

Astromers in the radioactive decay of r -process nucleiG. WENDELL MISCH ^{1,2,3} T. M. SPROUSE ¹ AND M. R. MUMPOWER ^{1,2,3}¹*Theoretical Division, Los Alamos National Laboratory, Los Alamos, NM, 87545, USA*²*Center for Theoretical Astrophysics, Los Alamos National Laboratory, Los Alamos, NM, 87545, USA*³*Joint Institute for Nuclear Astrophysics - Center for the Evolution of the Elements, USA*

ABSTRACT

We study the impact of astrophysically relevant nuclear isomers (astromers) in the context of the rapid neutron capture process (r -process) nucleosynthesis. We compute thermally mediated transition rates between long-lived isomers and the corresponding ground states in neutron-rich nuclei. We calculate the temperature-dependent β -decay feeding factors which represent the fraction of material going to each of the isomer and ground state daughter species from the β -decay parent species. We simulate nucleosynthesis by including as separate species nuclear excited states with measured terrestrial half-lives greater than 100 μ s. We find a variety of isomers throughout the chart of nuclides are populated, and we identify those most likely to be influential. We comment on the capacity of isomer production to alter radioactive heating in an r -process environment.

Keywords: r -process — astromers — isomers — nucleosynthesis

1. INTRODUCTION

The existence of long-lived excited (metastable) states of atomic nuclei, known as nuclear isomers, was proposed by Soddy (1917) and verified by Hahn (1921) a century ago (Walker & Podolyák 2020). The study of nuclear isomers has since been a focus of experimental efforts throughout the chart of nuclides, often requiring challenging measurements (Raut et al. 2013; Watanabe et al. 2014; Simpson et al. 2014; Patel et al. 2014; Svirikhin et al. 2017; Andreev et al. 2019; Sikorsky et al. 2020). Over 700 nuclei have been identified as possessing isomeric states with half-lives greater than 100 μ s (Brown et al. 2018). Theoretical nuclear structure methods (Brown & Rae 2014) are able to characterize some of these isomers, which can arise due to large spin differences (spin traps), differences in nuclear deformation (shape coexistence/shape isomers), and large differences in the projection of spin along the axis of symmetry in a deformed nucleus (K isomers) (Dracoulis et al. 2016).

Despite the experimental and theoretical progress in terrestrial studies, much remains uncertain regarding the population of nuclear isomers in astrophysical en-

vironments (Arahamian & Sun 2005; Hayakawa et al. 2005, 2009). The most well studied cases in nuclear astrophysics are in relatively light nuclei. For instance, ²⁶Al can be used as a tracer of star formation due to the long half-life of the ground state (Mahoney et al. 1982; Diehl et al. 1995), but its in-situ production is complicated by a low-lying isomer (Gupta & Meyer 2001; Runkle et al. 2001; Banerjee et al. 2018). The production of ³⁴Cl may be observable immediately after a nova (Coc et al. 1999), while ⁸⁵Kr is a branch point in the slow neutron capture (s) process (Abia et al. 2001) with implications for cosmochronometry (Ward 1977); both of these nuclei are influenced by isomers.

In stark contrast, while mechanisms for inclusion of these additional states in nucleosynthesis codes exist (Reifarth et al. 2018), the population of astrophysically relevant nuclear isomers (astromers) is a missing component in the simulation of r -process nucleosynthesis. Exploration of astromers should impact searches for the most influential nuclei to measure at radioactive beam facilities (Mumpower et al. 2016; Horowitz et al. 2019), and it overlaps with modern multi-messenger astronomy (Abbott et al. 2017; Tanvir et al. 2017; Troja et al. 2017; Wang et al. 2020).

In environments where the r process may operate, nuclear isomers can be populated thermally then frozen out as the temperature drops rapidly. They can also

be fed by β -decay of more neutron-rich isotopes, as well as through other nuclear reactions such as neutron capture and fission (Wisshak et al. 2006; Okumura et al. 2018). The direct population of nuclear isomers has recently been shown to be potentially influential in the radioactive heating of r -process events by Fujimoto & Hashimoto (2020). They compared a no-isomer simulation against two which included several hand-picked isomers; both isomer models essentially replaced the ground-state properties of the selected nuclei with the isomer properties.

In this work, we explore for the first time dynamical freeze-out as well as the thermal and β -decay population mechanisms of nuclear isomers in the r process. We demonstrate that a range of nuclear isomers are significantly populated in the r process between first peak (mass number, $A \sim 80$) and third ($A \sim 195$) peak elements. We introduce an Astromer Importance Rating (AIR) to identify astromers among our included isomers, and we show the effects of our careful treatment of known nuclear isomers on an r -process heating curve. By dynamically including both nuclear isomers and their corresponding ground states, we account for a previously unaddressed non-equilibrium process in simulations of heavy element nucleosynthesis.

2. NUCLEAR ISOMERS & NUCLEOSYNTHESIS

We treat isomers with half-lives greater than 100 μ s and ground states as distinct “long-lived” species. We use the method of Misch et al. (2020) to calculate the thermally mediated transition rates between long-lived states via intermediate states. This method assumes the states communicate through a thermal photon bath and does not rely on any approximations for the solution to the linear system of equations. The temperature-dependent β -decay rates for these long-lived states is computed using the ensemble formalism of Gupta & Meyer (2001).

To account for the feeding of β -decay daughter nuclei with isomers, we combine laboratory β -decay $\log(ft)$ values to specific daughter states with that state’s probability to contribute to each daughter ensemble; we use available laboratory β -decay rates and measured β intensities to obtain the β feeding when $\log(ft)$ values are unavailable. This yields temperature-dependent decay rates into each of the daughter “species.” When β intensities are unavailable in the literature, we assume that all decays go to the daughter ground state.

Our isomer calculations use experimental level energies, half-lives, spin-parity assignments, γ intensities,

and $\log(ft)$ values/ β intensities from ENSDF¹. We include all experimental levels if there are fewer than 30 reported; if there are more than 30 levels, we include all levels up to the 10 levels above the highest-lying isomer (minimum 30); the analysis of Coc et al. (1999) shows that this selection is more than adequate to accurately compute transition rates. For unmeasured γ transitions, we use the Weisskopf approximation (Weisskopf & Wigner 1930).

We simulate nucleosynthesis with the Jade nuclear reaction network of Sprouse et al. (2020). Jade uses a matrix exponential solver that enables the inclusion of temperature-dependent effects on nuclear transmutation rates. This allows us to dynamically track the population and de-population of the long-lived states of individual isotopes as a function of time. For nuclei with isomers, we use the aforementioned treatment. Otherwise, we take laboratory ground-state-to-ground-state half-lives from evaluated data² (Brown et al. 2018; Audi et al. 2017) and supplement with theoretical predictions for yet-to-be-measured nuclei (Möller et al. 2019).

A species in our nucleosynthesis network is referenced by two indices: 1) i maps between a tuple consisting of proton number Z and neutron number N , and 2) j indicates the individual long-lived state of the nucleus. For the sake of this work, we use g and m as values of j to denote the ground state and isomer, respectively. Our network explicitly includes all isomeric states of each isotope, but for clarity, our arguments are presented as though each isotope has at most one isomer.

We focus this work on the population of nuclear isomers in the r process after ~ 15 minutes. In this epoch, all relevant neutron captures have finished, and β -decay, thermally induced excitation, and radiative/thermally stimulated de-excitation are the primary nuclear reaction channels. We assume an initial solar-like distribution of r -process material (Arnould et al. 2007) situated far from stability; the distribution consists of all three major r -process abundance peaks (Sprouse et al. 2020) as motivated by recent observations (Abbott et al. 2017; Watson et al. 2019). We have studied the production of nuclear isomers in a variety of astrophysical conditions; here we take the temperature evolution used in Zhu et al. (2018), which is based on a homologous expansion into free space (Lippuner & Roberts 2015). We stop our calculations after $\sim 3,600$ days (10 years).

¹ From ENSDF database as of June 29th, 2020. Version available at <http://www.nndc.bnl.gov/ensarchivals/>

² <https://www.nds.iaea.org/public/download-endf/ENDF-B-VIII.0/>

3. AIR: ASTROMER IMPORTANCE RATING

We define the population ratio $R_{i,m}$ of the isomer species of isotope i as

$$R_{i,m} = \frac{Y_{i,m}}{\sum_j Y_{i,j}}. \quad (1)$$

The abundance $Y_{i,j}$ is the number of isotope i in state j per baryon of material. Thus $R_{i,m}$ is the ratio of the isomer abundance to the total abundance of the isotope. For the sake of brevity and legibility, we suppress the indices on R from here on.

By finding the maximum value of R over time for each isomer, we can identify which are populated in a nucleosynthesis event. Figure 1 shows this value for the 36 isotopes which at some time in our simulation had $R \geq 0.1$. Many of the most populated isomers lie in one of the three *r*-process abundance peaks located at $A \sim 80$, $A \sim 130$, and $A \sim 195$. The isomers in these three peaks may be especially significant to the *r* process, as they not only represent a significant fraction of their respective isotopes, but also these isotopes are among the most abundantly populated during an *r* process event.

However, not all highly populated isomers are astromers. They may not freeze out before the isotope has decayed away, or they may not decay at a rate appreciably different from the ground state. And even if an isomer is an astromer, the nuclear isotope may not be sufficiently abundant to influence *r*-process evolution. Therefore, we develop a metric to assist in identifying important astromers.

A species affects evolution through its destruction (decay or de-excitation). For example, consider an abundant ground state of a stable isotope in a cold environment. This species does not participate in the evolution since it neither affects heating nor drives change in abundance. Similarly, a rapidly decaying isotope with low abundance will not affect the environment. We calculate the specific activity a of each species to measure its influence.

$$a_{i,j} = \lambda_{i,j} Y_{i,j} \quad (2)$$

$$\lambda_{i,j} \equiv \sum_{\text{channels}} \lambda_{i,j}^{\text{channel}} \quad (3)$$

The λ s are destruction rates (s^{-1}), and the indices i and j are as in equation 1; we will suppress the a indices i and j going forward, taking j to always be the isomer. In this work, our channels consist exclusively of β decays and thermally-mediated internal transitions. Activity then quantifies the number of actions taken by an isotope (transitions and/or decays) per baryon of material per

second. The most active isotopes at a given time drive the evolution, provided that that activity is not simply internal transitions back and forth between the ground state and isomer. This brings us to the rate imbalance.

To be an astromer, we also require that the isomer m be destroyed at a rate different from the ground state g (the $\lambda_{i,j}$ in Equation 3); otherwise, it does not meaningfully change the evolution as nucleosynthesis proceeds (compared to g only). This is especially true at high temperature, where most of the activity could be due to rapid back-and-forth internal transitions. To quantify how λ_g and λ_m are different, we use (the absolute value of) their imbalance I .

$$I(\lambda_g, \lambda_m) = \frac{|\lambda_g - \lambda_m|}{\lambda_g + \lambda_m} \quad (4)$$

If the destruction rates are equal, the imbalance is zero, and as the rates diverge, the imbalance tends toward unity. Imbalance has the helpful feature of being independent of the scale of the rates without going to infinity the way a ratio of the two quantities might; this second point is important when e.g. the ground state is stable. Furthermore, unlike a ratio, the imbalance will be large for very different rates regardless of which rate is greater.

Taken as a product, the activity a , imbalance I , and population ratio R give a useful metric—the Astromer Importance Rating AIR—to help identify populated and influential astromers.

$$\begin{aligned} \text{AIR} &= \text{Activity} \times \text{Imbalance} \times \text{population Ratio} \\ &= a \times I \times R \end{aligned} \quad (5)$$

AIR selects isomers which 1) have a high activity, 2) behave distinctly from the ground state, and 3) have a significant population relative to ground. These are the necessary ingredients for an influential astromer. Note that it is possible for a slow-decaying astromer to effectively reduce a . However, at later times, everything with a high a will have decayed away, leaving the slower astromer to dominate the AIR then. We search AIR over a broad range of times to maximize our ability to find important astromers.

4. RESULTS

We computed the AIR for every nuclear isotope in our simulation to isolate influential astromers. Figure 2 shows the AIR for the 24 isotopes which, at some simulation timestep, are among the top 5 most important astromers as ranked by AIR.

Figure 2 shows that the most dominant first peak astromers (blue) are relevant for about 1 day with only

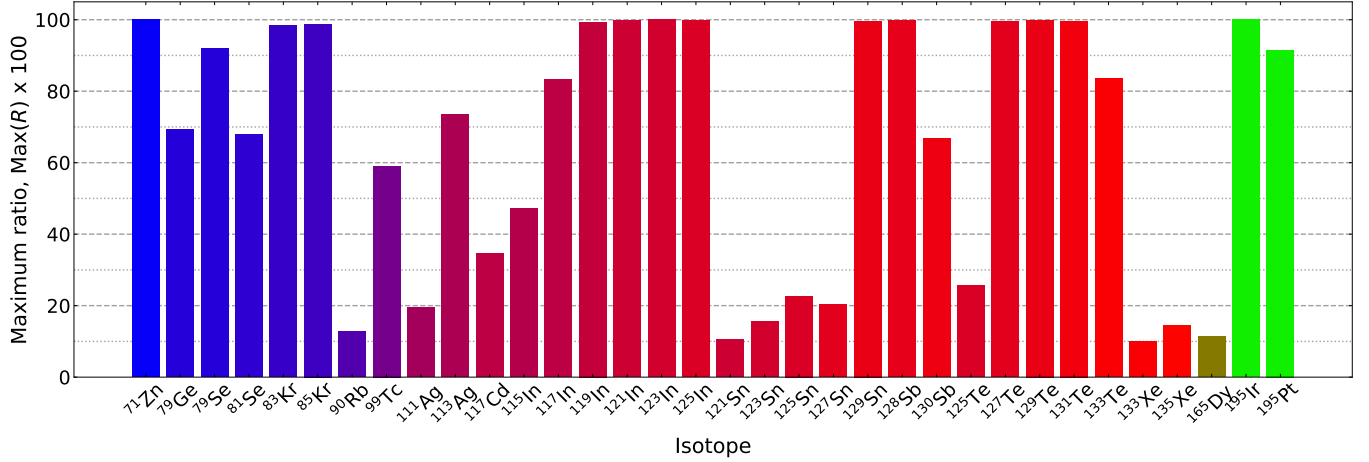


Figure 1. The maximum isomer population ratio, $\text{Max}(R)$, over the course of our simulation for isomers populated above 10%. Color indicates approximately the r -process abundance peak the nucleus resides in; see figure 2.

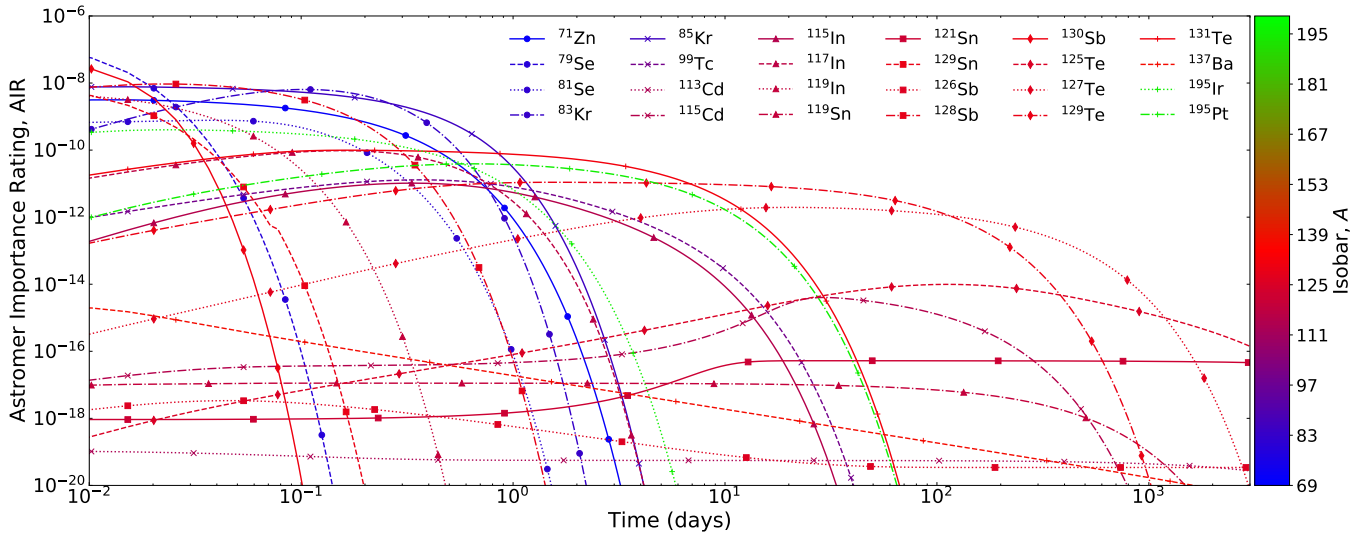


Figure 2. Influential r -process astromers ranked by their Astromer Importance Rating, AIR. We show isotopes with AIR values that appear among the top five at some point in our simulation. Color indicates approximately the r -process abundance peak the nucleus resides in: 1st (blue), 2nd (red), or 3rd (green).

^{95}Nb contributing after 100 days. A surprising contender on the timescale of a day is ^{85}Kr , which is most famous as an s -process branch point. In the r -process, it has the highest AIR from ~ 0.5 to 1 day due to the $> 90\%$ feeding of the isomer by ^{85}Br . This greatly accelerates the decay to ^{85}Rb , and it may produce an electromagnetic signal.

In contrast with the first peak astromers, those located in the second peak (red) persist over much of our simulation, ranging from 0.01 to 10,000 days. As a result, they have the potential to affect many aspects of the evolution throughout a range of observational timescales. At the earliest times in our simulation, around 0.01 days, we note the formation and large AIR of $^{128,130}\text{Sb}$ and ^{131}Te ,

which contribute alongside a number of additional isomers located in the first and third peaks. By 30 days, all remaining isomers are located among the second peak nuclei, the most notable of which include $^{125,127}\text{Te}$. Finally, we identify the particularly interesting isomer in ^{129}Te . It remains important across all epochs of our simulation, from 0.01 days to 1 year, reflecting a complex interplay between the β -decay feeding parent, thermal excitation, and de-excitation.

We identify ^{195}Pt as the dominant astromer located in the third r process peak (green). In contrast with the conventional assumption that this isotope is stable upon population via the ground state β decay of ^{195}Ir , our calculations show that the isomer of ^{195}Pt may be

populated directly from the β decay of the isomer of ^{195}Ir . The ^{195}Pt isomer de-excites on a timescale of ~ 4 days, which is relevant to the study of electromagnetic signals associated with the *r* process.

Notably, there are species in Figure 2 that do *not* appear in Figure 1. This is because while the isomer might not be heavily populated (low R), the isotope might have comparatively high activity a and a large destruction rate imbalance I . The isotopes with a maximum R less than 10% but a high AIR include ^{95}Nb , ^{113}Cd , ^{115}Cd , ^{119}Sn , ^{126}Sb , and ^{144}Pr .

In the interest of readability, Figure 2 shows only those astromers which rank in the top 5 by AIR. When we expand our enumeration of *r*-process astromers to include the top 10, we identify only 12 additional astromers. This highlights the effectiveness of AIR as a filter for distinguishing astromers from the broader population of isomers: the list of isomers identified as *astromers* is relatively insensitive to the particular choice of AIR top- N . We summarize our expanded (top 10) list of astromers in Table 1.

The astromers identified in Table 1 that change β -decay rates will affect *r*-process heating. Some (e.g. ^{115}Cd , $^{127,129,131}\text{Te}$) will defer heating by delaying decay, while others (e.g. ^{85}Kr , $^{128,130}\text{Sb}$) partially counteract this trend by accelerating decay.

Furthermore, astromers may generate identifiable x-ray or γ -ray signals either directly through de-excitation (^{85}Kr , ^{115}In , ^{119}Sn , $^{131,133}\text{Xe}$, ^{195}Pt) or indirectly subsequent to β decay (any with $T_{1/2} \gtrsim 1$ day). Such isomers could be used to directly associate future *r* process observations with the production of specific nuclei.

We emphasize two points about the relationship between available nuclear data and our astromer calculations. First, more complete excited-state data would improve our ground state \leftrightarrow isomer transition rates. Specifically, measurements of intermediate state half-lives and γ intensities enable more reliable calculations of thermally mediated transition rates. This is exemplified by ^{128}Sb , whose isomer dramatically accelerates the decay of the $A = 128$ mass chain. All measured excited states in this nucleus connect to the isomer—which itself has an unknown energy—and the missing links to ground suppress our calculated transition rate.

Second, we need more complete β intensities to compute reliable feeding factors. Because we assume that decays with unpublished intensities always go to ground in the daughter, we preclude the possibility that such decays populate a daughter isomer. The production of ^{170}Ho is one example of this issue: the β intensities of ^{170}Dy are unknown, leading to our inability to effectively calculate the feeding factors into the ^{170}Ho isomer

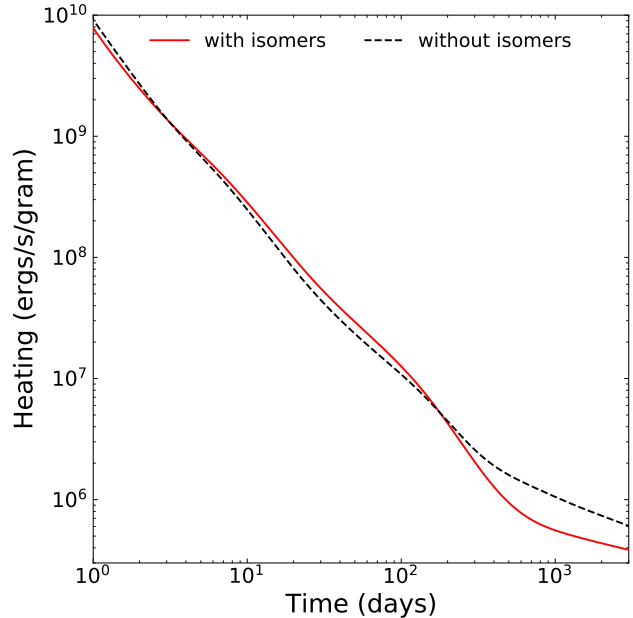


Figure 3. Radioactive heating from β -decay with and without isomers. We find a 25% increase in heating from the inclusion of isomers between three and one hundred days.

(Misch et al. 2020). The $A = 115$ and $A = 129$ mass chains are particularly striking examples of missing β -decay information. While the feeding factors from the ^{115}Cd and ^{129}Sn parents are measured, we lack data on how multiple earlier ancestors populate isomers in those parents. Experimental and theoretical efforts to quantify unknown β intensities would improve understanding of the population of astromers in *r*-process events.

One of the key roles that isomers have recently been proposed to play is in adjusting the timescale on which nuclear energy is released during the radioactive decay towards stability immediately following *r*-process nucleosynthesis (Fujimoto & Hashimoto 2020). To investigate the possibility of these effects, we performed calculations for the radioactive heating of *r*-process material both with and without the consideration of isomers. We show our calculations of these two heating rates as a function of time in Figure 3.

Within the first day, astromers effectively store energy by slowing radioactive decay towards stability as a result of their (on average) comparatively longer half-lives. On longer timescales (10-100 days), the slower decay of these astromers leads to a gradual release of the energy they had previously retained, with the effect of boosting the heating by about 25%. Analogous cycles of energy retention and release by astromers tend to repeat on longer timescales, the next of which begins around 200 days. We find the inclusion of temperature-dependent

Table 1. Astromer importance rating (AIR) top 10 selected at each time point $t \gtrsim 15$ min. We use g and m to indicate the ground state and isomer, respectively. The isomer energy is E_m , and the J^π are the spin and parity of the respective levels (parentheses denote uncertain J^π). The half-lives ($T_{1/2}$) and β -decay branching for the isomer ($B_{m\beta}$) are as measured in the laboratory; $B_{m\beta}$ is the percent of isomer decays which are β decays rather than internal transitions to another nuclear state. T_{pop} is the approximate timescale on which the isotope is populated, and Notes gives some brief comments on the nucleus.

Isotope	E_m (keV)	J_g^π	J_m^π	$T_{1/2\ g}$ (s)	$T_{1/2\ m}$ (s)	$B_{m\beta}$ (%)	T_{pop}	Notes
⁶⁹ Zn	438.636	1/2 ⁻	9/2 ⁺	3.38×10^3	4.95×10^4	0.033	3 min	λ^β slowed for 14 hours, EM signal ^a
⁷¹ Zn	157.7	1/2 ⁻	9/2 ⁺	1.47×10^2	1.43×10^4	100	20 s	λ^β slowed for 4 hours
⁷⁹ Se	95.77	7/2 ⁺	1/2 ⁻	1.03×10^{13}	2.35×10^2	0.056	9 min	$T_{1/2\ m} < T_{pop}$, no new effect
⁸¹ Se	103.00	1/2 ⁻	7/2 ⁺	1.11×10^3	3.44×10^3	0.051	30 s	λ^β slowed for 1 hour
⁸³ Kr	41.5575	9/2 ⁺	1/2 ⁻	stable	6.59×10^3	0	2.5 h	γ only, likely no effect
⁸⁵ Kr	304.871	9/2 ⁺	1/2 ⁻	3.39×10^8	1.61×10^4	78.8	3 min	λ^β accelerated for 5 hr ^b , EM signal ^a
⁹³ Nb	30.77	9/2 ⁺	1/2 ⁻	stable	5.09×10^8	0	1.6 Myr	$T_{1/2} < T_{pop}$, no new effect
⁹⁵ Nb	235.69	9/2 ⁺	1/2 ⁻	3.02×10^6	3.12×10^5	5.6	64 d	$T_{1/2} < T_{pop}$, no new effect
⁹⁷ Nb	743.35	9/2 ⁺	1/2 ⁻	4.33×10^3	5.87×10^1	0	17 h	$T_{1/2} < T_{pop}$, no new effect
⁹⁹ Tc	142.684	9/2 ⁺	1/2 ⁻	6.66×10^{12}	2.16×10^4	0.0037	66 h	$T_{1/2\ m} < T_{pop}$, no new effect
¹¹³ Cd	263.54	1/2 ⁺	11/2 ⁻	2.54×10^{23}	4.45×10^8	99.86	6 h	low Y_m , long $T_{1/2}$, likely unobservable
¹¹⁵ Cd	181.0	1/2 ⁺	(11/2) ⁻	1.92×10^5	3.85×10^6	100	20 min	λ^β slowed for 45 d
¹¹⁵ In	336.244	9/2 ⁺	1/2 ⁻	1.39×10^{22}	1.61×10^4	5.0	54 h	¹¹⁵ Sn production boosted, EM signal ^a
¹¹⁷ In	315.303	9/2 ⁺	1/2 ⁻	2.59×10^3	6.97×10^3	52.9	3 h	$T_{1/2} < T_{pop}$, no new effect
¹¹⁹ In	311.37	9/2 ⁺	1/2 ⁻	1.44×10^2	1.08×10^3	95.6	3 min	λ^β slowed for 18 min
¹²¹ In	313.68	9/2 ⁺	1/2 ⁻	2.31×10^1	2.33×10^2	98.8	15 s	feeds ¹²¹ Sn isomer
¹¹⁹ Sn	89.531	1/2 ⁺	11/2 ⁻	stable	2.53×10^7	0	3-18 min	EM signal ^a
¹²¹ Sn	6.31	3/2 ⁺	11/2 ⁻	9.73×10^4	1.39×10^9	22.4	4 min	λ^β slowed for 44 yr, EM signal ^a
¹²⁹ Sn	35.15	3/2 ⁺	11/2 ⁻	1.34×10^2	4.14×10^2	100	1 s	λ^β slowed for 7 min
¹²⁶ Sb	17.7	(8 ⁻)	(5 ⁺)	1.07×10^6	1.15×10^3	86	230 kyr	$T_{1/2} < T_{pop}$, no new effect
¹²⁸ Sb	0.0+X	8 ⁻	5 ⁺	3.26×10^4	6.25×10^2	96.4	1 h	λ^β accelerated to 11 min $T_{1/2}$ ^b
¹³⁰ Sb	4.8	(8 ⁻)	(4,5) ⁺	2.37×10^3	3.78×10^2	100	4 min	λ^β accelerated to 6.3 min $T_{1/2}$ ^b
¹²⁵ Te	144.775	1/2 ⁺	11/2 ⁻	stable	4.96×10^6	0	3 yr	$T_{1/2\ m} < T_{pop}$, no new effect
¹²⁷ Te	88.23	3/2 ⁺	11/2 ⁻	3.37×10^4	9.17×10^6	2.4	4 d	λ^β slowed for 100 d, EM signal ^a
¹²⁹ Te	105.51	3/2 ⁺	11/2 ⁻	4.18×10^3	2.9×10^6	36	4.5 h	λ^β slowed for 34 d, EM signal ^a
¹³¹ Te	182.258	3/2 ⁺	11/2 ⁻	1.5×10^3	1.2×10^5	74.1	23 min	λ^β slowed for 33 h, EM signal ^a
¹³³ Te	334.26	(3/2 ⁺)	(11/2 ⁻)	7.5×10^2	3.32×10^3	83.5	2.5 min	λ^β slowed for 1 h
¹³¹ Xe	163.930	3/2 ⁺	11/2 ⁻	stable	1.02×10^6	0	8 d	EM signal ^a
¹³³ Xe	233.221	3/2 ⁺	11/2 ⁻	4.53×10^5	1.9×10^5	0	21 h	EM signal ^a
¹³⁷ Ba	661.659	3/2 ⁺	11/2 ⁻	stable	1.53×10^2	0	30 yr	$T_{1/2} < T_{pop}$, no new effect
¹⁴⁴ Pr	59.03	0 ⁻	3 ⁻	1.04×10^3	4.32×10^2	0.07	285 d	$T_{1/2} < T_{pop}$, no new effect
¹⁶⁶ Ho	5.969	0 ⁻	7 ⁻	9.66×10^4	3.79×10^{10}	100	82 h	λ^β slowed for 1200 y
¹⁸⁹ Os	30.82	3/2 ⁻	9/2 ⁻	stable	2.09×10^4	0	1 d	$T_{1/2} < T_{pop}$, no new effect
¹⁹¹ Ir	171.29	3/2 ⁺	11/2 ⁻	stable	4.90×10^0	0	16 d	$T_{1/2\ m} < T_{pop}$, no new effect
¹⁹⁵ Ir	100	3/2 ⁺	11/2 ⁻	8.24×10^3	1.32×10^4	95	7 min	feeds ¹⁹⁵ Pt isomer ^c
¹⁹⁵ Pt	259.077	1/2 ⁻	13/2 ⁺	stable	3.46×10^5	0	4 h	EM signal ^a

^aEM signals (detectable x-rays and γ -rays) are *possible*, but we do not study them carefully here.

^bNearly all parent β decay feeds the isomer, effectively bypassing the longer-lived ground state.

^cTo explore ¹⁹⁵Pt isomer population, we assume all ¹⁹⁵Os β decay feeds the ¹⁹⁵Ir isomer (actual feeding unknown).

β -decay rates, as well as thermally induced excitation and spontaneous/thermally stimulated de-excitation between long-lived nuclear states, tempers the dramatic effects suggested by Fujimoto & Hashimoto (2020).

5. CONCLUSIONS

We have calculated temperature-dependent production and destruction rates (β decay and internal transitions) using available data for all known neutron-rich isomers that may participate in the rapid neutron capture process. We show for the first time that nuclear isomers are dynamically populated in an r -process event.

We focus on those populated isomers of notable astrophysical importance, and refer to these as astromers. Our nucleosynthesis simulations for the radioactive decay of r process nuclei, in combination with our newly defined Astromer Importance Rating, AIR, facilitates the identification of influential r -process astromers.

Our work highlights the need for careful treatment of isomeric states when simulating the many and varied aspects of r -process nucleosynthesis. This is especially true when attempting to incorporate astromers that may originate from fission deposition or be dynamically produced and/or destroyed at early times when neutron capture is still active.

Current (Zhang et al. 2019; Orford et al. 2020; Nesterenko et al. 2020; Walker et al. 2020) and future studies at radioactive beam facilities that focus on elucidating the properties of nuclear isomers will help to refine our astromer predictions and strengthen our understanding of the creation of the heavy elements. This work provides fresh motivation for experimental campaigns to measure the beyond-ground-state properties—such as excited state half-lives and γ -ray intensities—that astromer evolution relies on.

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