huge changes in the oxygen isotopic composition of the envelope, starting from FDU. Their effects include a drastic shrinking of the mass interval in which AGB stars can produce oxide grains like those recovered in pristine meteorites. Most such grains appear to come from stars below 1.5 \( M_\odot \). Another important revision of previous knowledge was the discovery that all \(^{18}\text{O}\)-poor grains (those with \(^{18}\text{O}/^{16}\text{O} < 0.0015\)) can now be explained, without the “forbidden regions” previously found for the lowest \(^{17}\text{O}/^{16}\text{O}\) ratios. Concerning the evolution of AGB stars of LM and solar metallicity, we suggested that most MS and S stars should have masses below 1.5–1.7 \( M_\odot \); in this mass interval they would never produce C-stars, also thanks to \(^{12}\text{C}\) processing in deep mixing.

Constraints coming from observations of CNO nuclei in both RGB and AGB stars were well reproduced by deep-mixing models with the new rates. In particular, for RGB stars we found that extra-mixing processing does not need to penetrate to very high temperatures, so that thermohaline diffusion might be an adequate scenario for those phases, if the problems recently mentioned in the literature for its too-low diffusion coefficients can be solved. We then confirmed that deep mixing is also required on the AGB. In this case, the abundances of \(^{26}\text{Al}\) in presolar grains indicate that the transport must reach much hotter layers than in RGB stages. This fact (and the previous consumption of \(^{3}\text{He}\)) induced us to conclude that thermonuclear diffusion is not a viable explanation for the transport processes occurring in the latest evolutionary stages. It was suggested that the existence of a small population of oxide grains with low \(^{17}\text{O}/^{16}\text{O}\) and high \(^{26}\text{Al}/^{27}\text{Al}\) ratios require some further revision of the rates for \(^{26}\text{Al}\) synthesis at low temperatures, probably an increase for the \(^{25}\text{Mg}(p, \gamma)^{26}\text{Al}\) rate.

The longstanding impossibility of accounting for the whole range of the nitrogen isotopic ratios in SiC grains remains unchanged after our work. It was argued that no stellar process occurring in LM stars can solve this problem and that SiC grains might be contaminated by cosmic ray spallation processes in the ISM.

We are indebted to many colleagues for helping us in understanding the subtleties of LM star nucleosynthesis. While recognizing the role of each of them is impossible, we cannot avoid acknowledging the fundamental lessons we learned from friends such as G. J. Wasserburg, R. Gallino, O. Straniero, U. Ott, and K. M. Nollett. M.L. acknowledges the support of the Italian Ministry of Education, University and Research under grant No. RBFR082838 (FIRB2008)

APPENDIX

THE CHOICE OF NUCLEAR REACTION RATES

A.1. The Reactions \(^{14}\text{N}(p, \gamma)^{15}\text{O}\) and \(^{15}\text{N}(p, \gamma)^{16}\text{O}\)

The \(^{14}\text{N}(p, \gamma)^{15}\text{O}\) reaction is the slowest reaction of the CN cycle, and hence the one determining the duration of the process. The low-energy cross section is dominated by a resonance at 259 keV. In particular, at the temperatures of interest here (\( T_\text{eff} \lesssim 0.06 \)), the reaction rate mainly depends on the resonance tail, the Gamow window extending (at most) to around 110 keV. Such a region is only partly covered by experimental data, spanning energies as low as 70 keV, so the \( S\)-factor is very sensitive to the extrapolating curve and can be severely affected by the \(-504\) keV subthreshold resonance. In the last years, direct measurements (in both surface and underground laboratories) and indirect measurements (Doppler shift attenuation, Coulomb excitation, and asymptotic normalization constant) have been performed, primarily triggered by the discovery that the contribution of the ground-state transition had been previously overestimated. From these measurements, ADE10 recommend \( S(0) = 1.66 \pm 0.12\) keV b, which is about half the NACRE value \( S(0) = 3.2 \pm 0.8\) keV b. This entails a significantly lower reaction rate in the range of our interest, as shown in Figure 13, left panel.

The \(^{15}\text{N}(p, \gamma)^{16}\text{O}\) reaction fixes the leak rate from the CN-cycle to the CNO bi-cycle. Besides the direct capture contribution, the cross section is characterized by the interference of two \( 1^-\) resonances at 312 and 964 keV. As for \(^{14}\text{N}(p, \gamma)^{15}\text{O}\), the largest energy of interest is \( \sim 110 \) keV, thus the extrapolation procedure is critical. In NACRE, a very large \( S(0) \) was obtained (\( 64\pm6\) keV b) while ADE10 recommend \( 36\pm6\) keV b following the results from an underground measurement (which does not cover the Gamow window anyway) and an asymptotic normalization coefficient measurement, yielding an absolute value of
the direct-capture contribution 10 times smaller than NACRE. Consequently, a 30%–40% smaller reaction rate is now obtained in the temperature region we are dealing with (Figure 13, right panel) by using the low-energy $S(E)$-factor recommended by ADE10.

A.2. The Reactions $^{16}\text{O}(p, \gamma)^{17}\text{F}$, $^{17}\text{O}(p, \gamma)^{18}\text{F}$, and $^{17}\text{O}(p, \alpha)^{14}\text{N}$

The operation of the $^{16}\text{O}(p, \gamma)^{17}\text{F}$ reaction contributes to the establishment of the ratios of oxygen isotopes in red giants and therefore plays a major role in the analysis of oxygen-rich presolar grains of circumstellar origin. At the energies of interest in this work, and up to $\sim 120$ keV, the cross section is determined by the direct capture to the ground and first excited states of $^{17}\text{F}$. Direct measurements exist down to about 100 keV, so extrapolations are needed. In the previous evaluation by NACRE, the greatest part of the error was attributed to the model assumed in the calculation. Improved calculations have now reduced the uncertainty affecting the $S(0)$ parameter, from 30% to 7.5% (see ADE10 for details), while the recommended value is in substantial agreement with the one from NACRE. Figure 14 (left panel) shows the resulting reaction rate.

Much like the previously mentioned reaction, $^{17}\text{O}(p, \gamma)^{18}\text{F}$ is also expected to modify the relative abundances of oxygen isotopes. In the $E_{\text{cm}} \lesssim 120$ keV energy range, the cross section shows a resonance at 65.1 keV that strongly influences the reaction rate above $T_9 = 0.02$. At lower temperatures, the dominant role is played by direct captures, while the subthreshold resonance at $-3.1$ keV does not provide any significant contribution. The 65.1 keV resonance and the direct capture term have been the subject of several improved calculations, leading to a change in the resonance energy (it was 66 keV in NACRE) and in its strength (which is a factor $\sim 3.7$ smaller than given in NACRE). Concerning the direct-capture contributions, the revised calculations yield a $\sim 30\%$ smaller $S(E)$ factor below 120 keV, which is responsible for the reduced reaction rate below $T_9 = 0.02$, as shown in Figure 14, central panel.

The $^{17}\text{O}(p, \alpha)^{14}\text{N}$ channel competes with the $^{17}\text{O}(p, \gamma)^{18}\text{F}$ channel, removing $^{17}\text{O}$ nuclei and turning the nucleosynthesis path back toward nitrogen. In contrast with the radiative capture
channel, the $-3.1$ keV subthreshold resonance gives a major contribution to the reaction rate for $T_9 < 0.02$. Updated calculations for these resonances are given in both the reviews we are considering. In particular, the $-3.1$ keV state proton width is lowered by a factor of $\sim 3$ as compared to NACRE. No expression for the reaction rate is given in ADE10; hence we use the reaction rate table in ILI10, extrapolating down to the $T_9 = 0.004-0.01$ temperature range. A remarkably lower reaction rate is thus obtained (see Figure 14, right panel). Regarding the $^{18}$O$(p, \alpha)^{18}$N reaction, despite the extensive experimental investigation in the past years (ADE10, ILI10, La Cognata et al. 2008a, 2008b) that has produced a significant change in the recommended reaction rate (about 30%) and a reduction of its uncertainty (about a factor of eight) at low temperatures, its effect on AGB nucleosynthesis turns out to be negligible. The updated reaction rate has been included anyway in the present work.

A.3. The $^{20}$Ne$(p, \gamma)^{21}$Na, $^{21}$Ne$(p, \gamma)^{22}$Na, and $^{22}$Ne$(p, \gamma)^{23}$Na Reactions

The $^{20}$Ne$(p, \gamma)^{21}$Na reaction is the starting point of the NeNa cycle and plays a key role, being the slowest reaction of the cycle. Inside the Gamow window, extending up to $\sim 130$ keV, the astrophysical factor is dominated by a subthreshold resonance at about $-7$ keV. An additional eight resonances show up at higher energies (up to about 2 MeV). In ADE10, both the resonance excitation energy and its strength have been re-evaluated using more recent measurements, resulting in a small increase of the $S(E)$ factor, by 2.9%. Moreover, a new fit of the energy trend for the $S(E)$ factor has been provided for the numerical integration of the reaction rate. The resulting rate is displayed in Figure 15, left panel, which shows a substantial agreement with the NACRE one and a slightly reduced uncertainty.

For the $^{22}$Ne$(p, \gamma)^{23}$Na reaction, we adopt the recommendation by ILI10. They make use of nuclear-physics input parameters from Iliadis et al. (2001). Resonances were measured for $E_{cm} \geq 121$ keV, corresponding to the upper edge of the Gamow window. Two threshold states, corresponding to resonance energies of $E_{cm} = 17$ and 95 keV, have been considered. While the latter (for which only an upper limit of the spectroscopic factor is known) adds a negligible uncertainty to the reaction rate, the former can influence it at low temperatures (below $T_9 \sim 0.02$).

Though the reaction-rate accuracy has been improved with respect to NACRE at around $T_9 \sim 0.03$, for $T_9 \lesssim 0.02$ the total rate remains uncertain by several orders of magnitude (see Figure 15, central panel).

The rate of the $^{22}$Ne$(p, \gamma)^{23}$Na reaction is taken from ADE10 and is shown in Figure 15, right panel. The main difference with respect to previous compilations is the treatment of the threshold states. Concerning the 151 keV resonance, its spectroscopic factor has been considered as a mean value rather than an upper limit. Lower energy resonances, whose existence had been considered as tentative, were disregarded by ADE10. As a result, the recommended rate is smaller than in NACRE above $T_9 \sim 0.03$, but remains in good agreement with the previous indications elsewhere. The error has been instead significantly reduced.

A.4. The $^{23}$Na$(p, \gamma)^{24}$Mg and $^{23}$Na$(p, \alpha)^{20}$Ne Reactions

The Gamow window of the $^{23}$Na$(p, \gamma)^{24}$Mg reaction extends up to about 140 keV. Therefore, the main contribution to the low-temperature reaction rate is due to the threshold states. ILI10 mostly make use of the updated nuclear physics input parameters by Hale et al. (2004) in this energy region, which have been indirectly measured by means of $(^3$He, $d)$ transfer reactions. The contribution of direct capture to the cross section has been taken from the same work. The resulting reaction rate is compared with the NACRE choice in Figure 16 (left panel). The updated value lays well within the upper and lower limits set by NACRE, although the uncertainty is greatly reduced.

Concerning the $^{23}$Na$(p, \alpha)^{24}$Mg reaction, the same considerations as for $^{21}$Na$(p, \gamma)^{20}$Ne can be repeated. The resulting reaction rate is given in Figure 16, right panel. The new rate is about 100 times smaller than the one recommended by NACRE at the lowest temperatures, while the uncertainty is significantly smaller.

A.5. The Reactions $^{24}$Mg$(p, \gamma)^{25}$Al, $^{25}$Mg$(p, \gamma)^{26}$Al$_{gs}$, $^{25}$Mg$(p, \gamma)^{26}$Al$_m$, and $^{26}$Mg$(p, \gamma)^{27}$Al

The $^{24}$Mg$(p, \gamma)^{25}$Al reaction influences the nucleosynthesis of magnesium and aluminum, which are key elements in the analysis of ancient meteorites and presolar grains. No experimental data are available inside the Gamow window (which extends up to $E_{cm} \sim 150$ keV). The most recent measurements provide the cross section down to 200 keV (see ILI10, and reference therein), where nine resonances occur (up to 2.3 MeV). In the review by ILI10, the reaction...
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Figure 16. Same comparison as in Figure 13, for the reactions $^{23}\text{Na}(p,\gamma)^{24}\text{Mg}$ and $^{23}\text{Na}(p,\alpha)^{20}\text{Ne}$. (A color version of this figure is available in the online journal.)

Figure 17. Same comparison as in Figure 13, for the reactions $^{24}\text{Mg}(p,\gamma)^{25}\text{Al}$ and $^{25}\text{Mg}(p,\gamma)^{26}\text{Al}_{gs}$. (A color version of this figure is available in the online journal.)

The reaction rate is calculated from these data, including re-normalizations of the strengths of higher-energy resonances and updated $Q$-values. Moreover, with respect to the NACRE compilation, a more reliable direct capture contribution to the $S(E)$ factor has been estimated, as well as the contribution of the 214 and 402 keV resonance tails, by means of numerical integrations. Despite all these improvements, the resulting reaction rate (Figure 17, left panel) is in fair agreement overall with the NACRE recommendations. It is shown for $T_9 > 0.01$, but the reaction rate there is $\sim 10^{-35}$ cm$^3$ mol$^{-1}$ s$^{-1}$, so that the $^{24}\text{Mg}(p,\gamma)^{25}\text{Al}$ reaction does not contribute to nucleosynthesis at very low energies.

As for the previous case, the $^{25}\text{Mg}(p,\gamma)^{26}\text{Al}_{gs}$ reaction (where “gs” stands for “ground state”) is important for us to understand the magnesium and aluminum nucleosynthesis in evolved red giants. Data exist in a wide energy range, starting from 37 keV and extensively overlapping with the Gamow peak. Concerning the nuclear physics inputs, the differences of the ILI10 review with respect to NACRE are (1) the use of re-normalized strengths for resonances above $E_{cm} = 189$ keV; (2) the adoption of the more accurate ground-state $\gamma$-ray branching ratios $f_0$ for the 189, 244, and 292 keV resonances; and (3) a new calculation of the contribution of the 37 keV resonance. The reaction rate is displayed in Figure 17, right panel: the recommended value is in good agreement with NACRE, but the uncertainty has been reduced by more than a factor of two at the lowest temperatures that are most important for us.

The same considerations done above apply to the $^{25}\text{Mg}(p,\gamma)^{26}\text{Al}_{m}$ reaction (where “$m$” indicates the isomeric meta-stable state), but in addition the isomeric-state $\gamma$-ray branching ratio $1 - f_0$ has been taken into account. Once again, while the recommended value by ILI10 is in overall agreement with NACRE, the uncertainties are reduced by more than a factor of two at the temperatures of AGB stars (Figure 18, left panel).

For the $^{26}\text{Mg}(p,\gamma)^{27}\text{Al}$ reaction, Figure 18 (right panel) shows a remarkable difference between the updated reaction rate by ILI10 and the NACRE value, ranging from a factor of two up to eight, while the error has been reduced, especially at high temperatures. Indeed, a different direct-capture component was adopted, the strengths of the measured resonances (above $E_{cm} \sim 100$ keV) were re-normalized, and the contributions of the lower-energy resonances at $E_{cm} = 53$ keV and $E_{cm} = 105$ keV were re-analyzed. One has to remark that some
A.6. The Reactions $^{26}\text{Al}(p,\gamma)^{27}\text{Si}$, $^{27}\text{Al}(p,\gamma)^{28}\text{Si}$, and $^{27}\text{Al}(p,\alpha)^{24}\text{Mg}$

The $^{26}\text{Al}(p,\gamma)^{27}\text{Si}$ reaction is important for establishing the equilibrium abundance of the radioactive isotope $^{26}\text{Al}$, playing a major role in presolar grain studies, γ-ray astronomy, and solar system formation (see, e.g., Busso 2010; Busso et al. 2003; Wasserburg et al. 2006 for this last point). Figure 19 (left panel) shows that the updated reaction rate by Iliadis et al. (2001) lies within the fiducial interval by NACRE, but at low temperatures, the recommended value is strongly decreased. Also, the uncertainty has been greatly reduced. In the estimate of the reaction rate, Iliadis et al. (2001) introduced a calculated direct-capture component and the contribution of 19 resonances. In particular, for almost all resonances below $E_{cm} \sim 500$ keV, the resonance energies were computed from updated excitation-energy determinations, while improved calculations of the proton widths of unobserved low-energy resonances were used to evaluate their contribution to the rate.

For the $^{27}\text{Al}(p,\gamma)^{28}\text{Si}$ reaction, Iliadis et al. (2001) made use of the nuclear physics input parameters reviewed in Iliadis et al. (2001). With respect to the NACRE compilation, the direct-capture contribution was recalculated, the strengths of the measured resonances were re-normalized (above $E_{cm} \sim 200$ keV, to be compared with the upper limit of the Gamow window, $\sim 160$ keV), and the contributions of four sub-threshold resonances were re-evaluated (the most important are at $E_{cm} = 72$ keV and $E_{cm} = 85$ keV). In particular, an improved determination of their spectroscopic factors was performed, based on the re-analysis of the $^{27}\text{Al}(^{3}\text{He},d)^{28}\text{Si}$ stripping data. Consequently, the recommended total rate deviates from NACRE by up to a factor of 20 in the energy range common to AGB stars (Figure 19, central panel), mainly because of improved estimates of spectroscopic factors for the threshold states.

A total of 91 resonances at energies of $E_{cm} = 72$–2966 keV were taken into account by Iliadis et al. (2001) for calculating the reaction rate for the $^{27}\text{Al}(p,\alpha)^{24}\text{Mg}$ process. For the measured resonances (for $E_{cm} \gtrsim 486$ keV), the same nuclear physics input parameters considered in Iliadis et al. (2001) were adopted. The same holds for threshold resonances, whose strengths were calculated from a re-analysis of the $^{27}\text{Al}(^{3}\text{He},d)^{28}\text{Si}$ stripping cross section and from spectroscopic data in the literature (see Iliadis et al., and references therein). The improved estimates of spectroscopic factors for the threshold states, as in the $^{27}\text{Al}(p,\gamma)^{28}\text{Si}$ case, determine an increase of the recommended total rate with respect to NACRE by a factor of 10 (Figure 19, right panel).
Finally, we recall that $(p, \gamma)$ and $(p, \alpha)$ captures on fluorine have not been addressed by either ADE10 or ILL10. However, recent measurements (Couture et al. 2008; Spyrou et al. 2000) only confirmed previous suggestions, so that the rates do not require updates, and we adopt them from NACRE.

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