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First Direct Measurement of ${}^{22}Mg(\alpha,p){}^{25}Al$ and Implications for X-Ray Burst Model-Observation Comparisons

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Type-I x-ray bursts can reveal the properties of an accreting neutron star system when compared with astrophysics model calculations. However, model results are sensitive to a handful of uncertain nuclear reaction rates, such as ${}^{22}Mg(\alpha, p)$. We report the first direct measurement of ${}^{22}Mg(\alpha, p)$, performed with the Active Target Time Projection Chamber. The corresponding astrophysical reaction rate is orders of magnitude larger than determined from a previous indirect measurement in a broad temperature range. Our new measurement suggests a less-compact neutron star in the source GS1826-24.

Type-I x-ray bursts (XRBs) are thermonuclear explosions on the surface of accreting neutron stars powered by nuclear burning [1-3]. In recent years, advances in XRB observations and modeling have opened a unique window to constrain the neutron star mass-radius relation and other underlying physics through comparisons between observations and models [4,5]. As XRB light curves are powered by nuclear reactions, XRB models are sensitive to various nuclear physics inputs (e.g., nuclear reaction rates) [6,7]. Models with reliable nuclear physics data are needed to validate the assumptions of astrophysical models through model-observation comparisons [8]. Accurate nuclear physics input plays an equally important role in predicting the burst ashes, which alter the composition of the crust of the neutron star. In mass-accreting systems, this crust is made in part or entirely out of XRB ashes. Various sensitivity studies conducted over years have shown that the ²²Mg(α , p)²⁵Al reaction rate is among the most significant reactions that directly impact light curves and burst ashes [6,7,9]. In a recent study, performed to assess the impact of uncertainties in nuclear inputs on the extraction of the neutron star mass-radius relation, the ²²Mg(α , p)²⁵Al reaction rate was shown to have a significant effect even when decreased by factor 10 [8].

Prior to an XRB, accreted hydrogen is compressed and burned via the hot CNO cycles, until the triple- α reaction is initiated and the thermonuclear runaway ensues. At this point the burst begins and the temperature rises rapidly. At temperatures around 0.5-0.6 GK, breakout from the hot CNO cycle via ¹⁵O(α, γ)¹⁹Ne and ¹⁸Ne(α, p)²¹Na becomes efficient. These breakout reactions are crucial as it takes the flow from hot CNO cycles to higher mass region. These breakout reactions open the door for the αp process, enabling the reaction flow to reach ²²Mg. At this branching point, ²²Mg(α , p)²⁵Al competes with the rather slow β^+ decay and with ${}^{22}Mg(p,\gamma){}^{23}A1$ [4,10]. The current experimental constraint on this reaction rate comes from an indirect measurement where resonant states in ²⁶Si were explored through the ${}^{28}\text{Si}(p, t){}^{26}\text{Si}$ reaction by Matic *et al.* [11]. This experimentally constrained reaction rate is more than a factor of 100 below Hauser-Feshbach (HF) predictions in the relevant XRB temperature range above 0.7 GK. The large deviation from the HF based model calculations was attributed to the lack of resonance data above 10 MeV excitation energy in ²⁶Si [11]; therefore their rate was considered to be a lower limit, and the HF based rate an upper limit. These two rates lead to significantly different results when used in XRB model calculations resulting in a significant uncertainty of model-observation comparisons. Since XRB ashes ultimately set the composition of the accreted neutron star crust, the discrepant ash results may also impact model-observation comparisons for neutron star crust cooling [4,12,13]. Therefore, it is important to directly measure this reaction to reduce this very large uncertainty and constrain the XRB model calculations. In this work, we report the first direct measurement of the ²²Mg(α , p)²⁵Al reaction using the Active-Target Time Projection Chamber (AT-TPC).

The ²²Mg(α , p)²⁵Al measurement was carried out at the National Superconducting Cyclotron Laboratory. ²²Mg was produced from the fragmentation of a ²⁴Mg primary beam accelerated by the coupled cyclotrons and selected by the A1900 fragment separator [14]. The ²²Mg fragments were stopped in a linear gas cell and transported to an electron beam ion source, where their charge state increased to 12^+ . Finally, the ions were injected into the ReA3 reaccelerator, accelerated to \sim 5 MeV/u with an average beam intensity of ~900 pps. A pure ²²Mg beam was delivered to the experiment station where it was first transported through a thin ionization chamber filled with isobutane at 10 torr. An ionization chamber recorded the beam intensity and identified beam contaminants [15]. Before the beam entered the active volume of the AT-TPC, it went through a 3.6- μ mthick aluminized para-aramid window of 1 cm diameter. The energy loss in the entrance window was 4.7 MeV. The AT-TPC active volume is a cylinder of length 1 m and of a radius 29.2 cm, placed in a uniform 1.9 T magnetic field generated by a solenoidal magnet. The AT-TPC was filled with 600 torr $\text{He:CO}_2(95\%:5\%)$ to stop the beam in the middle. The sensor plane consists of a mosaic of 10240 equilateral triangle pads that provides x and y information of the tracks of the particles. Ionization electrons drift time provides the longitudinal component of the track. Typical drift times through the detector were around 40 μ s. The ionization chamber signal was used to retain the arrival of beam particles through the window as a time reference. which is pivotal to determine the reaction vertex position along the beam axis. The AT-TPC is equipped with a hierarchical digital data acquisition system. At the bottom of this hierarchy is the AGET chip, which controls the sampling and shaping of the signals and compares them to a threshold to generate a channel level trigger. Each AGET outputs a multiplicity signal which is derived from the number of channels that are above a preset threshold amplitude. Multiplicities from each AGET (ASIC for General Electronics for TPCs) are further collected by their controlling CoBos (concentration boards) to produce a CoBo-level multiplicity signal [16]. In addition, a logical trigger signal from the ion chamber is delayed and put in coincidence with a large window opened by the CoBo multiplicity trigger. If the delayed ion chamber signal falls within this time window, a trigger signal will be generated. The AT-TPC electronics system allows the readout and trigger attributes of each channel to be set individually [15]. Among the dominating channels open at this energy are (α, α) and (α, p) . From kinematics, protons are the only particles that were emitted at backward angles, i.e., $\theta_{\text{lab}} > 90^{\circ}$. Therefore, protons were identified by selecting backscattered particle tracks. Another possible source of backscattered protons are fusion-evaporation reactions on carbon and oxygen. Any background contribution from reactions on carbon in this angular domain were estimated (using PACE4 [17]) to be less than 0.4%. The backscattered proton tracks were analyzed using the random sample consensus (RANSAC) method [18,19]. Figure 1 shows an example of a proton track in 3D (top panel). The middle and lower panels show a 2D projection of the same track on the pad plane as well as the RANSAC analysis, respectively. The RANSAC algorithm provides the best fit of the spiral to a circle and its radius. Then, we calculate the arc length starting between the point of closest approach, which in this case is the very first point of the track (isolated using our clustering algorithm) and each point of the hit pattern. From the arc length versus z-vertex plot [Fig. 1(c)] we can extract the angle and the interaction vertex. Tracks for the (α, p) reaction channel were simulated using the AT-TPC simulation package. The AT-TPC simulation uses the virtual Monte Carlo package which relies on the GEANT4 engine. The amount of primary ionization is estimated by dividing the energy lost by the tracked particle in each time step of the simulation. The energy loss in the gas is converted into electrons that are drifted to the pad plane which also takes into account lateral and longitudinal diffusion of the ionization electrons. Pulses are generated taking into account the response function of the pad and the electronics [15]. The data are then transformed into a three-dimensional hit pattern that represents the position of each simulated interaction point convoluted with the size of the pads and the sampling rate of the electronics. More about the AT-TPC simulation package and digitization can be found in Ayyad et al. [18]. Simulated tracks were also analyzed as described above. The top panel of Fig. 2 shows the event-by-event reaction vertex as a function of laboratory angle for the experimental and simulated data. It is evident from the plot that the experimentally accessible angular domain is limited from 90° to 130° in the laboratory frame and agrees with the simulated data. As we are analyzing only the backscattered protons, the higher angles are accessible only when the reaction vertex is at some distance from the entrance window. The detected angular range during the experiment depends on the geometrical acceptance of the detector and the threshold of the multiplicity trigger. Figure 2(b) shows the simulated acceptance as a function of laboratory angle without and with multiplicity threshold. The inset of Fig. 2(b) shows the number of pads hit as function of the laboratory angle. It is worth pointing out that the geometrical acceptance of the detector covers up to 160°

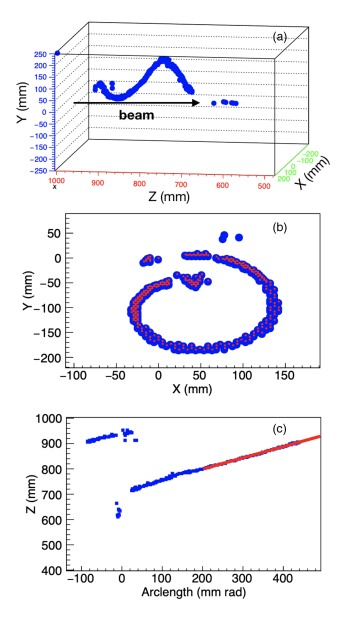


FIG. 1. Panel (a) shows a three-dimensional view of an example proton track and the arrow shows the beam direction. Panel (b) shows the projection of a proton track on the pad plane where blue dots are data points and red indicates the data points chosen for RANSAC analysis. Panel (c) shows the arclength of each hit pattern point as a function of the z coordinate. The red line is the least-squares fit performed to extract the scattering angle from the slope.

(red dots). The experimental data (blue triangles) show a sharp cutoff at ~80 hits per event. Higher laboratory angles were cut off due to this multiplicity threshold in the current experiment. To obtain the angle-integrated cross sections, the proton distribution in the laboratory frame was calculated at different beam energies using PACE4 [17]. The ratio of counts from 0° to 180° to counts in the angular region covered in this study are shown in Fig. 2(c). This energy-dependent ratio was used to obtain the angle-integrated cross section.

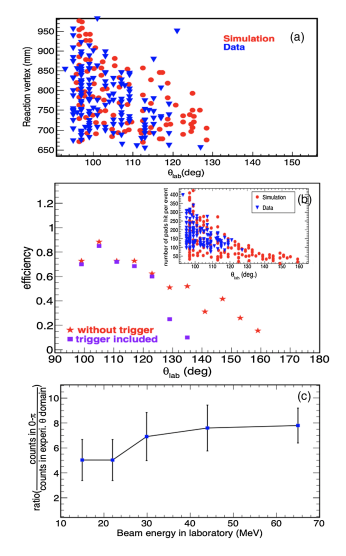


FIG. 2. Panel (a) shows the reaction vertex as a function of laboratory angle for experimental data (blue inverted triangle) overlaid on simulation data (red filled circles). Panel (b) shows the efficiency as a function of laboratory angle and the inset shows the number of pads hit per event as a function of the laboratory angle. Panel (c) shows the ratio of calculated (using PACE4) total counts over the counts in the experimental angular domain as a function of the beam energy.

The obtained excitation function is shown in Fig. 3(a) (and also in tabular form in Table I) in comparison to HF calculations. Data points are shown at the weighted energy. Here, the weighted energy is defined as $\int \sigma(E)EdE / \int \sigma(E)dE$, where the energy dependence of HF based NON-SMOKER cross sections was used [6]. Vertical error bars include contributions from statistical and systematic uncertainties. Systematic error bars include a 5% uncertainty in the number of target atoms based on energy loss tables uncertainties, 5% in the incident beam counts, and 35% error when accounting for counts outside the angular domain covered in this study. Our estimation of 35% uncertainty is based on any uncertainty originating from the model

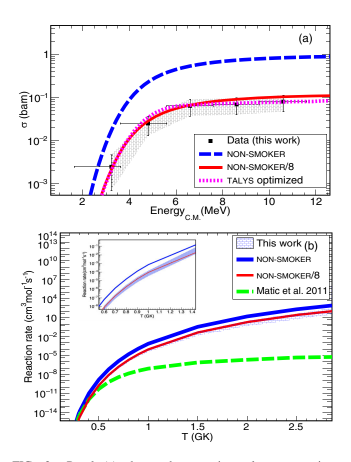


FIG. 3. Panel (a) shows the experimental cross sections obtained in the present work over a range of center-of-mass energies covered (black). For all the points, the cross section weighted energy is shown, which is the reason why horizontal error bars for the two lowest energy points are asymmetric. Panel (b) shows the reaction rate comparison of the current work to different model predictions and to the previous measurement by Matic *et al.* [11].

prediction of the proton angular distribution. Horizontal error bars reflect the bin size in center-of-mass energies. The cross sections obtained in the current work are a factor of 8–10 lower than the HF calculation results. The lowest center-of-mass energy of our measurement is located near the upper end of the astrophysical Gamow window for a 2 GK temperature [20]. Extrapolation to lower center-of-mass energies is required to obtain the cross section and reaction rate in the relevant temperature range. For this, a combination of input parameters for the TALYS code was identified to best reproduce the experimental data within error bars. In TALYS, we used the following combination of models:

TABLE I. Cross sections obtained in the present work as a function of (weighted) center-of-mass energy. Errors are shown in parentheses.

Weighted energy (MeV)	Protons detected per energy bin	Cross section (mb)	Error (stat, syst)	
3.2	3	$2.5(^{+2.3}_{-1.8})$	$\begin{pmatrix} 2.1, 0.9\\ 1.6, 0.9 \end{pmatrix}$	
4.8	18	26(12)	(6,11)	
6.6	41	64(27)	(10,25)	
8.6	40	69(29)	(11,26)	
10.6	45	80(33)	(12,31)	

semimicroscopic optical model potential, dispersive alpha optical potential from Demetriou et al. [21], microscopic level densities from Goriely's tables, and γ -ray strength functions from the Brink-Axel Lorentzian model [22]. The best fit to the experimental cross section was achieved after multiplying the default normalization factor for the shape of the double-folding α potential by 0.884 and the default level density of the compound nucleus by a factor of 2.60. The optimized TALYS reaction rate (this work) is shown in Fig. 3(b), lower panel, where the shaded area shows the estimated reaction rate uncertainty. The reaction rate uncertainty was estimated using the maximum and minimum cross sections obtained in this work indicated by vertical error bars and shaded regions in Fig. 3(a). The horizontal error bars in the cross-section measurement in Fig. 3(a) represents the energy range over which given cross sections are energy averaged. The astrophysical reaction rate in Fig. 3 (b) is energy averaged and hence does not reflect the effect of horizontal error bars of Fig. 3(a). The dramatic disagreement between the reaction rates from Matic et al. [11] and HF model calculations above 0.4 GK deduced in the previous measurement was not observed in the current work. Above 1 GK, the reaction rate reported in this work follows a similar trend, within the error bars, as the one inferred from NON-SMOKER but notably smaller in magnitude. With the current measurement the reaction rate in the critical temperature range around 0.7-1 GK is determined by using the best-fit TALYS model to experimental data just above the Gamow window. This new reaction rate was fit to the standard seven parameter REACLIB database form and best-fit parameters in the temperature range 0.3-3.5 GK are provided in Table II.

To assess the impact of our measurement on the XRB light curve, we performed multizone XRB calculations with the code MESA, following those described in Refs. [5,8].

TABLE II. Best-fit parameters for new rate as a fit to standard seven parameter REACLIB form, i.e., $\exp[a_0 + \sum_{i=1}^{5} a_i T_9^{(2i-5)/3} + a_6 \ln(T_9)]$.

Parameter	a_0	a_1	a_2	<i>a</i> ₃	a_4	a_5	<i>a</i> ₆
Best-fit value	43.939	0.0	-5.31×10	6.35×10^{-1}	-8.29×10^{-1}	6.5×10^{-2}	-6.6×10^{-1}

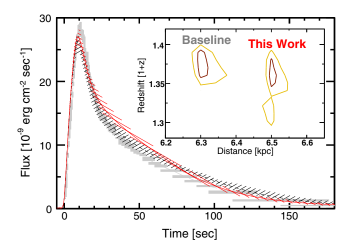


FIG. 4. XRB light curve comparison for GS1826-24 observations (gray boxes), MESA calculations using the NON-SMOKER rate for ${}^{22}Mg(\alpha, p)$ (black band), and the NON-SMOKER rate reduced by /8 (red band). The inset shows the corresponding 68% (red) and 95% (yellow) confidence intervals for the distance and redshift, which were determined as described in Ref. [8].

We employed the REACLIBv2.2 nuclear reaction rate library, where the NON-SMOKER HF rate of Ref. [6] is the default for ${}^{22}Mg(\alpha, p){}^{25}Al$. We used the astrophysical conditions that were found to best reproduce observables from the year 2000 bursting epoch of the source GS1826-24 [5,23]. Figure 4 compares MESA results to astronomical observations. We use a distance of 6.3 kpc and redshift (1 + z) of 1.38, which provide the best fit between the observed light curve and our baseline calculation, to mimic the x-ray detection solid angle and neutron star surface gravitational redshift that modify the observed light curve.

Figure 4 shows that the calculations from Refs. [5,8] (black band) generally reproduce the observed light curve (gray boxes), including the recurrence time between bursts (which is not shown). However, this agreement is substantially diminished when reducing the NON-SMOKER rate for ${}^{22}Mg(\alpha, p)$ by a factor of 8 (red band). In particular, the tail of the XRB light curve is substantially modified because a reduced ²²Mg(α , p) rate effectively enhances hydrogen burning early in the burst by making the path $^{22}Mg(p,\gamma)^{23}Al(p,\gamma)^{24}Si$ [24] more competitive. Therefore, less hydrogen is available to be burned at later times following the light curve peak, resulting in a more rapid decline of the light curve tail. The implication of our result is that an alternative distance and surface gravitational redshift are needed in order to reproduce observed features of GS1826-24, as shown in the Fig. 4 inset. This implies a less-compact neutron star in GS1826-24 than determined by Ref. [8]. Our results bring constraints to an important uncertainty in XRB model calculations which will improve the extraction of neutron star compactness via XRB light curve model-observation comparisons.

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