

ABSTRACT

FOX, WILLIAM CARR. Investigating Nucleosynthesis in Massive AGB Stars with Transfer Reactions. (Under the direction of Richard Longland).

Abstract text ...

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Investigating Nucleosynthesis in Massive AGB Stars with Transfer Reactions

by
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BIOGRAPHY

The author was born in a small town ...

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CHAPTER

1

INTRODUCTION

Understanding the origin of the elements has been the primary goal of nuclear astrophysics since its conception. Through the work of Burbidge, Burbidge, Fowler, and Hoyle [Bur57], it was established that the chemical elements observed in our solar system were synthesized through nuclear burning in stars. Indeed, all but the smallest elements in our observable universe (hydrogen through boron) are the ashes of stellar burning. Consequently, in the quest to discover the nature of the elemental origins, we must understand the nature of stellar processes. From birth to death, stars undergo nuclear reactions in their interiors. With sufficient temperatures, the reaction products themselves undergo subsequent nuclear reactions. An evolutionary stage is reached at which point the star either explodes or otherwise ejects its material outward. This nuclear-processed material is mixed together with the ashes from neighboring stars, forming new stars with new compositions and therefore new nucleosynthesis. One of the most complex evolutionary stages of a star is known as the asymptotic giant branch (AGB) phase, and stars that we currently observe on this phase are known as AGB stars. This thesis focuses on these complex nucleosynthesis processes in massive AGB stars, including s-process nucleosynthesis resulting from thermal pulses, as well as hot-bottom burning at the base of the convective hydrogen envelope. These stellar processes are investigated through laboratory nuclear reaction experiments here on earth. In particular, transfer reaction experiments are used in this thesis as a surrogate for the direct reactions occurring in the stellar processes. These experiments provide information on the nuclear structure of the nuclei involved in the key reactions, and therefore they constrain the reaction rates that are crucial to nucleosynthesis.

The present chapter will provide an overview of the nuclear astrophysics topics addressed in this thesis. Chapter 2 will introduce the mathematical formalism for quantifying nucleosynthesis with reaction rates, as well as the formalism for transfer reactions. Chapter 3 will address the “rubidium problem” in massive AGB stars, a discrepancy between observed and theoretical rubidium abundances from s-process nucleosynthesis. The s-process branching at ^{86}Rb is investigated through a Monte Carlo reaction network approach, identifying the most important reactions to rubidium abundance. Chapter 4 will introduce the experimental techniques that were used in this thesis to successfully perform transfer reaction experiments. Chapter 5 will introduce the development of a new digital data acquisition system (DAQ) for the focal-plane detector package at the Enge Split-Pole Spectrograph, which will replace the current analog system for future transfer reaction experiments. Finally, Chapter 6 will address the Mg–K abundance anomaly observed in globular cluster NGC 2419, which may have originated from hot-bottom burning in massive AGB stars. The $^{39}\text{K}({}^3\text{He}, d)^{40}\text{Ca}$ proton-transfer reaction is measured at high resolution, and new constraints are placed on the key potassium-destroying reaction $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$.

1.1 Asymptotic Giant Branch Stars

Asymptotic giant branch (AGB) stars produce about half of all elements heavier than iron in our galaxy [Bus99], and they are one of the leading candidates for the origin of abundance anomalies in globular clusters [Pra07]. An understanding of these stars therefore provides insight into the origin of the elements. The asymptotic giant branch refers to the evolutionary track that these stars follow on a typical Hertzsprung-Russell (HR) diagram, which asymptotically approaches that of the red giant branch. HR diagrams show the luminosity of a star as a function of its effective temperature. These properties are often cataloged for each star observed in a globular cluster, for example, and the current evolutionary phase of a star can be determined from its relative position on the diagram. A theoretical example of the evolutionary track of a $5 M_{\odot}$ star is represented on the HR diagram of Fig. 1.1. Starting from the zero-age main sequence (ZAMS), where the star first begins to burn hydrogen, the evolutionary track passes through the sub-giant branch (SGB), the red-giant branch (RGB), the horizontal branch (HB), the early (E-) and thermally-pulsing (TP-) asymptotic giant branch (AGB) phases, and the post-AGB branch. It then undergoes planetary nebula (PN) formation, before eventually becoming a white dwarf. Each of these branches represents a unique evolutionary phase of the star, many of which are described in the figure at key transitional points. As mentioned above, this thesis focuses on AGB stars, the evolutionary track of which is highlighted in red in the figure.

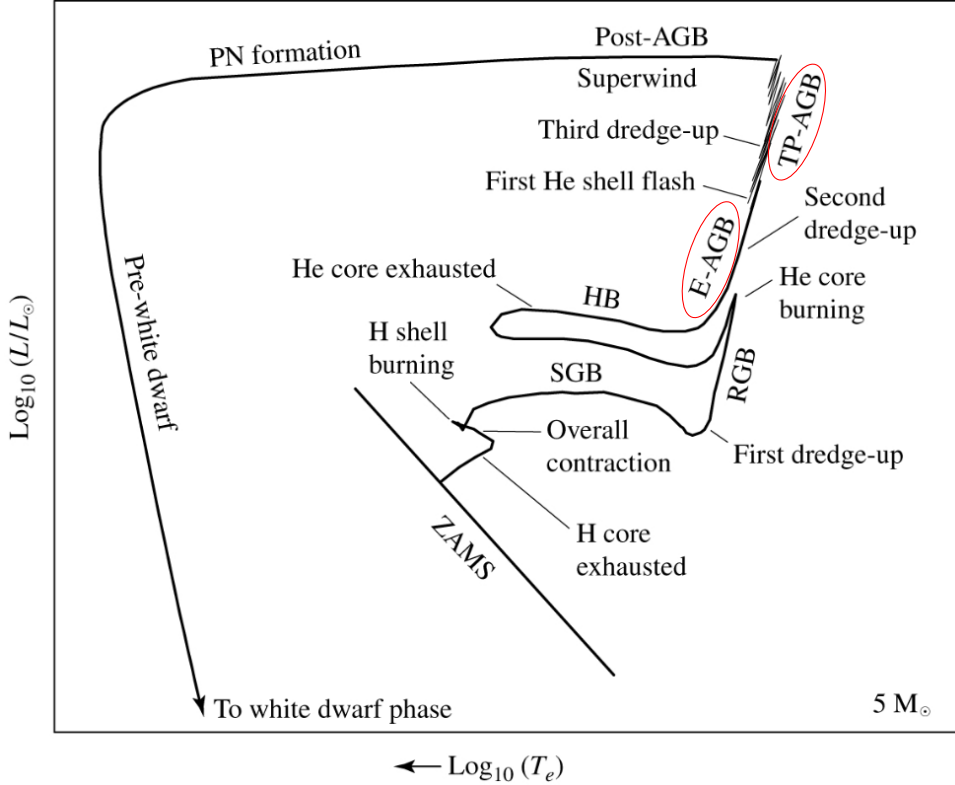


Figure 1.1: Evolution of a $5 M_{\odot}$ star represented on the Hertzsprung-Russell (HR) diagram. The two stages of the asymptotic giant branch (AGB) are highlighted in red. See text for definitions of acronyms. Figure adapted from Ref. [Car07].

It is clear from Figure 1.1 that AGB stellar evolution is separated into two stages. The *early-AGB* (e-AGB) stage occurs after both hydrogen and helium have been exhausted in the core of a star, leaving behind mostly inert carbon and oxygen. Hydrogen and helium burning is still active in separate shells around the core, with a helium-rich intershell region in between. The far more active hydrogen-burning shell perpetually increases the density and temperature of the intershell region, until the rate of energy generated by the helium-burning shell is greater than the rate at which energy can be transported outward through radiative diffusion [Ili15]. A thermal instability, called a *thermal pulse*, occurs as a result, initiating the *thermally pulsing AGB* (TP-AGB) stage, shown in Figure 1.2. This pulse causes the helium-burning shell to extend into the intershell region and ignite the hydrogen-burning shell, rendering it inactive for a brief period. The nuclei inside the intershell region get carried outward through the convective hydrogen envelope toward the surface of the star, in what is known as a *third dredge-up* (3DUP) event¹, where they are ejected by stellar winds. The

¹The first two dredge-ups are depicted in Figure 1.1, occurring at the start of the RGB and during the

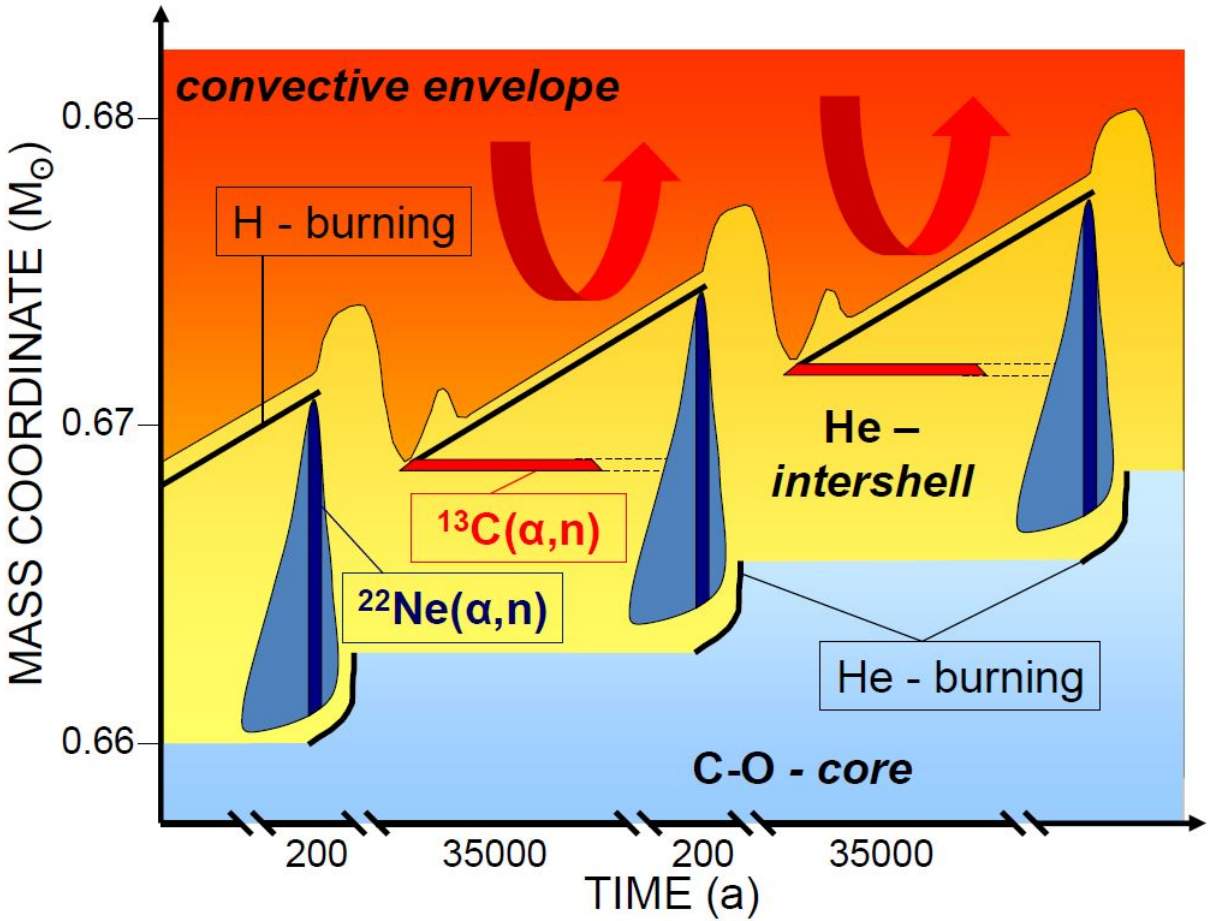


Figure 1.2: The evolution and structure of a thermally pulsing AGB star. Three thermal pulses are depicted in blue, extinguishing the hydrogen burning shell. The neutron sources $^{13}\text{C}(\alpha, n)^{16}\text{O}$ and $^{22}\text{Ne}(\alpha, n)^{25}\text{Mg}$ are shown to be active between pulses and during pulses, respectively. Figure adapted from Ref. [Rei14].

hydrogen-burning shell takes over once again as the dominant energy-producer after the thermal pulse, and this cycle repeats for on the order of tens to hundreds of pulses [Hab04].

Helium burning during each thermal pulse produces ^{12}C as a result of the triple- α reaction, followed by ^{16}O from $^{12}\text{C}(\alpha, \gamma)^{16}\text{O}$, which get accumulated mostly in the core, increasing its mass. However, some ^{12}C accumulates in the intershell region. After a 3DUP event resulting from a thermal pulse, some protons from the convective hydrogen envelope mix into the intershell region, initiating the reaction sequence $^{12}\text{C}(p, \gamma)^{13}\text{N}(\beta^+ \nu)^{13}\text{Ca}(p, \gamma)^{14}\text{N}$. The effect of these additional proton-capture reactions is the formation of a ^{13}C pocket and a ^{14}N pocket at the top of the intershell region. During the interpulse period, when temperatures

e-AGB stage, respectively. For stars with $M \lesssim 4 M_\odot$, the second dredge-up does not occur because the convective hydrogen envelope cannot penetrate the intershell region at this point.

are between $100 \text{ MK} \lesssim T \lesssim 300 \text{ MK}$, the $^{13}\text{C}(\alpha, n)^{16}\text{O}$ reaction is activated, producing a large neutron density of $N_n \approx 10^7 \text{ cm}^{-3}$. During thermal pulses, when temperatures are $T \gtrsim 300 \text{ MK}$, the reaction sequence $^{14}\text{N}(\alpha, \gamma)^{18}\text{F}(\beta^+\nu)^{18}\text{O}(\alpha, \gamma)^{22}\text{Ne}$ proceeds. As a result, the $^{22}\text{Ne}(\alpha, n)^{25}\text{Mg}$ reaction is activated, producing an even larger neutron density of $N_n \approx 10^{10} \text{ cm}^{-3}$. Nucleosynthesis occurs in the intershell region from these two neutron sources via a series of neutron-captures and β -decays, known as the s-process, described in Section 1.2. The heavy nuclei produced from the s-process are brought to the surface by subsequent thermal pulses and 3DUP events.

In the case of AGB stars with $M \gtrsim 4 M_\odot$, the convective hydrogen envelope reaches down to the top of the hydrogen-burning shell during the interpulse period, where temperatures are $T \gtrsim 50 \text{ MK}$ [Ili15], but can theoretically reach as high as $T \approx 150 \text{ MK}$ for *Super-AGB* (SAGB) stars with $M \gtrsim 7 M_\odot$ [Ven12]. The resulting hydrogen burning nucleosynthesis is known as hot-bottom burning, detailed in Section 1.3, and the products of this burning are brought to the surface by the convective hydrogen envelope. This process is one of the leading candidates for the recent abundance anticorrelations observed in globular clusters, detailed in Section 1.3.1.

1.2 s-Process Nucleosynthesis

Most light nuclei below the iron-peak at mass number $A \approx 60$ are formed through charged-particle fusion reactions in stars, where temperatures are sufficient for transmission through the repulsive Coulomb barrier via quantum tunneling. The synthesis of nuclei heavier than the iron-peak, however, cannot be explained by these charged-particle fusion reactions. At low temperatures, the transmission probability through the large Coulomb barrier of heavy nuclei becomes prohibitively small, while at high temperatures, nuclear statistical equilibrium favors the production of iron-peak nuclei due to their large binding energy per nucleon. Nuclei with $A \gtrsim 60$ must be formed instead by neutron capture, where Coulomb repulsion is no longer a factor. Neutron capture will generate neutron-rich isotopes of a given element, some of which eventually undergo β -decay if they are unstable. The stable daughter nuclei will then capture neutrons until another unstable isotope is reached, and the process continues this way up to some of the heaviest nuclei.

When the neutron density in the astrophysical environment is sufficiently small such that unstable nuclei preferentially β -decay, the ensuing nucleosynthesis is referred to as the *s*-process. This is a relatively slow process (hence the *s*-), as the nucleosynthesis can only proceed based on the half-lives of the unstable nuclei. If, on the other hand, the neutron density is sufficiently large such that unstable nuclei preferentially neutron-capture, the

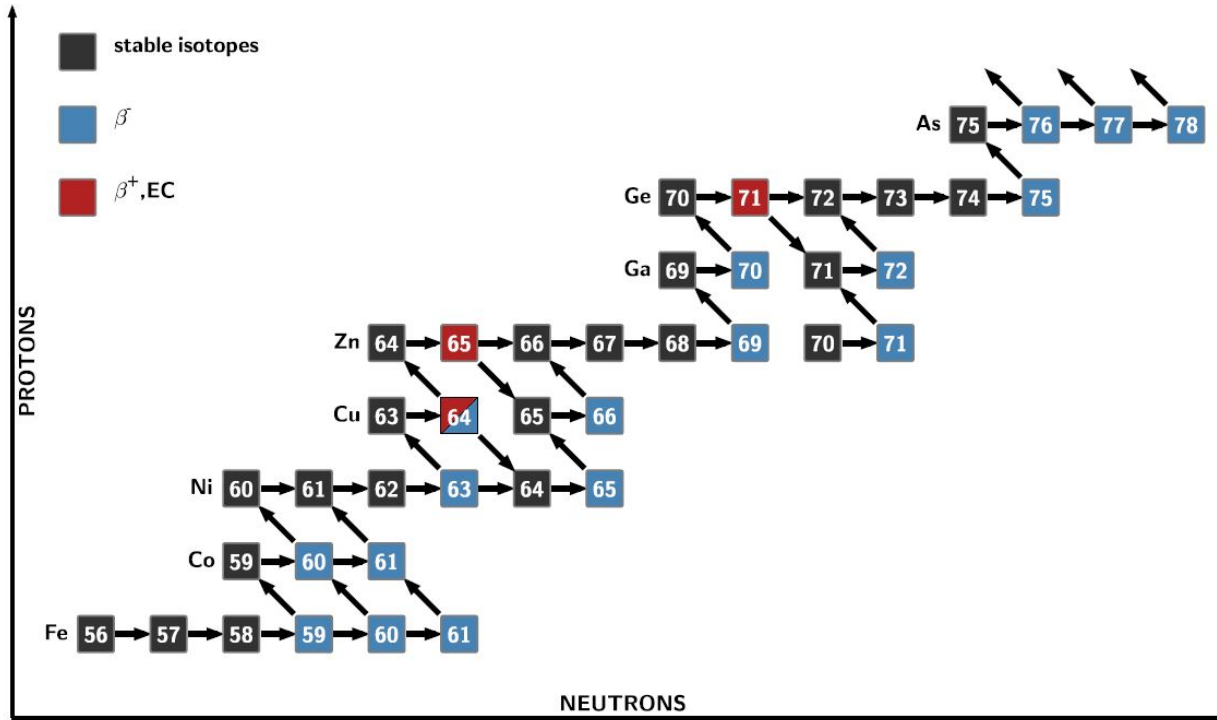


Figure 1.3: The s-process path from iron to arsenic. The stable nuclei in black undergo neutron capture, while the unstable nuclei in blue (β^- -decay) and red (β^+ -decay or electron capture) either decay or undergo neutron-capture depending on their half-lives and the neutron density N_n . At s-process branchings, e.g. ^{59}Fe , the probability for neutron-capture and decaying is approximately equal, hence the path is split into two branches. Figure adapted from Ref. [Rei14].

ensuing nucleosynthesis is referred to as the *r*-process because this happens at a much more (*r*-)apid pace. These two processes occur in different astrophysical sites and are equally responsible for almost all of the heavy nuclei in the observable universe. Some heavy nuclei are referred to as s-only or r-only nuclei because only one of these processes accounts for their abundance. The research presented in Chapter 3 is based on s-process nucleosynthesis occurring in AGB stars.

To illustrate how the s-process works, consider the chain of reactions shown in Figure 1.3, starting with the most abundant stable nucleus formed at the end of the stellar fusion reaction chains, ^{56}Fe . This is referred to as a *seed nucleus* because it is the nucleus that initiates the s-process reaction chain. Stable nuclei are shown in black in the figure, nuclei unstable to β^- -decay are shown in blue, and nuclei unstable to β^+ -decay or electron capture (EC) are shown in red. As is typical of a chart of the nuclides, neutron number increases to the right and proton number increases upward. Neutron-capture is therefore represented by an arrow to the right, while β -decays and electron capture are represented by diagonal arrows

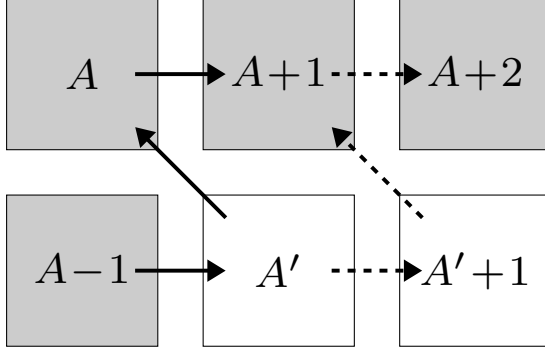


Figure 1.4: A simple s-process path initiating from the nucleus with mass number $A - 1$, where shaded nuclei are stable and unshaded nuclei are unstable. The simplest case, where only β -decays occur on unstable nuclei, is shown by the solid arrows. The added complication of a branching at A' is indicated by dashed arrows.

due to the conversion of a neutron to a proton, or vice versa. If there is a sufficient neutron density N_n in the astrophysical environment, the neutron-capture of ^{56}Fe , $^{56}\text{Fe}(n, \gamma)^{57}\text{Fe}$, leads to the stable nucleus ^{57}Fe . The two subsequent neutron-captures, $^{57}\text{Fe}(n, \gamma)^{58}\text{Fe}$ and $^{58}\text{Fe}(n, \gamma)^{59}\text{Fe}$, on the stable ^{57}Fe and ^{58}Fe nuclei, respectively, leads to the unstable ^{59}Fe nucleus. The nucleosynthesis path has now lead to an *s-process branching*, and ^{59}Fe is referred to as a *branching point nucleus*. At these branchings, the decay constant for β -decay, λ_β , is approximately equal to the decay constant for neutron capture, $\lambda_{n\gamma}$. That is, $\lambda_\beta \approx \lambda_{n\gamma}$, and the path splits into two branches with approximately equal probability. These branchings are sensitive to neutron density N_n and the neutron-capture cross section $\sigma_{n\gamma}$ of the unstable branching point nucleus, described below. After the s-process branching, the path continues in this way up to some of the heaviest nuclei.

Now consider the simple s-process path shown in Figure 1.4, starting from a stable nucleus with mass number $A - 1$. In the simplest case represented by solid arrows, the unstable nucleus with mass number A' will β -decay to nucleus A , which will then neutron-capture to nucleus $A + 1$, and so on. The more complicated case is represented by dashed arrows, where the nucleus A' is at a branching, and it can undergo neutron-capture and β -decay at roughly equal probability. We will first assume that $\lambda_\beta \gg \lambda_{n\gamma}$, such that the path does not lead through the dashed arrows. In this case, the abundance at each mass number exists in only one (unprimed) nucleus. Therefore the abundance evolution of any stable nucleus A only depends on the neutron-capture reaction rates of the $A - 1$ and A nuclei, as in

$$\frac{dN_s(A)}{dt} = N_n(t)N_s(A - 1)\langle\sigma v\rangle_{A-1} - N_n(t)N_s(A)\langle\sigma v\rangle_A, \quad (1.1)$$

where $N_s(A)$ and $N_n(t)$ are the time-dependent number densities of the A nucleus and the neutron, respectively, and $\langle\sigma v\rangle_A$ is the neutron-capture reaction rate per particle pair of the A nucleus. The first term in Eqn 1.1 represents the rate of production of the A nucleus, while the second term represents its rate of destruction. We will assume that the temperature is constant while the neutron source is active, so that the neutron-capture reaction rates are also constant. The neutron-capture reaction rate is frequently expressed in terms of a Maxwellian-averaged cross section $\langle\sigma\rangle_A \equiv \langle\sigma v\rangle_A/v_T$, where $v_T = \sqrt{2kT/\mu_{An}}$ is the thermal velocity, or in other words, the maximum of the velocity distribution; μ_{An} is the reduced mass between the A nucleus and the neutron, which is approximately equal to the neutron mass. That is, $\mu_{An} = M_A M_n / (M_A + M_n) \approx M_n$, which makes v_T approximately independent of target mass. The abundance evolution of the nucleus A can then be written as

$$\frac{dN_s(A)}{dt} = v_T N_n(t) [N_s(A-1)\langle\sigma\rangle_{A-1} - N_s(A)\langle\sigma\rangle_A]. \quad (1.2)$$

Far from any s-process branchings or closed neutron-shell nuclei, an approximate equilibrium is obtained where the abundance of the A nucleus increases until its production rate and destruction rate are approximately equal for a given neutron flux. This is known as the *local equilibrium approximation*, and it is expressed as $N_s(A, \Phi)\langle\sigma\rangle_A \approx N_s(A-1, \Phi)\langle\sigma\rangle_{A-1}$, where Φ is the time-integrated neutron flux. The approximation has been shown to hold for the s-only isotopes of tellurium ($Z = 52$), as well as many other s-only nuclei with adjacent mass numbers between $90 \lesssim A \lesssim 205$ [Ili15]. However, there is a distinct separation of observed solar system $N_\odot(A)\langle\sigma\rangle_A$ values at $A \approx 90$, and the approximation does not hold at all near the closed neutron-shell nuclei. Refs. [Cla61, See65] have shown that better agreement with solar system $N_\odot(A)\langle\sigma\rangle_A$ values is achieved when considering not just a single (time-integrated) neutron flux Φ , but an exponential distribution of Φ -values. Indeed, it is reasonable to assume that the number of ^{56}Fe seed nuclei decreases with increased exposure to neutrons. With this additional constraint, the solar system $N_\odot(A)\langle\sigma\rangle_A$ values are reproduced to a remarkable accuracy for $90 \lesssim A \lesssim 205$, when also accounting for s-process branchings, discussed below. The region between $60 \lesssim A \lesssim 90$ is reproduced only when adding a second exponential distribution of neutron fluxes. These two components are referred to as the *main s-process component* ($90 \lesssim A \lesssim 205$) and the *weak s-process component* ($60 \lesssim A \lesssim 90$). A third component, called the *strong s-process component* is needed to explain the solar system $N_\odot(A)\langle\sigma\rangle_A$ values at $A \gtrsim 205$.

These components originate from different astrophysical environments. The main s-process component is from thermally-pulsing AGB stars with $1.5M_\odot \lesssim M \lesssim 3M_\odot$, while most of the weak s-process component is thought to originate from core helium burning in massive stars

with $M \gtrsim 13M_{\odot}$ [Ili15]. The primary neutron source for the main component is $^{13}\text{C}(\alpha, n)^{16}\text{O}$, with the $^{22}\text{Ne}(\alpha, n)^{25}\text{Mg}$ neutron source only marginally activated. This latter source acts to enhance the efficiency of s-process branchings. The weak s-process component, however, relies on $^{22}\text{Ne}(\alpha, n)^{25}\text{Mg}$ as its primary neutron source, since temperatures are significantly higher for these massive stars.

If we now consider the nucleus A' in Figure 1.4 as an s-process branching, we can see that only a fraction of the path reaches the nucleus A , but the entire path reaches the nucleus $A+1$. In this case, the local equilibrium approximation becomes $N_s(A, \Phi)\langle\sigma\rangle_A + N_s(A', \Phi)\langle\sigma\rangle_{A'} \approx N_s(A+1, \Phi)\langle\sigma\rangle_{A+1}$. We can now define the *branching ratio* B as

$$\begin{aligned} B &\equiv \frac{N_s(A, \Phi)\langle\sigma\rangle_A}{N_s(A+1, \Phi)\langle\sigma\rangle_{A+1}} = \frac{\lambda_\beta(A')}{\lambda_\beta(A') + \lambda_{n\gamma}(A')} \\ &= \frac{\ln 2/T_{1/2}(A')}{\ln 2/T_{1/2}(A') + N_n\langle\sigma\rangle_{A'}v_T}. \end{aligned} \quad (1.3)$$

Solving for the neutron number density N_n , we obtain

$$N_n = \left(\frac{1-B}{B} \right) \frac{1}{\langle\sigma\rangle_{A'}v_T} \frac{\ln 2}{T_{1/2}(A')}. \quad (1.4)$$

Even though Eqn. 1.4 is just one of many examples of possible s-process branching configurations, it shows the only inputs necessary to calculate the neutron density, which is a key parameter in stellar models of the s-process. From this equation, a connection is made between the study of s-process branchings and the neutron density.

In order to extract the neutron density with a reasonable uncertainty, it is crucial to have precise experimental neutron-capture cross sections of the A , $A+1$, and A' nuclei. Since the A and $A+1$ nuclei are stable, these are usually readily available. However, A' is unstable, and it is typically not feasible to perform neutron-capture cross section measurements on unstable nuclei. In cases where data is not available, the cross sections have to be estimated using Hauser-Feshbach theory. It is possible to experimentally constrain neutron-capture cross sections on unstable nuclei by performing photoabsorption reactions on the recoil nucleus, if it is stable. The dipole response of the recoil nucleus via nuclear resonance fluorescence (NRF) and the (γ, n) reaction can be used to determine the photoabsorption cross section of the recoil nucleus, which can in turn be used to determine its photon strength function (PSF). The PSF is a crucial input for statistical model calculations constraining Maxwellian-averaged cross sections of neutron-capture on unstable nuclei [Wil20].

1.3 Hot-Bottom Burning

1.3.1 Candidate for Globular Cluster Abundance Anomalies

CHAPTER

2

NUCLEAR REACTIONS IN STARS

2.1 Nucleosynthesis

2.2 Reaction Rates

The reaction rate per particle pair in a stellar medium is given by

$$\langle \sigma v \rangle_{01} = \sqrt{\frac{8}{\pi \mu_{01}}} \frac{1}{(kT)^{3/2}} \int_0^\infty E \sigma(E) e^{-E/kT} dE, \quad (2.1)$$

where μ_{01} is the reduced mass of particle 0 and particle 1, $\mu = M_0 M_1 / (M_0 + M_1)$; k is the Boltzmann constant; T is the stellar temperature; E is the center-of-mass energy between the particles; and $\sigma(E)$ is the reaction cross section evaluated at E . The energy dependence of the cross section determines whether numerical integration must be performed. The formalism for reaction rate calculations therefore depends on whether $\sigma(E)$ is smoothly varying over energy (for non-resonant reactions) or if it can be described by isolated resonances (for resonant reactions). It can be described by both in principle, in which case the cross section is separated into non-resonant and resonant components, assuming interference effects are negligible. Regardless of the nature of $\sigma(E)$, it can be separated into a Coulomb factor and a

factor resulting only from nuclear structure. That is, the cross section can be rewritten as

$$\sigma(E) = \frac{1}{E} e^{-2\pi\eta} S(E), \quad (2.2)$$

where $S(E)$ is the astrophysical S-factor, governed by nuclear effects alone, and η is the Sommerfeld parameter defined by $2\pi\eta = \sqrt{2\mu_{01}/E} Z_0 Z_1 e^2 / \hbar$, where Z_0 and Z_1 are the atomic numbers of the nuclei. The $1/E$ factor is included to cancel with the E factor in the reaction rate. Substituting Eqn. 2.2 into Eqn. 2.1, we have

$$\langle \sigma v \rangle_{01} = \sqrt{\frac{8}{\pi\mu_{01}}} \frac{1}{(kT)^{3/2}} \int_0^\infty e^{-2\pi\eta} e^{-E/kT} S(E) dE. \quad (2.3)$$

The integrand in Eqn. 2.3 is composed of the Gamow factor $e^{-2\pi\eta}$, the Maxwell-Boltzmann factor $e^{-E/kT}$, and the astrophysical S-factor $S(E)$. The former two factors have a combined energy dependence of $e^{-\sqrt{1/E}} e^{-E}$, while the $S(E)$ energy dependence is based on the nuclear structure of the specific reaction. The overlap between the Gamow and Maxwell-Boltzmann factors is called the *Gamow peak*, $e^{-2\pi\eta} e^{-E/kT}$. This peak determines the energies at which the reaction will proceed in the stellar environment at the given temperature¹. It has a maximum at

$$E_0 = \left[\left(\frac{\pi}{\hbar} \right)^2 (Z_0 Z_1 e^2)^2 \left(\frac{m_{01}}{2} \right) (kT)^2 \right]^{1/3} \\ = 0.1220 (Z_0^2 Z_1^2 \mu_{01} T_9^2)^{1/3} [\text{MeV}] \quad (2.4)$$

where T_9 is the temperature in units of GK. The energy E_0 is the most probable energy for non-resonant reactions, where $S(E)$ varies smoothly. For resonant reactions, where $S(E)$ is described by a sharp Lorentzian function, the Gamow peak can still be a useful indicator of which resonances contribute significantly to the total reaction rate, particularly at lower resonance energies ($E_r^{\text{c.m.}} \lesssim 0.5$ MeV). The Gamow peak can be approximated by a Gaussian function with a mean of E_0 and a $1/e$ width of

$$\Delta = \frac{4}{\sqrt{3}} \sqrt{E_0 kT} = 0.2368 (Z_0^2 Z_1^2 \mu_{01} T_9^5)^{1/6} [\text{MeV}]. \quad (2.5)$$

Thermonuclear reactions therefore mainly occur in the energy window from $E_0 - \Delta/2$ to $E_0 + \Delta/2$, known as the *Gamow window*.

For the ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ reaction, discussed in Chapter 6, the Gamow window at temperatures

¹This is a slight oversimplification. For resonant reactions at high stellar temperatures, the resonances that contribute significantly to the reaction rate may occur below the Gamow peak [Ili15].

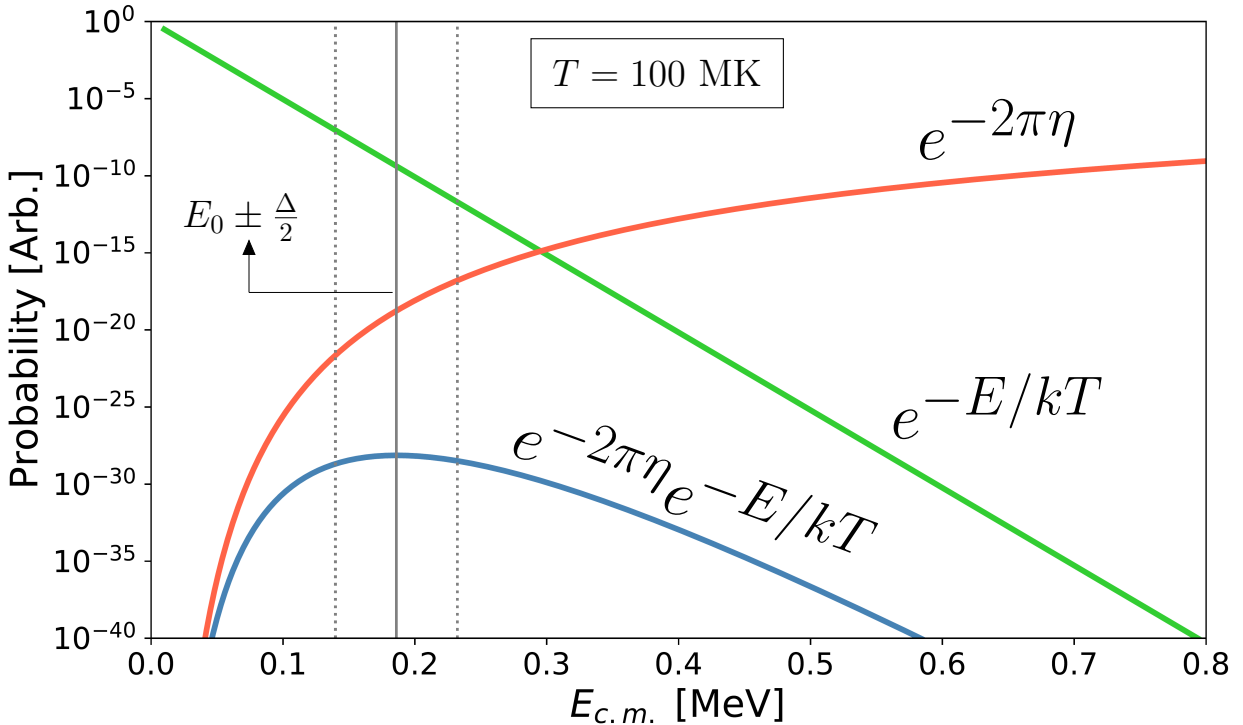


Figure 2.1: The Gamow window, $E_0 \pm \Delta/2$, for the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction at 100 MK. The Gamow factor $e^{-2\pi\eta}$ is shown in red; the Maxwell-Boltzmann factor $e^{-E/kT}$ is shown in green, and the Gamow peak $e^{-2\pi\eta}e^{-E/kT}$ is shown in blue. The resonances that contribute significantly to the reaction rate at this temperature will have energies within the Gamow window, between about 140 keV and 230 keV.

of 100 MK occurs between about 140 keV and 230 keV, centered on about 185 keV. This situation is depicted in Figure 2.1. The Gamow and Maxwell-Boltzmann factors are shown in red and green, respectively, while the combined Gamow peak is shown in blue. The Gamow window centered on E_0 is shown between the vertical dotted lines. As will be clear in Section 6.9.1, the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate is dominated by isolated resonances in the temperature range of astrophysical interest, 80–260 MK, occurring between about 120 keV and 450 keV.

2.2.1 Narrow Resonances

An isolated resonance at the resonance energy E_r is described by the Breit-Wigner cross-section [Bre36]

$$\sigma_{\text{BW}}(E) = \frac{\lambda^2}{4\pi} \omega \frac{\Gamma_a(E) \Gamma_b(E)}{(E - E_r)^2 + \Gamma(E)^2/4}, \quad (2.6)$$

where λ is the deBroglie wavelength, defined by $\lambda^2/2 = (\pi\hbar)^2/(\mu_{01}E)$; $\Gamma_a(E)$ and $\Gamma_b(E)$ are the entrance channel and exit channel partial widths, respectively; $\Gamma(E)$ is the total width, and ω is the spin factor, defined as

$$\omega = \frac{(2J+1)}{(2J_0+1)(2J_1+1)}, \quad (2.7)$$

where J is the resonance spin, and J_0 and J_1 are the spins of the interacting particles. Substituting Eqn. 2.6 into Eqn. 2.1, the reaction rate per particle pair arising from an isolated resonance becomes

$$\langle\sigma v\rangle_{01} = \frac{\sqrt{2\pi}\hbar^2}{(\mu_{01}kT)^{3/2}} \omega \int_0^\infty \frac{\Gamma_a(E)\Gamma_b(E)}{(E-E_r)^2 + \Gamma(E)^2/4} e^{-E/kT} dE. \quad (2.8)$$

Consider the partial widths Γ_i and the Maxwell-Boltzmann factor $e^{-E/kT}$ varying slowly with energy over the width of the resonance. In this situation, the resonance is considered narrow, and these factors can be treated as being independent of energy over this energy range. The reaction rate is then written as

$$\begin{aligned} \langle\sigma v\rangle_{01} &= \frac{\sqrt{2\pi}\hbar^2}{(\mu_{01}kT)^{3/2}} \omega \Gamma_a \Gamma_b e^{-E_r/kT} \int_0^\infty \frac{dE}{(E-E_r)^2 + \Gamma(E)^2/4} \\ &= \frac{\sqrt{2\pi}\hbar^2}{(\mu_{01}kT)^{3/2}} \omega \Gamma_a \Gamma_b e^{-E_r/kT} \frac{2}{\Gamma} \arctan\left(\frac{E-E_r}{\Gamma/2}\right) \Big|_0^\infty \\ &= \left(\frac{2\pi}{\mu_{01}kT}\right)^{3/2} \hbar^2 \omega \frac{\Gamma_a \Gamma_b}{\Gamma} e^{-E_r/kT}, \end{aligned} \quad (2.9)$$

where the last line results from the further approximation of narrow resonances, $E_r \gg \Gamma$, which makes the lower bound of integration effectively $-\infty$. The quantity

$$\omega\gamma \equiv \omega \frac{\Gamma_a \Gamma_b}{\Gamma} \quad (2.10)$$

is defined as the *resonance strength* because it is proportional to the maximum cross section multiplied by the total resonance width, $\omega\gamma \propto \sigma_{\max}\Gamma$. The reaction rate per particle pair for multiple isolated, narrow resonances forms an incoherent sum,

$$\langle\sigma v\rangle_{01} = \left(\frac{2\pi}{\mu_{01}kT}\right)^{3/2} \hbar^2 \sum_i (\omega\gamma)_i e^{-E_{r,i}/kT}. \quad (2.11)$$

The calculation of partial widths differs between particles and γ -rays. The following discussion will consider charged-particles and γ -rays only. For a discussion of neutron partial widths,

see Ref. [Ili15].

2.2.2 Particle Partial Widths

The charged-particle partial width for channel c is

$$\Gamma_c(E) = \frac{2\hbar^2}{\mu_{01}R^2} P_c(E) C^2 S \theta_{\text{sp},c}, \quad (2.12)$$

where the channel radius R is defined by $R = R_0 (A_0^{1/3} + A_1^{1/3})$; the penetration factor $P_c(E)$ can be calculated as

$$P_c(E) = \frac{\rho}{F^2(E) + G^2(E)}, \quad (2.13)$$

where $\rho = 0.21874 R \sqrt{\mu_{01}E}$ and F and G are the energy dependent Coulomb wave functions; $C^2 S$ is the spectroscopic factor, sometimes simply referred to as S , where C is the isospin Clebsch-Gordon coefficient, and $\theta_{\text{sp},c}$ is the dimensionless single-particle reduced width. The charged-particle partial width can be thought of as being proportional to the probability of the particle being emitted from the compound nucleus. Each of the latter three quantities in Eqn. 2.12 contribute to this probability. The penetration factor $P_c(E)$ represents the probability of overcoming or tunneling through the Coulomb barrier to reach the nuclear surface. The dimensionless single-particle reduced width $\theta_{\text{sp},c}$ represents the probability that the particle occupies the exact single-particle shell-model state of the resonance at the nuclear surface. Finally, the spectroscopic factor $C^2 S$ is a measure of how much the populated state resembles that single-particle shell-model state. Although $C^2 S$ is model-dependent, it is the only quantity in Eqn. 2.12 that is based on measurement, as will be clear in Section 2.3.2.

The dimensionless single-particle reduced width is given by

$$\theta_{\text{sp},c} = \frac{R}{2} |u_{\text{sp},c}(R)|^2, \quad (2.14)$$

where $u_{\text{sp},c}(R)$ is the single-particle radial wavefunction evaluated at the channel radius, which is normalized to unity as in

$$\int_0^R |u_{\text{sp},c}(r)|^2 r^2 dr = 1. \quad (2.15)$$

One method of numerically calculating $\theta_{\text{sp},c}$ is to vary the parameters of the single-particle potential until a resonance is produced at the binding energy. This is performed, for example, by the code BIND, detailed in Ref. [Ili97]. Another procedure, performed for ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ in Sec. 6.8.5, is to use the weak binding approximation. This approximation is required if the

nuclear reaction code used in the determination of C^2S only handles bound states, as is the case with **Fresco** [Tho88, Tho06]. This method assumes unbound states are weakly bound (by ~ 1 keV), and therefore $u_{\text{sp},c}(R)$ is the bound state radial wavefunction evaluated at the channel radius. The channel radius R is selected to represent the nuclear surface, where the wavefunctions for the bound state and unbound state are equivalent.

A fact that appears troubling for the weak binding approximation at first glance is that both C^2S and $\theta_{\text{sp},c}$ depend linearly on the binding energy, assuming R is chosen at the nuclear surface. Therefore, both quantities will diverge from unbound calculations at high binding energies. However, as Ref. [HAR23] has shown for the proton-transfer reaction $^{30}\text{Si}(^{3}\text{He}, d)^{31}\text{P}$, C^2S decreases as a function of binding energy at roughly the same rate as $\theta_{\text{sp},c}$ increases, such that their product is balanced. The only significant energy dependence from Eqn. 2.12 therefore comes from the penetration factor $P_c(E)$, which can easily be evaluated for unbound energies without the use of the weak binding approximation.

2.2.3 γ -ray Partial Widths

The γ -ray partial width for a single transition is

$$\Gamma_\gamma(\bar{\omega}, E_\gamma) = \frac{8\pi(L+1)}{L [(2L+1)!!]^2} \left(\frac{E_\gamma}{\hbar c} \right)^{2L+1} B(\bar{\omega}L), \quad (2.16)$$

where $\bar{\omega}$ represents either electric or magnetic radiation; E_γ is the γ -ray transition energy; L is the γ -ray multipolarity, and B is the branching ratio. The γ -ray partial width at the incoming particle energy E can be scaled with respect to that of the resonance energy E_r as in

$$\Gamma_\gamma(E) = \Gamma_\gamma(E_r) \left(\frac{E + Q - E_f}{E_r + Q - E_f} \right)^{2L+1}, \quad (2.17)$$

where Q is the Q -value, or equivalently, the particle separation energy, and E_f is the final excitation energy of the γ -decay.

2.3 Transfer Reactions

2.3.1 Optical Model

2.3.2 Distorted Wave Born Approximation

CHAPTER

3

RUBIDIUM PRODUCTION IN MASSIVE AGB STARS

3.1 Introduction

3.2 Rubidium Overabundance

3.3 s-Process Reaction Network Methodology

3.3.1 The Reaction Network

3.3.2 Reaction Rate Sensitivities

Method 1: Manually Varying the $^{22}\text{Ne}(\text{a},\text{n})$ Rate

Method 2: Monte Carlo

3.4 Re

3.4.1 Abundance Evolution

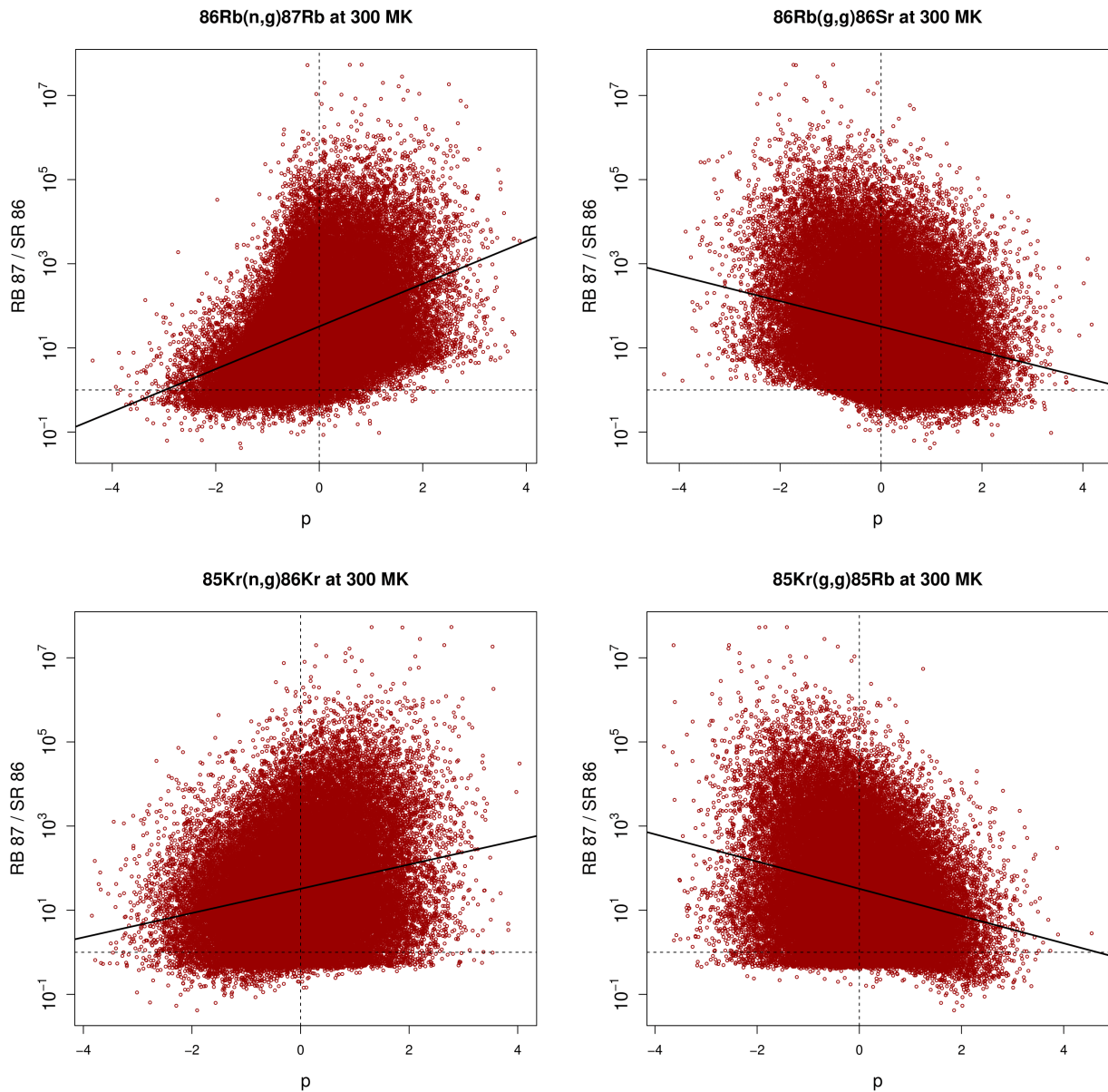


Figure 3.1: Here's my caption!

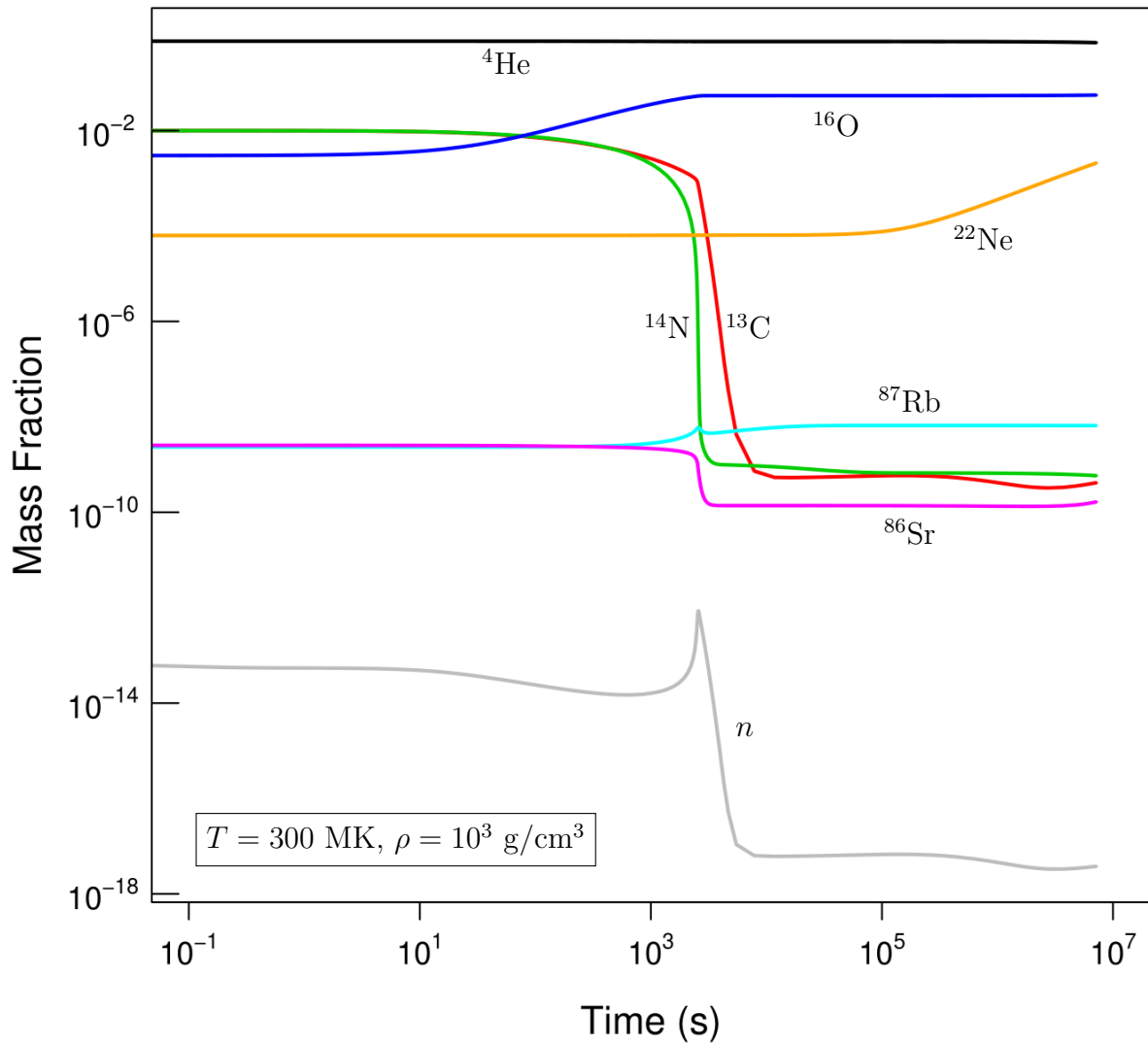


Figure 3.2: The abundance evolution of key nuclei related to the s-process branching at ^{86}Rb for a single network calculation simulating the conditions during a single AGB thermal pulse. The T , ρ , and $X_{\text{last}}({}^4\text{He})$ parameters are 300 MK, 10^3 g/cm^3 , and 0.7, respectively, where the initial mass fraction of ${}^4\text{He}$ was 0.75.

CHAPTER

4

**EXPERIMENTAL NUCLEAR
TECHNIQUES**

CHAPTER

5

DATA ACQUISITION SYSTEM FOR THE FOCAL-PLANE DETECTOR PACKAGE

5.1 Introduction

A new digital data acquisition system (DAQ) and graphical user interface (GUI) have been developed for use with the focal-plane detector package [Mar19] of the Enge Split-Pole Spectrograph at the Triangle Universities Nuclear Laboratory (TUNL). The CAEN V1730 digitizer is meant to replace the traditional analog modules and analog-to-digital converter (ADC) that have previously been used for the focal-plane detector.

The purpose of this document is twofold. Firstly, it serves as an operation manual on the components of our upgraded system - the CAEN V1730 digitizer (Section 3), our custom frontend DAQ software (Section 4), and our custom GUI: EngeSpec (Section 5). Secondly, it provides a record of the author's contributions to the development of the DAQ (Section 4) and the GUI (Section 5) and to testing the new system with live data (Section 6). Section 2 provides a brief background and motivates the need for the upgrades.

5.2 Background and Motivation

Nuclear physics experiments involving the detection of radiation (γ -rays, neutrons, protons, α -particles, etc.) require a method of converting detected radiation into numerical data. Figure 5.1 illustrates the components of the traditional data acquisition method. Detectors first convert each radiation event into an analog electrical signal, or pulse. Pulse processing then alters the detector signal so as to reduce noise and amplify the pulse to sufficiently measurable levels. At this point, the energy and timing of the event can be determined by further processing. This processing is traditionally performed by a network of analog modules, where the processed pulse is then converted to a digital pulse via an analog-to-digital converter (ADC). The digital pulse can finally be read by a computer and collected by a data acquisition (DAQ) software. The stored data can be visualized by a graphical user interface (GUI), which sorts all of the data based on the corresponding detectors and observables, displays histograms, and can perform a number of functions, such as curve fitting and gating.

The network of analog pulse-processing modules needed for nuclear physics experiments can be incredibly complex. Input and output cables connect modules together, and the vast number of cables can be cumbersome. Rearranging the network of modules between experiments amounts to physically altering the cable setup, which can lead to cable-mapping issues. The modules themselves also take up a significant amount of space. Hence, analog modules have an inherent portability problem.

A network of traditional analog modules can be replaced by a single digitizer, a module that performs the same pulse processing digitally after it samples the detector signals itself. Figure 5.2 illustrates the new data acquisition method using a digitizer instead of analog

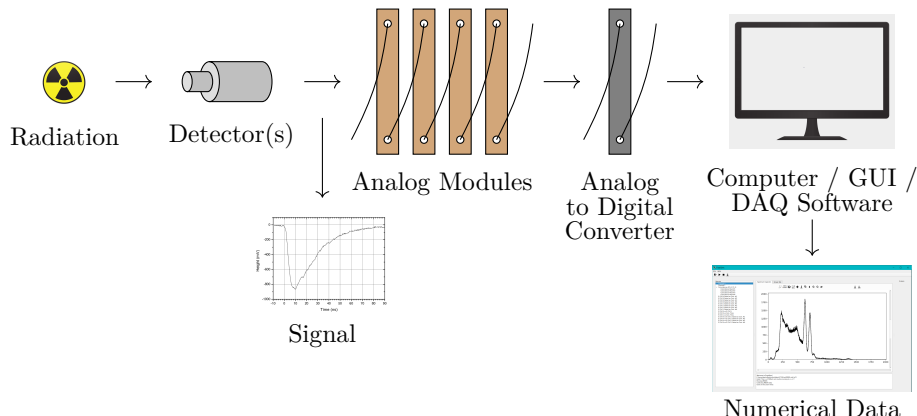


Figure 5.1: The traditional method of converting detected radiation into numerical data using analog pulse-processing modules and an analog-to-digital converter (ADC).

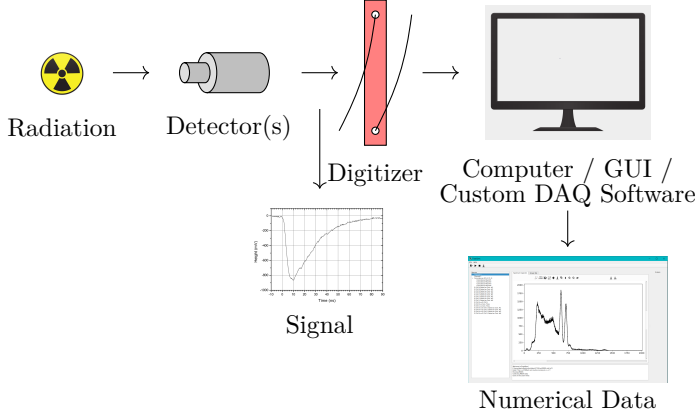


Figure 5.2: The new method of converting detected radiation into numerical data with a digitizer, which replaces analog modules.

modules and an ADC. Digitizers use firmware algorithms to emulate the functions of analog modules. Custom software is written by the user to perform specific functions that are needed for a given experiment. Changing the setup between experiments now amounts to altering existing code, rather than moving physical equipment.

At TUNL, the Longland group is replacing the focal-plane detector's analog modules with the CAEN V1730 digitizer and its associated Digital Pulse Processing (DPP) firmware algorithms. Figure 5.3 shows the difference between the old analog module setup versus the digitizer. Portability will now be significantly improved, and a large amount of physical space will be freed up for other purposes. The 14-bit digitizer will also provide enhanced energy and timing resolutions compared to the previous 12-bit ADC.

We have written a custom frontend DAQ software to accompany the digitizer, which uses MIDAS (Maximum Integrated Data Acquistion System) as our event-based DAQ. The DAQ software we used for our old system, NSCLDAQ, will no longer be supported soon, which is another incentive for the upgrades. In order to visualize the data from our new digitizer and DAQ, we have developed a GUI, called EngeSpec, that is connected with MIDAS to provide live data visualization during experiments. It also has an offline mode for analysis with previously stored data. EngeSpec has several data analysis tools like curve fitting for Gaussian and double-Gaussian peaks, peak area calculation, and gating functionality. A sort routine, called EngeSort, that was used with the old GUI has been revamped for use with EngeSpec. As before, it sorts the data by detector channel and by observable into each given histogram. However, the new EngeSort includes trigger and coincidence logic for the Enge focal-plane detector package, something that was previously performed by analog modules.

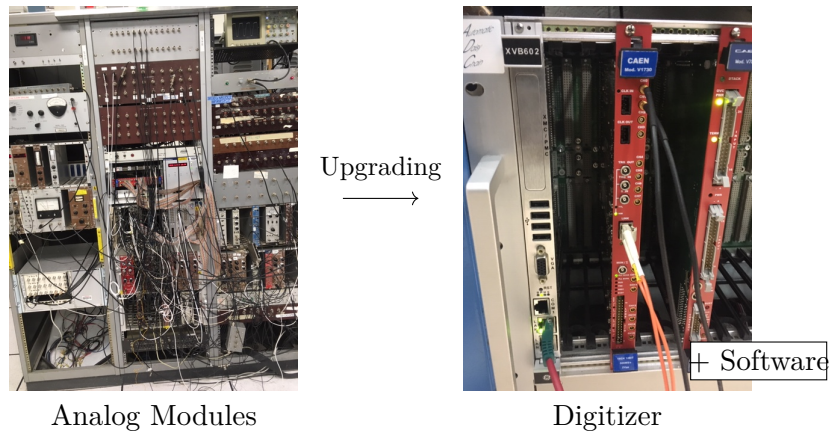


Figure 5.3: The CAEN V1730 Digitizer (pictured on the right), along with its associated DPP firmware and custom frontend DAQ software, is replacing the need for the unportable analog pulse-processing modules (pictured on the left) associated with the Enge focal-plane detector package at TUNL.

5.3 Overview of the CAEN V1730 Digitizer

This section provides a brief overview of how the CAEN V1730 digitizer functions, where the most important aspects of its Digital Pulse Processing (DPP) algorithms are discussed. Section 5.4 covers how to implement the algorithms via our frontend DAQ software. Our version of the digitizer firmware is known as DPP-PSD, or Digital Pulse Processing for Pulse Shape Discrimination. The official CAEN DPP-PSD user manual [Ref] is a comprehensive reference for details not covered in this document.

5.3.1 Digital Pulse-Processing Parameters

The V1730 digitizer is capable of sampling signals directly from detectors at a rate of 500 MS/s for each of its 16 input channels, corresponding to a time resolution of 2 ns per sample. Those samples are used by the DPP firmware in calculations of important quantities associated with a given signal, such as its baseline, energy, and time stamp. A typical sampled signal is shown in Figure 5.4, where parameters used by the DPP algorithms are summarized. These parameters are all user-defined and depend on the shape of the detector signals. One can use an oscilloscope or any waveform display software, such as CoMPASS, to approximate the parameters. For optimal resolution, it may be necessary to perform resolution tests for

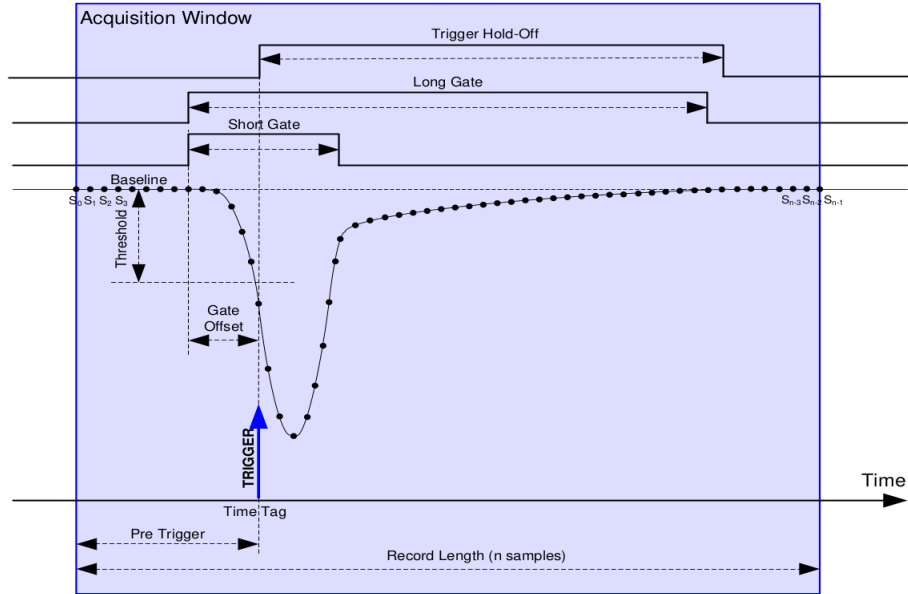


Figure 5.4: A signal sampled by the V1730 digitizer, summarizing the parameters used by the DPP algorithms. When the signal crosses the threshold value, the trigger fires, which primes the system for data acquisition.

certain parameters over a range of values (see Section 5.6.1 for an example).

A signal is triggered for data acquisition when it crosses the *threshold* voltage, which should be set high enough to filter out low-energy noise. When the trigger fires, the signal is delayed by the *pre-trigger* width to prime for the energy calculation. Since the energy of a radiation event is proportional to the charge that it deposits, the digitizer calculates this charge, and therefore determines the (uncalibrated) energy, by integrating the signal voltage over time. The length of time used in the integration is a user-defined parameter known as the *long gate*, and the integrated charge (i.e. uncalibrated energy) is denoted by Q_{Long} . The energy integration begins a length of time before the trigger fires, known as the *gate offset*, which must be less than the pre-trigger width. The purpose of the gate offset is to include the portion of the signal before the trigger in the energy integration. After the trigger fires, all other triggers from the same channel are inhibited during the *trigger hold-off* width, which must last at least until the end of the long gate. After this width, the channel can accept new triggers.

One of the most important quantities associated with the DPP algorithms is the baseline of the signal. The baseline is the reference value used for the energy calculation and to determine the threshold. This can either be a user-defined quantity, in which case it retains a fixed value, or the baseline can be dynamically calculated based on the average value of a user-defined number of samples during a moving time window (see Section 5.4.2 for the specific options). In the latter case, as shown in Figure 5.5, the baseline freezes during energy

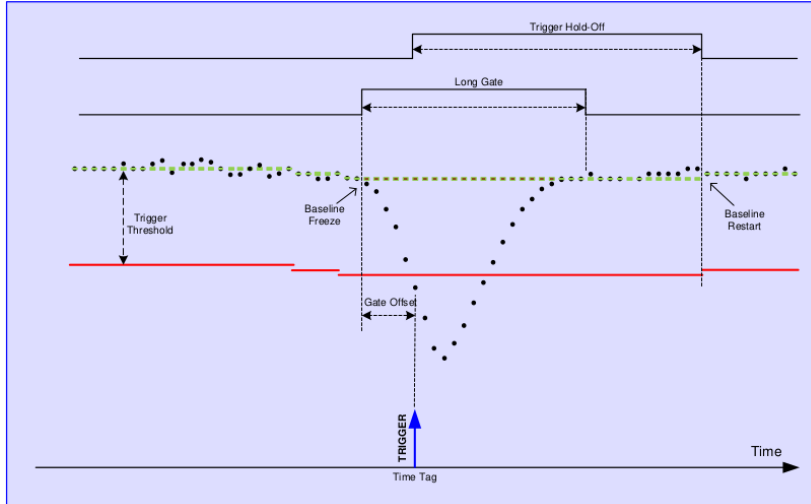


Figure 5.5: The dynamic baseline calculation using the mean value of a user-defined number of samples in a moving time window. The baseline freezes during the energy integration of a signal that has been triggered.

integration, and the baseline calculation restarts after the trigger hold-off width, using the samples both before and after the freeze. This means the baseline calculation contributes almost no dead-time. Note that the threshold follows the baseline variations.

The *short gate* is a measure of the fast portion of the signal. This is used, in conjunction with the long gate, to discern between different types of particles, e.g. between gamma-rays and neutrons. The pulse shape discrimination (PSD) is a quantity that measures the ratio between the area of the tail of the signal and the area of the total signal, i.e.

$$\text{PSD} = \frac{Q_{\text{Long}} - Q_{\text{Short}}}{Q_{\text{Long}}},$$

where Q_{Long} and Q_{Short} are the charge collected in the long gate and short gate during charge integration, respectively. Neutron events tend to have a distinctly higher PSD than gamma-ray events because they have larger tails. The option to collect the PSD for each event is not yet available in our frontend software, but it will be added soon.

The only parameter yet to be discussed in Figure 5.4, the *record length*, is not used by our digitizer. The V1730 only operates in List mode, where the event data is stored, but the samples themselves are not. Some other CAEN digitizers can operate in Waveform mode, where the samples in the record length that make up the *acquisition window* are stored to visualize the signals and the baseline with the appropriate CAEN software. However, the reduced amount of data transferred in List mode compared to Waveform mode keeps the dead-time of the system very low during runs.

5.3.2 Time Stamp Determination

By default, the timing of events in the DPP firmware is performed by *leading edge discrimination* (LED), where a time stamp is given to each event the moment it crosses the trigger threshold. However, threshold triggering on the leading edge of signals causes well-known issues with timing, known as *time walk*. Two signals exactly coincident, but with different amplitudes, will register different time tags using LED mode because the leading edge of the larger signal rises sooner.

A more precise way of determining signal time is by using *constant fraction discrimination* (CFD), which can be enabled by the DAQ user instead of LED mode (see Section 5.4.2). This method assigns the time stamp to the moment the signal amplitude reaches a constant fraction of its total amplitude, allowing for timing to be independent of signal amplitude. As in traditional analog CFD modules, the digitizer takes the input signal and produces two new signals, one attenuated by a factor f of the total amplitude and the other inverted and delayed by a time d . The two signals are summed together to form a bipolar pulse with a

single *zero-crossing*, which is taken as the time stamp of the event. Because of the finite sampling resolution of the digitizer, the exact zero-crossing location will always be located somewhere between two samples of the signal. The DPP firmware, by default, assigns the sample before the zero-crossing as the time stamp of the event, called the *coarse time stamp*. This is usually sufficient, but for more precise timing it is also possible to assign the time stamp as the interpolated zero-crossing. The width between the coarse time stamp and the interpolated zero-crossing is called the *fine time stamp*. If the user enables zero-crossing interpolation, the data for each event will take up more memory. Note that in the case of CFD mode, the threshold trigger serves the purpose of priming the system for CFD implementation rather than assigning a time stamp.

The time stamp for LED mode and the coarse time stamp for CFD mode are both referred to as the *trigger time tag* in the DPP firmware, a 31-bit unsigned integer that has units equal to the time resolution of the digitizer, 2 ns. In CFD mode, the fine time stamp is a 10-bit unsigned integer, meaning the interpolated zero-crossing time stamp is a 41-bit unsigned integer (with a maximum value of 2^{31}). The trigger time tag is stored by default, but the fine time stamp can be retrieved separately. See Section 5.4.2 to learn how to retrieve the fine time stamp and/or the interpolated zero-crossing time stamp. Both the interpolated zero-crossing time stamp and the trigger time tag roll back to zero once they reach their maximum value of 2^{31} , or 2×2^{31} ns \approx 4.295 s. It is also possible to extend the rollback of the time stamps by an extra 16-bits by enabling the *extended time stamp*. ***CHECK THIS PARAGRAPH***

5.3.3 Event Readout Format

Each of the 16 V1730 digitizer channels uses RAM to temporarily store event data. This memory is divided into *memory buffers*, or *aggregates*, that each contain data for up to 1023 events. The number of events per aggregate is programmable, but we have set this number to its maximum of 1023 as to optimize readout bandwidth. The drawback of this is that events are not available for readout until all 1023 events have been stored in an aggregate, unless the run is stopped before an aggregate becomes full. The total number of aggregates contained in the RAM is also programmable, but we have set this number to 8 due to the large size of each aggregate. The RAM for each aggregate is cleared after all aggregates have been read to make room for upcoming events. ***CHECK THIS LAST PART***

In reality, there are two types of aggregates for the V1730 memory organization - *channel aggregates*, which are what have been described so far, and *board aggregates*. Each channel aggregate is shared by two consecutive channels, i.e. channel 0 and channel 1, channel 2 and

channel 3, etc. The sample of all channel aggregates is a board aggregate. The sample of all board aggregates is called the *data block*. Since there are 16 channels in the V1730, there are up to 8 channel aggregates that comprise a board aggregate. If a channel aggregate contains no data, it is not included in the board aggregate. When data readout is performed, each successive channel aggregate is read at a time in a given board aggregate, and each successive board aggregate is read from the data block via a *block transfer*.

Channel Aggregate Event Format

The channel aggregates are formatted for readout as in Figure 5.6. Each line represents a *memory location*, consisting of a 32-bit integer (each bit is either a 0 or 1). Note that only 2 events are shown, and a full channel aggregate will contain 1023 events. The first two memory locations, SIZE and FORMAT, comprise the channel aggregate *header*, which is common to all events in the aggregate. Beyond the header is data for each event in the aggregate. Our V1730 digitizer does not save individual samples, so every memory location that contains a sample, i.e. S_1 , S_2 , etc., does not appear for readout. Therefore, each event contains a maximum of 3 memory locations - the trigger time tag, EXTRAS, and the charges from the long and short gates (Q_{Long} and Q_{Short} , respectively). EXTRAS contains extra information associated with CFD zero crossing interpolation. It is therefore omitted by default, unless the user has enabled CFD zero crossing interpolation (see Section 5.4.2 for EXTRAS options). Note that if CFD mode is enabled, the trigger time tag is the coarse time stamp, associated with the sample before the zero crossing.

Not every bit in Figure 5.6 will be discussed here. See the CAEN DPP-PSD manual for detailed information [Ref]. However, the data most frequently accessed are as follows (from top to bottom in Figure 5.6). In the SIZE memory location, the CHANNEL AGGREGATE SIZE is the number of memory locations that are contained in the given channel aggregate, including itself and FORMAT. This can be useful for debugging and determining how many events are contained in the aggregate. In the FORMAT memory location, EX contains 3 bits that the user inputs (through functions in the frontend DAQ) to specify what the EXTRAS memory location outputs. See Section 5.4.2 for the specific options. If all 3 of these bits are 0, the EXTRAS memory location is omitted. In the event memory locations, CH specifies which channel the event came from in the couple (0 for even or 1 for odd). PUR is a flag that detects events whose energy was not evaluated correctly (e.g. from pile-up or event saturation).

Board Aggregate Event Format

The board aggregates are formatted for readout as in Figure 5.7. Each board aggregate contains up to 8 channel aggregates and has 4 memory locations making up the header. Each channel aggregate that has data contributes all of its memory locations to the board aggregate in the order shown in Figure 5.7. Channel aggregates with no data do not contribute to the board aggregate.

Again, only the most important values in the board aggregate will be discussed here. The BOARD AGGREGATE SIZE is analogous to the CHANNEL AGGREGATE SIZE in that it is the total number of memory locations contained in the board aggregate, including all of the memory locations in the channel aggregates that comprise it. The DUAL CHANNEL MASK corresponds to the channel couples participating to the board aggregate. THE BOARD AGGREGATE COUNTER gives the current board aggregate count. The BOARD AGGREGATE TIME TAG gives the time that the board aggregate was created. This is not a physical quantity, but can be useful for debugging.

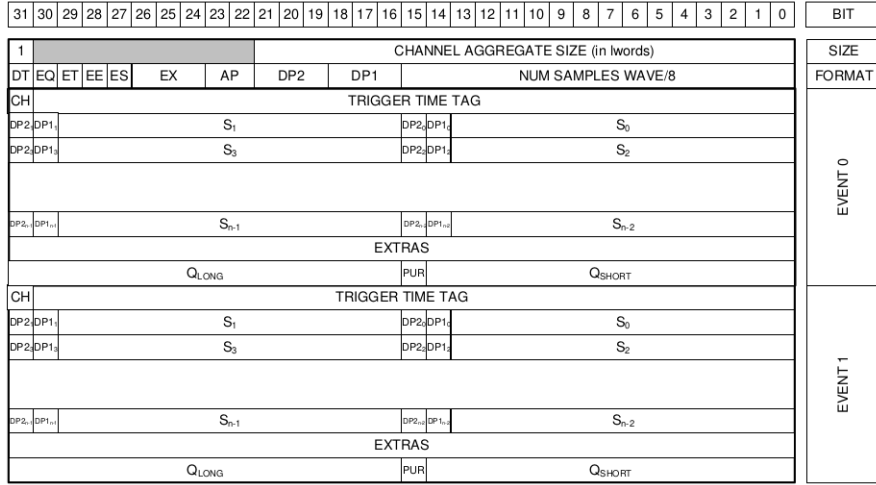


Figure 5.6: The data format for the first two events in a channel aggregate. Each memory location is 32-bits and is accessed during readout. Individual samples are not saved by our digitizer, so every 32-bit line that contains a sample, i.e. S₁, S₂, etc., is omitted. Each event therefore contains up to three memory locations. The channel aggregate itself has a header consisting of the first two memory locations - SIZE and FORMAT.

5.4 Frontend DAQ Software

5.4.1 V1730 Registers

The CAEN V1730 digitizer firmware uses Digital Pulse Processing (DPP) algorithms, which are implemented in the Field-Programmable Gate Array (FPGA) of the digitizer board. The DPP algorithms and settings can therefore be programmed by the user. This is performed by accessing and writing values to 32-bit wide *registers*, which have distinct hexadecimal addresses. For example, setting the width of the long gate for channel 0 amounts to writing the long gate value (as a 16-bit integer) into the Long Gate Width register specific to channel 0, which has the address 0x1058. All of the registers that the user can access are documented in the official V1730 DPP-PSD Registers manual [Ref]. We will highlight some of the most important ones in the next section.

An important aspect of our custom DAQ software is to provide a simple way of changing user-defined settings without the user having to manually change register values. In the DAQ software, the different settings are implemented via C++ functions that write the given parameter values into the appropriate register when executed. The function definitions are stored in the C file v1730DPP.c, while the functions are executed in the frontend C++ file fev1730-DPP.hxx. The most common settings, however, can be applied via the settings file, settings-DPP.dat.

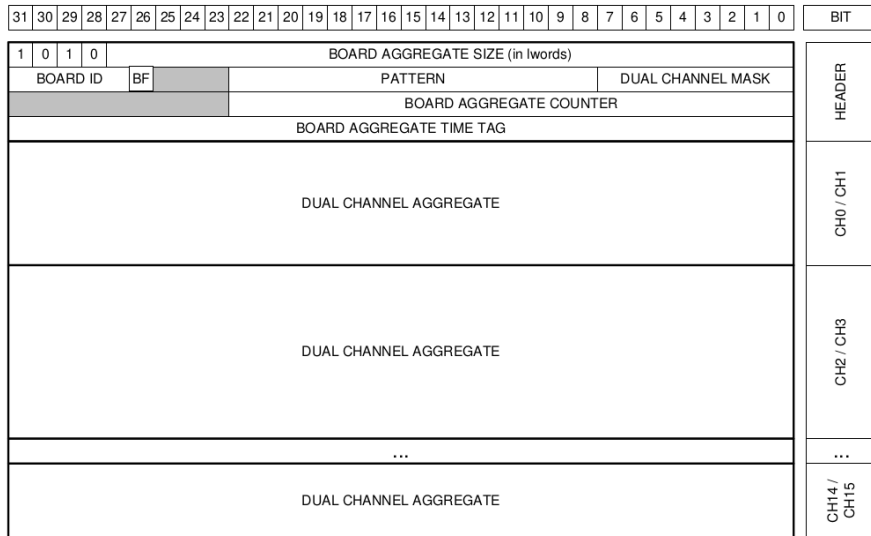


Figure 5.7: The data format for a board aggregate, consisting of up to 8 channel aggregates and 4 memory locations making up the header. Each memory location is 32-bits and is accessed during readout.

5.4.2 User Settings

Settings Determined from Oscilloscope Signals

LED vs. CFD Mode

Gain

Mean Baseline Calculation

Input Smoothing

Coincidences

Other Settings

5.4.3 Making Changes and Starting Runs

```
{0,1}      ## Enable Channels (0-15) e.g. --> {0,1,2,3}
700      ## Long gate (ns)
400      ## Short gate (ns)
110      ## Gate offset (ns)
0       ## Trigger Hold-off (ns, multiple of 8 - If too low, automatically adjusts to lowest possible value)
0       ## Pre Trigger (ns, multiple of 8, must be > Gate Offset - If too low, automatically adjusts to lowest possible value)
1       ## Gain (0-5)
1       ## negSignals (1=yes, 0=no)
32767    ## DC Offset (in DAC LSB Units)
3       ## Trigger threshold (in mV)
40      ## CFD Delay (ns)
75      ## CFD Fraction (%)

0       ## Dynamic Range (0 = 2 Vpp, 1 = 0.5 Vpp)
## Input Smoothing (0 = disabled, 1 = input pulse averaged over 2 samples, 2 = 4 samples, 3 = 8 samples, 4 = 16 samples)
## Mean Baseline Calc (0 = fixed baseline, 1 = 16 samples for mean baseline calculation, 2 = 64 samples, 3 = 256 samples, 4 = 1024 samples)
## Fixed Baseline (in mV, 0 = disabled, Mean Baseline Calc must be 0 if enabled)
## Baseline Calc Restart after Long Gate (0 = disabled (default), 1 = enabled)
## Trigger Mode (0 = LED mode, 1 = CFD mode)
## Trigger Counting Mode (0 = only accepted self-triggers (default), 1 = all self-triggers)
## Trigger Pile Up (0 = disabled (default), 1 = enabled)
## Detect Signals with Opposite Polarity (0 = enabled (default), 1 = disabled)
## Charge Zero Suppression Threshold (in LSB, 0 = disabled (default), cuts Q_Long < Q_thresh)
```

Figure 5.8: The settings file, settings-DPP.dat, where the most common user-defined DPP parameters are adjusted. Other common V1730 register settings can be adjusted here as well.

5.5 EngeSpec GUI

5.5.1 Overview of EngeSpec Operation

Sort Routine: EngeSort

Online MIDAS Runs

Offline MIDAS Runs

5.5.2 GUI Features

Histogram Display

Curve Fitting

Coincidences

Gating

5.6 Examples of DAQ/EngeSpec Operation

5.6.1 Resolution Tests with a NaI Detector

Energy Resolution

Timing Resolution (Coincidence Mode)

5.6.2 Coincidences with NaI Detectors

LED Mode

Timing Spectrum

2D Energy Spectrum

2D Energy vs. Coincidence Time Spectrum

Gating on Timing Spectrum

CFD Mode

Timing Spectrum

2D Energy Spectrum

2D Energy vs. Coincidence Time Spectrum

5.6.3 Dead Time Tests with Fixed Energy Pulser

CHAPTER

6

$^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ CONSTRAINTS FOR GLOBULAR CLUSTER NGC 2419

6.1 Introduction

This chapter details the original research constraining hydrogen-burning conditions responsible for abundance anomalies in globular clusters. In particular, the observed potassium enrichment and magnesium depletion in red giants of the globular cluster NGC 2419 is reproduced with nuclear reaction network calculations using new nuclear physics information presented in this chapter. A new reaction rate of the key potassium-destroying reaction, $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$, is calculated from the new proton partial-widths and resonance energies of astrophysically-important, proton-unbound ^{40}Ca states measured in the proton-transfer reaction $^{39}\text{K}({^3\text{He}}, d)^{40}\text{Ca}$. The reaction rate is significantly constrained for $T \lesssim 110$ MK. Monte Carlo network calculations will show new temperature-density constraints for hydrogen-burning, compared with that of the previously calculated reaction rate. Hot-bottom burning in Super-AGB stars is investigated as a polluter candidate driving the Mg–K anticorrelation in NGC 2419, as well as other abundance patterns involving potassium.

6.2 Globular Clusters

Globular clusters are compact conglomerates of stars associated with all types of galaxies. They are typically on the order of 1 pc to a few tens of pc in radius, and they are very old,

in most cases having an age of about 10 Gyr. Peak globular cluster formation is thought to pre-date most stellar formation in galaxies, and they may have played a crucial role in early galaxy formation [Gra19]. In the Milky Way galaxy, they often have low metallicity and are distributed throughout the halo, the thick disk, and the bulge. These properties, while interesting, are not what have made globular clusters garner considerable interest in the last few decades.

A characteristic recently attributed to globular clusters, which distinguishes them from similar objects such as open clusters, is the presence of chemical inhomogeneities among their low-mass stars. Once labeled *abundance anomalies*, these inhomogeneities take the form of anticorrelations among the abundances of light element pairs, such as O–Na and Mg–Al. A new definition of globular clusters that includes this characteristic was suggested by Ref. [Car10], since almost every globular cluster in which abundances have been observed has exhibited an O–Na anticorrelation. The only exceptions are Terzan 7 and Pal 12, where the simultaneous Na and O abundances of only 7 [Sbo07] and 4 [Coh04] stars were observed, respectively. Other anticorrelations and correlations are exhibited in some, but not all globular clusters, making each cluster distinguishable by the combination and extent of their abundance inhomogeneities. These cluster-to-cluster differences are primarily driven by differences in luminosity and metallicity. Meanwhile, no abundance anticorrelations have ever been observed in open clusters [Gra19].

Fig. 6.1 shows the ubiquitous O–Na anticorrelation in 1,200 red giants among all 19 globular clusters sampled by Ref. [Car10]. The red giants with an enhanced Na abundance are associated with a depletion in O abundance, and vice versa. The populations of red giants are split into 3 categories based on the extent of their Na enrichment, and associated O depletion. These populations are the primordial (P), extreme (E), and intermediate (I) components, as indicated in the first panel of Fig. 6.1. The primordial population is Na-poor/O-rich, the extreme population is Na-rich/O-poor, and the intermediate population is in-between. The primordial population is so-called because the same abundance pattern is typical of field stars with a similar metallicity. However, the Na and O abundances of the intermediate and extreme populations indicates that these red giants went through an unknown process. Low-mass stars like red giants and main sequence stars do not reach the high temperatures necessary for the nucleosynthesis chains that produce the observed Na and O abundances. This reasoning also applies to the other observed anticorrelations in globular clusters [Pra07].

The most likely scenario explaining these inhomogeneities is that the currently observed stars are part of a second generation which formed from the ashes of an older, first generation of stars. The material ejected from these more-massive first generation stars upon their death likely polluted the inter-cluster medium where the second generation stars began to form.

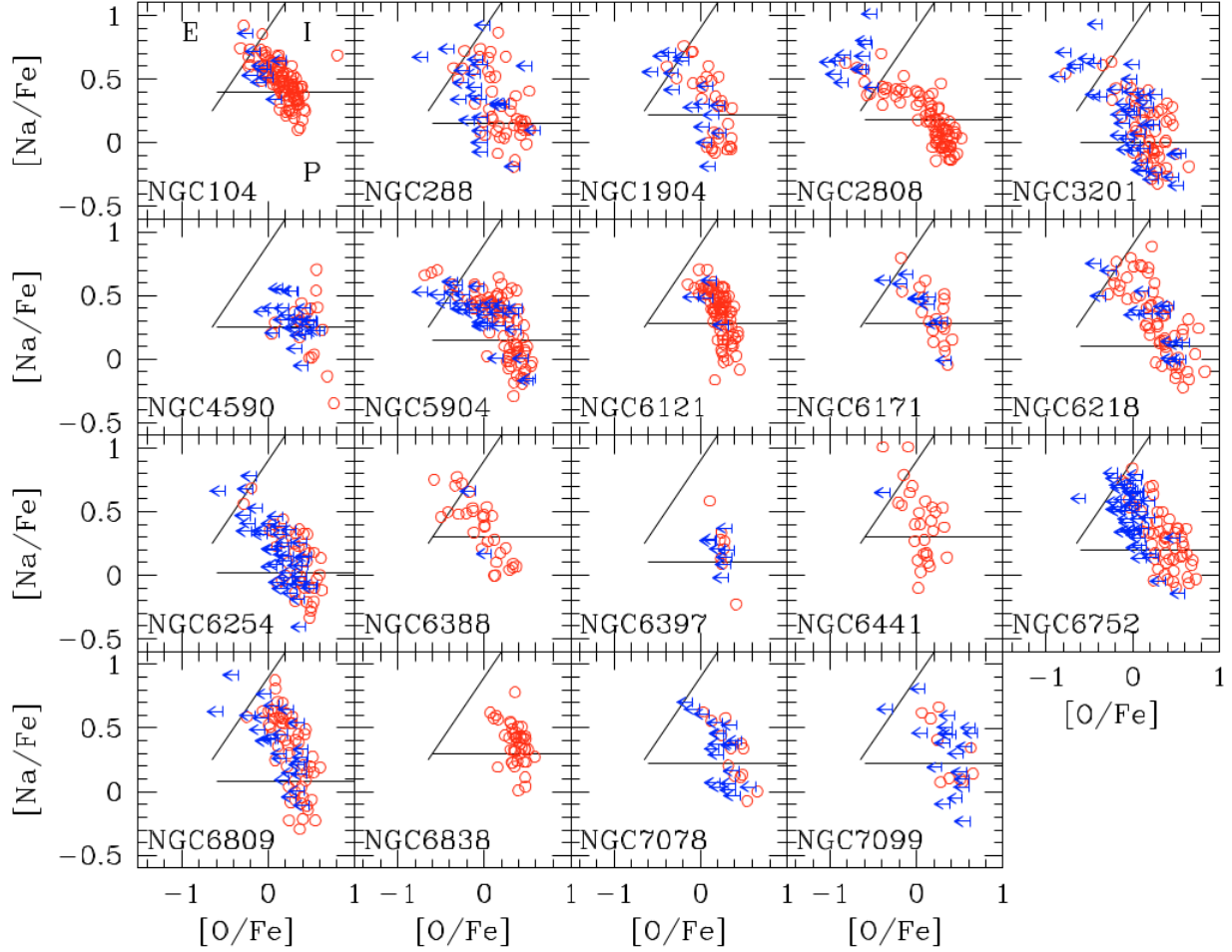


Figure 6.1: The O–Na anticorrelation of 1,200 red giants among all 19 globular clusters observed in the survey of Ref. [Car10]. Blue arrows indicate upper limits in O abundances, while lines separate different populations of red giants based on their relative O and Na richness. Primordial (P), extreme (E), and intermediate (I) populations are labeled in the first panel. See text for definitions of these populations. Adapted from [Car10].

Hence, these first generation stars are sometimes called *polluter stars*, a class of massive, extinct stars that are the origin of the apparent abundance anomalies. Different generations of stars existing in a given globular cluster would provide evidence for *multiple stellar populations*. This premise has been widely debated over the last few decades, with most research supporting it at present [Gra04, Gra12, Gra19].

Polluter stars are likely the site of the nucleosynthesis that gives rise to the currently observed abundance patterns in globular clusters. Their exact nature is unknown, but the nucleosynthesis mechanism driving the inhomogeneities in the vast majority of globular clusters is well established as proton-capture reactions in high temperature hydrogen-burning

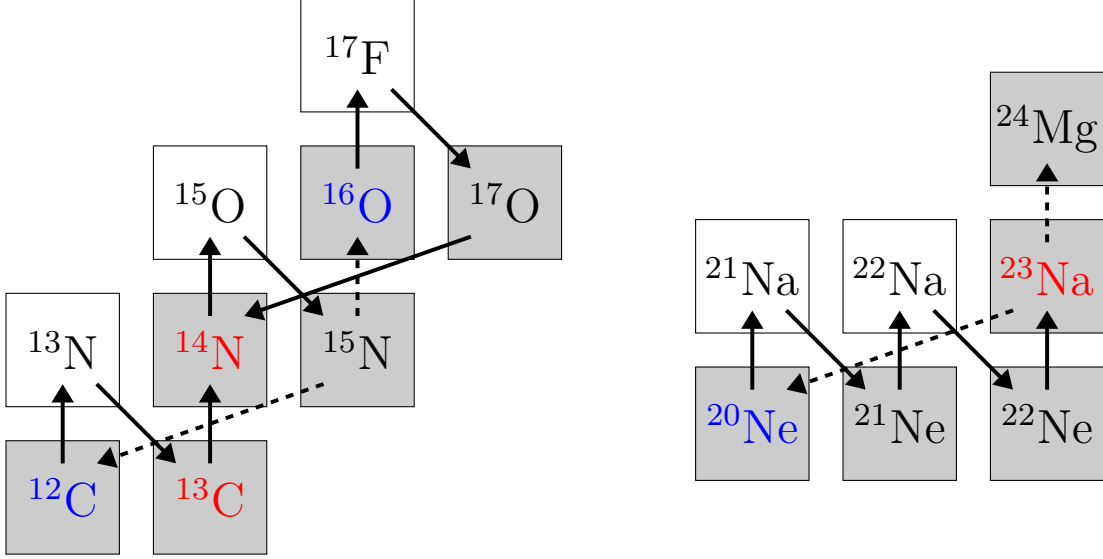


Figure 6.2: The hydrogen burning mechanism driving the O–Na anticorrelation. Nuclei in blue are destroyed overall in the various cycles, while nuclei in red are produced. Shaded nuclei are stable. The first two stages of the CNO cycle (left), distinguished by the temperature-dependent branch at ^{15}N , lead to an overall depletion of oxygen. The $^{15}\text{N}(p, \gamma)^{16}\text{O}$ reaction is activated at about 40 MK. The Ne–Na cycle (right), occurring simultaneously at about the same activation temperature, leads to an overall production of sodium. The $^{23}\text{Na}(p, \gamma)^{24}\text{Mg}$ reaction activates at about 70 MK.

environments [Den89, Lan93]. The ubiquitous O–Na anticorrelation, for example, is exhibited in the hydrogen burning of the CNO and Ne–Na cycles at about 40 MK [Gra19]. Fig. 6.2 shows the nucleosynthesis chains that lead to an overall oxygen depletion and sodium production. The first stage of the CNO cycle involves a series of (p, γ) and $(\beta^+\nu)$ reactions on stable and radioactive nuclei, respectively, starting from ^{12}C . This stage occurs at fusion temperatures of about 10 MK. Once ^{15}N is reached, $^{15}\text{N}(p, \alpha)^{12}\text{C}$ closes the cycle if the temperature is less than about 40 MK. Otherwise, the second stage of the CNO cycle will be activated from $^{15}\text{N}(p, \gamma)^{16}\text{O}$. This eventually leads to ^{17}O , which closes the second stage of the cycle via $^{17}\text{O}(p, \alpha)^{14}\text{N}$. This second stage leads to an overall destruction of oxygen. Meanwhile, the Ne–Na cycle activates at about 40 MK as well, starting from $^{20}\text{Ne}(p, \gamma)^{21}\text{Na}$. Once ^{23}Na is reached, it will close the cycle via $^{23}\text{Na}(p, \alpha)^{20}\text{Ne}$ if temperatures are less than about 70 MK, otherwise $^{23}\text{Na}(p, \gamma)^{24}\text{Mg}$ is activated. These destruction reactions on ^{23}Na are slow enough such that sodium is produced overall.

Although nucleosynthesis via hydrogen burning is what drives the observed abundance patterns, it is not the only mechanism responsible for them. The majority of second generation

stars must be composed of a mixture of nuclear-processed ejecta and matter with a pristine composition, in order for nuclear reaction network models to reproduce the observed globular cluster abundance patterns [Pra07]. The nuclear-processed ejecta is the matter resulting from nucleosynthesis via hydrogen burning in the first generation polluter stars, while pristine matter refers to the unprocessed gas left behind from the first burst of star formation. Many polluter star sites such as AGB stars, fast-rotating massive stars, and supermassive stars require dilution with unprocessed gas in order to convert their correlated O and Na abundances into the observed anticorrelation [Erc10, Erc11, Erc12]. The observed, mixed abundances in a typical dilution model are obtained by diluting one part of processed matter with f parts of pristine matter, as in

$$X_{\text{mix}} = \frac{X_{\text{proc}} + f X_{\text{pris}}}{1 + f}, \quad (6.1)$$

where X is the mass fraction of a given nuclide among its specific composition and f is the dilution factor [Pra07, Car09]. A very small dilution factor represents nearly pure processed matter, while a very large dilution factor results in mostly pristine matter. The dilution model works very well for most globular clusters, but often a single dilution model does not simultaneously reproduce the abundances of the extreme and intermediate populations. In these cases, more than one class of polluter stars is required to match the observations, suggesting that the multiple populations premise may be more complex than previously considered [Gra19].

6.3 NGC 2419 and the Mg–K Anticorrelation

While the O–Na anticorrelation is ubiquitous in globular clusters, there are other abundance patterns that are unique to individual or a small subset of globular clusters. The globular cluster NGC 2419 was recently found to exhibit some puzzling abundance patterns. About 30% of the red giants observed show a strong K enrichment correlated with a considerable Mg depletion [Muc12, Coh12]. Fig. 6.3 shows the observed elemental potassium and magnesium abundances, with respect to iron, for the red giants in NGC 2419 sampled by Ref. [Muc12] in red and Ref. [Coh12] in blue. This was the first discovery of a Mg–K anticorrelation in any globular cluster. Only one other cluster to date, NGC 2808, has exhibited such an anticorrelation in a small portion of its stars [Muc15, Muc17], but the extent of the K enrichment is about 7 times less than that of NGC 2419. Meanwhile, the extent of the Mg depletion is about 3 times less. The evolutionary history of NGC 2419 must therefore be rather unique for it to exhibit such abundance patterns not seen in other globular clusters.

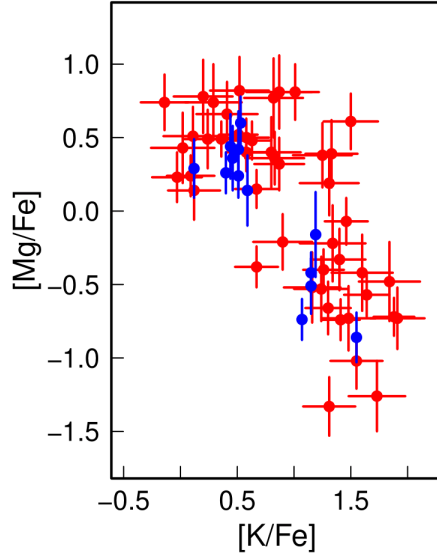


Figure 6.3: The observed Mg and K elemental abundances of red giants in NGC 2419 from Refs. [Muc12] (red) and [Coh12] (blue). Figure adapted from [Ili16].

Just as the nucleosynthesis mechanism for the O–Na anticorrelation involves a series of (p, γ) reactions and radioactive decays, the same is true for the Mg–K anticorrelation. The hydrogen burning schemes are illustrated in Fig. 6.4. The Mg–Al cycle, shown on the left-hand side of the figure, starts from ^{24}Mg at about 70 MK. After the subsequent $^{24}\text{Mg}(p, \gamma)^{25}\text{Al}$ and $^{25}\text{Al}(\beta^+\nu)^{25}\text{Mg}$ reactions, there is a chance the $^{26}\text{Al}^m$ isomer is populated. The isomer preferentially β^+ decays to ^{26}Mg , while the ground state $^{26}\text{Al}^g$ preferentially captures a proton. Either scenario eventually synthesizes ^{27}Al , which can repeat the cycle with $^{27}\text{Al}(p, \alpha)^{24}\text{Mg}$ or proceed with $^{27}\text{Al}(p, \gamma)^{28}\text{Si}$ if the temperature is more than about 80 MK. The overall effect is a depletion of magnesium and an enrichment of aluminum and silicon.

The hydrogen burning scheme for the K enrichment in NGC 2419 is illustrated on the right-hand side of Fig. 6.4. It has been shown to start from ^{36}Ar , an α -nucleus and the most abundant argon isotope at the low metallicities of the first stellar generation [Ven12]. Some authors [Ven12, Muc15] have incorrectly claimed that the main K enrichment chain involves the decay of ^{37}Ar to ^{37}Cl , then the subsequent $^{37}\text{Cl}(p, \gamma)^{38}\text{Ar}$ reaction, indicated by the dashed arrows in Fig. 6.4. However, as Ref. [Ili16] indicates, the $^{37}\text{Ar}(p, \gamma)^{38}\text{K}$ reaction has a much larger decay constant than the electron capture reaction $^{37}\text{Ar}(e^-, \nu)^{37}\text{Cl}$ at the stellar densities of interest. Therefore, the main chain must proceed to ^{38}K before the subsequent $^{38}\text{K}(\beta^+\nu)^{38}\text{Ar}$ and $^{38}\text{Ar}(p, \gamma)^{39}\text{K}$ reactions. The former chain is expected to contribute only marginally to ^{39}K nucleosynthesis. Additionally, unlike the mechanisms

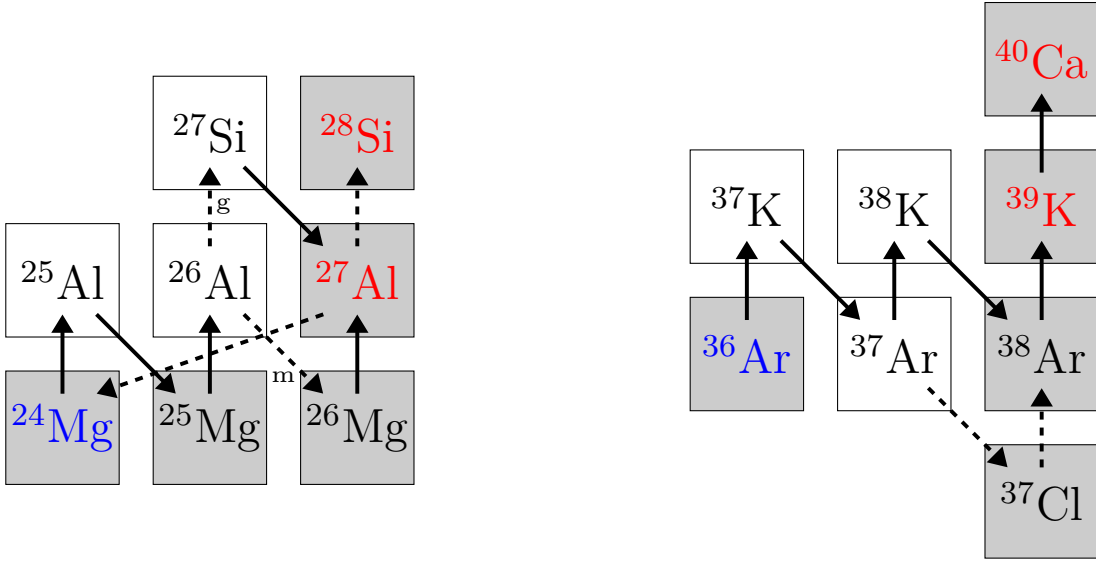


Figure 6.4: The hydrogen burning mechanism driving the Mg–K anticorrelation in the globular cluster NGC 2419. The nucleosynthesis chain for the Mg depletion is the Mg–Al cycle, shown on the left-hand side. Whether the $^{26}\text{Al}^m$ isomer or the $^{26}\text{Al}^g$ ground state is populated, the result is ^{27}Al production, which repeats the cycle with $^{27}\text{Al}(p, \alpha)^{24}\text{Mg}$. The main nucleosynthesis chain for the K enrichment is shown on the right-hand side with solid arrows. The $^{37}\text{Ar}(p, \gamma)^{38}\text{K}$ reaction proceeds at a much higher rate than $^{37}\text{Ar}(e^-, \nu)^{37}\text{Cl}$ electron capture at the stellar densities of interest. Nuclides in blue are destroyed and nuclides in red are produced in the overall nucleosynthesis. Shaded nuclides are stable.

for the O–Na anticorrelation and the Mg depletion, the sequence leading from ^{36}Ar to ^{39}K is not a cycle because the $^{39}\text{K}(p, \alpha)^{36}\text{Ar}$ reaction has a much smaller decay constant than $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$. However, as it will become clear in Section 6.3, the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate is currently very uncertain in the astrophysical temperature range of interest.

The nucleosynthesis mechanism for the Mg–K anticorrelation may be well-established, but the site of this nucleosynthesis is much more uncertain. Ref. [Ven12] were the first to propose polluter star candidates for the Mg–K anticorrelation in NGC 2419, as well as the other observed abundance patterns. Of consideration were massive AGB stars and Super-AGB (SAGB) stars of $\sim 6 M_\odot$, where hydrogen burning occurs during hot-bottom burning (HBB) at the base of the convective hydrogen envelope. Agreement was found with the observed intermediate K-enriched population only in an artificially optimized scenario. The $^{38}\text{Ar}(p, \gamma)^{39}\text{K}$ reaction rate would need to increase by a factor of 100, and the poorly known mass loss rate in the AGB models would need to simultaneously decrease by a factor of 4. However, Ref. [Ven12] does not investigate the effect of varying other relevant reaction

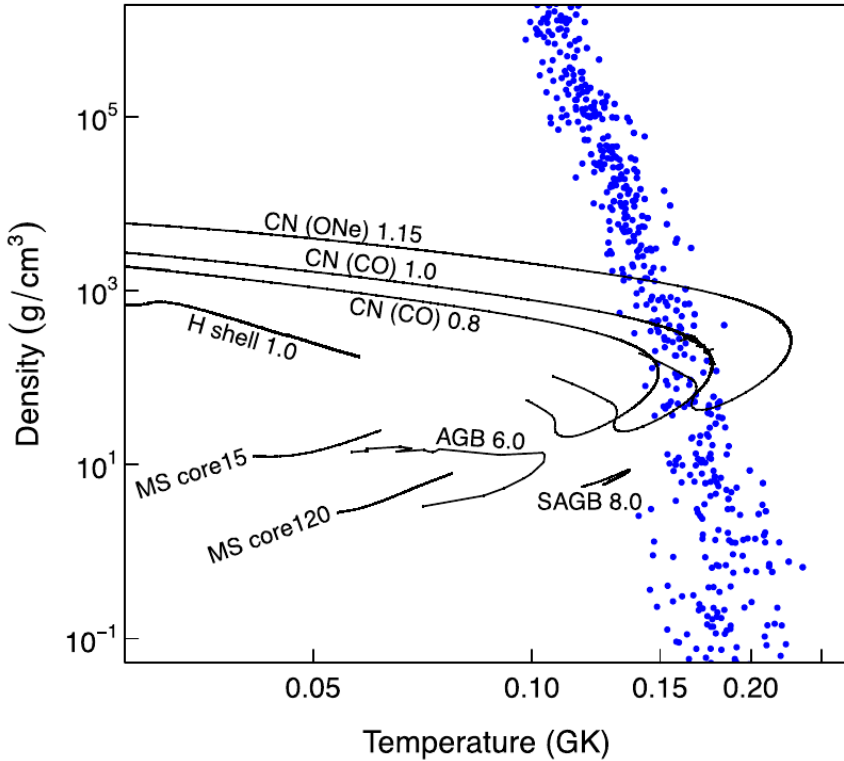


Figure 6.5: Temperature-density conditions reproducing the Mg–K anticorrelation and other abundance patterns in NGC 2419, obtained by sampling T , ρ , X_H , reaction rate probability densities, and initial abundances from a nuclear reaction network (see text). The T – ρ conditions for several polluter star candidates are represented by the black curves. Adapted from Ref. [Ili16].

rates, which could have the same effect as increasing the $^{38}\text{Ar}(p, \gamma)^{39}\text{K}$ rate.

Ref. [Ili16] expanded the search for polluter candidates by performing a Monte Carlo nuclear reaction network calculation that reproduced all of the observed abundances in NGC 2419, including the Mg–K anticorrelation. The parameters that were randomly sampled include the temperature T , density ρ , final hydrogen mass fraction X_H , all reaction rate probability densities in the network, and initial abundances. T and ρ were held constant for each network calculation. The reaction rates were obtained by truncating the Starlib [Sal13] library for the relevant rates. The resulting T – ρ conditions that matched all of the observed abundances are shown by the blue dots in Fig. 6.5. A narrow band of possible T – ρ conditions was found between $\approx 80 - 260$ MK and $\approx 10^{-4} - 10^8$ g/cm 3 , with additional solutions not shown in the figure up to 10 11 g/cm 3 . The black curves represent the T – ρ hydrogen burning conditions of several polluter star candidates. These include the cores of 15 M $_\odot$ and 120 M $_\odot$ main sequence (MS) stars, the hydrogen-burning (H) shell of a 1 M $_\odot$

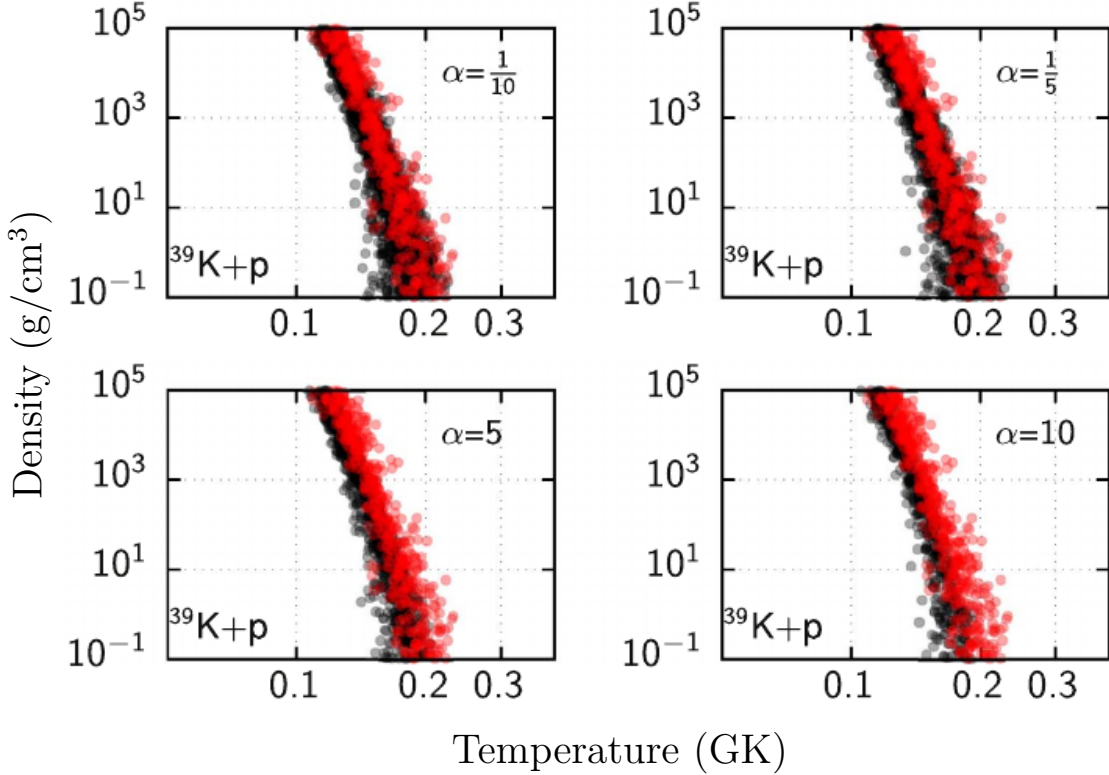


Figure 6.6: Systematic effects of the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate influencing temperature-density conditions. The indicated variation factors ($\alpha = 1/10, 1/5, 5, 10$) are applied to the reaction rate in each panel, and the black dots show the resulting temperature-density conditions that provide an acceptable match with observed abundances. The red dots represent the case where no systematic effects ($\alpha = 1$) have been added. Figure adapted from [Der17].

red giant star, hot-bottom burning of a $6 M_{\odot}$ and $8 M_{\odot}$ thermally-pulsing AGB and SAGB star, respectively, a $1.15 M_{\odot}$ ONe classical nova (CN), and $0.8 M_{\odot}$ and $1.0 M_{\odot}$ CO classical novae. The only candidates that nominally overlap with this band are ONe and CO classical novae. However, another candidate Ref. [Ili16] considers SAGB stars because of the large uncertainty in AGB stellar model parameters, such as the mass loss rate and the prescription of convective mixing. Adjusting these parameters within their current uncertainties could increase the temperature of HBB enough to match the observed abundances.

While Ref. [Ili16] simultaneously sampled the reaction rate probability densities of all the reactions in the network, they did not investigate the effect of individual reaction rates. Ref. [Der17] investigated the sensitivity of the acceptable $T-\rho$ conditions for NGC 2419 polluter stars on unknown systematic effects of individual reaction rates. Only 4 reactions were found to make a significant impact on the acceptable $T-\rho$ conditions, $^{30}\text{Si}(p, \gamma)^{31}\text{P}$, $^{37}\text{Ar}(p, \gamma)^{38}\text{K}$, $^{38}\text{Ar}(p, \gamma)^{39}\text{K}$, and $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$. If the reaction rate of $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$, for example, were

systematically larger than its recommended rate, by even as little as a factor of 5, the acceptable T - ρ band would have significantly reduced scatter and would be constrained to the low-temperature side for all densities. This scenario is shown in Fig. 6.6 on the bottom-left panel, where the black dots represent the new rate multiplied by the indicated variation factor α , and the red dots represent the recommended rate ($\alpha = 1$). This makes sense in theory because potassium abundance would be reduced if the rate of its primary destruction reaction were increased. The high temperature conditions would therefore destroy too much potassium compared to its observed abundance. In contrast, if the rate were systematically smaller by a factor of 5, the solutions interestingly only have increased scatter. These same effects are also true for $^{30}\text{Si}(p, \gamma)^{31}\text{P}$, but the opposite is true for $^{37}\text{Ar}(p, \gamma)^{38}\text{K}$ and $^{38}\text{Ar}(p, \gamma)^{39}\text{K}$, as these latter reactions increase the potassium abundance.

6.4 The Previous $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ Reaction Rate

The remainder of this chapter will focus specifically on the key potassium-destroying reaction $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$, found by Ref. [Der17] to be one of the few important reactions able to constrain the potassium abundance in NGC 2419 and therefore constrain the possible T - ρ conditions of its polluter stars.

6.4.1 Longland *et al.* (2018) Evaluation

The most recent $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate evaluation is that of Ref. [Lon18b], which Ref. [Der17] used while it was in preparation. Its probability density was calculated using a Monte Carlo reaction rate formalism [Lon10] with resonance strengths and resonance energies provided by the direct measurements of Refs. [Kik90, Che81, Lee66]. The lowest directly measured resonance strength from these experiments is for the resonance at $E_r^{\text{c.m.}} = 606$ keV. For resonances below this, indirect $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ [Cag71] and $^{39}\text{K}(d, n)^{40}\text{Ca}$ [Fuc69] proton-transfer measurements provided proton partial-widths, when available. Additionally, the indirect $^{36}\text{Ar}(^6\text{Li}, d)^{40}\text{Ca}$ α -transfer measurement of Ref. [Yam94] provided α -particle partial widths for the (p, α) channel, when available. Otherwise, measured or theoretical upper limits were given to the single particle reduced widths of the remaining resonances. These upper limits are described by the Porter-Thomas distribution [Por56, Wei09], truncated at the upper limit value in the case of measured upper limits. This provided a physically motivated probability density from which to sample unknown or upper-limit partial widths.

The reaction rate probability density evaluated by Ref. [Lon18b] is shown in Fig. 6.7. The reaction rate uncertainty is represented on the y -axis as a factor of the mean, recommended

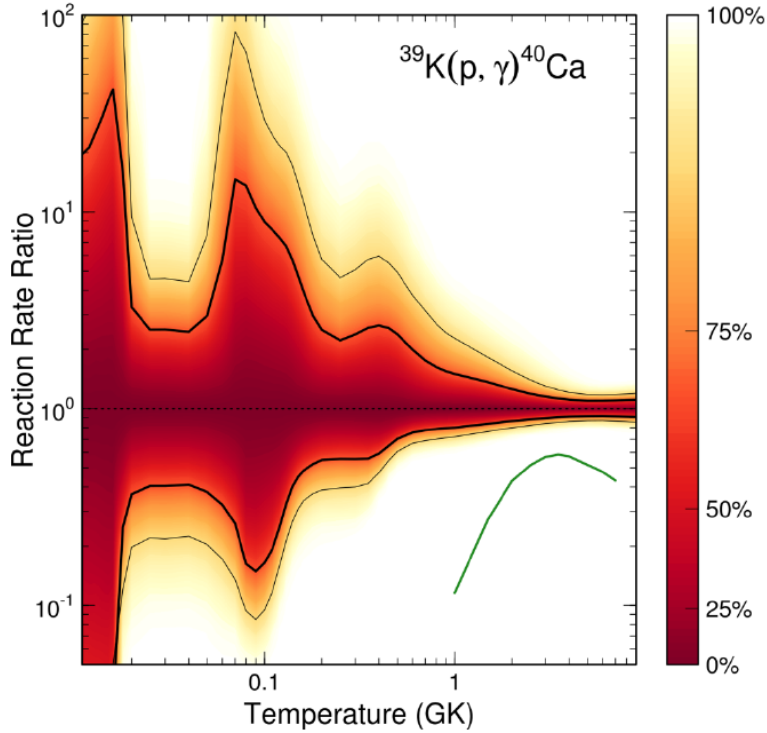


Figure 6.7: The recent $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate probability density calculation of [Lon18b] as a function of temperature. The median, recommended rate is shown as the dotted normalization line. The thick and thin black lines represent the 68% (1σ) and 95% (2σ) uncertainty bands, respectively. The color scale shows the continuous nature of the probability density, with darker red colors closer to the recommended rate. The green line represents the previous calculation of Ref [Che81].

rate, which has been normalized to unity. The 68% (1σ) and 95% (2σ) uncertainty bands are represented by the thick and thin black lines, respectively, while the color scale indicates the continuity of the probability density. Clearly, the rate uncertainty is very large between about 0.05 – 0.2 GK (50 – 200 MK), where the total 1σ width peaks at a factor of 84 at about 80 MK. Recall from Section 6.3 that the astrophysically relevant temperature range spanning the $T-\rho$ band in Figs. 6.5 and 6.6 is $\approx 80 - 260$ MK. The most recent $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate is therefore very uncertain in the astrophysical temperature range of interest. This is concerning, given that Ref. [Der17] found that the $T-\rho$ conditions are sensitive to systematic variations in the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ rate, by even as little as a factor of 5. This motivates the necessity of constraining the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ rate, particularly for the contributing resonances between $\approx 50 - 260$ MK.

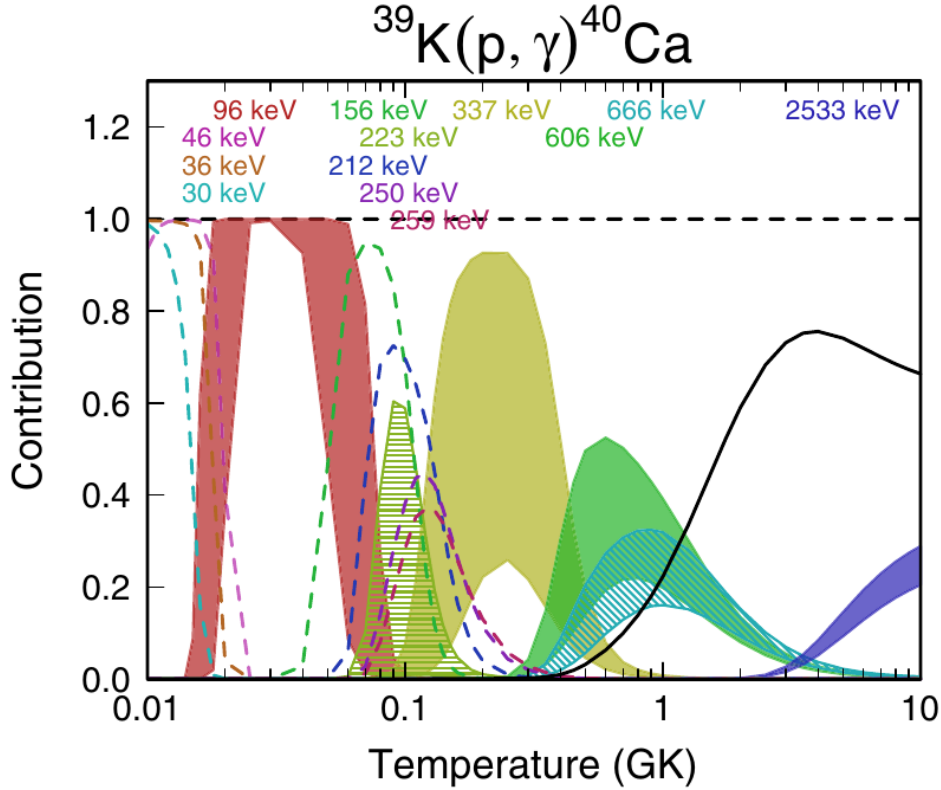


Figure 6.8: The individual resonance contributions to the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate from Ref. [Lon18b] as a fraction of unity. Measured resonances are represented with shading or hatched lines and are shown with their 1σ uncertainty bands. Resonances from upper limits are represented by dashed lines and show their 84% ($+1\sigma$) value. The black line represents the total resonance contribution of all resonances that individually contribute less than 20% to the total reaction rate.

6.4.2 Contributing Resonances

The $^{39}\text{K} + p$ resonances that contribute to the reaction rate in the $\approx 50 - 260$ MK range span resonance energies of $E_r^{\text{c.m.}} \approx 90 - 450$ keV (See Section 2.2) and ^{40}Ca excited state energies of about $E_x \approx 8400 - 8800$ keV, where the most recent ^{40}Ca proton separation energy measurement is $S_p = 8328.18(2)$ keV from Ref. [Wan21]. This resonance energy range is below the lowest directly measured resonance at $E_r^{\text{c.m.}} = 606$ keV ($E_x = 8935$ keV), indicating that this region is entirely described by inferred transfer reaction partial widths or upper limits at present.

The contributing resonances from the reaction rate evaluation of Ref. [Lon18b] are shown in Fig. 6.8. The individual, fractional resonance contribution for each resonance is shown as a function of temperature. Measured resonances are shown with their 1σ width, representative of the Monte Carlo formalism [Lon10] and are shaded or have hatched lines for clarity. Upper

limit resonances are represented by dashed lines at their 84% ($+1\sigma$) value. All resonances that individually contribute at least 20% to the total reaction rate are shown in the figure with corresponding resonance energy labels, while the remaining resonance contributions sum to the black line. Of note are the 96 keV, 156 keV, and 337 keV resonances, which overwhelmingly contribute to the reaction rate at their respective temperatures. The 337 keV resonance, in particular, is the most important resonance between $\approx 0.15 - 0.4$ GK (150 – 400 MK), which makes up most of the astrophysical temperature range of interest, 80 – 260 MK. However, the discrepancy between the 84% ($+1\sigma$) contribution and the 16% (-1σ) contribution is very large. The large uncertainty in the reaction rate around 50 – 200 MK in Fig. 6.7 is partly due to the uncertainty in this resonance contribution, but it is mostly a result of the many unknown resonances with only upper limit values between the 96 keV resonance and the 337 keV resonance.

6.4.3 Previous Proton-Transfer Measurements

The previous ${}^{39}\text{K}({}^3\text{He}, d){}^{40}\text{Ca}$ [Ers66, Set67, For70, Cag71] and ${}^{39}\text{K}(d, n){}^{40}\text{Ca}$ [Fuc69] proton-transfer measurements were not capable of achieving high resolution, particularly for unbound ${}^{40}\text{Ca}$ states, i.e. at energies above the proton separation energy. The $({}^3\text{He}, d)$ measurements of Refs. [Ers66, Set67, Cag71] obtained spectroscopic factors (C^2S) for only 2 unbound ${}^{40}\text{Ca}$ states, with ENSDF [Che17] excitation energies of $E_x = 8425$ keV ($E_r^{\text{c.m.}} = 96$ keV) and 8551 keV ($E_r^{\text{c.m.}} = 223$ keV), which are both below the lowest directly measured resonance at $E_x = 8935$ ($E_r^{\text{c.m.}} = 606$ keV). Meanwhile, Ref. [For70] did not observe unbound ${}^{40}\text{Ca}$ states due to their focus on lower energy excited states. The (d, n) measurement of Ref. [Fuc69] obtained C^2S for 14 unbound ${}^{40}\text{Ca}$ states, with only 4 of these states below the lowest directly measured resonance, including 8359 keV ($E_r^{\text{c.m.}} = 30$ keV), 8425 keV ($E_r^{\text{c.m.}} = 96$ keV), 8551 keV ($E_r^{\text{c.m.}} = 223$ keV), and 8665 keV ($E_r^{\text{c.m.}} = 337$ keV). Hence, there is an overlap between the $({}^3\text{He}, d)$ and (d, n) measurements of 2 unbound ${}^{40}\text{Ca}$ states, 8425 keV ($E_r^{\text{c.m.}} = 96$ keV) and 8551 keV ($E_r^{\text{c.m.}} = 223$ keV). The ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ reaction rate calculation of Ref. [Lon18b] used the C^2S measurements of Ref. [Cag71] for these 2 states, due to their more sophisticated coupled-channel DWBA calculations. They used the C^2S measurement of Ref. [Fuc69] for the 8665 keV ($E_r^{\text{c.m.}} = 337$ keV) state. However, the reaction rate calculation did not include the 8359 keV ($E_r^{\text{c.m.}} = 30$ keV) C^2S measurement of Ref. [Fuc69], as this state has an energy discrepancy with ENSDF [Che17] of 12 keV. This resonance was instead given an upper limit. Several unknown resonances remain below $E_x = 8935$ keV ($E_r^{\text{c.m.}} = 606$ keV). Ref. [Lon18b] uses upper limits for 18 resonances in this region. Hence, this was strong motivation to revisit the proton-transfer reaction at modern resolution to potentially resolve the currently

unknown resonances.

6.5 The $^{39}\text{K}(\text{He}, d)^{40}\text{Ca}$ Experiment

The $^{39}\text{K}(\text{He}, d)^{40}\text{Ca}$ experiment was performed in 2019 over 5 days using the Enge Split-Pole Spectrograph at the Triangle Universities Nuclear Laboratory (TUNL). The 10 MV FN tandem Van de Graaff accelerator at TUNL accelerated a fully-ionized ^3He beam to $E_{\text{lab}} = 21$ MeV, where the energy was stabilized using a pair of high-resolution slits between two 90° dipole magnets (see Section ?? for experimental details). The $^{39}\text{K}(\text{He}, d)^{40}\text{Ca}$ proton transfer and $^{39}\text{K}(\text{He}, \text{He})^{39}\text{K}$ elastic scattering reactions were measured separately between $\theta_{\text{lab}} = 5 - 20^\circ$ and $\theta_{\text{lab}} = 15 - 59^\circ$, respectively, using the focal plane detector package of Ref. [Mar19]. The d and ^3He particles that were ejected from the target at the angle θ_{lab} passed through the magnetic field B of the Enge Split-Pole Spectrograph, where they were then focused at a position along the focal plane, based on their magnetic rigidity $B\rho$ (See Sec. ??). The focal-plane detector was preemptively moved via dual motors to align itself with the focal plane. Meanwhile, for every $^{39}\text{K}(\text{He}, d)^{40}\text{Ca}$ or $^{39}\text{K}(\text{He}, \text{He})^{39}\text{K}$ focal-plane measurement, the $^{39}\text{K}(\text{He}, \text{He})^{39}\text{K}$ reaction was simultaneously measured with a silicon detector telescope inside the target chamber, positioned at a constant 45° from the beamline to monitor potential target degradation and other target properties.

6.5.1 Targets

A total of 7 natural KI (93.26% ^{39}K , 6.73% ^{41}K) targets were made throughout the course of the experiment, in 2 separate evaporation batches (see Section ?? for details on the evaporation procedure). The targets labeled KI #1, KI #2, and KI #3 were made in the first batch, while those labeled KI #4, KI #5, KI #6, and KI #7 were made in the second batch, on the second day of the experiment. For each batch, the targets were simultaneously produced by evaporating approximately $75 \mu\text{g}/\text{cm}^2$ of natural KI onto aluminum target frames, each with a $21 \mu\text{g}/\text{cm}^2$ natural carbon (98.84% ^{12}C , 0.96% ^{13}C) foil backing. The KI target thickness was measured by a thickness monitor inside the evaporation chamber, placed at roughly the same distance from the evaporation material as were the aluminum target frames with carbon backings. A separate $21 \mu\text{g}/\text{cm}^2$ natural carbon target was also used in the experiment for calibrations with the $^{12}\text{C}(\text{He}, d)^{13}\text{N}$, $^{13}\text{C}(\text{He}, d)^{14}\text{N}$, and $^{16}\text{O}(\text{He}, d)^{17}\text{F}$ reactions, among other contaminants.

Potassium iodide is hygroscopic, meaning it will eventually oxidize at atmospheric pressure. To prevent this, care was taken to transport the targets from the evaporation chamber to the

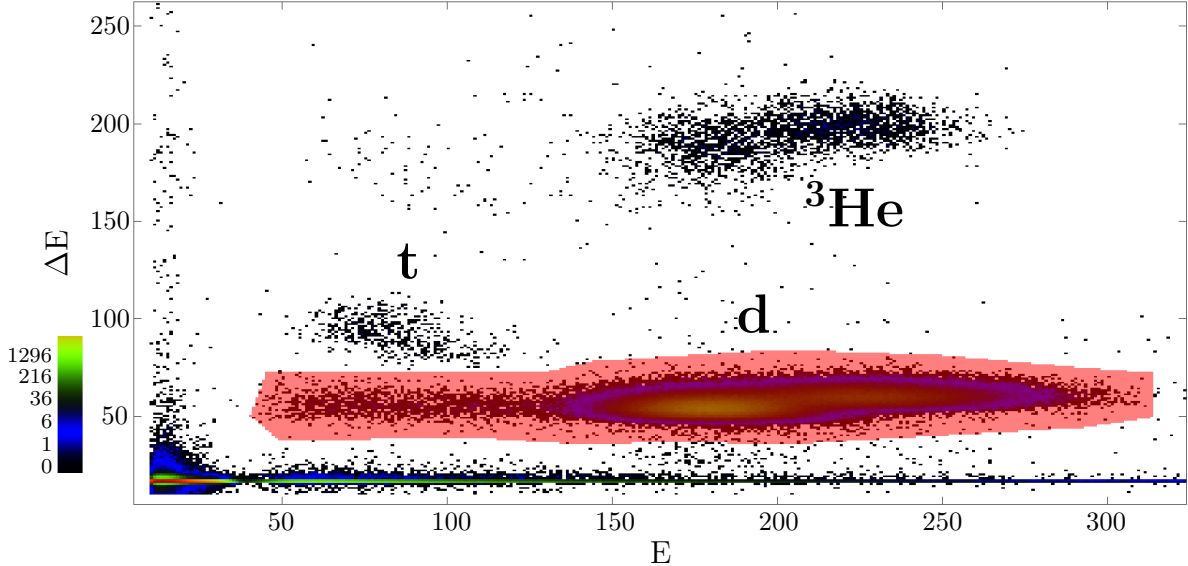


Figure 6.9: A 2D histogram of the focal-plane ΔE vs E detector signals. Different groups correspond to different particles depending on their mass and charge. The deuteron group, highlighted in red, is being gated on in the figure.

target chamber in a vacuum-sealed box at rough vacuum ($\sim 10^{-2}$ torr), where they were then exposed to high vacuum ($\sim 10^{-6}$ torr) for the remainder of the experiment. However, as will be described later, the first batch of targets (KI #1, KI #2, and KI #3) oxidized at some point during the first day of the experiment, for the $\theta_{\text{lab}} = 5^\circ$ and 7° (${}^3\text{He}, d$) measurements. The focal-plane spectra for these runs, exclusively using the KI #1 target, consisted of peaks with high-energy tails, expected of targets that undergo oxidation [Lan44]. These angles were measured again toward the end of the experiment with the non-oxidized KI #6 target, but the oxidized KI #1 data was not discarded nevertheless. Section 6.6.2 details a novel technique for fitting these spectra, an example of which is shown in Section 6.5.5.

6.5.2 Focal-Plane Spectra

The focal-plane detector package (see Section ?? and Ref. [Mar19]) consists of two position-sensitive avalanche counters, one located in the front of the detector (P1 section) and one located near the back (P2 section), a gas proportionality counter (ΔE section), and a residual energy scintillator (E section). The data acquisition system (DAQ) triggers on the E section, located at the back of the detector, so that coincidences with all sections are established (see Section ??). The combination of the ΔE and E sections enables particle discrimination based on mass and charge, allowing a given particle to be gated on and therefore filtering out undesired particles and detector noise in other spectra. In the ${}^{39}\text{K}({}^3\text{He}, d){}^{40}\text{Ca}$ experiment,

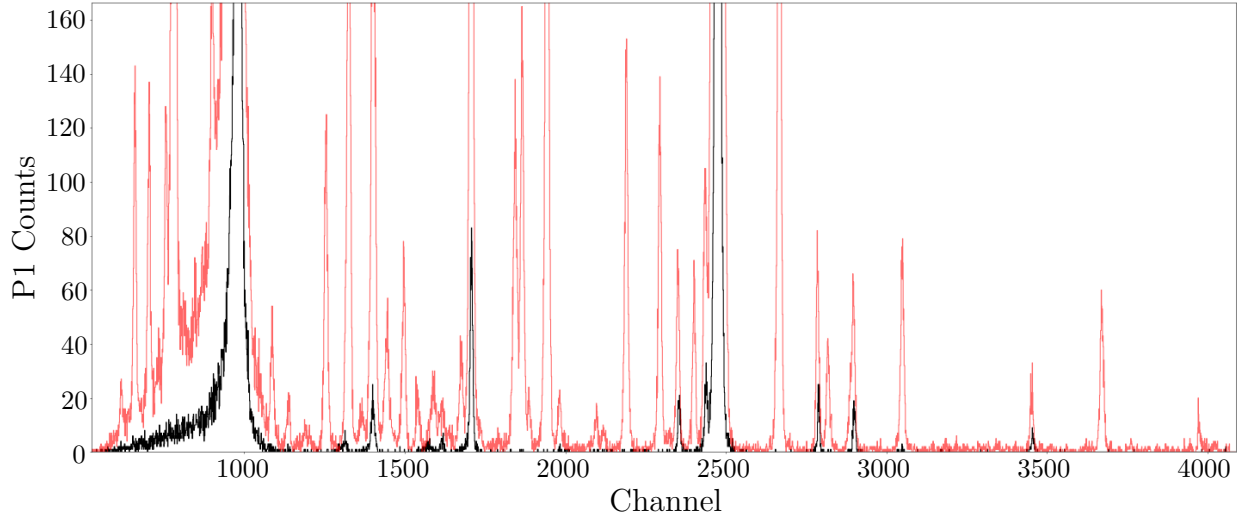


Figure 6.10: A histogram spectrum of the front (P1) focal-plane detector position section at $\theta_{\text{lab}} = 5^\circ$ and $E_{\text{lab}} = 21$ MeV with the KI #6 target (in red) and the carbon target (in black), both gated on the deuteron group from the the ΔE vs. E spectrum. Overlapping peaks indicate contaminants.

the $\Delta E/E$ spectrum was used to gate on the deuteron group for $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ and the ^3He group for $^{39}\text{K}(^3\text{He}, ^3\text{He})^{39}\text{K}$. Figure 6.9 shows an example 2D histogram of ΔE vs E in the offline **Jam** software package [Swa01, Swa02] for the KI #6 target at $\theta_{\text{lab}} = 5^\circ$ and an Enge magnetic field of $B = 1.14$ T. The d , t , and ^3He particles appear in groups, with the d group highlighted in red to represent a gate. The p group is joined by noise in the ΔE detector below the ΔE threshold and are filtered out along with the other particle groups. Both the ΔE and E signals are compressed from their original 4096 channels to 512 channels in the 2D spectrum.

The 1D histogram of the P1 section, gated on the deuteron group of Fig 6.9, is shown in Fig. 6.10 in red. The overlayed black spectrum is from a carbon target to show overlapping peaks, indicating contaminants in the KI target. The deuteron peaks shown only in red were ejected from $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ reactions in the KI target, where different excited states of the ^{40}Ca recoil nuclei were populated, resulting in different kinematics for each deuteron depending on the excited state. High-energy deuterons are associated with high-energy ^{40}Ca excited states, and vice versa. The deuterons are momentum-separated by the Enge Split-Pole Spectrograph, focusing the higher energy ones on the left side of the position sections and the lower energy ones on the right side. Thus, the P1 channel number in Fig. 6.10 is inversely related to the ^{40}Ca excited state energies. The same is true of the excited states from the contaminants.

6.5.3 Energy Shifts

The P1 peak centroid positions typically remain constant over time. However, certain changes in the beam can introduce shifts in these positions, broadening the peaks over more channels than usual. For instance, adjusting the beam steerers or focusing elements at any point along the beamline can cause very slight kinematic differences to occur in the target. This is because the target material is never perfectly uniformly-distributed. Sometimes these differences in kinematics are large enough to cause a noticeable drift in the P1 spectrum. Retuning the beam between runs at the same Enge angle θ_{lab} is the most common example of energy shifts, followed by switching between targets from a different evaporation batch. Any unintentional beam drift can also cause them, albeit usually to a lesser degree.

To visualize these energy shifts, the P1 spectrum must be traced over time. This feature was recently added to the offline version of the `EngeSpec` [Lon19] graphical user interface (GUI), and I implemented the same feature in my local `Jam` source code. For each event, the P1 data is collected and the event number is recorded during the sort routine. The total number of events is unknown until the end of the sort routine, which unfortunately makes an online version of this feature nearly impossible. The 4096-channel P1 data and the corresponding event numbers are compressed into 512 bins, where they each get incremented into a 2D histogram.

There were a few instances of energy shifts during the $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ experiment. One happened due to a necessary beam-retune after a tour group was scheduled to view the tandem accelerator. The low-energy beamline Faraday cup was put in place for about an hour, and tuning was required to get the beam back on target. The runs before and after this retune are represented by the 2D histogram of P1 vs event number in Fig. 6.11. Depicted in the figure are the runs #23 – #27 at $\theta_{\text{lab}} = 9^\circ$ with the KI #5 target. The P1 section is zoomed in on the ^{12}C ground state, as it had the most counts and is therefore easiest to visualize energy shifts. The ^{12}C ground state centroid starts at P1 bin number 296, and a clear shift to bin number 297 is seen at event number 320. A 1-bin shift in the 512-bin 2D spectrum corresponds to an 8-channel shift in the 4096-channel P1 spectrum, which was noticeable even during the experiment. Based on the BCI of each run, it was easy to determine that the shift happened between runs #25 and #26, exactly when the retune happened. This is also evident from the shift in counts at this point because more beam current was acquired as a result of the retune. There is also an unknown phenomenon just after event bin number 100, occurring during run #23. The peaks do not appear to be shifted from it, at least not as dramatically as from the retune, but it is clear that the beam current changed abruptly.

It would not be appropriate to sort all runs at $\theta_{\text{lab}} = 9^\circ$ together for the $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ energy calibration and yield analysis. Because of the presence of energy shifts at some angles,

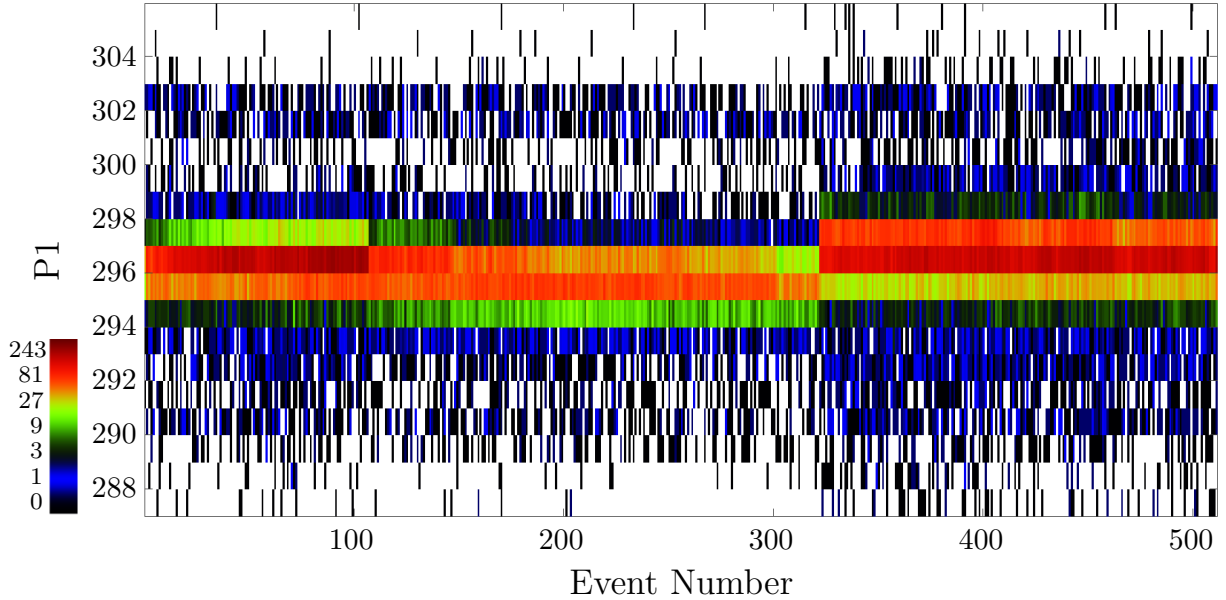


Figure 6.11: A 2D histogram of the front (P1) focal-plane position section vs event number over all $^{39}\text{K}(\text{He}^3, d)^{40}\text{Ca}$ runs (#23 – #27) at $\theta_{\text{lab}} = 9^\circ$ with the KI #5 target. Both axes are compressed to 512 channels. The P1 section is zoomed in on the ^{12}C ground state to show the effect of energy shifts due to beam retunes. A beam retune after run #25 caused the shift in the P1 spectrum shown at event (bin) number 320. The unknown phenomenon at event (bin) number 100 occurred during run #23, which caused a reduction in counts but no energy shift.

it was necessary to group runs together based on where the energy shifts occurred. I refer to these groups as *sort groups*. All of the runs in the $^{39}\text{K}(\text{He}^3, d)^{40}\text{Ca}$ experiment, as well as their sort groups, are shown in Table 6.1. Because there were 7 angles measured in this experiment ($\theta_{\text{lab}} = 5^\circ, 7^\circ, 9^\circ, 11^\circ, 13^\circ, 15^\circ$ and 20°), the ideal scenario would be to have 7 sort groups, 1 for each angle. However, there are 16 sort groups for various reasons, including beam retunes, switching targets, and the occasional beam noise from beamline element power supplies failing. In the case of sort group #13 at $\theta_{\text{lab}} = 13^\circ$, consisting of runs #45 and #47, 2 different targets are used between them, but they are from the same evaporation batch and therefore did not cause an energy shift. The $^{39}\text{K}(\text{He}^3, \text{He})^{39}\text{K}$ P1 spectra fortunately did not contain energy shifts because only a single run was needed per angle, and the same target (KI #7) was used for each run.

In the case of the $^{39}\text{K}(\text{He}^3, d)^{40}\text{Ca}$ run #23 at $\theta_{\text{lab}} = 9^\circ$, and for a few other runs, the energy shift occurred during the run, not between runs. For this reason, it was also necessary to have the ability to gate on the P1 vs Event Number histogram for the entire area before or after a shift. I added this feature to both my local `EngeSpec` and `Jam` source codes. Only the events occurring in the gate are used in a new, event-gated P1 spectrum. Unfortunately, this

Table 6.1: Information on each run of the $^{39}\text{K}(\text{He}, d)^{40}\text{Ca}$ experiment, including the angle of the Enge, the target used, the magnetic field of the Enge, and the sort group (see text).

Sort Group	θ_{lab} [°]	Run Numbers	Target	Enge Field [T]
1	5	9, 10	KI #1	1.14
2	5	11	KI #1	1.14
3	5	83 – 87	KI #6	1.14
4	7	15 – 20	KI #1	1.14
5	7	72, 75 – 78	KI #6	1.14
6	9	23	KI #5	1.145
7	9	24, 25	KI #5	1.145
8	9	26, 27	KI #5	1.145
9	11	31 – 33	KI #5	1.145
10	13	39 – 41	KI #5	1.145
11	13	42	KI #5	1.145
12	13	43, 44	KI #5	1.145
13	13	45, 47	KI #5, KI #6	1.145
14	15	55 – 59	KI #6	1.145
15	20	60, 61, 64	KI #6	1.145
16	20	65, 66	KI #6	1.145

means that the recorded BCI is no longer accurate. An approximation to the new, reduced BCI is to scale the original BCI by the fraction of events in the gate. This is not exact due to the loss of resolution when compressing the events into 512 bins, but it is more than sufficient considering the much larger uncertainty from the BCI measurement itself. The same fraction can be applied to the time-keeping scalars to correct for the new deadtime. The exact same gate is also used in these cases for the simultaneous silicon detector telescope spectra, as introduced in the next section, to ensure consistency with the reduced BCI and deadtime between the focal-plane and silicon detector spectra. This is essential for a relative measurement between the two detectors, as discussed in Section 6.7.4.

6.5.4 Silicon Detector Telescope

To minimize the effects of uncertainty in the target thickness, non-uniformity in the target, and target degradation after exposure to the beam, the $^{39}\text{K}(\text{He}, \text{He})^{39}\text{K}$ elastic scattering and $^{39}\text{K}(\text{He}, d)^{40}\text{Ca}$ proton-transfer yields from the focal-plane P1 spectra were normalized to the simultaneous $^{39}\text{K}(\text{He}, \text{He})^{39}\text{K}$ elastic scattering yield of a Si detector telescope positioned at a constant $\theta_{\text{lab}} = 45^\circ$ inside the target chamber. The scale for the Si-normalized focal-plane differential cross-section measurements was then corrected using the global ^3He optical model

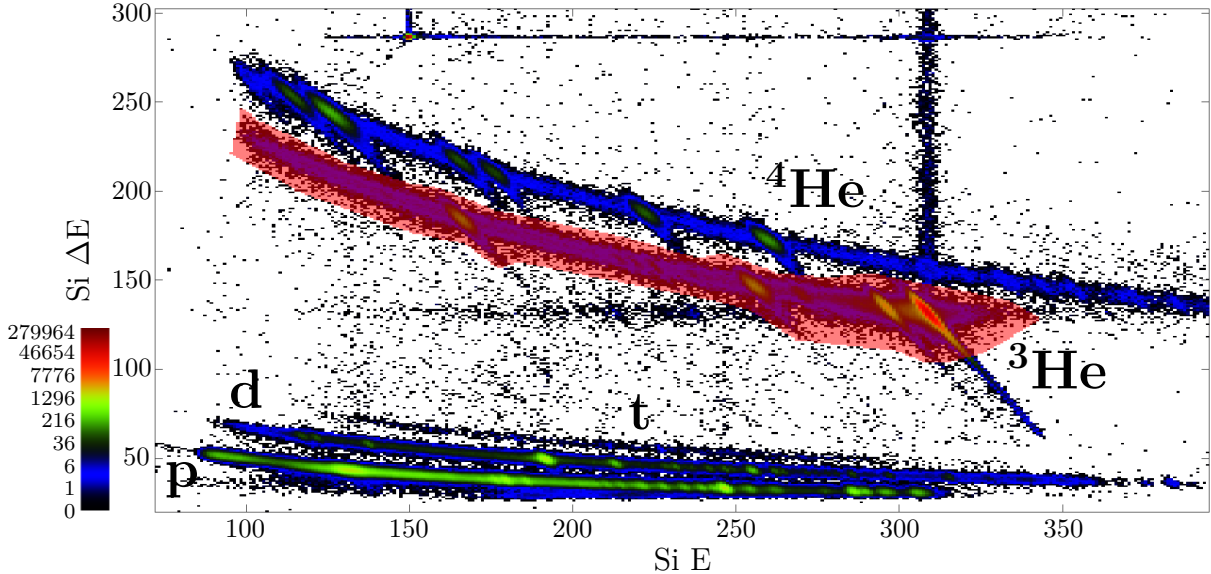


Figure 6.12: A 2D histogram of the Si ΔE vs Si E detector signals. Different groups correspond to different particles depending on their mass and charge. The ^3He group, highlighted in red, is being gated on in the figure.

potential (OMP) differential cross-section for ^{39}K from Ref. [Lia09] (see Sections 6.7.4 and 6.7.5). The scaling factor was the ratio between the differential cross-section of the global OMP and that of the Si-normalized focal-plane $^{39}\text{K}(^3\text{He}, ^3\text{He})^{39}\text{K}$ measurements.

The silicon detector telescope consists of a ΔE detector and a thicker, residual energy E detector, used in coincidence for particle discrimination in the same way as the ΔE and E focal-plane detectors. A representative 2D ΔE vs E histogram for the silicon detectors at $\theta_{\text{lab}} = 45^\circ$ with the KI #6 target is shown in Fig. 6.12. As before, different groups correspond to different particles depending on mass and charge. The highlighted gate is around the ^3He group. It is clear that there are 4 regions in this group that are more dense than the background and that have an even more negative slope than the trend of the overall group. These different regions correspond to scattering from different elements in the target. From left to right, they are carbon, oxygen, potassium, and iodine, as the more massive elements will deposit more energy in the Si E detector.

To obtain the $^{39}\text{K}(^3\text{He}, ^3\text{He})^{39}\text{K}$ Si detector yield, the gated Si ΔE vs Si E spectrum is first transformed. The regions corresponding to the individual scattering elements all have the same slope, and by rotating them so that they vertically align with the y -axis, they can be projected onto the x -axis to create a new 1D histogram with narrow peaks. This

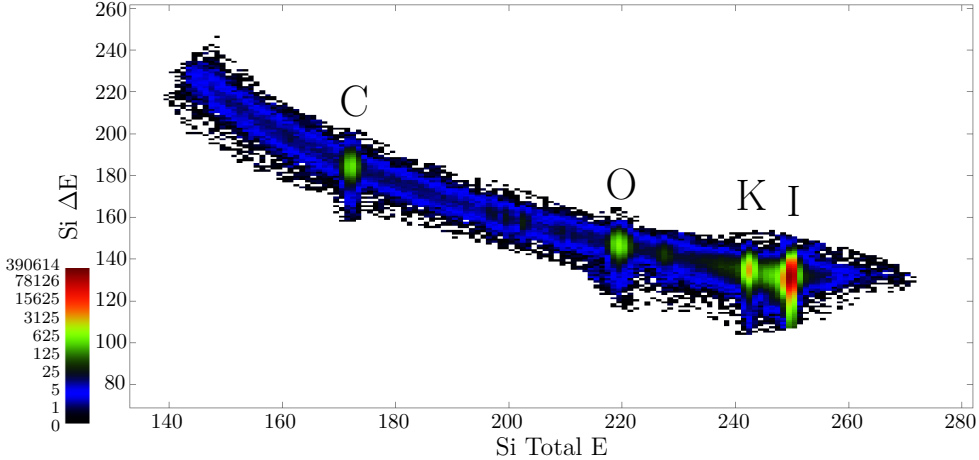


Figure 6.13: A 2D histogram of the Si ΔE vs Si total E detector signals, gated on the ${}^3\text{He}$ group and at $\theta_{\text{lab}} = 45^\circ$. This is similar to Fig. 6.12, except the total Si energy has been calculated based on the slope of the peaks in the Si ΔE vs Si E spectrum (see Eqn. 6.2). Each peak is labeled with its corresponding elastic scattering element.

transformation is equivalent to finding the total energy deposited in the silicon detectors,

$$E^{\text{tot}} = \frac{E + p_{\text{Si}}\Delta E}{1 + p_{\text{Si}}}, \quad (6.2)$$

where p_{Si} is the Si slope parameter defined by the user in either `EngeSpec` or `Jam`. The transformed 2D Si ΔE vs Si total E spectrum, gated on the ${}^3\text{He}$ group, is shown in Fig. 6.13. The scattering elements are labeled with their element symbols. Finally, the projected 1D histogram for Si total E is presented in Fig. 6.14, where the potassium (K) peak can be seen on the shoulder of the iodine (I) peak. The yield of the potassium peak ideally remains constant throughout the experiment, for a given target, since the Si detector telescope is positioned at a constant $\theta_{\text{lab}} = 45^\circ$. However, target degradation occurs naturally over time, decreasing the yield of both focal-plane and Si detector measurements. Hence, the normalization corrects for this, as well as the other unknown target properties.

One complication is the poor resolution between different isotopes for elastic scattering, especially at low angles. In the Si detectors, the potassium peak always consisted of ${}^{41}\text{K}$ in addition to ${}^{39}\text{K}$, since a natural KI target was used. In the focal-plane P1 detector, which has better energy resolution, the elastic scattering ${}^{39}\text{K}$ peak could be resolved only for $\theta_{\text{lab}} \geq 40^\circ$. The presence of ${}^{41}\text{K}$ was corrected for in the differential cross-section calculations of ${}^{39}\text{K}({}^3\text{He}, {}^3\text{He}){}^{39}\text{K}$ by taking into account both the isotopic abundance ratio of potassium (93.26% ${}^{39}\text{K}$, 6.73% ${}^{41}\text{K}$) and the relative differential cross-sections of global ${}^3\text{He}$ OMPs for ${}^{39}\text{K}$ and ${}^{41}\text{K}$. The former contribution cancels out in the Si detector normalization for

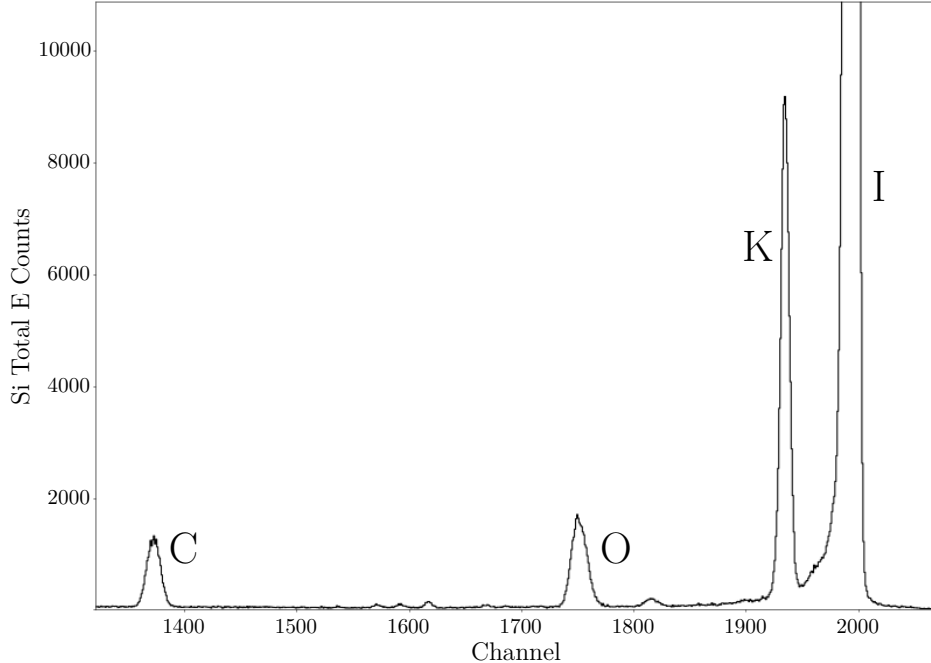


Figure 6.14: A histogram spectrum of Si Total E gated on the ${}^3\text{He}$ group at $\theta_{\text{lab}} = 45^\circ$, obtained by projecting Fig. 6.13 onto its x -axis. Each peak is labeled with its corresponding elastic scattering element. The individual ${}^{39}\text{K}$ and ${}^{41}\text{K}$ isotopes are unresolvable.

$\theta_{\text{lab}} < 40^\circ$, but provides a $\sim 7\%$ increase in the differential cross section ratio for $\theta_{\text{lab}} \geq 40^\circ$. The latter contribution is almost negligible due to the similar theoretical angular distributions of both differential cross sections over the angles of interest.

6.5.5 Oxidized Spectra

The KI #1 target was the first one to be used in the ${}^{39}\text{K}({}^3\text{He}, d){}^{40}\text{Ca}$ experiment at $\theta_{\text{lab}} = 5^\circ$ and 7° . It was immediately clear that this target was producing peaks that were not gaussian in the focal-plane P1 spectra, as were the other targets in the first evaporation batch, KI #2 and KI #3. They were transported together and all placed in the target chamber simultaneously. At some point, they were presumably exposed to atmosphere suddenly or for an extended period of time, despite the care taken in transporting them to the target chamber in a sealed box at rough vacuum. The oxidized focal-plane P1 spectrum for all KI #1 runs (#15 – #20) at $\theta_{\text{lab}} = 7^\circ$, gated on the deuteron group, are shown sorted together in Fig. 6.15. The low-channel (high-energy) side of each peak has a tail that is potentially smeared over other nearby peaks. This makes fitting peaks very challenging for yield measurements, when high resolution is required. However, Bayesian Monte Carlo techniques were implemented to obtain yield measurements with realistic uncertainties from these runs, as presented in Section 6.6.2.

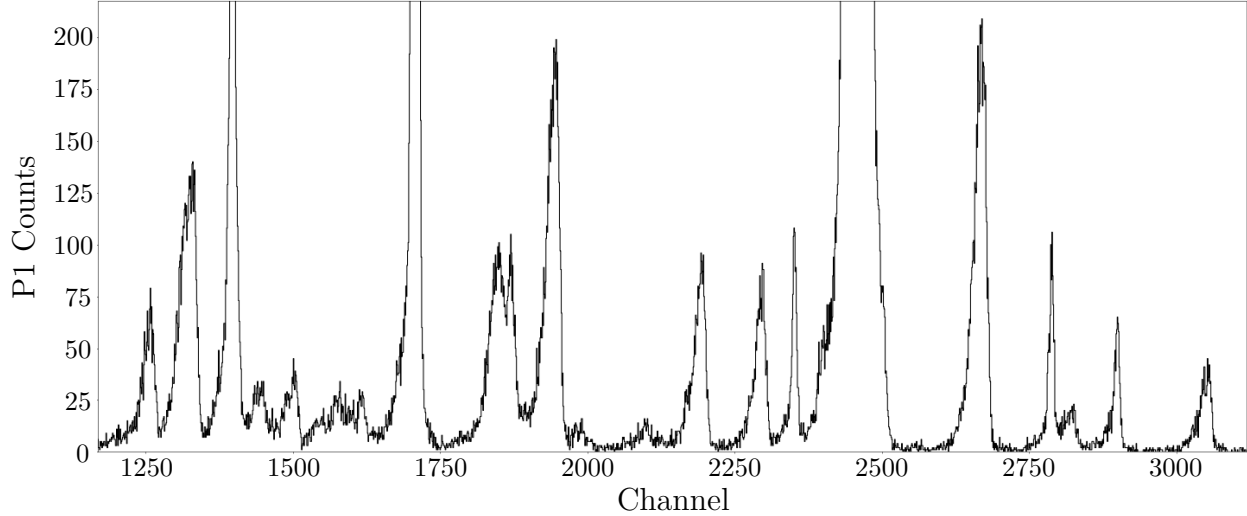


Figure 6.15: A histogram spectrum of the focal-plane P1 section at $\theta_{\text{lab}} = 7^\circ$ with the KI #1 target, gated on the deuteron group. The peaks are clearly not gaussian because of the presence of high-energy (low-channel) tails, a result of oxidation in the KI #1 target.

6.6 Bayesian Peak Fitting and Target Oxidation

The energy loss of light, charged particles through a medium is a statistical process. Particles with the same initial energy will traverse the same length in the medium with a distribution of final energies. This phenomenon is known as energy straggling and was theoretically described by Landau [Lan44]. Thin-film targets are ideal for high resolution spectral analysis because the energy loss distribution is approximately gaussian. Thick targets, however, alter the energy loss distribution in ways that present challenges to fitting peaks in a spectrum. Section 6.6.1 presents the Bayesian peak fitting procedure implemented in the analysis of the $^{39}\text{K}(\text{He}, d)^{40}\text{Ca}$ experiment for the case of thin targets, and Section 6.6.2 presents a modified procedure for the case of thick targets, resulting from target oxidation which produces tails in the energy loss distribution.

6.6.1 Bayesian Peak Fitting

Transfer reaction analysis requires precise experimental cross section measurements for each relevant excited state of the residual nucleus. Each cross section is proportional to the yield of the transfer reaction populating the given excited state. An essential ingredient in the yield measurement, described in more detail in Section 6.7, is the number of counts of ejectile particles measured by the focal-plane detector. Each count contributes to a peak along the focal-plane spectrum corresponding to the given excited state with a finite width. The

peaks are distinguished by their excitation energies, which are determined through an energy calibration based on their relative centroid positions along the focal-plane, as is done in Section 6.8.1. A typical transfer reaction spectrum will include many such peaks, as well as peaks from reactions in the target not necessarily of interest, known as contaminants. As mentioned in the introduction of this section, excited state peaks are typically gaussian-distributed, assuming a thin-film target is used. The spectrum will also include background counts, represented by a straight line. A precise determination of the number of counts in a given peak, in the simplest case, is therefore done by fitting a gaussian with a background line.

The target for the $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ experiment was natural potassium iodine on a natural carbon film backing. The most prominent contaminants were from $(^3\text{He}, d)$ reactions on ^{12}C , ^{13}C , ^{14}N , and ^{16}O , producing excited states from the residual nuclei ^{13}N , ^{14}N , ^{15}O , and ^{17}F , respectively. The chosen magnetic field of the Enge Split-Pole Spectrograph was such that deuterons from $(^3\text{He}, d)$ reactions with iodine were not at all present along the focal-plane, and deuterons from $^{41}\text{K}(^3\text{He}, d)$ were only minimally present. However, given the presence of the many other contaminants, and to distinguish doublets, triplets, and other mutliplets, it was deemed necessary to use a more sophisticated technique to fit the peaks from this experiment than a simple chi-squared minimization. Bayesian Monte Carlo sampling was a natural choice, considering its realistic uncertainty handling and flexibility when dealing with non-gaussian fits, presented in Section 6.6.2. I used the `BayeSpec` [Lon18a] graphical user interface (GUI), along with custom Bayesian sampling routines described below, to acquire fits to the $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ data.

Most peaks from transfer reactions on a thin-film target are gaussian-distributed,

$$f(x; A, \mu, \sigma) = A \exp\left(-\frac{(x - \mu)^2}{2\sigma^2}\right), \quad (6.3)$$

where A is the peak intensity, μ is its mean, and σ is its standard deviation. Bayesian sampling can be used to optimize these 3 parameters to match the focal-plane data if appropriate prior distributions are known. In this case, each prior distribution is approximately normal, $\mathcal{N}(\mu_0, \sigma_0^2)$, with its own mean μ_0 and standard deviation σ_0 . Posterior distributions can then be computed from these priors to achieve parameter values with statistically realistic uncertainties that match the data. These more realistic uncertainties make it possible to handle difficult fitting scenarios that would normally cause problems for chi-square minimizations, such as distinguishing peaks in a multiplet with appropriate uncertainties.

Prior means μ_0 for A and μ are particularly simple to obtain if the peak apex is visible. Even if it is not visible, as in the case of a multiplet, a guess can be made for each peak apex

location, (x, y) , based on the curvature of their sum, where x is the mean guess and y is the intensity guess. Each guess is established by selecting a point using the **BayeSpec** GUI. The prior standard deviations σ_0 for A and μ are given constant values conservative enough for sampling to be successful in even the most uncertain cases. These can be adjusted by the user, but are typically left as their defaults, listed below.

The prior mean μ_0 and standard deviation σ_0 for σ are ideally given constant values for a given reaction representative of most peaks from that reaction in focal-plane spectra. This is because energy loss is nearly equivalent for peaks with similar excitation energies from the same reaction, but it is not equivalent for peaks from different reactions. Therefore, peaks from each reaction should ideally have a shared σ prior and a shared σ posterior in a single fit if multiple peaks are fit simultaneously. In practice, a single σ prior can be used for different reactions if it is conservative enough for sampling to be successful, but the σ posteriors should remain different for different reactions.

A background line must also be included in the model, where its intensity y_{bg} and slope m_{bg} have their own $\mathcal{N}(\mu_0, \sigma_0^2)$ priors. Put together, the prior distributions for gaussian peaks on a background line in a typical focal-plane spectrum are

$$\begin{aligned} A_i &\sim \mathcal{N}(y_i, 10.0^2) \\ \mu_i &\sim \mathcal{N}(x_i, 1.0^2) \\ \sigma &\sim \mathcal{N}(5.0, 1.0^2) \\ y_{\text{bg}} &\sim \mathcal{N}(\max(1, \min(d)), 0.1^2) \\ m_{\text{bg}} &\sim \mathcal{N}(0, 0.01), \end{aligned} \tag{6.4}$$

where σ is a constant prior for all reactions, but it can easily be adjusted by the user for different reactions if needed, and d refers to the focal-plane data in the specified fit range. The prior for m_{bg} can also be adjusted if the background line is clearly not horizontal. The model function to be fit is

$$f(x; A, \mu, \sigma, y_{\text{bg}}, m_{\text{bg}}) = \sum_i^N A_i \exp \left(-\frac{(x - \mu_i)^2}{2\sigma_j^2} \right) + \left(y_{\text{bg}} + m_{\text{bg}}(x - \min(x)) \right), \tag{6.5}$$

where N is the total number of gaussian peaks to be fit and σ_j refers to the shared standard deviation for peaks from reaction j .

The posterior computation in the custom Bayesian sampling routine uses the **quap** function in **R**, part of the Bayesian **rethinking** package [McE13]. This function finds a quadratic approximation to each full posterior distribution at its mode. It is less sophisticated than Markov Chain Monte Carlo, but its relative simplicity makes it a more efficient option to use

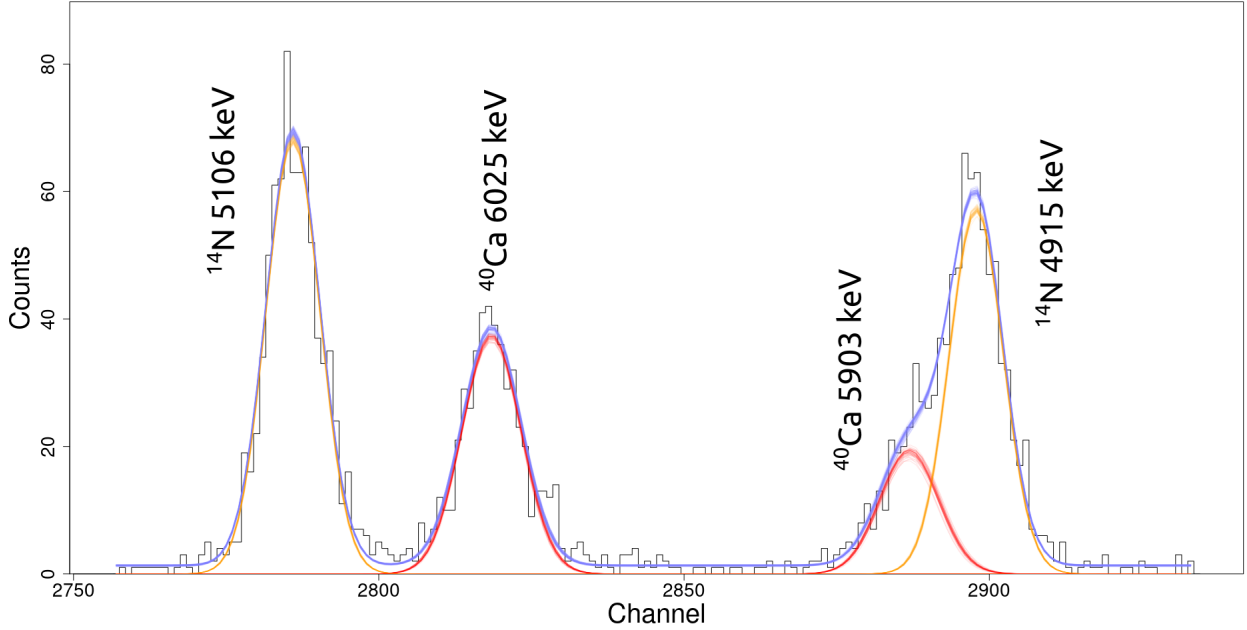


Figure 6.16: A Bayesian multi-gaussian fit with `BayeSpec` for the ^{40}Ca excited states (in red) 6025 keV and 5903 keV from $^{39}\text{K}(\text{He}, d)^{40}\text{Ca}$ and the ^{14}N excited states (in orange) 5106 keV and 4915 keV from $^{13}\text{C}(\text{He}, d)^{14}\text{N}$ at $\theta_{\text{lab}} = 5^\circ$. In red and orange are 50 random samples of the gaussian distributions from the σ , μ , and A posteriors for each peak. The ^{40}Ca peaks share an identical σ posterior, while the ^{14}N peaks share their own as well. In blue are the sums of the peaks plus the background line for each of those 50 samples.

with a GUI. The model composed of Eqns. 6.4 and 6.5 is provided to `quap`, which returns the posterior quadratic approximations with built-in uncertainties. Samples from these posteriors are then used to compute the area of each gaussian, given by $\sqrt{2\pi} \sigma_j A_i$, along with their uncertainties. A small number of posterior samples are also used to graphically display the gaussian fits, with different colors representing different reactions.

Figure 6.16 shows an example of this Bayesian fitting procedure for a $^{39}\text{K}(\text{He}, d)^{40}\text{Ca}$ focal-plane spectrum at $\theta_{\text{lab}} = 5^\circ$. Excited states of ^{40}Ca are shown in red and ^{14}N excited state contaminants are shown in orange. In blue is the sum of the gaussians and the background line, given by Eqn. 6.5. Each gaussian, as well as the blue curve, consists of 50 individual curves that were sampled from the posteriors to represent the uncertainties. The reason it is useful to combine these 4 peaks into one fit in this example is to better resolve the 5903 keV ^{40}Ca state in red from the 4915 keV ^{14}N state in orange in the double gaussian on the right side of the spectrum. The addition of the 6025 keV ^{40}Ca peak in red and the 5106 keV ^{14}N peak in orange on the left constrains the σ posteriors for the double gaussian. While this may seem like a minor correction, and it is minor in the present example, it is crucial in situations where the double gaussian or multiplet is otherwise impossible to resolve.

6.6.2 Fitting Peaks with Target Oxidation

Potassium iodine is hygroscopic, meaning it easily absorbs moisture from the environment. When exposed to atmospheric pressure, the salt slowly oxidizes, forming potassium carbonate and molecular iodine. This oxidation increases the thickness of thin-film targets. The probability of particles traversing the target with large final energies is decreased, introducing a high energy tail in the ejected particle spectrum. The Landau distribution describes this energy loss, but it is computationally challenging to implement. An approximation that has been found to fit oxidized-target spectra well is the exponentially-modified gaussian (EMG) distribution [Bab16]. The power of the EMG approximation is that its area calculation is the exact same as that of a gaussian distribution, $\sqrt{2\pi}\sigma A$, where σ and A are parameters of both gaussians and EMGs, but they have slightly different definitions as it will become clear below. This makes it very simple to determine the number of counts for EMG distributions, unlike Landau distributions, where the full area integral must be calculated.

The probability density function (PDF) of an EMG distribution f is a convolution of exponential g and gaussian h PDFs,

$$\begin{aligned} f(x; \sigma, \lambda, \mu, A) &= (g * h)(x) = \int_{-\infty}^{\infty} g(x')h(x - x') dx' \\ &= \int_0^{\infty} \lambda \exp(-\lambda x') A \exp\left(-\frac{1}{2}\left(\frac{x - x' - \mu}{\sigma}\right)^2\right) dx' \\ &= A \sigma \lambda \sqrt{\frac{\pi}{2}} \exp\left(\frac{1}{2}(\sigma\lambda)^2 - (x - \mu)\lambda\right) \operatorname{erfc}\left(\frac{1}{\sqrt{2}}\left(\sigma\lambda - \frac{x - \mu}{\sigma}\right)\right), \end{aligned} \quad (6.6)$$

where λ is the exponential-component rate, μ , σ , and A are the gaussian-component mean, standard deviation, and peak intensity, respectively, and erfc is the complimentary error function, defined as $\operatorname{erfc} z = 1 - \operatorname{erf} z$ for the complex variable z , where erf is the error function,

$$\operatorname{erf} z = \frac{2}{\sqrt{\pi}} \int_0^z \exp(-t^2) dt. \quad (6.7)$$

The standard EMG distribution, as a function of x , has a high- x tail. Since the focal-plane position spectrum channel number is inversely related to energy, the EMG must be modified to have a low- x tail. This is done by simply replacing $x - \mu$ with $\mu - x$ to reflect the standard EMG distribution about its gaussian-component mean, μ . The following discussion of EMG distributions assumes this reflection has been performed.

Previously it was shown for non-oxidized targets that the Bayesian fitting procedure for a gaussian peak requires gaussian mean μ and peak intensity A priors. The means of these priors are provided by the user when selecting an estimate at the peak apex with the BayeSpec GUI. However, this presents a problem when extending the procedure to EMG fits.

An EMG is defined by the μ and A parameters of its gaussian component, not the mean and peak intensity of the EMG itself. To obtain reasonable priors for the μ and A parameters, they must be derived from the attributes of the EMG peak apex, where the user selects. This apex defines the mode x_m and peak intensity y_m of the EMG, which can both be calculated by determining the coordinates where the derivative of the PDF is equal to zero. These are

$$\begin{aligned} x_m &= \mu - \sigma^2 \lambda + \sqrt{2} \sigma z, \\ y_m &= A \exp \left(-\frac{1}{2} \left(\frac{\mu - x_m}{\sigma} \right)^2 \right), \end{aligned} \quad (6.8)$$

where z is defined such that

$$\exp(z^2) \operatorname{erfc}(z) = \sqrt{\frac{2}{\pi}} \frac{1}{\sigma \lambda}, \quad (6.9)$$

which can be solved numerically. With x_m and y_m provided by the user, μ and A become

$$\mu(\sigma, \lambda) = x_m + \sigma^2 \lambda - \sqrt{2} \sigma z, \quad (6.10)$$

$$A(\sigma, \lambda) = y_m \exp \left(\frac{1}{2} \left(\frac{\mu - x_m}{\sigma} \right)^2 \right). \quad (6.11)$$

The priors would be fully determined if σ and λ , the parameters related to the EMG width and skewness, can be constrained. Fortunately, for a given transfer reaction observed in a focal-plane spectrum, there is little variation between these properties, much like the gaussian σ parameter of Section 6.6.1. From a sample of EMG fits for $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ peaks, reasonable priors for σ and λ were determined to be

$$\sigma \sim \mathcal{N}(5.0, 1.0^2), \quad (6.12)$$

$$\lambda \sim \mathcal{N}(0.09, 0.02^2), \quad (6.13)$$

where these are also usually sufficient for other reactions. Priors for Eqns. 6.10 and 6.11 can then be constructed by sampling from Eqns. 6.12 and 6.13. The resulting prior distributions for $\mu - x_m$ and A/y_m are shown in Figure 6.17 in black after taking 10,000 samples from Eqns. 6.12 and 6.13. The gaussian approximations (blue) were constructed from the mean and standard deviation of the $\mu - x_m$ and A/y_m samples, while the lognormal approximations (red) were similarly constructed from the mean and standard deviation of the natural logarithm of those samples. Note that the priors are not gaussian. In fact, the prior for A/y_m is neither gaussian nor lognormal. However, a lognormal approximation suffices for both in practice and is preferable to the actual distributions, which are either non-analytical or at least exceedingly complicated. The Bayesian quadratic approximation function, `quap`, used with `BayeSpec`

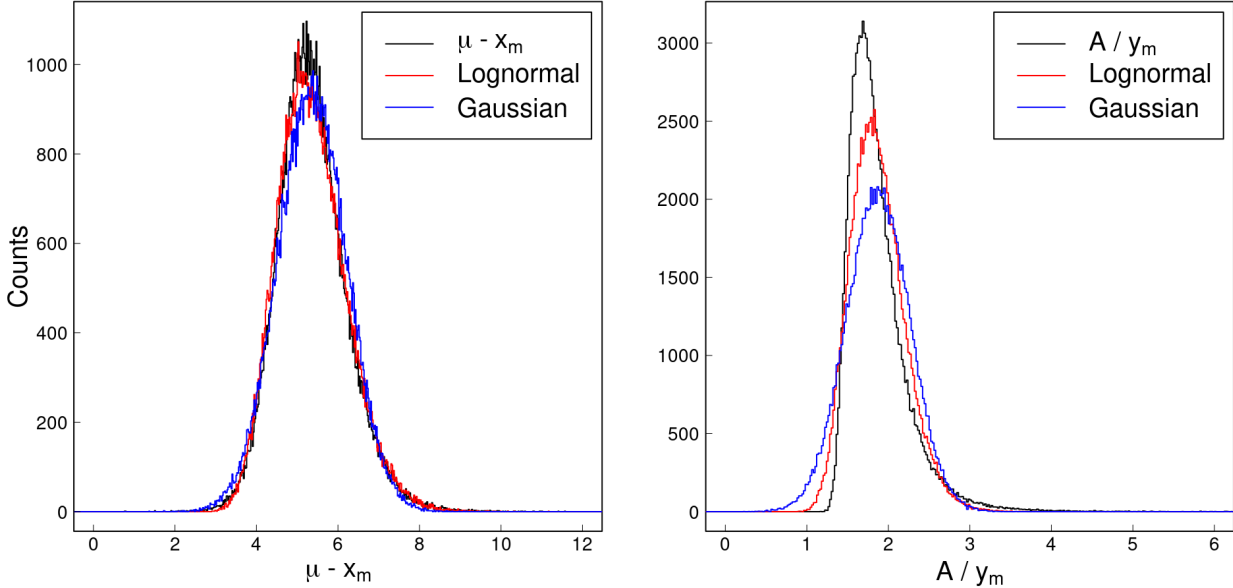


Figure 6.17: The constructed priors for $\mu - x_m$ (left) and A/y_m (right) from 10,000 samples of the σ and λ priors of Eqns. 6.12 and 6.13. The gaussian (blue) approximations were derived from the mean and standard deviation of the samples. The lognormal (red) approximations were derived from the mean and standard deviation of the natural logarithm of those samples.

requires named prior distributions, forbidding the use of non-analytical distributions. The lognormal approximations also correctly prevent negative sample values and are therefore taken as the priors for μ and A .

As in Section 6.6.1, once the priors have been determined, the quadratic approximations to the posteriors are computed with `quap`, and the fit is constructed from samples of the EMG distribution with those posteriors. An example of this Bayesian EMG fitting method is shown in Figure 6.18, where the 6285 keV ${}^{40}\text{Ca}$ state from ${}^{39}\text{K}({}^3\text{He}, d){}^{40}\text{Ca}$ at $\theta_{\text{lab}} = 5^\circ$ is fit with an EMG distribution because the potassium iodine target for this run (KI #1) had undergone oxidation. The closely-packed blue lines are 50 representative samples of the EMG posteriors, plus that of the background line, while the green lines are the corresponding gaussian components of those EMG samples with mean μ , standard deviation σ , and peak intensity A . The new mode and peak intensity from Eqn 6.8, obtained by sampling from the posteriors, are shown by the black lines, where the solid line represents its mean and the dashed lines represent its standard deviation. Note that the (x_m, y_m) coordinate lies along its gaussian component, a ubiquitous property of EMG distributions. The number of counts in the EMG distribution is simply equivalent to the area of its gaussian component, $\sqrt{2\pi} \sigma A$, constructed by sampling from the σ and A posteriors.

Figure 6.19 shows how powerful this method can be for high resolution spectral analysis

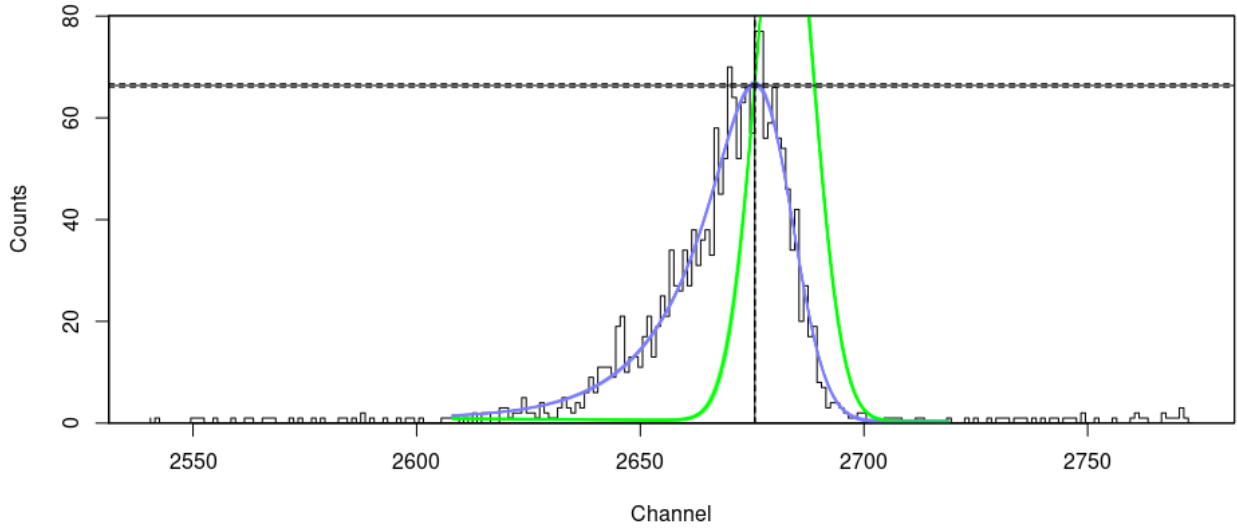


Figure 6.18: A Bayesian exponentially-modified gaussian fit with BayeSpec for the 6285 keV ^{40}Ca state from $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ at $\theta_{\text{lab}} = 5^\circ$ with an oxidized potassium iodide target (KI #1). In blue are 50 random samples of the exponentially-modified gaussian distribution from the σ , λ , μ , and A posteriors, plus the background line. In green are the gaussian components of the exponentially-modified gaussian samples. The new mode and peak intensity are represented by the black lines, where the solid line represents their mean and the dashed lines represent their standard deviation.

with an oxidized target. The same focal-plane spectrum as Figure 6.18 is shown here, but it is focused on a region with multiple peaks. The blue summed fit consists of 5 EMG distributions, shown individually by the red and orange lines, and a small, virtually horizontal background line. As before, each distribution shows 50 lines drawn from samples of the EMG posteriors. From a simple energy calibration, the red peaks were found to be ^{40}Ca excited states with excitation energies 7694 keV, 7658 keV, 7623 keV, and 7532 keV, from left to right, and the orange peak corresponds to the ^{14}N 6446 keV excited state. Because we expect energy loss to be nearly equivalent for peaks with similar energies from the same reaction, the σ and λ width and skewness parameter posteriors are fixed between the ^{40}Ca peaks, while the ^{14}N peak has its own σ and λ posteriors. The custom Bayesian sampling routines for both gaussian and EMG fits can be used for a simultaneous fit of any number of peaks from up to 3 different reactions at present, and this could easily be extended to any number of reactions when the need arises.

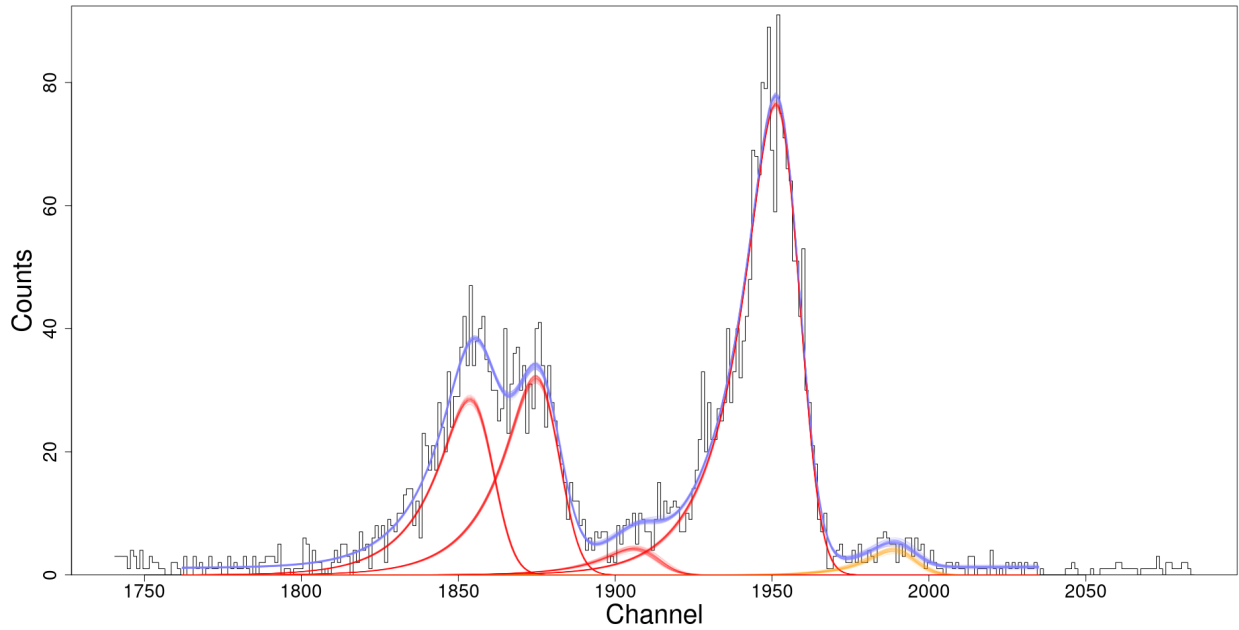


Figure 6.19: A Bayesian exponentially-modified gaussian fit with BayeSpec for the ^{40}Ca excited states (left to right, in red) 7694 keV, 7658 keV, 7623 keV, and 7532 keV from $^{39}\text{K}(\text{He}^3, d)^{40}\text{Ca}$ and the ^{14}N excited state (in orange) 6446 keV from $^{13}\text{C}(\text{He}^3, d)^{14}\text{N}$ at $\theta_{\text{lab}} = 5^\circ$ with an oxidized potassium iodine target (KI #1). In red and orange are 50 random samples of the exponentially-modified gaussian distributions from the σ , λ , μ , and A posteriors for each peak. The ^{40}Ca peaks share identical σ and λ posteriors, whereas the ^{14}N peak has its own. In blue are the sums of the peaks plus the background line for each of those 50 samples.

6.7 Cross Sections

An essential ingredient in the calculation of thermonuclear reaction rates is the nuclear reaction cross section σ , introduced in Section 2.2. The reaction cross section is an imaginary, effective area around the target nucleus where the reaction proceeds with a probability of unity if an incident particle passes through it. It is therefore a measure of the probability for a reaction to occur, varying with the incident particle energy. The cross section is closely related to the yield Y of the reaction

$$Y = \frac{N_R}{N_b}, \quad (6.14)$$

where N_R is the total number of reactions and N_b is the total number of incident particles. The total number of reactions N_R that occur is related to the number of counts N_c , or the area, of a given peak in a detector. However, this peak count is affected by the deadtime t_{dead} of the data acquisition system. To correct for the deadtime, and therefore get an accurate measure of the number of reactions, the livetime $t_{\text{live}} = 1 - t_{\text{dead}}$ is used as in

$$N_R = \frac{N_c}{t_{\text{live}}}. \quad (6.15)$$

The total number of incident particles N_b is

$$N_b = \frac{q}{Z_p e}, \quad (6.16)$$

where q is the total charge deposited by the incident particles, Z_p is the unit charge of the incident particles, and e is the electronic charge (1.6×10^{-19} C). The total charge q is derived from the beam-integrated current (BCI) scalar value collected for each pulse during a given run. The BCI has an associated $\text{BCI}_{\text{scale}}$ setting that determines the scale of the deposited charge, which can be adjusted on the beam current readout module. For the $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ experiment, the $\text{BCI}_{\text{scale}}$ setting was 10^{-10} C/pulse. The number of incident particles N_b is therefore

$$N_b = \frac{\text{BCI} \times \text{BCI}_{\text{scale}}}{Z_p e}. \quad (6.17)$$

As described in Section 6.7.1, the conversion between the yield and the cross section depends on the target thickness ΔE (in energy units) and the stopping power $\mathcal{E} = -(1/N_t)dE/dx$ (in units of eV cm²/atom), where N_t denotes the number density of target nuclei.

A measurement of the total cross section or total yield would require the full 4π sr solid angle coverage of detectors around the target to ensure that no emitted particles from the

reaction are missed. Since this is usually not feasible, the alternative is the measurement of a differential cross section $d\sigma/d\Omega_\theta$ or differential yield $dY/d\Omega_\theta$ from a detector covering a solid angle Ω collecting the particles ejected at an angle θ from the incident beam. The differential cross section can be measured at several angles to determine the angular distribution of the reaction. In the case of charged-particle reactions, such as $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$, each excited state of the recoil nucleus ^{40}Ca has its own angular distribution that depends theoretically on the single-particle quantum numbers associated with the quantum mechanical selection rules for the transition, presented in Section 6.8.4.

6.7.1 Yield to Cross Section Conversion

For thin targets, where the cross section σ and stopping power \mathcal{E} are approximately constant over the target thickness ΔE , the yield Y is proportional to σ , with a proportionality constant $n_X = \Delta E / \mathcal{E}$ representing the number of active target nuclei per cm^2 in the target compound X_aY_b [Ili15]. The term *active nuclei* in this sense means the nuclei of interest X in the target, whereas the term *inactive nuclei* refers to the nuclei not of interest Y in the target. The number of inactive target nuclei per unit area is defined as n_Y , and the ratio n_Y/n_X is equivalent to the target molecule stoichiometry, b/a . For compound targets, the effective stopping power $\mathcal{E}_{\text{eff}} = \mathcal{E}_X + n_Y \mathcal{E}_Y / n_X$ must be used. The same proportionality constant

$$n_X = \frac{\Delta E}{\mathcal{E}_{\text{eff}}} \quad (6.18)$$

also applies to the differential yield and differential cross section. Additionally, as some of the energy of the incident particles E_0 is lost in the target, the cross section is evaluated at an effective, reduced energy $E_{\text{eff}} = E_0 - \Delta E/2$. The conversion between the differential yield and differential cross section therefore becomes

$$\left[\frac{dY(E_0)}{d\Omega} \right]_\theta = n_X \left[\frac{d\sigma(E_{\text{eff}})}{d\Omega} \right]_\theta. \quad (6.19)$$

The number of active nuclei per cm^2 , n_X , can also be expressed in a form more tractable for experiments in the lab system as [Rol88, Set11]¹

$$n_X = \frac{\nu N_A \Delta x}{A_{t,\text{mol}}} \quad [\text{in cm}^{-2}], \quad (6.20)$$

where ν is the number of X atoms per molecule in the target material, N_A is Avogadro's

¹The “yield” Y defined in Eqns. 3.30 and 3.31 of Ref. [Set11] is N_R , not N_R/N_b , even though it describes it as such in the text.

number (6.023×10^{23} atoms/mole), Δx is the thickness of the target (in g/cm²), and $A_{t,\text{mol}}$ is the molecular mass of the target material (in grams).

Combining everything, the differential cross section in units of mb/sr, where $1 \text{ b} = 10^{-24} \text{ cm}^{-2}$, measured in the lab system is calculated as

$$\left[\frac{d\sigma}{d\Omega} \right]_\theta = \frac{N_c Z_p A_{t,\text{mol}}}{(3.75 \times 10^9) \Omega t_{\text{live}} \text{BCI} \times \text{BCI}_{\text{scale}} \nu \Delta x} \quad [\text{in mb/sr}], \quad (6.21)$$

where the solid angle Ω is in units of msr, all other variables are defined as before, and the factor e/N_A has been evaluated and converged into the numerical factor in the denominator.

6.7.2 Rutherford Scattering

For elastic scattering cross section measurements, it is typical to report the cross section as a ratio to the theoretical Rutherford scattering cross section at the same angle. Theoretical elastic scattering cross sections from optical model potentials (OMPs) are also typically given in terms of the Rutherford ratio, depending on the nuclear reaction code. In the coupled-reaction channels code Fresco [Tho06] for example, the Rutherford ratio is the default output of the elastic scattering cross section.

In the center-of-mass system, the Rutherford scattering cross section is [Ili15]

$$\begin{aligned} \left[\frac{d\sigma}{d\Omega} \right]_\theta^{\text{Ruth}} &= \left(\frac{Z_p Z_t e^2}{4E_{\text{c.m.}}} \right)^2 \frac{1}{\sin^4(\theta_{\text{c.m.}}/2)} \\ &= 1.296 \left(\frac{Z_p Z_t}{E_{\text{c.m.}}} \right)^2 \frac{1}{\sin^4(\theta_{\text{c.m.}}/2)} \quad [\text{in mb/sr}], \end{aligned} \quad (6.22)$$

where Z_p is the unit charge of the incident particles, as before, Z_t is the unit charge of the active target nuclei, $E_{\text{c.m.}}$ is the total kinetic energy in the center-of-mass system, in units of MeV, and $\theta_{\text{c.m.}}$ is the angle measured in the center-of-mass system. The elastic scattering cross section measured in the lab system must first be converted to the center-of-mass system before taking the ratio to Rutherford scattering. This conversion is described in Section 6.7.3.

6.7.3 Cross Sections in the Center-of-Mass System

The incident beam energy E_{lab} , scattering angle θ_{lab} , and the differential cross section itself $d\sigma/d\Omega_{\theta_{\text{lab}}}$ must be converted to the center-of-mass frame to apply the Rutherford ratio and to compare with theoretical differential cross sections, which are usually given in the center-of-mass frame.

The conversion from the incident beam energy in the lab system E_{lab} to the total kinetic energy in the center-of-mass system $E_{\text{c.m.}}$ is [Ili15]

$$E_{\text{c.m.}} = E_{\text{lab}} \frac{A_t}{A_t + A_p}, \quad (6.23)$$

where A_t and A_p are the nuclear masses of the active target nucleus and the incident particle nucleus, respectively. However, mass evaluations typically report atomic masses, not nuclear masses. In general, the nuclear masses A_{Nuc} can be calculated from the reported atomic masses A_{Atom} as [Wan21]

$$A_{\text{Nuc}} = A_{\text{Atom}} - Z m_e + B_e(Z), \quad (6.24)$$

where Z is the unit charge of the nucleus, m_e is the electron mass ($548579.909065(16) \times 10^{-9}$ g/mole [Hua21]), and B_e is the electron binding energy, found from Ref. [Hua21] to be calculated as

$$B_e(Z) = 14.4381 Z^{2.39} + 1.55468 \times 10^{-8} Z^{5.35} \text{ eV}. \quad (6.25)$$

The lab angle θ_{lab} can be converted to the center-of-mass angle $\theta_{\text{c.m.}}$ using the kinematics relation [Ili15]

$$\cos \theta_{\text{lab}} = \frac{\gamma + \cos \theta_{\text{c.m.}}}{\sqrt{1 + \gamma^2 + 2\gamma \cos \theta_{\text{c.m.}}}}, \quad (6.26)$$

where the γ parameter is defined as

$$\gamma = \sqrt{\frac{A_p A_e E_{\text{lab}}}{A_r (A_e + A_r) Q + A_r (A_r + A_e - A_p) E_{\text{lab}}}}, \quad (6.27)$$

with A_e and A_r representing the nuclear masses of the ejected particle and recoil nucleus, respectively, and Q representing the Q -value of the reaction. For elastic scattering, $A_e = A_p$, $A_r = A_t$, and $Q = 0$, so that $\gamma = A_p/A_t$. Eqn. 6.26 can be solved numerically or it can be rewritten as a quadratic equation for $\theta_{\text{c.m.}}$, using the positive solution for forward scattering.

The differential cross section is defined such that the same number of ejected particles cross the solid angle $d\Omega_{\text{lab}}$ in the direction θ_{lab} as those crossing the solid angle $d\Omega_{\text{c.m.}}$ in the direction $\theta_{\text{c.m.}}$. That is,

$$\left(\frac{d\sigma}{d\Omega} \right)_{\theta_{\text{lab}}}^{\text{lab}} d\Omega_{\text{lab}} = \left(\frac{d\sigma}{d\Omega} \right)_{\theta_{\text{c.m.}}}^{\text{c.m.}} d\Omega_{\text{c.m.}}. \quad (6.28)$$

Assuming the cross section does not depend on the azimuthal angle, the conversion between the lab and center-of-mass systems is therefore

$$\frac{(d\sigma/d\Omega)_{\theta_{\text{c.m.}}}^{\text{c.m.}}}{(d\sigma/d\Omega)_{\theta_{\text{lab}}}^{\text{lab}}} = \frac{d\Omega_{\text{lab}}}{d\Omega_{\text{c.m.}}} = \frac{d(\cos \theta_{\text{lab}})}{d(\cos \theta_{\text{c.m.}})} = \frac{1 + \gamma \cos \theta_{\text{c.m.}}}{(1 + \gamma^2 + 2\gamma \cos \theta_{\text{c.m.}})^{3/2}}. \quad (6.29)$$

6.7.4 Si Detector Normalization

As mentioned in Section 6.5.4, a Si detector telescope was used to measure the $^{39}\text{K}(^3\text{He}, ^3\text{He})^{39}\text{K}$ elastic scattering differential cross section at a constant $\theta_{\text{lab}} = 45^\circ$, while the focal-plane detector was simultaneously measuring the differential cross sections of both $^{39}\text{K}(^3\text{He}, ^3\text{He})^{39}\text{K}$ and $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ over multiple θ_{lab} angles. The Si detector elastic scattering cross section was used as a normalization to correct for systematic uncertainties associated with unknown target effects during the focal-plane measurements. Although this is a relative measurement, an absolute scale can be established by comparing to optical model predictions, described in Section 6.7.5.

The reason the relative measurement is effective is due to the conversion factor n_X between the yield and cross section in Eqn. 6.19. Since n_X only depends on the target properties, taking a relative cross section is equivalent to taking a relative yield, where the target properties cancel out. Because of the angular dependence, however, each differential yield (or equivalently, each differential cross section) must first be converted into the center-of-mass system before taking the ratio. The final form of the differential cross-section ratio R in the center-of-mass system becomes

$$R \equiv \frac{(d\sigma/d\Omega_{\text{FP}})_{\theta_{\text{c.m.}}}^{\text{FP}}}{(d\sigma/d\Omega_{\text{Si}})_{\theta_{\text{Si}}}^{\text{Si}}} = \frac{N_c^{\text{FP}} \Omega_{\text{Si}}}{\Omega_{\text{FP}} N_c^{\text{Si}}} \frac{1 + \gamma \cos \theta_{\text{c.m.}}}{(1 + \gamma^2 + 2\gamma \cos \theta_{\text{c.m.}})^{3/2}} \frac{(1 + \gamma^2 + 2\gamma \cos \theta_{\text{Si}})^{3/2}}{1 + \gamma \cos \theta_{\text{Si}}}, \quad (6.30)$$

where $\theta_{\text{c.m.}}$ is the angle for the focal plane measurement, $\theta_{\text{Si}} = 48.14^\circ$ is the center-of-mass angle for the Si measurement ($\theta_{\text{lab}} = 45^\circ$), $\Omega_{\text{Si}} = 4.23(4)$ msr, and $\Omega_{\text{FP}} = 1.00(4)$ msr for all runs of the transfer reaction and at angles $\theta_{\text{lab}} \leq 35^\circ$ for elastic scattering; otherwise, $\Omega_{\text{FP}} = 0.50(4)$ msr for elastic scattering. Note that the only experimental quantities required with a relative measurement are the number of detector counts N_c and the solid angle Ω that each detector covers, in addition to the quantities making up the center-of-mass conversion. The u_R uncertainty is found to be

$$u_R = R \sqrt{\left(\frac{u_{N_c}}{N_c}\right)_{\text{FP}}^2 + \left(\frac{u_\Omega}{\Omega}\right)_{\text{FP}}^2 + \left(\frac{u_{N_c}}{N_c}\right)_{\text{Si}}^2 + \left(\frac{u_\Omega}{\Omega}\right)_{\text{Si}}^2}, \quad (6.31)$$

where the uncertainties in the angle measurement and γ are sufficiently small to be neglected.

Due to the presence of energy shifts in the $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ experiment (see Section 6.5.3), the runs usually could not all be sorted together at a given scattering angle. Therefore, there were multiple relative cross sections that needed to be calculated at these angles, one per sort group, and a method of averaging these measurements was necessary. A weighted average over each sort group i at a given angle was deemed appropriate, where the weight is given in

terms of the individual relative cross section uncertainties u_R , as in

$$\bar{R} = \frac{\sum_i^n R_i / u_{R,i}^2}{\sum_i^n 1 / u_{R,i}^2}, \quad (6.32)$$

for n total sort groups at the given angle. This ensures that the more precise measurements are weighted more. The uncertainty in this weighted average is then

$$u_{\bar{R}} = \sqrt{\frac{1}{\sum_i^n 1 / u_{R,i}^2}}. \quad (6.33)$$

In calculating the individual relative cross section uncertainties u_R , it was clear that an additional scatter uncertainty u_s was required to accurately account for the variations from the average R . The goal was to apply the same fractional scatter uncertainty for all measurements of a given angle, based on the typical deviation from the average R over all ^{40}Ca states. An example of this is depicted in Fig. 6.20, where the variations from the average R , normalized to unity, at $\theta_{\text{lab}} = 13^\circ$ are shown over all sort groups and for all ^{40}Ca states, up to $E_x = 8935$ keV. Each color represents a different ^{40}Ca state. It is clear that the variations from the average are typically no more than about 10%, which is also true for the other angles. The variation from the average R is written as

$$\text{var} = \frac{R}{R_{\text{avg}}}, \quad (6.34)$$

where each fractional deviation is found from

$$f_{\text{dev}} = |\text{var} - 1|. \quad (6.35)$$

The average of these fractional deviations $f_{\text{dev, avg}}$ over all the ^{40}Ca states and all sort groups for a given angle is used as the fractional uncertainty applied to the additional scatter,

$$u_s = f_{\text{dev, avg}} R. \quad (6.36)$$

In cases where only one sort group was needed for an angle, the fractional uncertainty was instead obtained by taking $f_{\text{dev, avg}}$ for each sort group and ^{40}Ca state over all angles.

For sort groups where an oxidized target (KI #1) was used, it was determined that the uncertainty was still underestimated, based on the tension between measurements at the same angle with targets that were not oxidized. An additional $f_{\text{ox}} = 20\%$ scatter was added in these cases to account for any unknown variability and to apply more weight to measurements

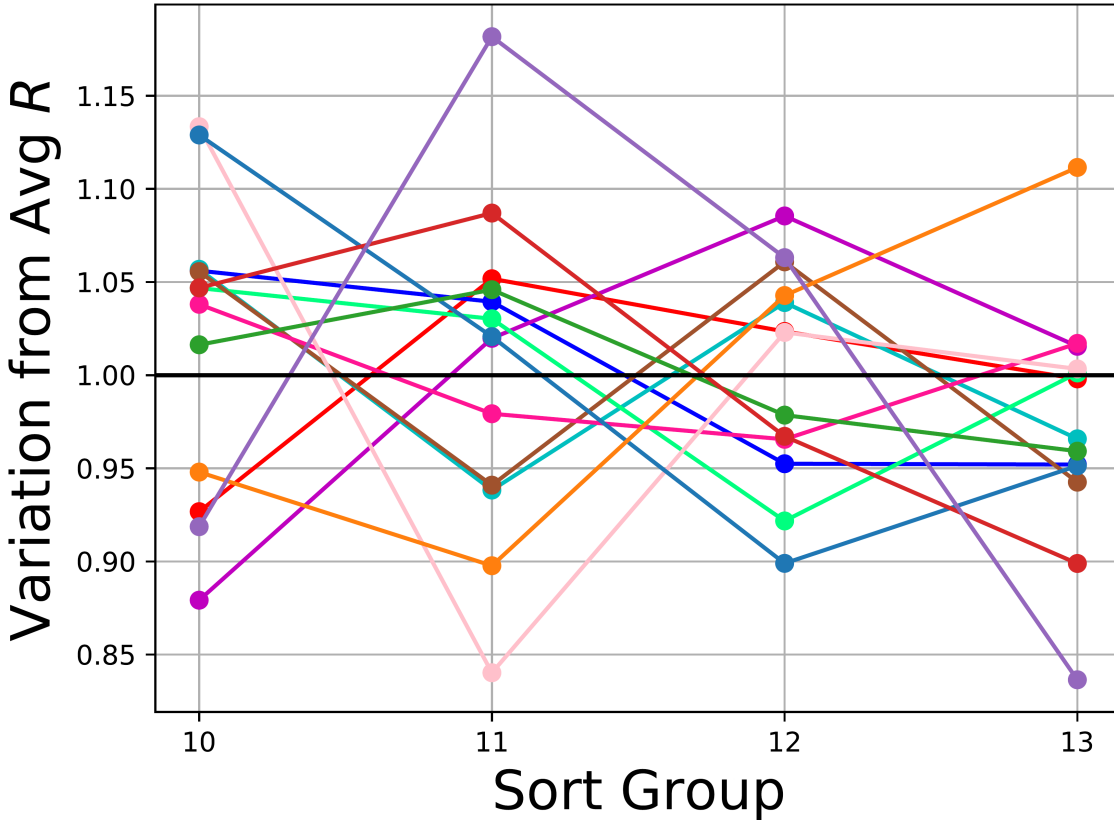


Figure 6.20: The variations from the average differential cross-section ratio R , normalized to unity, for each sort group over all ^{40}Ca states measured at $\theta_{\text{lab}} = 13^\circ$. The colored lines correspond to each ^{40}Ca state up to $E_x = 8935$ keV. The variations are typically within 10% of the average, which is true for the other angles as well. The average deviation for a given angle is used as an additional scatter uncertainty, as described in the text.

with non-oxidized targets. The resulting uncertainty in each R measurement with the added scatter is thus

$$u'_R^2 = u_R^2 + u_s^2 + u_{\text{ox}}^2 = u_R^2 + (f_{\text{dev, avg}}^2 + f_{\text{ox}}^2) R^2, \quad (6.37)$$

where $f_{\text{dev, avg}}$ covers all angles in cases where only one sort group was needed for an angle and $f_{\text{ox}} = 0$ for non-oxidized targets.

6.7.5 Optical Model Normalization

An absolute scale for the relative transfer differential cross section is established by applying a normalization factor N_{ES} obtained from the comparison between the relative elastic scattering differential cross section and the absolute differential cross section from a global ^3He optical model potential (OMP) for ^{39}K (see Section 2.3.1). The parameters of a global OMP are

Table 6.2: Global optical model potential parameters for $^{39}\text{K} + ^3\text{He}$ at $E_{\text{lab}} = 21$ MeV from Ref. [Lia09].

V_r	r_r	a_r	W_v	r_v	a_v	W_s
117.881	1.178	0.768	-0.646	1.415	0.847	20.665
r_s	a_s	V_{so}	r_{so}	a_{so}	W_{so}	r_c
1.198	0.852	2.083	0.738	0.946	-1.159	1.289

found through an optimization procedure, fitting them to an exhaustive list of experimental elastic scattering data for several target nuclei. Both the elastic scattering and global OMP cross sections are given in terms of a ratio to Rutherford scattering, which is theoretically unity at $\theta_{\text{c.m.}} = 0^\circ$.

The global ^3He OMP of Ref. [Lia09] was chosen as the reference for the absolute scale. This potential includes real and imaginary volume terms, an imaginary surface term, real and imaginary spin-orbit terms, and a Coulomb potential, where the Woods-Saxon form factor $f_i(r)$ and its derivative are used, as described in Section 2.3.1. A total of 148 sets of elastic scattering data for 52 target nuclei were used in the parametrization. The potential parameters are expressed as functions of beam energy and target mass. For $^{39}\text{K} + ^3\text{He}$ at $E_{\text{lab}} = 21$ MeV, the potential parameters of Ref. [Lia09] are given in Table 6.2, which are defined in the same manner as in Section 2.3.1.

The nuclear reaction code **Fresco** [Tho88, Tho06] was used to calculate the $^{39}\text{K}(^3\text{He}, ^3\text{He})^{39}\text{K}$ differential cross section shown in Fig. 6.21 using the OMP parameters in Table 6.2 at $E_{\text{lab}} = 21$ MeV. This differential cross section, represented as the black curve, was calculated in steps of 1° from $\theta_{\text{c.m.}} = 0^\circ - 80^\circ$. The present experimental differential cross section is shown in red at the angles $\theta_{\text{lab}} = 15^\circ - 55^\circ$ in steps of 5° , and 59° , converted to $\theta_{\text{c.m.}}$. The experimental points have been corrected for the presence of ^{41}K in all of the elastic scattering peaks of the Si spectra and only at $\theta_{\text{lab}} < 40^\circ$ in the focal-plane P1 spectra, described in Section 6.5.4. Both the data and the model are given in terms of the ratio to Rutherford scattering. The difference in scale is attributed to the relative measurement of the focal-plane and the Si detectors.

The elastic scattering scale factor N_{ES} was calculated by taking the average ratio between each experimental point and the global model,

$$N_{\text{ES}} = \frac{1}{N} \sum_{i=1}^N \frac{R_{\text{ES},i}}{(d\sigma/d\Omega_{\text{ES,global}})_i}, \quad (6.38)$$

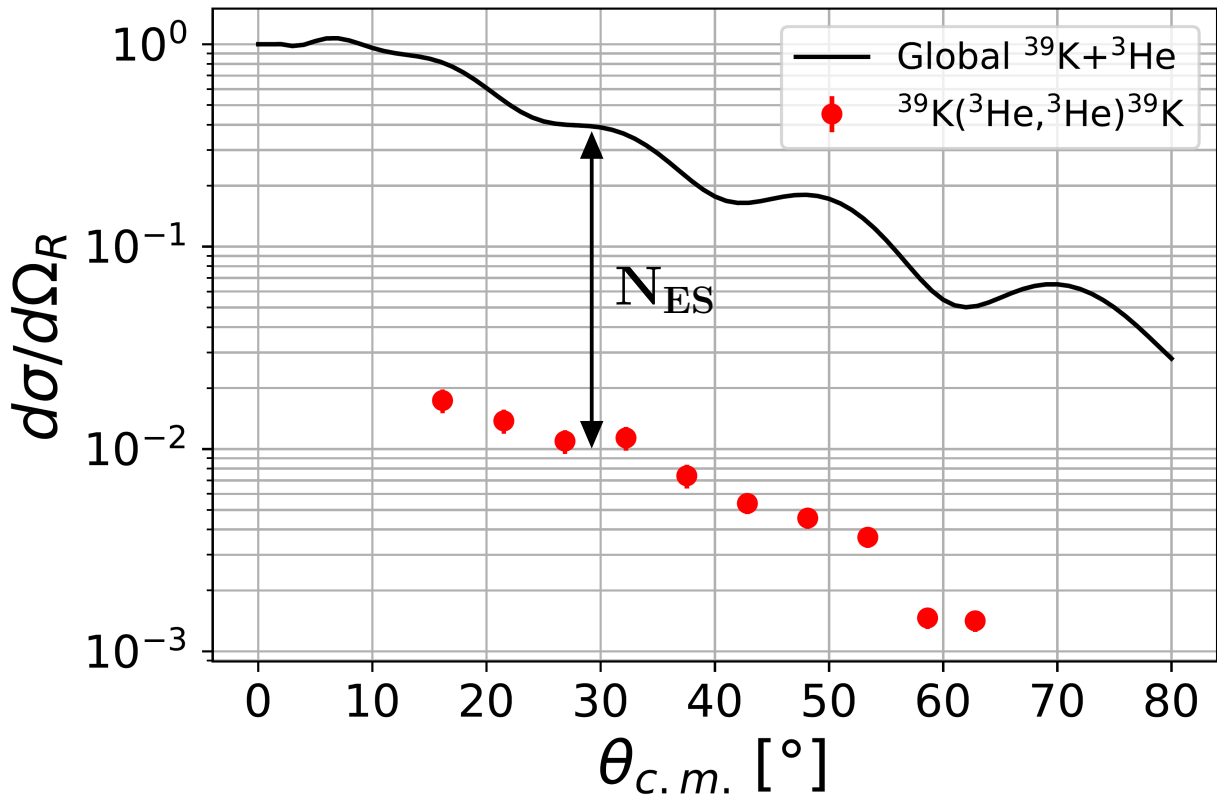


Figure 6.21: The differential cross-section, as a ratio to Rutherford scattering, of the global ${}^3\text{He}$ optical model potential of Ref. [Lia09] for ${}^{39}\text{K}$ (black line) and that of the present relative ${}^{39}\text{K}({}^3\text{He}, {}^3\text{He}){}^{39}\text{K}$ measurements (red data). The scaling factor N_{ES} is illustrated, which is also applied to the transfer data to establish an absolute scale.

where $R_{\text{ES},i}$ is defined by Eqn. 6.30 for the simultaneous ${}^{39}\text{K}({}^3\text{He}, {}^3\text{He}){}^{39}\text{K}$ measurements of the focal-plane and Si detectors, $(d\sigma/d\Omega_{\text{ES,global}})_i$ is the global OMP differential cross section evaluated at the experimental angle $\theta_{\text{c.m.,}i}$, and N is the number of experimental points. Since the global OMP differential cross section is not continuous, its value at each experimental center-of-mass angle must be interpolated. This is performed by cubic spline interpolation, where each pair of neighboring $d\sigma/d\Omega_{\text{ES,global}}$ points defines a polynomial that ensures its first and second derivatives at each point are equal to that of neighboring polynomials. After interpolating the N points, the scale factor becomes $N_{\text{ES}} = 0.0277(11) \approx 1/36$. Fig. 6.22 shows the global model scaled by N_{ES} to match the data. No χ^2 -minimization or Bayesian Monte Carlo algorithm has been performed on the OMP potential parameters. The global ${}^3\text{He}$ OMP parameters themselves sufficiently reproduce the angular distribution of the experimental points. However, the OMP parameters give rise to large systematic uncertainties in the spectroscopic factor calculations for the transfer reaction, as demonstrated in Section 6.8.4.

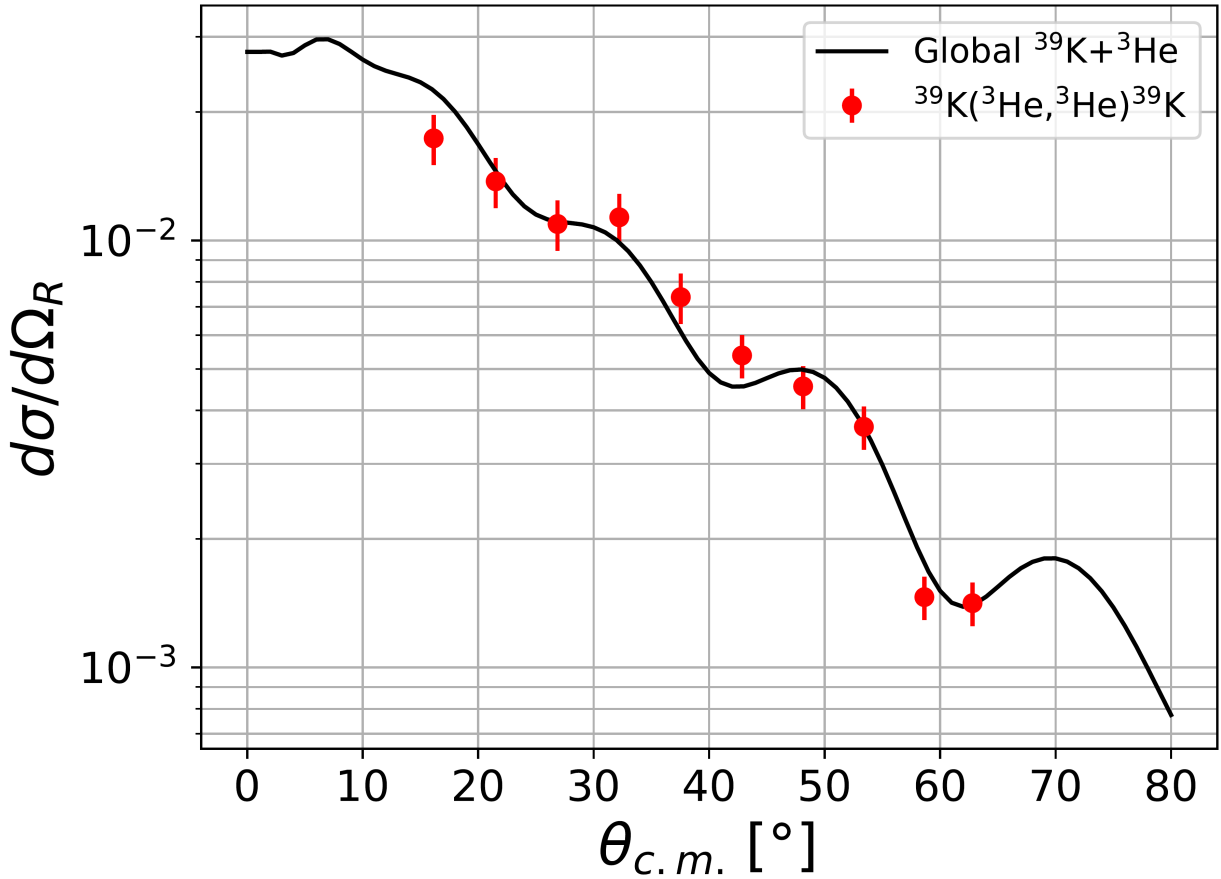


Figure 6.22: The same as Fig. 6.21, except the differential cross-section of the global ${}^3\text{He}$ optical model potential has been scaled by N_{ES} to match the experimental measurements, as described in the text.

The same N_{ES} normalization is applied to the transfer differential cross sections, since they were also measured relative to the elastic scattering of the Si detector. The ratio between this corrected ${}^{39}\text{K}({}^3\text{He}, d){}^{40}\text{Ca}$ differential cross section and that of an appropriate distorted-wave Born approximation (DWBA) model is the spectroscopic factor C^2S (see Sections 2.2.1 and 2.3.2). This ratio is performed in the exact same way as the global OMP normalization, including the cubic spline interpolation. The transfer differential cross sections will be presented in Section 6.8.2.

6.8 $^{39}\text{K}(\text{He}^3, d)^{40}\text{Ca}$ Analysis

6.8.1 Energy Calibrations

The centroid positions μ of the ^{40}Ca states along the P1 focal-plane spectra were used to determine their excitation energies, given several known calibration states. These μ were obtained with high precision through the Bayesian peak fitting procedure, detailed in Section 6.6. However, the centroid uncertainty obtained from the fit does not represent the true channel uncertainty. The fit underestimates the uncertainty, as it does not take into account scatter associated with the known focal-plane detector response. An additional scatter uncertainty of $u_s \sim 0.6$ channels was added to rectify this, corresponding to roughly 1 keV in excitation energy. The new channel uncertainty associated with each centroid value is

$$u_{\text{ch}}^2 = u_\mu^2 + u_s^2. \quad (6.39)$$

The oxidized target fits were also not used in the energy calibration. It is not clear whether the mode of the exponentially-modified gaussian (EMG) fits properly represents their energies. The inconsistency between using gaussian means and EMG modes could cause problems. However, the only angles that have oxidized target data, $\theta_{\text{lab}} = 5^\circ$ and 7° , also have data from a target that was not oxidized. The EMG fit area was still used for the yield determination because the total number of counts should not be affected by target oxidation. Scatter was added to these yields regardless to put more weight on those from non-oxidized targets in the weighted averages.

The ^{40}Ca calibration states that were used in the energy calibration are highlighted in red in Fig. 6.23 for the $\theta_{\text{lab}} = 5^\circ$ focal plane position spectrum. These are the 4491 keV, 5614 keV, 6025 keV, 7532 keV, 8425 keV, and 9454 keV ^{40}Ca states. The energies are provided by the ENSDF evaluation of Ref. [Che17], except the 8425 keV state uses the more recent energy measurement of Ref. [Gri22]. Unfortunately, not all of these states were unobscured throughout the course of the experiment, which made it necessary to use alternative calibration states in those cases. In total, of the 13 sort groups with non-oxidized targets in the $^{39}\text{K}(\text{He}^3, d)^{40}\text{Ca}$ experiment (see Section 6.5.3), an alternative calibration state had to be used for 4 of them. At both the $\theta_{\text{lab}} = 5^\circ$ (sort group #3) and $\theta_{\text{lab}} = 7^\circ$ (sort group #5) angles, the 8425 keV state was obscured by the first excited state from $^{16}\text{O}(\text{He}^3, d)^{17}\text{F}$. The alternative calibration state for these sort groups was the 8484 keV state, as it is closest in energy and was clearly resolved. Similarly, for both sort groups (#15 – #16) at $\theta_{\text{lab}} = 20^\circ$, the 9454 keV calibration state did not have enough counts to properly resolve it from the neighboring 9405 – 9432 keV multiplet. In these cases, the 9136 keV state was used as the alternative calibration state.

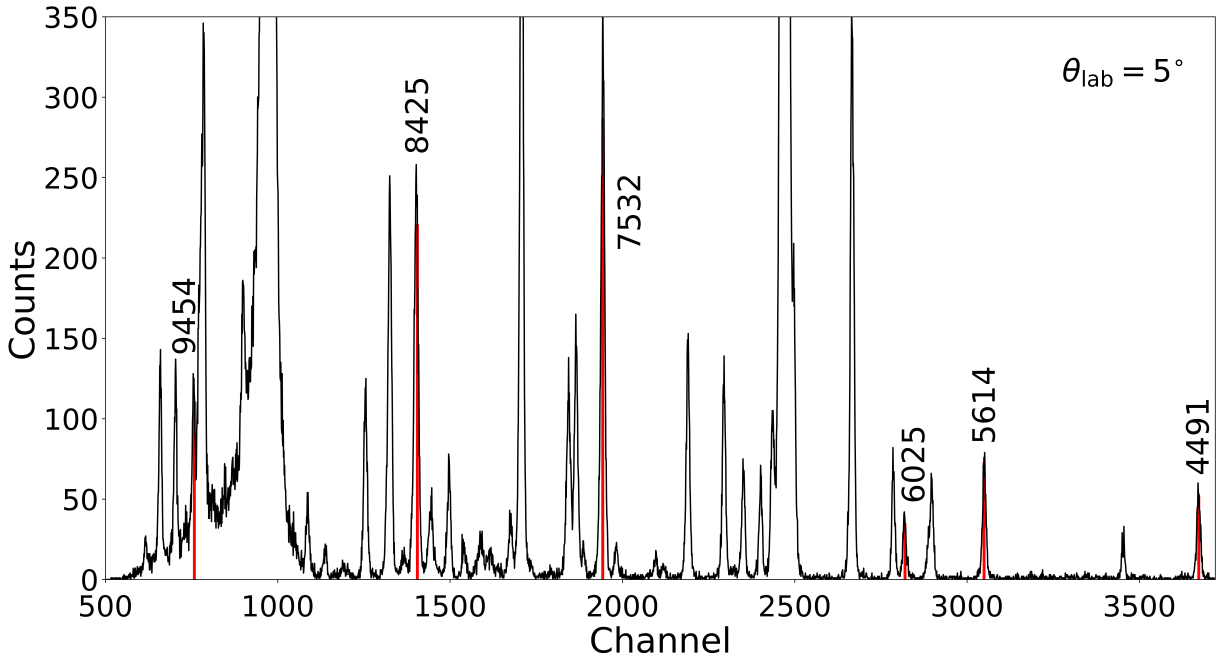


Figure 6.23: The calibration states for the MCMC energy calibration, represented by red vertical lines for the $\theta_{\text{lab}} = 5^\circ$ focal plane position spectrum. Their ENSDF [Che17] energies are labeled in keV. The 8425 keV state uses the energy of the recent measurement from Ref. [Gri22] instead. The 8484 keV and 9136 keV alternative calibration states are used when the 8425 keV and 9454 keV states were obscured at $\theta_{\text{lab}} = 5, 7^\circ$ and 20° , respectively. The ^{17}F contaminant appears in exactly the same position as the 8425 keV state in the figure.

The closer 9538 keV state was determined to be too near the edge of the detector to have a reliable centroid and yield measurement. The 9227 keV state was also ruled out due to it being impossible to determine which of the 9226.69 keV or 9227.43 keV states, or both, from ENSDF it belongs to, as they both have a wide range of possible J^π assignments. It was still possible to report energies for these alternative calibration states over the sort groups where they were not involved in the calibration.

The energy calibration was performed with a quadratic fit that represents the magnetic rigidity $B\rho$ of the Enge Split-Pole Spectrograph as a function of the focal-plane detector channels x_{ch} ,

$$B\rho = a + b_1 x_{\text{ch}} + b_2 x_{\text{ch}}^2. \quad (6.40)$$

The calibration uses x_{ch} and $B\rho$ from the known calibration states to predict $B\rho$ for all other states, known as test states. The excitation energies of the calibration states are first converted to $B\rho$ by interpolation from the kinematics code `jrelkin` [Vis03]. After the calibration, the predicted $B\rho$ are converted back to energy in the same manner, with uncertainties propagated

through both the calibration and the energy conversion.

Markov chain Monte Carlo (MCMC) fits were performed with Eqn 6.40 using the Hamiltonian Monte Carlo and Metropolis algorithms built into `rstan` [Tea11, Guo20]. This allowed for Bayesian statistical inference of the fit parameters from MCMC samples. The parameters, including an extra scatter parameter σ added in quadrature to each $B\rho$ calibration state uncertainty, were given the prior distributions

$$\begin{aligned} a &\sim \mathcal{N}(0, 1^2), \\ b_1 &\sim \mathcal{N}(0, 1^2), \\ b_2 &\sim \mathcal{N}(0, 1^2), \\ \sigma &\sim \text{Cauchy}(0, 1^2). \end{aligned} \quad (6.41)$$

A total of 5×10^4 iterations were performed for each of the four chains that are used by `rstan`. Samples were extracted from the last 2.5×10^4 iterations, and the first half were used as warmup. Using the new posteriors, $B\rho$ was sampled over the 4096 channels and converted to ^{40}Ca excitation energy. Energy residuals from the median calibration line were also sampled over the 4096 channels.

The energy residuals for $\theta_{\text{lab}} = 5^\circ$ (sort group #3) are shown in Fig. 6.24. The 68% (1σ) and 95% (2σ) confidence intervals from the MCMC samples are shown in dark and light shading, respectively, centered on the median calibration line. The red points represent the calibration states, and the black points represent the test states. Uncertainty bars shown for the test states are simply their input centroid uncertainties u_{ch} converted to energy from the fit, $u_{\text{ch-to-E}}$, added in quadrature with the ENSDF uncertainties. The uncertainties reported for each test state among each sort group are $u_{\text{ch-to-E}}$ added in quadrature with half the 1σ width from the fit at the given energy. That is,

$$u_E^2 = u_{\text{ch-to-E}}^2 + \left(\frac{u_{\text{MCMC,upper}} - u_{\text{MCMC,lower}}}{2} \right)^2, \quad (6.42)$$

where upper and lower refer to the 84th and 16th percentile of the MCMC samples, respectively.

The calibration was performed for every sort group, except for those where an oxidized target was used (sort groups #1, #2 and #4). Each test state therefore resulted in up to 13 reported energies, 1 for each sort group where the state was resolvable. Several averaging methods were attempted to obtain a final reported energy for each test state. A weighted average \bar{E}_{wavg} was the first attempt, where the weights were $1/u_E^2$, and the uncertainties were propagated through the weighted average. However, it was clear that for many test states, the scatter was not properly accounted for with the weighted average and its uncertainty.

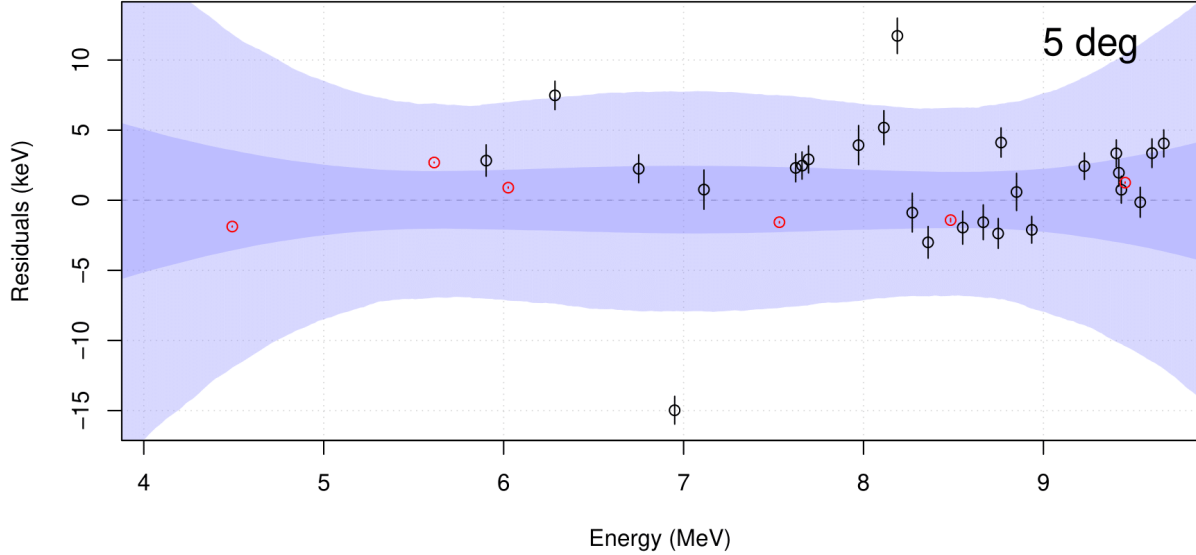


Figure 6.24: Residuals between the excitation energies predicted from the MCMC energy calibration and that of the median calibration line at $\theta_{\text{lab}} = 5^\circ$ (sort group #3) for the calibration states (in red) and the test states (in black). The 68% (1σ) and 95% (2σ) confidence intervals for the fit are represented by the dark and light shading, respectively. See text for details.

The few sort groups with smaller uncertainties shifted the average unreasonably far from that of the other sort groups. A simple average \bar{E} was then attempted, which seemed to be better suited for the scatter. However, the uncertainty propagated through the simple average also did not reproduce the scatter appropriately. What finally captured the appropriate scatter was taking a simple average of the reported uncertainties. That is,

$$\bar{E} \pm u_{\bar{E}} = \frac{1}{N} \sum_i^N E_i \pm \frac{1}{N} \sum_i^N u_{E,i}, \quad (6.43)$$

was determined to be the appropriate final energy and uncertainty for each state.

To illustrate these different averaging methods, the resulting distribution of calibrated energies for the representative 8551 keV state is shown by the black data in Fig. 6.25. This is an example where the scatter was minimized, potentially due to its relatively large yield, but the different averaging methods are clearly distinguished. The black line represents the simple average, while the orange line represents the weighted average. The propagated uncertainty from each of these averages are the blue and orange bands, respectively. The green band

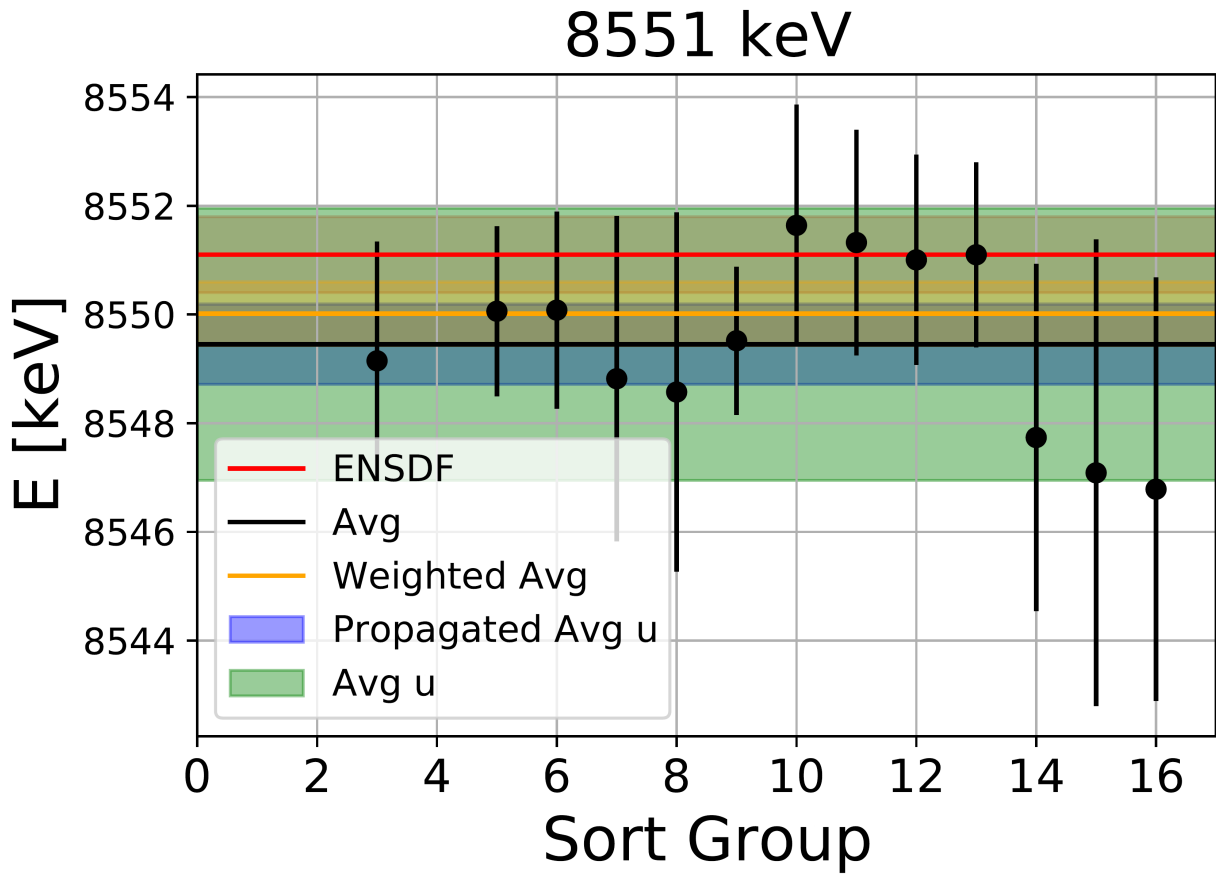


Figure 6.25: Excitation energies and their uncertainties of the 8551 keV state from the MCMC energy calibration for each sort group, excluding sort groups #1 and #2 that used an oxidized target. The ENSDF energy and its uncertainty are given as the red line and red band, respectively. A simple average and a weighted average of the energies are given by the black and orange lines, respectively. The uncertainty propagated from the simple average and from the weighted average are given by the blue and orange bands, respectively. The simple average of the MCMC uncertainties is represented by the green band, which best captures the range of the individual uncertainties for each state.

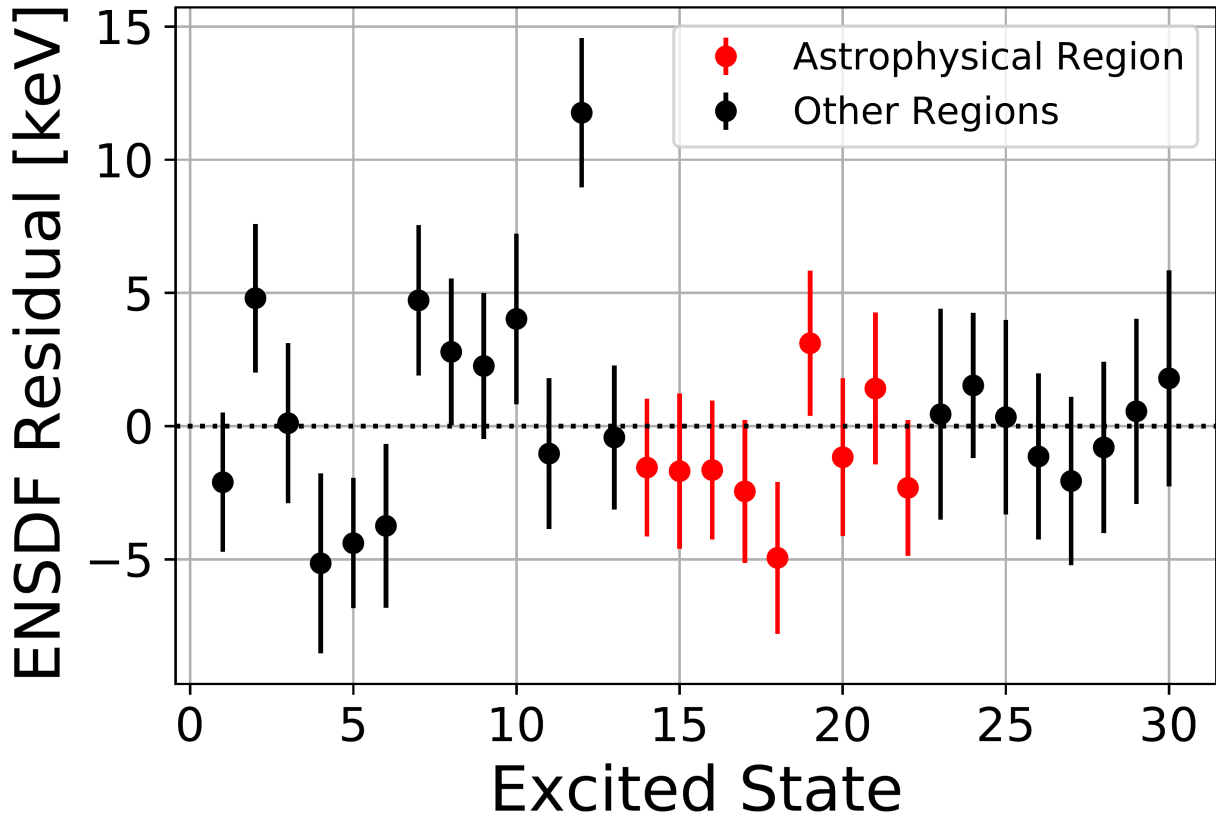


Figure 6.26: Residuals between the final reported energies for each state from the MCMC energy calibration and that of ENSDF [Che17]. The residuals and uncertainties for each state result from a simple average of the MCMC energies and uncertainties of each sort group, respectively. The residuals in red represent the astrophysical region, from 8359 keV to 8995 keV. The x -axis values correspond to the following states in keV; 1: 5903, 2: 6285, 3: 6582, 4: 6750, 5: 6950, 6: 7113, 7: 7623, 8: 7658, 9: 7694, 10: 7973, 11: 8113, 12: 8188, 13: 8271, 14: 8359, 15: 8484, 16: 8551, 17: 8665, 18: 8748, 19: 8764, 20: 8851, 21: 8935, 22: 8995, 23: 9092, 24: 9136, 25: 9227, 26: 9405, 27: 9419, 28: 9432, 29: 9538, 30: 9603.

shows the simple average uncertainty of Eqn. 6.43, coming from the black line. The energy from ENSDF is shown in red, with its uncertainty band also in red. Note that the blue band does not agree with ENSDF, and there is also tension between the orange band and ENSDF.

The final residuals between the energy-calibrated states using Eqn. 6.43 and their corresponding ENSDF state are shown in Fig. 6.26. It is clear that the typical energy uncertainty is in the range of 2 – 3 keV. The states highlighted in red are the ones that contribute to the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate and are therefore of astrophysical interest. These range from 8359 keV to 8995 keV. The excited state labels are provided in the figure capture. There is decent agreement with ENSDF for most states at relatively high excitation energy, but there is considerable scatter at lower energies. In particular, the state that most disagrees with ENSDF is the 8188 keV state, which has a calibrated energy of 8199.3(27) keV, or a residual of 11.8(28) keV. This may correspond to the 8196 keV state in ENSDF instead, but there is a considerable lack of experimental evidence for this state. This hypothesis will be discussed further in Section 6.8.2.

The final energies from the MCMC calibration are reported in Table 6.3. Calibration states are shown in italics, except for the alternative calibration states 8484 keV and 9136 keV that were still able to have their energies determined. Cases where the observed state corresponds to an unresolvable doublet according to ENSDF are represented with a bracket around the doublet. The assigned J^π values from ENSDF are also given. Fig. 6.27 shows an energy calibrated focal plane spectrum of the proton-unbound ^{40}Ca states labeled with their ENSDF [Che17] energies, where the proton separation energy $S_p = 8328(18)$ keV is indicated by the red line.

Table 6.3: Excitation energies E_x determined through a Markov Chain Monte Carlo (MCMC) energy calibration for the ^{40}Ca states observed in the present experiment. For comparison, the ENSDF [Che17] excitation energies and J^π values are also given, unless otherwise indicated. Brackets represent states from ENSDF too close in energy to resolve via the energy calibration alone. Calibration states are shown in italics.

E_x^{a} [keV]	E_x^{b} [keV]	J^π ^a	E_x^{a} [keV]	E_x^{b} [keV]	J^π ^a
4491.43(4)	<i>4491</i> ^c	5^-	8850.6(9)	8849.5(27)	$6^-, 7^-, 8^-$
5613.52(3)	<i>5614</i> ^c	4^-			
5902.63(7)	5900.5(26)	1^-	$\begin{cases} 8934.81(7) \\ 8938.4(9) \end{cases}$	8936.3(27)	2^+
6025.47(5)	<i>6025</i> ^c	2^-			0^+
6285.15(4)	6290.0(28)	3^-			
6582.47(10)	6582.6(30)	3^-	8994.5(11)	8992.2(25)	$(1^-, 2^+)$
6750.41(7)	6745.3(34)	2^-	9091.7(6)	9092.2(36)	3^-
6950.48(7)	6946.1(24)	1^-	9135.66(5)	9137.8(34) ^c	$2^-, 3^-$
$\begin{cases} 7113.1(10) \\ 7113.73(5) \end{cases}$	7109.4(29)	1^-	$\begin{cases} 9226.69(5) \\ 9227.43(7) \end{cases}$	9227.8(34)	$(1^-, 2, 3^-)$
		4^-			$(1, 2^+)$
7532.26(5)	<i>7532</i> ^c	2^-	$\begin{cases} 9404.85(19) \\ 9406.3(6) \end{cases}$	9403.7(31)	2^-
7623.11(8)	7627.8(28)	$(2^-, 3, 4^+)$			0^+
7658.23(5)	7661.0(28)	4^-			
7694.08(4)	7696.3(27)	3^-	$\begin{cases} 9412.3(2) \\ 9418.8(2) \end{cases}$	9416.7(32)	
7972.5	7976.5(30)	3^-			3^-
8113.2(5)	8112.2(28)	1^-			
8187.5(8)	8199.3(27)	$3, 4, 5^-$	$\begin{cases} 9429.11(5) \\ 9432.46(18) \end{cases}$	9431.7(32)	$(3, 4)^-$
8271(1)	8270.6(25)	3^-			1^-
8358.9(6)	8357.3(25)	$(0, 1, 2)^-$			
8424.35(31) ^d	<i>8425</i> ^c	2^-	9453.95(5)	<i>9454</i> ^c	3^-
8484.02(13)	8482.3(29) ^c	$(1^-, 2^-, 3^-)$	9537.8(5)	9538.4(34)	1^-
8551.1(7)	8549.5(25)	5^-			
8665.3(8)	8662.9(25)	1^-	$\begin{cases} 9603.0(4) \\ 9604.6(4) \end{cases}$	9605.1(40)	3^-
8748.59(19) ^d	8743.6(28)	2^+			1^-
8764.18(6)	8767.3(27)	3^-			

^aFrom the ENSDF evaluation of Ref. [Che17], unless otherwise indicated.

^bPresent experiment.

^cLevel used as a calibration point for at least one angle in this work. See text for details.

^dEnergy from the nuclear resonance fluorescence measurement of Ref. [Gri22].

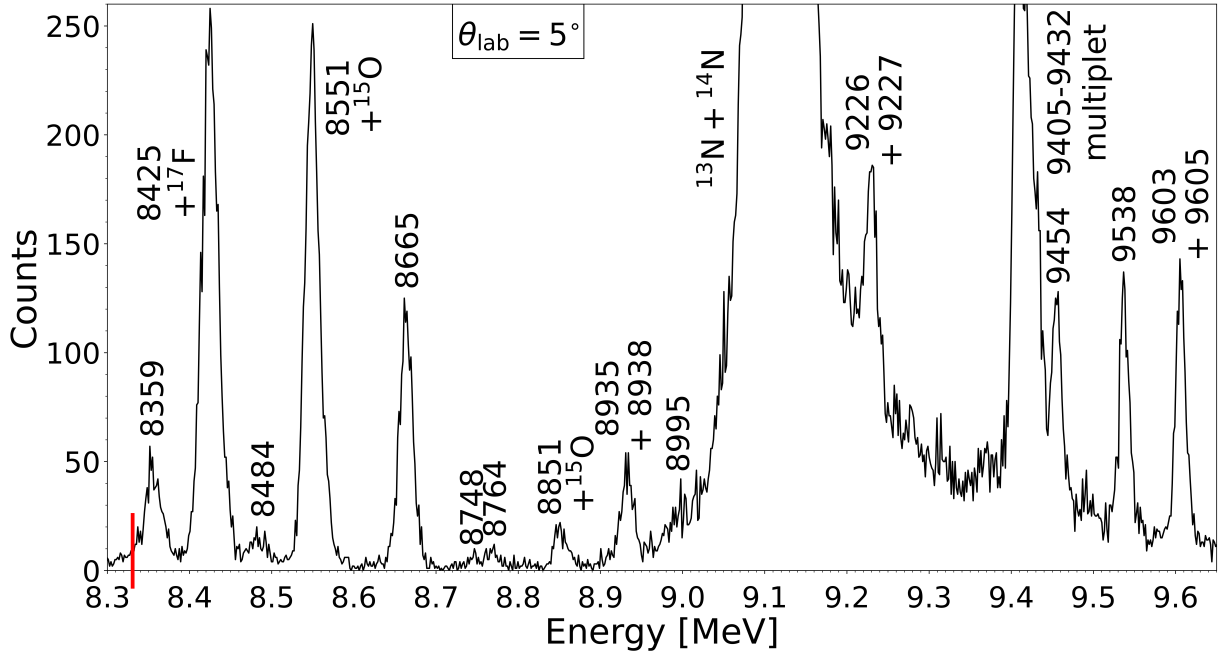


Figure 6.27: The energy-calibrated focal plane position spectrum, focused on the ^{40}Ca states above the proton separation energy at $\theta_{\text{lab}} = 5^\circ$ (sort group #3). The ^{40}Ca states are labeled with their ENSDF [Che17] energies in keV, and contaminants at this angle are indicated. The red vertical line indicates the proton separation energy, $S_p = 8328.18(2)$ keV from Ref. [Wan21].

6.8.2 DWBA Calculations

The $^{39}\text{K}(\text{He}^3, d)^{40}\text{Ca}$ differential cross sections were normalized by the same amount N_{ES} as that of the focal plane $^{39}\text{K}(\text{He}^3, \text{He})^{39}\text{K}$ measurements, since both were measured relative to the elastic scattering of the Si detector. The remaining normalization,

$$\frac{d\sigma}{d\Omega_{\text{Exp}}} = C^2 S \frac{d\sigma}{d\Omega_{\text{DWBA}}}, \quad (6.44)$$

between the scale-corrected transfer differential cross section $d\sigma/d\Omega_{\text{Exp}}$ and that of an appropriate distorted-wave Born approximation (DWBA) $d\sigma/d\Omega_{\text{DWBA}}$ defines the spectroscopic factor $C^2 S$. The spectroscopic factor for $^{39}\text{K}(\text{He}^3, d)^{40}\text{Ca}$ is broken into two parts, corresponding to the $^{39}\text{K} + p$ and $d + p$ overlaps (see Section 2.3.2),

$$C^2 S = C^2 S_{^{39}\text{K}+p} C^2 S_{d+p}. \quad (6.45)$$

The C^2S_{d+p} spectroscopic factor is frequently omitted in the literature, as it is a constant factor applied to all C^2S for $(^3\text{He}, d)$ reactions. It can be approximated as $A/2$ for $A \leq 4$ nuclei [Sat88], which implies $C^2S_{d+p} = 1.5$. However, *ab-initio* calculations such as those from Ref. [Bri11] can be more precise, reducing systematic uncertainties associated with $C^2S_{^{39}\text{K}+p}$. Ref. [Bri11] calculates $C^2S_{d+p} = 1.32(1)$, which is adopted in the present analysis. The $C^2S_{^{39}\text{K}+p}$ spectroscopic factor for proton-unbound states, when converted to a proton partial-width in Section 6.8.5, is a direct input in the ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ reaction rate. This is the main quantity of astrophysical interest in transfer reactions.

DWBA transfer reaction calculations require optical model potentials (OMPs) for both the incoming ${}^{39}\text{K} + {}^3\text{He}$ and the outgoing $d + {}^{40}\text{Ca}$ channels. The potential for the incoming channel is evaluated at the lab beam energy E_{lab} , whereas the outgoing channel potential is evaluated at $E_{\text{lab}} - Q$, where Q is the Q -value of the transfer reaction. The global ${}^3\text{He}$ OMP of Ref. [Lia09] from the elastic scattering normalization is retained for the incoming channel. The parameters of this OMP were given in Table 6.2. For the outgoing channel, the global d OMP of Ref. [An06] is used in the present analysis, where the potential is defined in the exact same way as that of Ref. [Lia09].

The proton bound-state potential must also be specified for DWBA calculations of proton-bound ${}^{40}\text{Ca}$ states. It consists of a real volume potential with a Woods-Saxon form and a Coulomb potential. The Woods-Saxon well depth V_r is automatically adjusted to reproduce the proton binding energy,

$$\text{BE}({}^{A-1}(X-1) + p) = S_p({}^A X) - E_x, \quad (6.46)$$

where S_p is the proton-separation energy of the compound nucleus ${}^A X$, and E_x is its excitation energy. However, many of the states observed in the present experiment, and indeed all of the states of astrophysical interest, are proton-unbound. In these cases, one could adjust the Woods-Saxon well depth until a resonance is formed at E_r . This is unfortunately not performed automatically in **Fresco** [Tho88, Tho06], the nuclear reaction code chosen for the present analysis. An alternative to this is to approximate the resonance scattering wave function as the wave function for a loosely-bound particle, with a binding energy of ~ 1 keV. This is known as the weak binding approximation, and it has been shown to reproduce unbound calculations to within 1% for states within about 500 keV of the particle threshold and 6% for states within about 1800 keV above the particle threshold [Kan16, Kah19]. The present analysis uses the weak binding approximation for unbound ${}^{40}\text{Ca}$ states. The proton bound-state parameters, along with the global OMP parameters are summarized in Table 6.4.

Another factor that must be considered in DWBA calculations is the transferred-particle

Table 6.4: Global optical model potential parameters for $^{39}\text{K} + ^3\text{He}$ and $^{40}\text{Ca} + d$ and the proton bound-state parameters.

particle	V_r	r_r	a_r	W_v	r_v	a_v	W_s	r_s	a_s	V_{so}	r_{so}	a_{so}	W_{so}	r_c
^3He	117.881	1.178	0.768	-0.646	1.415	0.847	20.665	1.198	0.852	2.083	0.738	0.946	-1.159	1.289
d	91.120	1.150	0.762	2.234	1.334	0.513	10.274	1.378	0.743	3.557	0.972	1.011		1.303
p	V_p^1	1.250	0.650											1.250

overlaps, in this case for $^{39}\text{K} + p$ and $d + p$. Each overlap is specified by a single-particle state $n\ell_j$ from the shell model with the quantum numbers n , ℓ , and j corresponding to the node, transferred orbital angular momentum, and total angular momentum ($j = \ell \pm 1/2$), respectively (see Ref. [Kra87]). Only a select few single-particle states can be populated from a given transition, based on quantum-mechanical selection rules for angular momentum and parity (see Ref. [Ili15]). For example, if the spin-parity J^π of the final ^{40}Ca state is known, the possible n , ℓ , and j single-particle quantum numbers can be extracted from the selection rules. This can also be done for final states where J^π is narrowed down to a few possibilities. This extraction was performed for each excited state of ^{40}Ca populated in the present experiment, given their known spin-parities, or spin-parity candidates, from the ENSDF evaluation of Ref. [Che17]. The $d + p$ overlap was treated as constant for each DWBA calculation, since it was assumed that it occupied only the ^3He ground state.

The nuclear reaction code **Fresco** [Tho88, Tho06] was used to calculate $^{39}\text{K}(^3\text{He}, d)^{40}\text{Ca}$ DWBA differential cross sections using the zero-range (ZR) approximation introduced in Section 2.3.2. Calculations were performed for the allowed $n\ell_j$ combinations of each excited state, based on their known J^π values from ENSDF. In cases where none of the angular distributions agreed with the experimental cross section, the $n\ell_j$ combinations were expanded to account for different final J^π values. An example of the normalization procedure and the different $n\ell_j$ combinations for the 7532 keV state is shown in Fig. 6.28. This state has a spin-parity of 2^- from ENSDF, and therefore allows $\ell = 1$ or $\ell = 3$ orbital angular momenta from selection rules. The most likely single-particle states corresponding to these ℓ -values are $2p_{3/2}$ or $2p_{1/2}$ for $\ell = 1$ and $1f_{7/2}$ or $1f_{5/2}$ for $\ell = 3$, as they are the lowest energy transitions from the ^{39}K ground state at $1d_{3/2}$. The higher n -values for a given ℓ transition have a much greater transition energy and are therefore less likely. For that reason they are typically omitted in the literature. However, they were included in the analysis for completion and to test their effects. From Fig. 6.28, it is clear that the DWBA differential cross sections increase with increasing n . It can also be seen that the n - ℓ pairs produce very similar differential cross sections at low angles. This is especially true at angles below the first minimum, where the DWBA differential cross sections are most reliable. It is common during normalization

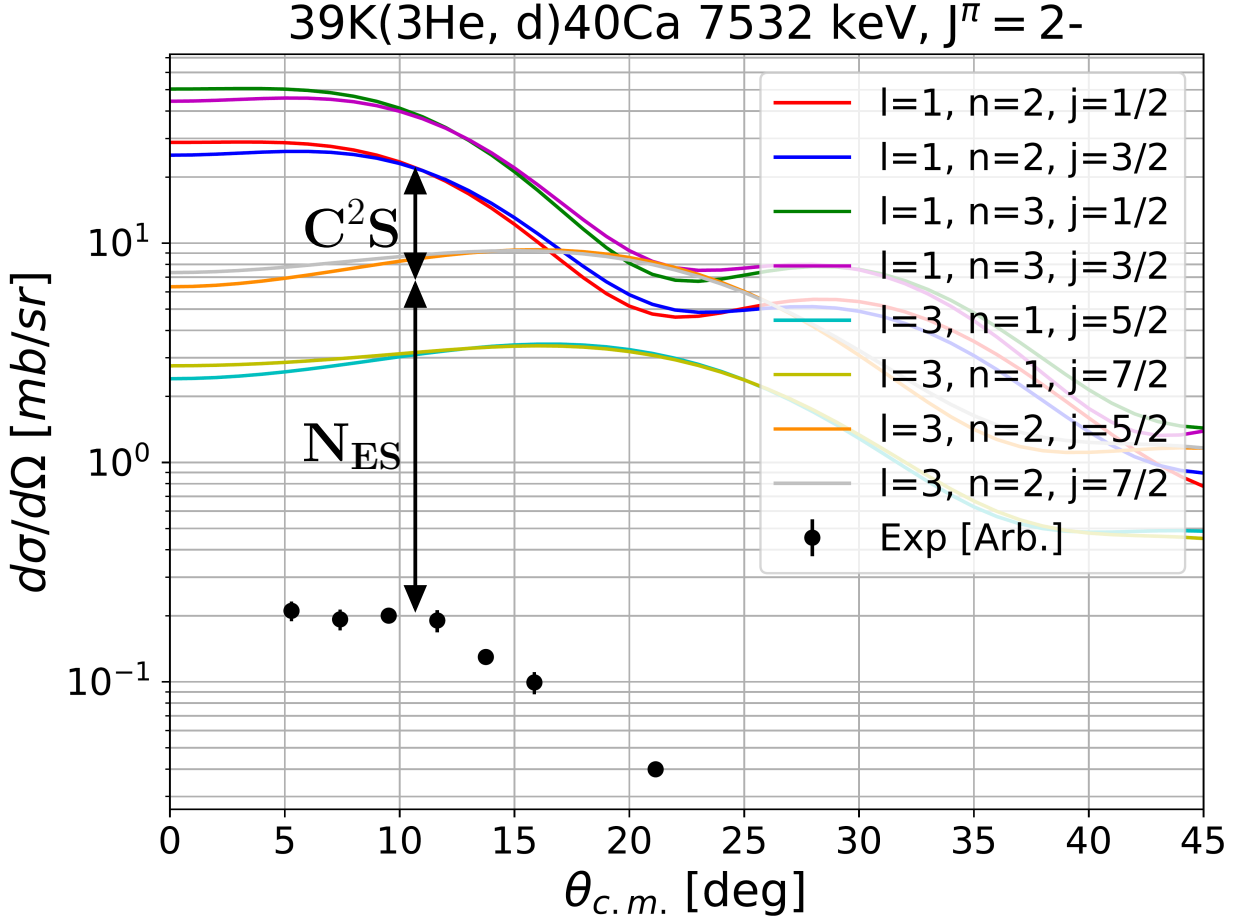


Figure 6.28: Normalization between the relative transfer differential cross section of the 7532 keV state and that of all possible DWBA models based on the current ENSDF $J^\pi = 2^-$ assignment. The normalization N_{ES} between elastic scattering and the global ${}^3\text{He}$ OMP is shown to scale, as is the C^2S normalization.

to omit angles greater than the first DWBA minimum because that is roughly where the distorted-wave Born approximation loses validity [??]. However, the present ${}^{39}\text{K}({}^3\text{He}, d){}^{40}\text{Ca}$ experiment did not obtain data beyond the first minimum, so normalization was performed with all experimental points.

Fig. 6.29 shows the same differential cross sections as in Fig. 6.28, but the DWBA models have been scaled down to the experimental points to show all normalizations simultaneously. It is clear that the $\ell = 1$ models match the angular distribution of the points. However, it is not possible to tell which of the $\ell = 1$ models is the best match. As is done in the literature [Ers66, Set67, For70, Cag71, Fuc69], the present analysis chooses the $n-j$ pair associated with the lowest energy single-particle state, unless it is forbidden by selection rules. That is, unless

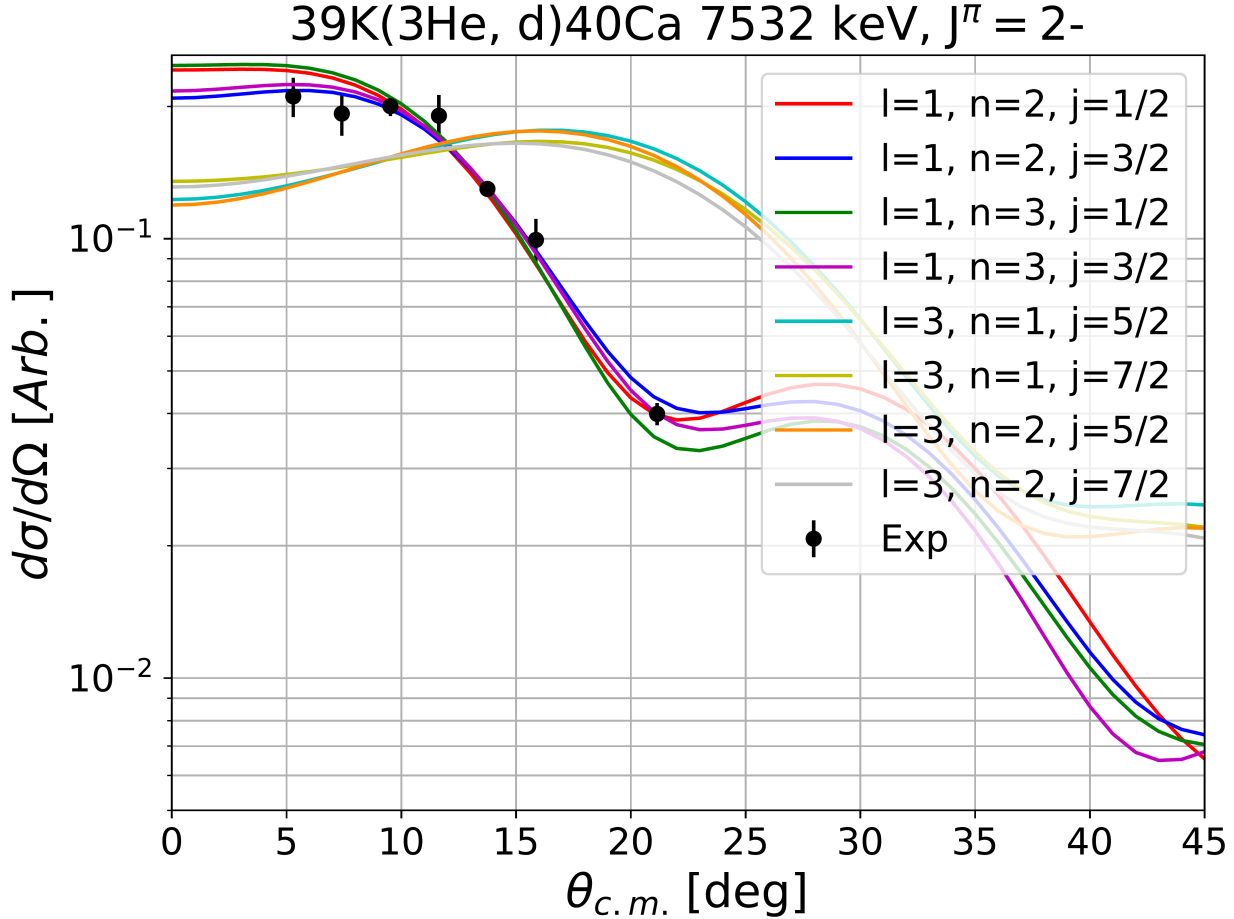


Figure 6.29: The DWBA models of Fig. 6.28 scaled down to the 7532 keV data to compare angular distributions. The $l = 1$ models are a clear match for this state.

they are forbidden, all $\ell = 0$, $\ell = 1$, $\ell = 2$, and $\ell = 3$ distributions will be associated with $3s_{1/2}$, $2p_{3/2}$, $2d_{5/2}$, and $1f_{7/2}$ single-particle states, respectively.

Finally, another transfer consideration comes from the fact that ^{39}K has a non-zero ground state ($J^\pi = 3/2^+$). It is possible in this case that more than one single-particle state contributes to the differential cross section, resulting in a mixed- ℓ angular distribution. Theoretically, all allowed ℓ transitions can contribute, but it is usually impossible to distinguish between more than the two lowest ℓ transitions. The differential cross section decreases with increasing ℓ , making each larger ℓ contribution more negligible [Hod71]. The mixed- ℓ angular distributions should then resemble a sum of two DWBA differential cross sections,

$$\frac{d\sigma}{d\Omega}_{\text{Exp}} = C^2 S_{\ell_1} \frac{d\sigma}{d\Omega}_{\text{DWBA}, \ell_1} + C^2 S_{\ell_2} \frac{d\sigma}{d\Omega}_{\text{DWBA}, \ell_2}. \quad (6.47)$$

The individual spectroscopic factors are given the common parameter α which determines their fractional contributions, as in

$$C^2 S_{\ell_1} = \alpha C^2 S, \quad (6.48)$$

$$C^2 S_{\ell_2} = (1 - \alpha) C^2 S, \quad (6.49)$$

where $C^2 S$ represents the joint spectroscopic factor. A χ^2 -minimization is performed in mixed- ℓ cases to determine the α and $C^2 S$ parameters, and the individual $C^2 S_{\ell_1}$ and $C^2 S_{\ell_2}$ are then reported. Their statistical uncertainties are obtained from the variance-covariance matrix elements added in quadrature to the $d\sigma/d\Omega_{\text{Exp}}$ uncertainties.

6.8.3 Angular Distribution Results

Figs. 6.30–6.34 show the angular distributions extracted for each ^{40}Ca state observed in the $^{39}\text{K}(^{3}\text{He}, d)^{40}\text{Ca}$ experiment, with a few exceptions. The states observed above 9227 keV (9405–9603 keV) did not match any DWBA prescription. The experimental differential cross sections for these states rapidly decreased as a function of angle faster than even the lowest possible ℓ transition. This is likely due to these peaks being too near the edge of the focal plane detector, which is exacerbated with increasing angle. The detector efficiency is significantly reduced at the fringes, which makes yield measurements unreliable. Another factor could be the breakdown of the weak binding approximation in DWBA models for states with energies this high above the proton separation energy, albeit likely to a lesser degree, considering the good agreement with the nearby 9227 keV differential cross section. Energies were still extracted for states up to 9603 keV with relatively small residuals from ENSDF [Che17] because detector efficiency does not affect centroid position, and DWBA calculations are not used in the energy calibration. Fig. 6.34 focuses on the proton-unbound ^{40}Ca states of astrophysical interest ($E_x = 8359$ –8851 keV), which are below the lowest directly measured $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ resonance at $E_x = 8935$ keV ($E_r^{\text{c.m.}} = 606$ keV). These are the states that are updated in the new reaction rate calculation, shown in Section 6.9.

The angular distribution of each state is described in detail in the following discussion, where each excited state energy is labeled by its ENSDF [Che17] energy for clarity. For proton-unbound states, the resonance energy $E_r^{\text{c.m.}}$ is also provided based on the energy calibration of the present analysis. Spectroscopic factors are compared to literature in Section 6.8.4, and the proton partial-widths of the states of astrophysical interest are discussed in Section 6.8.5.

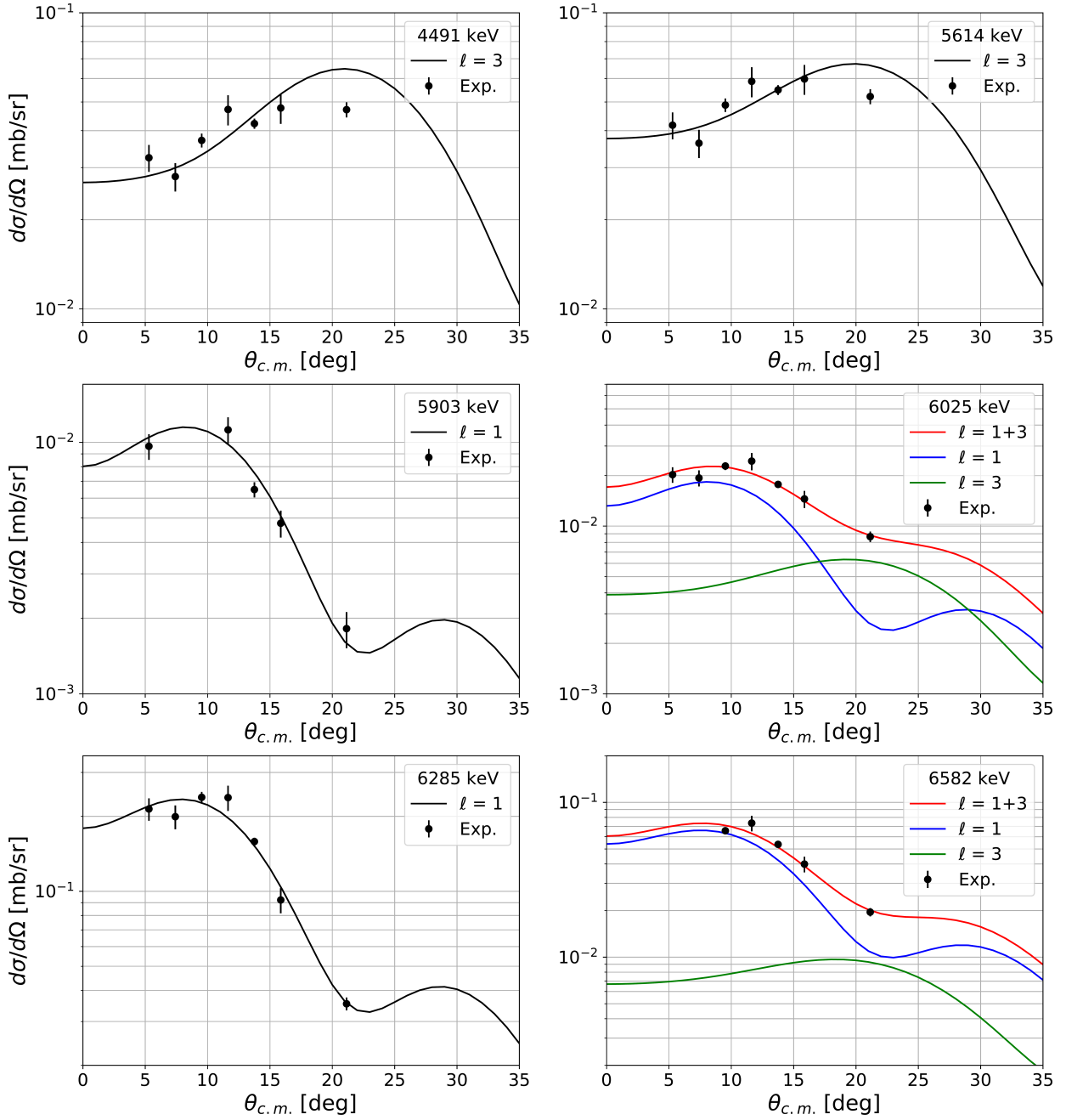


Figure 6.30: Differential cross sections for 4491 keV through 6582 keV. See text for details.

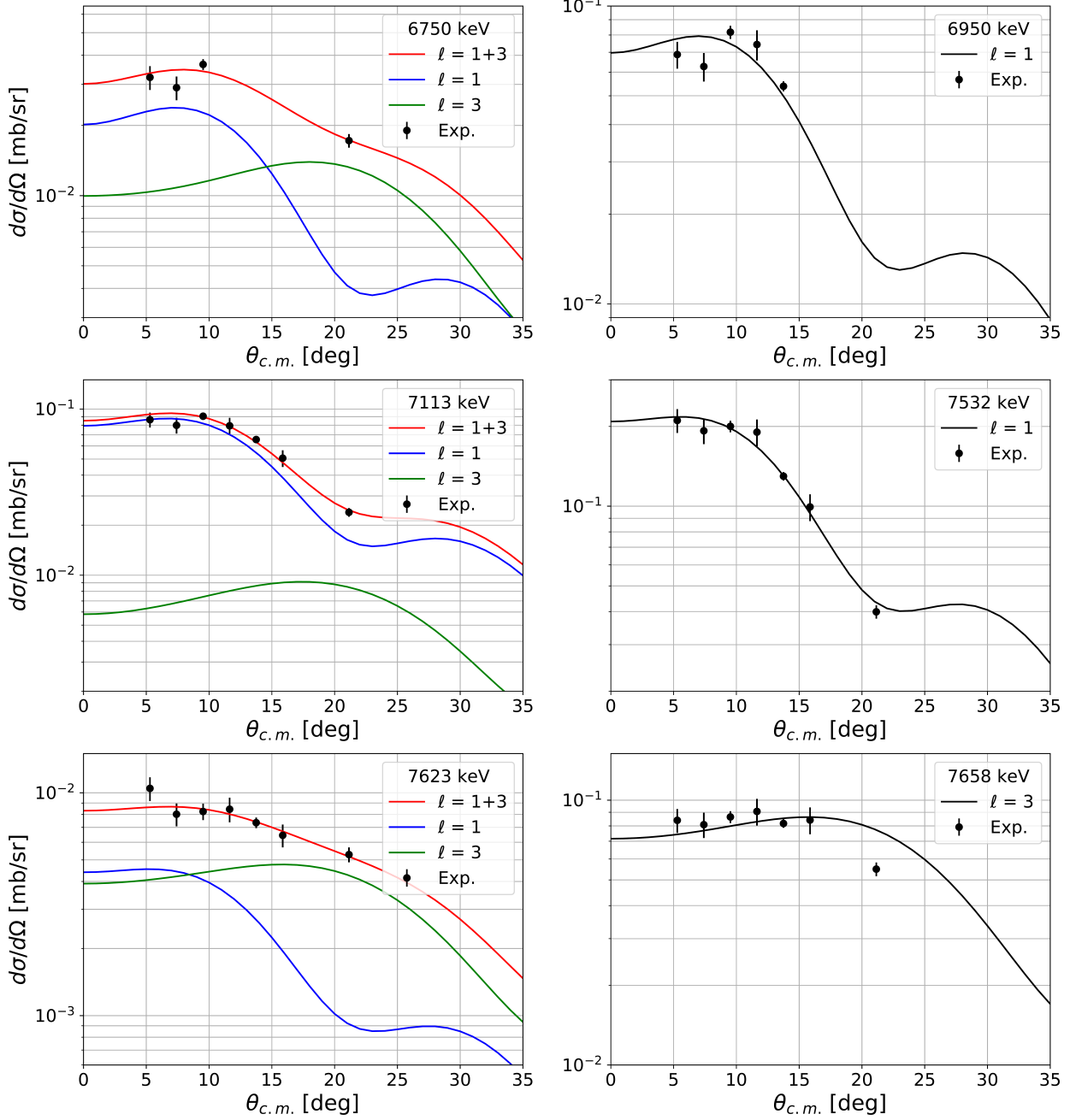


Figure 6.31: Differential cross sections for 6750 keV through 7658 keV. See text for details.

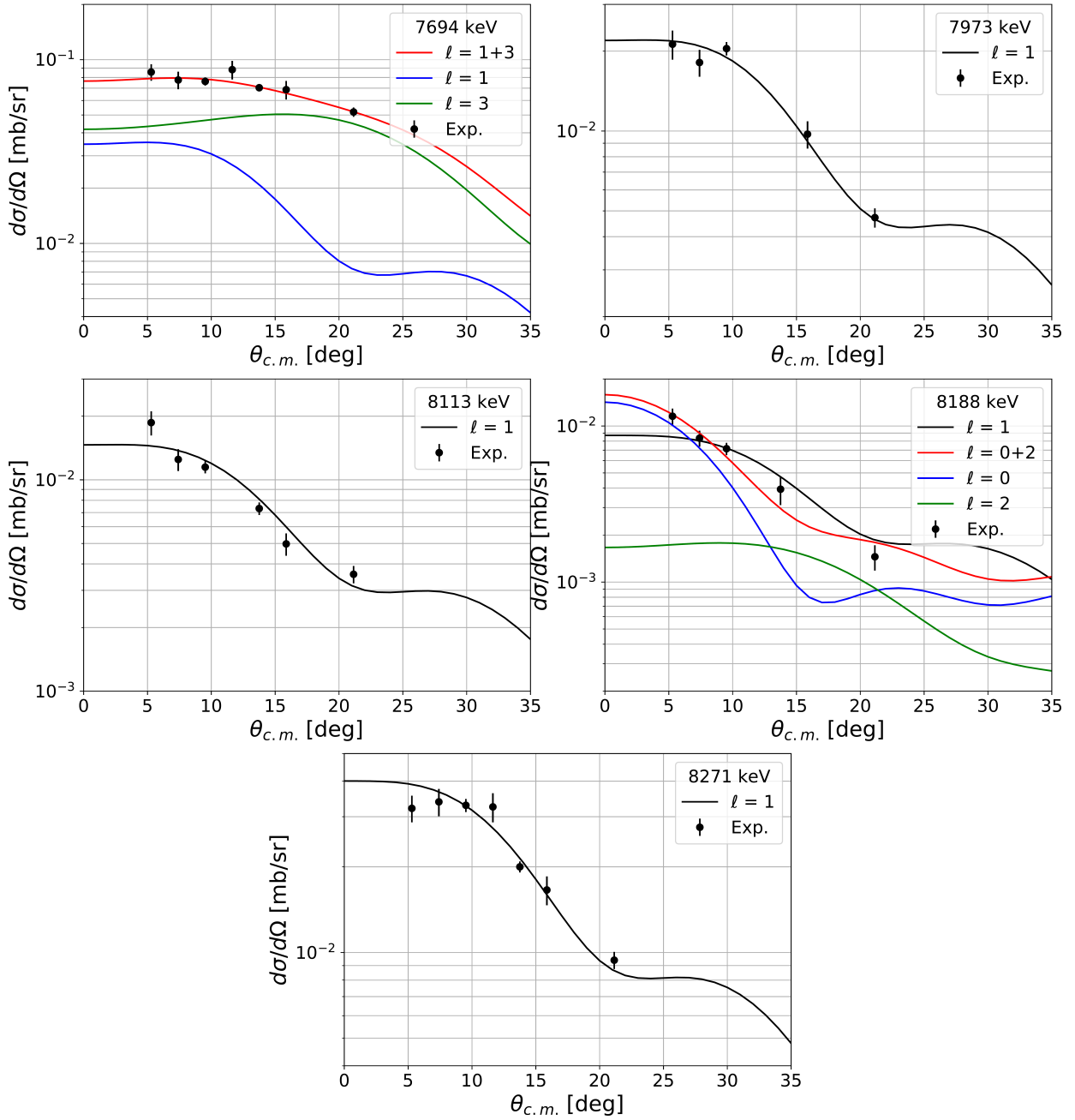


Figure 6.32: Differential cross sections for 7694 keV through 8271 keV. See text for details.

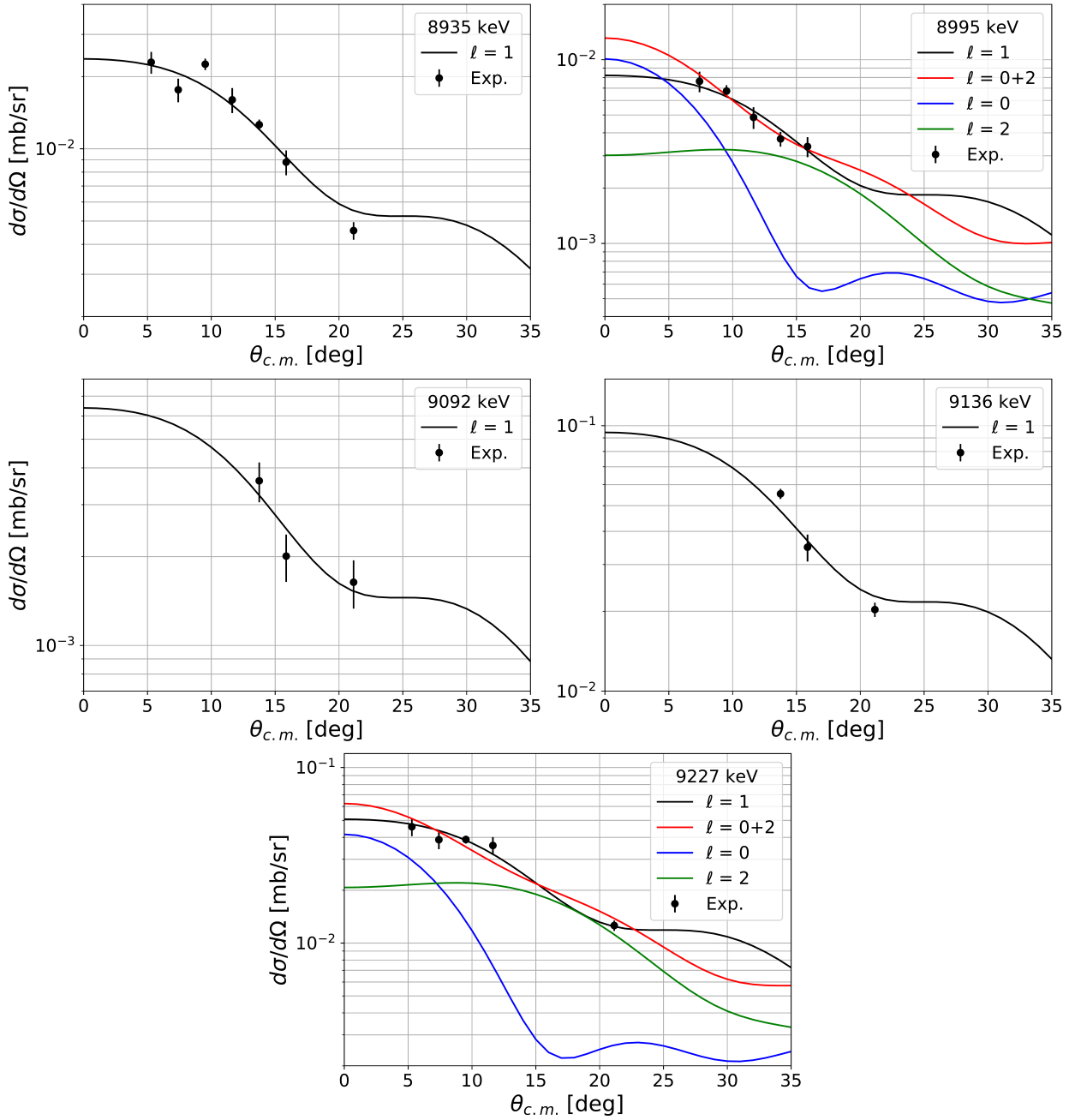


Figure 6.33: Differential cross sections for 8935 keV through 9227 keV. See text for details.

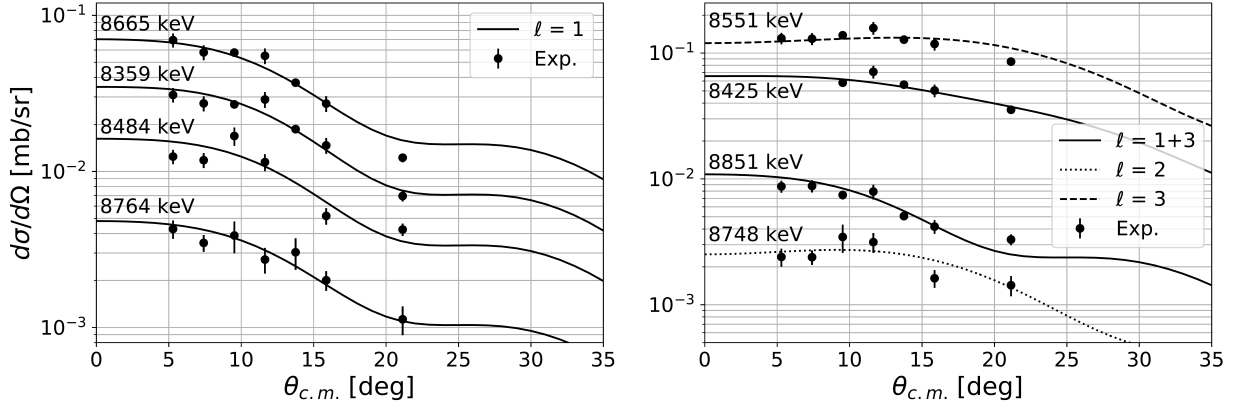


Figure 6.34: Differential cross-sections of proton-unbound ^{40}Ca states observed in the present experiment below the lowest directly measured (p, γ) resonance at 8935 keV. The left panel shows the $l = 1$ distributions, while the right panel shows all other distributions.

$E_x = 4491$ keV: The spin-parity of this state is assigned $J^\pi = 5^-$ by ENSDF [Che17], and its angular distribution is assigned $\ell = 3$ from the present analysis. This is in agreement with all of the $(^3\text{He}, d)$ measurements from Refs. [Ers66, Set67, For70, Cag71] and the only (d, n) measurement of Ref. [Fuc69] in the literature.

$E_x = 5614$ keV: The spin-parity of this state is assigned $J^\pi = 4^-$ by ENSDF. Its angular distribution is assigned $\ell = 3$ from the present analysis, in agreement with the literature [Ers66, Set67, For70, Cag71, Fuc69].

$E_x = 5903$ keV: The spin-parity of this state is assigned $J^\pi = 1^-$ by ENSDF. Its angular distribution is assigned $\ell = 1$ from the present analysis, despite it being obscured by the 4915 keV ^{14}N contaminant at $\theta_{\text{lab}} = 7^\circ$ and 9° . This is in agreement with Refs. [Ers66, Set67, Cag71] but in disagreement with (d, n) [Fuc69], where $\ell = 1 + 3$ is assigned. It was not resolved by Ref. [For70].

$E_x = 6025$ keV: The spin-parity of this state is assigned $J^\pi = 2^-$ by ENSDF, and its angular distribution is assigned $\ell = 1 + 3$ from the present analysis. This is in agreement with Refs. [Ers66, Set67] and (d, n) [Fuc69], but in disagreement with Ref. [Cag71], where $\ell = 3$ is assigned. It was not resolved by Ref. [For70].

$E_x = 6285$ keV: The spin-parity of this state is assigned $J^\pi = 3^-$ by ENSDF, and its angular distribution is assigned $\ell = 1$ from the present analysis. This is in agreement with Refs. [Ers66, For70, Cag71], but in disagreement with Ref. [Set67] and (d, n) [Fuc69] where $\ell = 1 + 3$ is assigned.

$E_x = 6582$ keV: The spin-parity of this state is assigned $J^\pi = 3^-$ by ENSDF, and its angular distribution is assigned $\ell = 1 + 3$ from the present analysis, with most of the contribution from $\ell = 1$. It is obscured by the ^{13}N ground state at $\theta_{\text{lab}} \leq 7^\circ$. This is in

agreement with (d, n) [Fuc69], but in disagreement with $(^3\text{He}, d)$ [Ers66, Set67, For70, Cag71], which assign it purely $\ell = 1$.

$E_x = 6750 \text{ keV}$: The spin-parity of this state is assigned $J^\pi = 2^-$ by ENSDF, and its angular distribution is assigned $\ell = 1 + 3$ from the present analysis. It is obscured by the ^{13}N ground state at $\theta_{\text{lab}} = 11^\circ, 13^\circ$, and 15° . This state is widely disputed in the literature. It is in agreement with (d, n) [Fuc69], but Refs. [Ers66, Set67, For70] assign it $\ell = 1$, and Ref. [Cag71] assigns it $\ell = 3$.

$E_x = 6950 \text{ keV}$: The spin-parity of this state is assigned $J^\pi = 1^-$ by ENSDF, and its angular distribution is assigned $\ell = 1$ from the present analysis. It is obscured by the 5834 keV ^{14}N state at $\theta_{\text{lab}} = 15^\circ$ and 20° . The energy residuals are about 13 keV lower than ENSDF at $\theta_{\text{lab}} = 5^\circ$ and 7° , which makes these measurements questionable, but the remaining angles show only a 2 keV residual from ENSDF. Even among the other angles alone, however, the angular distribution is clearly $\ell = 1$ and not $\ell = 3$. This assignment is in agreement with $(^3\text{He}, d)$ [Ers66, Set67, For70, Cag71], but in disagreement with (d, n) [Fuc69] where $\ell = 1 + 3$ is assigned.

$E_x = 7113 \text{ keV}$: In ENSDF, the 7113.1(10) keV and 7113.73(5) keV states have spin-parities of $J^\pi = 1^-$ and $J^\pi = 4^-$, respectively. The present energy calibration yielded a state at 7109.4(29) keV, within the uncertainty of the 1^- state, but not the 4^- state. An angular distribution of $\ell = 1 + 3$ is assigned in the present analysis, with the main contribution from $\ell = 1$. This is impossible for a 4^- state, where only $\ell = 3$ and/or $\ell = 5$ can be observed. Both of these measurements indicate that this corresponds with the 7113.1(10) keV 1^- state, with possibly a small contribution from the 7113.73(5) keV 4^- state at $\ell = 3$. This assignment is in agreement with (d, n) [Fuc69], but in disagreement with $(^3\text{He}, d)$ [Ers66, Set67, For70, Cag71], where purely $\ell = 1$ is assigned. This was the highest energy state measured by Ref. [For70], so this reference is omitted from the remaining discussion.

$E_x = 7532 \text{ keV}$: The spin-parity of this state is assigned $J^\pi = 2^-$ by ENSDF, and its angular distribution is assigned $\ell = 1$ from the present analysis. This is in agreement with Refs. [Ers66, Cag71], but in disagreement with Ref. [Set67] and (d, n) [Fuc69] where $\ell = 1 + 3$ is assigned.

$E_x = 7623 \text{ keV}$: The spin-parity of this state is assigned $J^\pi = (2^-, 3, 4^+)$ by ENSDF and was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71] or (d, n) [Fuc69]. This experiment has resolved this state for the first time through proton-transfer. Its angular distribution is assigned $\ell = 1 + 3$ in the present analysis. This implies a J^π assignment of $1^-, 2^-, 3^-$. Comparing with ENSDF, the $J^\pi = 3^+$ and 4^+ assignments can be ruled out with this measurement.

$E_x = 7658 \text{ keV}$: The spin-parity of this state is assigned $J^\pi = 4^-$ by ENSDF, and its

angular distribution is assigned $\ell = 3$ in the present analysis, in agreement with the literature [Ers66, Set67, Cag71, Fuc69].

$E_x = 7694$ keV: The spin-parity of this state is assigned $J^\pi = 3^-$ by ENSDF, and its angular distribution is assigned $\ell = 1+3$ in the present analysis. This state is disputed across the literature. The present assignment is in agreement with Ref. [Ers66], but in disagreement with Refs. [Set67, Cag71] and (d, n) [Fuc69], where $\ell = 3$ and $\ell = 1$ are assigned, respectively.

$E_x = 7973$ keV: The spin-parity of this state is assigned $J^\pi = (\text{LE } 3)^-$ by ENSDF based only on (d, n) [Fuc69]. It was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71]. This state was obscured in this experiment by the ^{17}F ground state at $\theta_{\text{lab}} = 11^\circ$ and 13° . The present analysis assigns $\ell = 1$, in agreement with (d, n) . This experiment alone suggests $J^\pi = 0^-, 1^-, 2^-, 3^-$.

$E_x = 8113$ keV: The spin-parity of this state is assigned $J^\pi = 1^-$ by ENSDF, and it was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71]. The present analysis assigns $\ell = 1$, in disagreement with (d, n) [Fuc69], where $\ell = 1+3$ is assigned. However, Ref. [Fuc69] observes a doublet at 8113 keV and 8135 keV, which is not observed in the present experiment. The presence of the 6859 keV ^{15}O contaminant complicates matters because it overlaps with where the 8135 keV state is expected, making this state impossible to resolve. The 7029 keV ^{14}N and 6793 keV ^{15}O contaminants obscure the $\theta_{\text{lab}} = 11^\circ$ measurement.

$E_x = 8188$ keV ($E_x = 8196$ keV): The spin-parity of the 8188 keV state is assigned $J^\pi = (3, 4, 5^-)$ in ENSDF, and it was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71]. No spectroscopic factor was reported from (d, n) [Fuc69] for a state observed at 8186 keV, but it was assigned a $\ell = 0$ transition nonetheless. This would imply $J^\pi = 1^+, 2^+$, which conflicts with ENSDF. The present energy calibration found a state at 8199.3(27) keV, or a residual from the 8188 keV state of 11.8(28) keV. There is in fact a state at 8196 keV reported in ENSDF, but it has only been observed in (p, t) [Set74, Set77] and (p, p') [Nol75], and no official J^π assignment has been made. However, the (p, t) measurement of Ref. [Set77] makes a weak assignment of $\ell = (2)$ for this state, which for (p, t) implies $J^\pi = (0, 1)^+$. In the present experiment, there is ambiguity in the angular momentum between $\ell = 0+2$ and $\ell = 1$, with the $\ell = 0+2$ transition being slightly more favorable. The spin-parities would be $J^\pi = 0^+, 1^+$ for the former transition and $J^\pi = 0^-, 1^-, 2^-, 3^-$ for the latter. It is possible that the (d, n) [Fuc69] assignment of $\ell = 0$ applies to the 8196 keV state, a 10 keV residual, which would be more in alignment with the present measurement and Refs. [Set74, Set77, Nol75]. This may even be likely, considering that Ref. [Fuc69] also determined the energy of a nearby state to be 8371 keV, which is interpreted by ENSDF as coinciding with the 8359 keV state, a 12 keV residual. Observation of this state in the present experiment was obscured at $\theta_{\text{lab}} = 11^\circ$ by the 6859 keV ^{15}O state and at 15° by the 7029 keV ^{14}N and 6793 keV ^{15}O states.

$E_x = 8271$ keV: The spin-parity of this state is assigned $J^\pi = (\text{LE } 3)^-$ by ENSDF, and it was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71]. The present analysis assigns $\ell = 1$, in agreement with (d, n) [Fuc69]. These experiments alone suggest $J^\pi = 0^-, 1^-, 2^-, 3^-$.

$E_x = 8359$ keV; $E_r^{\text{c.m.}} = 29$ keV: This is the lowest energy proton-unbound state observed in the present experiment. The spin-parity of this state is assigned $J^\pi = (0, 1, 2)^-$ in ENSDF, and it was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71]. The present analysis assigns $\ell = 1$, in agreement with (d, n) [Fuc69]. The recent ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ reaction rate evaluation from Ref. [Lon18b] did not use the (d, n) [Fuc69] C^2S measurement for a state at 8371 keV in their analysis because of the 12 keV energy discrepancy with ENSDF. This state was instead assigned a proton partial-width upper limit in the reaction rate calculation which the present work now replaces.

$E_x = 8425$ keV; $E_r^{\text{c.m.}} = 96$ keV: The spin-parity of this state is assigned $J^\pi = 2^-$ in ENSDF, and it has also since been confirmed via NRF [Gri22]. The present work assigns $\ell = 1 + 3$, in agreement with (d, n) [Fuc69], but in disagreement with $(^3\text{He}, d)$ [Ers66, Set67, Cag71], where $\ell = 3$ is assigned. Ref. [Lon18b] uses the $(^3\text{He}, d)$ C^2S measurement of Ref. [Cag71] in their reaction rate calculation. This state was obscured in the present experiment at $\theta_{\text{lab}} = 5^\circ$ and 7° by the ${}^{17}\text{F}$ first excited state.

$E_x = 8484$ keV; $E_r^{\text{c.m.}} = 154$ keV: This weakly-populated state has not been observed in $(^3\text{He}, d)$ [Ers66, Set67, Cag71] or (d, n) [Fuc69]. It has therefore been resolved for the first time in ${}^{39}\text{K} + p$ from this experiment because the lowest energy resonance resolved by (p, γ) [Kik90, Che81, Lee66] is for $E_r^{\text{c.m.}} = 606$ keV. Its spin-parity is assigned $J^\pi = (1^-, 2^-, 3^-)$ in ENSDF. The present work assigns $\ell = 1$ in support of the ENSDF J^π assignment. This measurement replaces the proton partial-width upper limit of Ref. [Lon18b] in the reaction rate calculation. This state was obscured in the present experiment at $\theta_{\text{lab}} = 13^\circ$ by the ${}^{17}\text{F}$ first excited state.

$E_x = 8551$ keV; $E_r^{\text{c.m.}} = 221$ keV: The spin-parity of this state is assigned $J^\pi = 5^-$ in ENSDF. The present work assigns $\ell = 3$, in agreement with $(^3\text{He}, d)$ [Ers66, Set67, Cag71] and (d, n) [Fuc69].

$E_x = 8665$ keV; $E_r^{\text{c.m.}} = 335$ keV: The spin-parity of this state is assigned $J^\pi = 1^-$ in ENSDF, and it was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71]. The present work assigns $\ell = 1$ in agreement with (d, n) [Fuc69].

$E_x = 8748$ keV; $E_r^{\text{c.m.}} = 415$ keV: This weakly-populated state was observed in (d, n) [Fuc69] as part of an unresolved doublet with the 8764 keV state, but no ℓ assignments were made. Neither states were resolved in $(^3\text{He}, d)$ [Ers66, Set67, Cag71] or (p, γ) [Kik90, Che81, Lee66], but they have been resolved in the present experiment, making this the first time they have been resolved in ${}^{39}\text{K} + p$ altogether. The spin-parity of the 8748 keV state

is assigned $J^\pi = 2^+$ in ENSDF and has since been confirmed by NRF [Gri22]. The present work assigns $\ell = 2$ in support of the ENSDF J^π assignment. This measurement replaces the proton partial-width upper limit of Ref. [Lon18b] in the reaction rate calculation. The statistics were not sufficient to obtain fits for $\theta_{\text{lab}} = 13^\circ$ for this state.

$E_x = 8764 \text{ keV}; E_r^{\text{c.m.}} = 439 \text{ keV}$: This state is part of a doublet with 8748 keV and was not resolved in $(^3\text{He}, d)$ [Ers66, Set67, Cag71], (d, n) [Fuc69], or (p, γ) [Kik90, Che81, Lee66]. Its spin-parity is assigned $J^\pi = 3^-$ in ENSDF, and the present work assigns $\ell = 1$, in support of this assignment. This experiment is therefore the first to resolve this state for $^{39}\text{K} + p$, and its measurement replaces the proton partial-width upper limit of Ref. [Lon18b] in the reaction rate calculation.

$E_x = 8851 \text{ keV}; E_r^{\text{c.m.}} = 521 \text{ keV}$: The spin-parity of this state is assigned $J^\pi = 6^-, 7^-, 8^-$ in ENSDF based on (p, p') [Gru72]. It was observed in (d, n) [Fuc69] with no ℓ assignment determined, and it was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71] or (p, γ) [Kik90, Che81, Lee66]. The present experiment assigns $\ell = 1 + 3$, making its resolution a first for $^{39}\text{K} + p$. The ℓ assignment suggests $J^\pi = (1, 2, 3)^-$, in stark disagreement with ENSDF. The upper limit from Ref. [Lon18b] for this state uses the ENSDF J^π assignment in the reaction rate calculation. For this reason, the $(2J + 1)\Gamma_p$ value from this work is about 2 orders of magnitude larger than the upper limit, as shown in Table ??.

$E_x = 8935 \text{ keV}; E_r^{\text{c.m.}} = 608 \text{ keV}$: In ENSDF, the 8934.81(7) keV and 8938.4(9) keV states have spin-parities of $J^\pi = 2^+$ and $J^\pi = 0^+$, respectively. The 8934.81(7) keV state is the lowest energy proton-unbound state measured by the direct (p, g) [Kik90] experiment, and they do not observe the 8938.4(9) keV state. The present energy calibration yielded a state at 8936.3(27) keV, within the uncertainty of both states. Neither state is resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71]. The present analysis assigns $\ell = 1$, in agreement with the 8931 keV assignment from (d, n) [Fuc69], and this joins Ref. [Fuc69] in disagreement with ENSDF. These measurements imply $J^\pi = 0^-, 1^-, 2^-, 3^-$. It is worth noting that a $\ell = 0 + 2$ assignment was considered in the present analysis, which would imply $J^\pi = 1^+, 2^+$, in agreement with (p, g) [Kik90]. However, the $\ell = 1$ transition is significantly more likely.

$E_x = 8995 \text{ keV}; E_r^{\text{c.m.}} = 664 \text{ keV}$: The spin-parity of this state is assigned $J^\pi = (1^-, 2^+)$ in ENSDF, and it was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71] or (d, n) [Fuc69]. The angular momentum assignment in the present analysis is ambiguous between $\ell = 0 + 2$ and $\ell = 1$, which does not narrow down the J^π assignment. Higher angle measurements of $\theta_{\text{lab}} \geq 30^\circ$ could resolve the angular momentum. The state was obscured at $\theta_{\text{lab}} \geq 5^\circ$ by the ^{13}N first excited state, and the statistics were insufficient at $\theta_{\text{lab}} \geq 20^\circ$.

$E_x = 9092 \text{ keV}; E_r^{\text{c.m.}} = 764 \text{ keV}$: The spin-parity of this state is assigned $J^\pi = 3^-$ in ENSDF, and it was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71] or (d, n) [Fuc69]. The

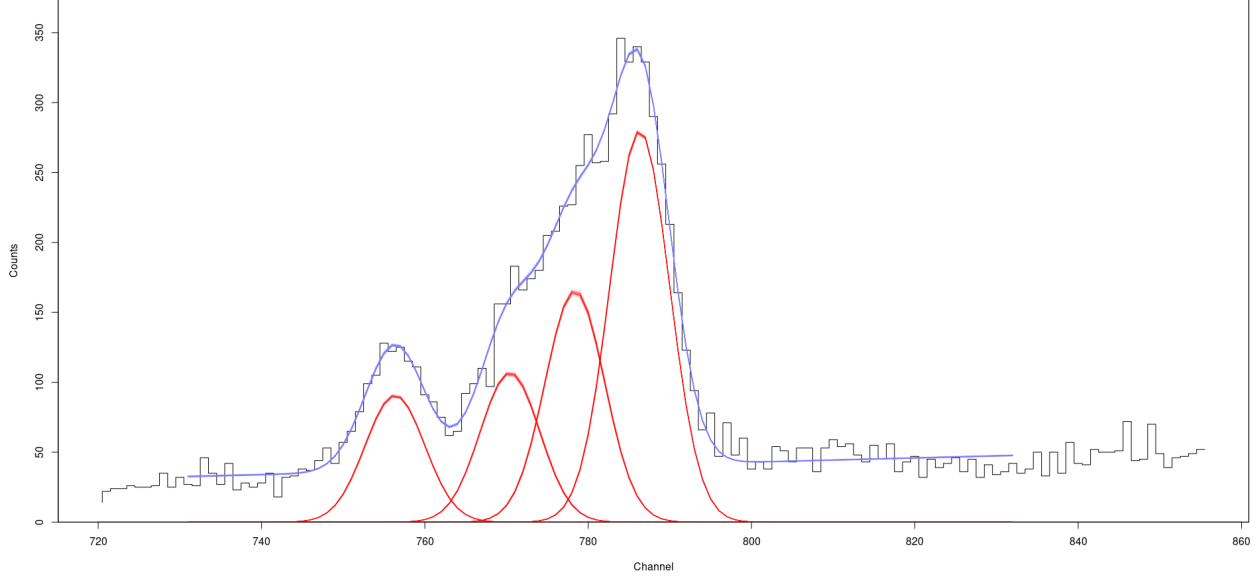


Figure 6.35: A Bayesian Monte Carlo fit for the 9405–9432 keV multiplet at $\theta_{\text{lab}} = 5^\circ$, including the clearly distinguishable 9454 keV state. The multiplet is fit with 3 peaks in this experiment, with calibrated energies of 9403.7(31) keV, 9416.7(32) keV, and 9431.7(32) keV. This is better resolution than (d, n) [Fuc69] was able to achieve, where only 2 peaks in the multiplet were fit at energies of 9408 keV and 9431 keV.

present analysis assigns $\ell = 1$, despite the state being obscured at $\theta_{\text{lab}} = 5\text{--}11^\circ$ by the ^{13}N first excited state. This is in agreement with the J^π assignment.

$E_x = 9136 \text{ keV}; E_r^{\text{c.m.}} = 810 \text{ keV}$: The spin-parity of this state is assigned $J^\pi = 2^-, 3^-$ in ENSDF, and it was not resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71]. The present analysis assigns $\ell = 1$, in agreement with (d, n) [Fuc69], despite the state being obscured at $\theta_{\text{lab}} = 5\text{--}11^\circ$ by the ^{13}N first excited state. This is in agreement with the J^π assignment.

$E_x = 9227 \text{ keV}; E_r^{\text{c.m.}} = 900 \text{ keV}$: In ENSDF, the 9226.69(5) keV and 9227.43(7) keV states have spin-parities of $J^\pi = (1^-, 2, 3^-)$ and $J^\pi = (1, 2^+)$, respectively. The present energy calibration yielded a state at 9227.8(34) keV, within the uncertainty of both states. Neither state is resolved by $(^3\text{He}, d)$ [Ers66, Set67, Cag71]. The angular momentum assignment in the present analysis is ambiguous between $\ell = 0 + 2$ and $\ell = 1$, which does not narrow down the J^π assignment in either case. The (d, n) measurement of Ref. [Fuc69] unambiguously assigns $\ell = 1$. The state was obscured at $\theta_{\text{lab}} = 13^\circ$ and 15° by the ^{13}N first excited state.

$E_x = 9405 - 9432 \text{ keV Multiplet}$: This multiplet consists of the states 9405 keV ($J^\pi = 2^-$), 9406 keV ($J^\pi = 0^+$), 9412 keV (no J^π assignment), 9419 keV ($J^\pi = 3^-$), 9429 keV ($J^\pi = (3, 4)^-$), and 9432 keV ($J^\pi = 1^-$) in ENSDF. Even though these individual states are unresolvable at the energy resolution attainable with the focal plane detector, it was possible

Table 6.5: The states resolved in the present experiment for the first time in the indicated regimes. Parentheses denote ambiguous ℓ assignments.

New for ${}^{39}\text{K}({}^3\text{He}, d){}^{40}\text{Ca}$	New for ${}^{39}\text{K}$ Proton-Transfer	New for ${}^{39}\text{K} + p$
7623 keV	7623 keV	
7973 keV		
8113 keV		
(8188 keV)	(8188 keV)	
8271 keV		
8359 keV		
8484 keV	8484 keV	8484 keV
8665 keV		
8748 keV	8748 keV	8748 keV
8764 keV	8764 keV	8764 keV
8851 keV	8851 keV	8851 keV
8935 keV		
(8995 keV)	(8995 keV)	
9092 keV	9092 keV	
9136 keV		
(9227 keV)		

to consistently fit 3 peaks in this region with the Bayesian Monte Carlo procedure of Section 6.6, as shown in Fig. 6.35. From the present energy calibration, the peaks result in energies of 9403.7(31) keV, 9416.7(32) keV, and 9431.7(32) keV. These can likely be assigned as 3 separate doublets from the 6 ENSDF states, 9405+9406 keV, 9412+9419 keV, and 9429+9432 keV. The (d, n) measurement of Ref. [Fuc69] obtained fits for only 2 peaks in this region, at 9408 keV and 9431 keV. Attempting fits with 2 peaks in this region resulted in much worse agreement with the present data. Angular distributions were not obtained for these states in the present analysis, or for any higher energy states, since the focal plane detector efficiency at the high energy fringe resulted in dramatic yield loss over increasing angle.

To summarize the results of this section, the present experiment has unambiguously resolved 13 new states in ${}^{39}\text{K}({}^3\text{He}, d){}^{40}\text{Ca}$, 6 new states in ${}^{39}\text{K}$ proton-transfer, and 4 new states in ${}^{39}\text{K} + p$ altogether. Table 6.5 lists the states that are newly resolved in these regimes. States in parentheses were given an ambiguous ℓ assignment in the present experiment.

6.8.4 Spectroscopic Factors

Spectroscopic factors $C^2 S_{39\text{K}+p}$ for each state were extracted from Eqns. 6.44 and 6.47, using the angular momentum assignments and DWBA calculations from the previous section.

These are summarized and compared with the $(^3\text{He}, d)$ [Ers66, Set67, For70, Cag71] and (d, n) [Fuc69] measurements in the literature in Table 6.6. Doublets unresolved in the present analysis are given brackets around their ENSDF [Che17] energies. Ambiguous ℓ assignments are also indicated with brackets. In mixed- ℓ or disputed- ℓ cases, each ℓ -component is shown in parentheses on separate rows, where the lowest ℓ is always the first row.

The J^π values shown in the table were used in the present DWBA models to calculate C^2S . They were also used to convert the other measurements from their reported $(2J+1)C^2S$ values to C^2S for consistency, since their chosen J^π sometimes differed. The J^π values that differ from ENDSF are indicated. This difference is usually an arbitrary choice of the lowest appropriate ENDSF J^π value that is consistent with the present ℓ -assignment. However, the choice sometimes genuinely conflicts with the ENDSF assignment, as is the case for the 8188 keV 1^+ ambiguity, 8851 keV, and the 8935+8938 keV doublet.

The statistical uncertainty of C^2S for the present experiment is also given in the table. An additional 30% systematic uncertainty in C^2S is assumed, which is typical of $(^3\text{He}, d)$ reactions in the literature [End77]. The main contribution to this systematic uncertainty is the choice of OMP parameters, especially for the ${}^3\text{He} + {}^{39}\text{K}$ incoming channel. This is due to the fact that the ${}^{39}\text{K}(^3\text{He}, d){}^{40}\text{Ca}$ differential cross section was normalized to the ${}^3\text{He} + {}^{39}\text{K}$ OMP prediction. The OMP parameters for the $d + {}^{40}\text{Ca}$ outgoing channel also contribute to the systematic uncertainty, albeit to a lesser degree. For example, the ${}^{23}\text{Na}(^3\text{He}, d){}^{24}\text{Mg}$ measurement of Ref. [Hal04] varied the choice of OMP parameters within a reasonable range. They found a 26% uncertainty associated with the ${}^3\text{He}$ parameters alone and a 11% uncertainty associated with the d parameters alone. Added in quadrature, this is a 28% uncertainty from the choice of OMP parameters, which agrees with the historical scatter between $(^3\text{He}, d)$ measurements of C^2S [End77].

6.8.5 Proton Partial Widths and Resonance Energies

The ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ reaction rate is dominated by narrow resonances. As described in Section 2.2.1, narrow resonances contribute individually to thermonuclear reaction rates. The resonance strength $\omega\gamma$ and resonance energy E_r of each resonance are the key experimental quantities that go into the reaction rate calculation of Eqn. ???. Typically, proton-transfer experiments report proton partial widths from Eqn. 2.12, rather than resonance strengths from Eqn. ???. The only difference is the knowledge of the γ -ray partial width and any other partial widths contributing to the total width, which arise from other measurements. The spectroscopic factors of the proton-unbound ${}^{40}\text{Ca}$ states from the previous section go directly into the proton partial width calculation, along with the penetration factor $P(E)$ and the dimensionless single-particle reduced width. The latter two quantities are calculated theoretically and are detailed in Section 2.2.1.

Center-of-mass resonance energies $E_r^{\text{c.m.}}$ are simply calculated from the measured excitation energies in Section 6.8.1 by adding the proton separation energy S_p of ${}^{40}\text{Ca}$, or equivalently, the Q -value of the ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ reaction. The present work uses $S_p = 8328.18(2)$ keV from Ref. [Wan21].

The results of the proton partial widths and the resonance energies are summarized in Table ??.

6.9 New ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ Reaction Rate

The Monte Carlo reaction rate code, **RatesMC** [Lon10, Lon23], was used to calculate a new ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ reaction rate probability density from the resonance energies and proton partial-widths reported in Sections 6.8.1 and 6.8.5, respectively. Among the states observed in the ${}^{39}\text{K}({}^3\text{He}, d){}^{40}\text{Ca}$ experiment, only those that do not already have a directly measured ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ resonance strength from Refs. [Kik90, Che81, Lee66] were modified from the most recent reaction rate evaluation of Ref. [Lon18b]. That is, the only resonance strengths and resonance energies that were updated from those in Ref. [Lon18b] are below the $E_x = 8935$ keV ($E_r^{\text{c.m.}} = 606$ keV) resonance. Otherwise, the expectation values and variances of the ${}^{39}\text{K}(p, \gamma){}^{40}\text{Ca}$ experiments [Kik90, Che81, Lee66] are adopted from Ref. [Lon18b].

The new reaction rate is compared with that of Ref. [Lon18b] in Fig. 6.36. The solid line and blue band represent the median, recommended rate and the 1σ confidence interval of this work, respectively. The dotted line and gray band represent that of Ref. [Lon18b], except with resonance energies calculated using $S_p = 8328.18(2)$ keV from Ref. [Wan21] for consistency, but with marginal effect. Both reaction rates are normalized to the median, recommended

Table 6.6: Spectroscopic factors compared with Refs. [Ers66, Set67, For70, Cag71, Fuc69].

E_x^a	J^π ^b	ℓ ^b	$C^2 S_{^{39}\text{K}+p}$ ^b	[Ers66]	[Set67]	[For70]	[Cag71] ^c	[Fuc69]
4491	5 ⁻	3	0.454(25)	0.609	0.382	0.364	0.418	0.340
5614	4 ⁻	3	0.461(25)	0.656	0.389	0.378	0.389	0.470
5903	1 ⁻	1	0.019(1)	0.037	0.025		0.015	0.027 (1) <0.067 (3)
6025	2 ⁻	1 + 3	0.018(1) (1) 0.072(6) (3)	0.027 (1) 0.180 (3)	0.020 (1) 0.100 (3)		0.105	0.030 (1) 0.096 (3)
6285	3 ⁻	1	0.167(9)	0.357	0.200 (1)	0.307	0.150	0.246 (1) <0.171 (3)
6582	3 ⁻	1 + 3	0.047(3) (1) 0.069(9) (3)	0.114	0.086	0.107	0.060	0.080 (1) <0.114 (3)
6750	2 ⁻	1 + 3	0.024(2) (1) 0.135(11) (3)	0.100	0.096	0.094		0.027 (1) 0.264 (3)
6950	1 ⁻	1	0.134(8)	0.275	0.200	0.215	0.080	0.227 (1) <0.267 (3)
7113	1 ⁻	1 + 3	0.150(8) (1) 0.132(23) (3)	0.317	0.257	0.233	0.257	0.240 (1) <0.133 (3)
7532	2 ⁻	1	0.227(12)	0.436	0.150 (1) 0.400 (3)		0.095	0.392 (1) <0.080 (3)
7623	2 ^{-d}	1 + 3	0.005(1) (1) 0.037(3) (3)					
7658	4 ⁻	3	0.375(20)	0.483	0.306		0.306	0.345
7694	3 ⁻	1 + 3	0.027(4) (1) 0.279(20) (3)	<0.057 (1) 0.584 (3)	0.414		0.393	0.029
7973	3 ⁻	1	0.017(1)					0.023
8113	1 ⁻	1	0.026(2)					0.033 (1) 0.160 (3)
8188	3 ^{-d} 1 ^{+e}	$\begin{cases} 1 \\ 0 + 2 \end{cases}$	0.007(1) 0.017(2) (0) 0.006(1) (2)					
8271	3 ⁻	1	0.031(2)					0.046
8359	0 ^{-d}	1	0.191(11)					0.320
8425	2 ⁻	1 + 3	0.037(6) (1) 0.246(21) (3)	0.320	0.500		0.280	0.008 (1) 0.288 (3)
8484	1 ^{-d}	1	0.031(2)					
8551	5 ⁻	3	0.416(23)	0.591	0.455		0.418	0.356
8665	1 ⁻	1	0.148(8)					0.187
8748	2 ⁺	2	0.004(1)					
8764	3 ⁻	1	0.005(1)					
8851	2 ^{-e}	1 + 3	0.011(1) (1) 0.011(1) (3)					
$\{8935$	1^{-e}	1	0.057(3)					0.147
$8938\}$								

Table 6.6: (continued)

E_x^a	J^π ^b	ℓ^b	$C^2 S_{^{39}\text{K}+p}^b$	[Ers66]	[Set67]	[For70]	[Cag71] ^c	[Fuc69]
8995	1 ⁻	1	0.020(1)					
	2 ⁺	$\begin{cases} 0 \\ + 2 \end{cases}$	0.006(1) (0) 0.006(1) (2)					
9092	3 ⁻	1	0.007(1)					
9136	2 ^{-d}	1	0.152(9)				0.136	
{9226.69	1 ^{-d}	1	0.143(9)				0.213	
{9227.43	1 ^{+d}	$\begin{cases} 0 \\ + 2 \end{cases}$	0.048(6) (0) 0.071(6) (2)					

^aFrom the ENSDF evaluation of Ref. [Che17]. Energy is given in keV.

^bPresent experiment. Statistical uncertainties are given. A 30% systematic uncertainty is assumed.

^cUsing the modified DWBA results of Ref. [Cag71].

^dThe lowest appropriate J^π from the ENSDF assignment possibilities.

^eDiffers from the ENSDF J^π assignment altogether.

rate of Ref. [Lon18b].

As mentioned previously, the large uncertainty in Ref. [Lon18b], between about 50 MK and 200 MK, corresponds to most of the relevant temperatures that reproduce the Mg–K anticorrelation in the globular cluster NGC 2419 [Ili16]. Fig. 6.36 illustrates that the new reaction rate increases significantly in this range, particularly at 70 MK where the median rate is a factor of 13 larger than the rate of Ref. [Lon18b]. The 1σ width is also significantly reduced in this region, from a factor of 84 at 80 MK to just a factor of 2, a reduction of a factor of 42.

Our new determination of $(2J + 1)\Gamma_p$ for the 154 keV resonance is primarily responsible for the increase in the rate and decrease in the uncertainty between about 55 MK and 110 MK. Similar effects occur between about 20 MK and 55 MK, primarily from our $l = 1 + 3$ assignment of the 96 keV resonance, which has replaced the $l = 3$ assignment by the other (${}^3\text{He}, d$) measurements in this calculation. The (d, n) measurement of Ref. [Fuc69] is in agreement with the $l = 1 + 3$ assignment for this resonance. Note that using the previous $l = 3$ assignment has a negligible effect on the results mentioned above at 70 and 80 MK. Our $(2J + 1)\Gamma_p$ determination for the 29 keV resonance is responsible for the rate increase below 20 MK. The smaller effects above 110 MK are from a combination of the 335 keV, 415 keV, 439 keV, and 521 keV resonances, the latter three of which have replaced $(2J + 1)\Gamma_p$ upper limits in Ref. [Lon18b].

6.9.1 Resonance Contributions

6.10 Potassium Abundance in Reaction Network Calculations

6.11 Conclusions

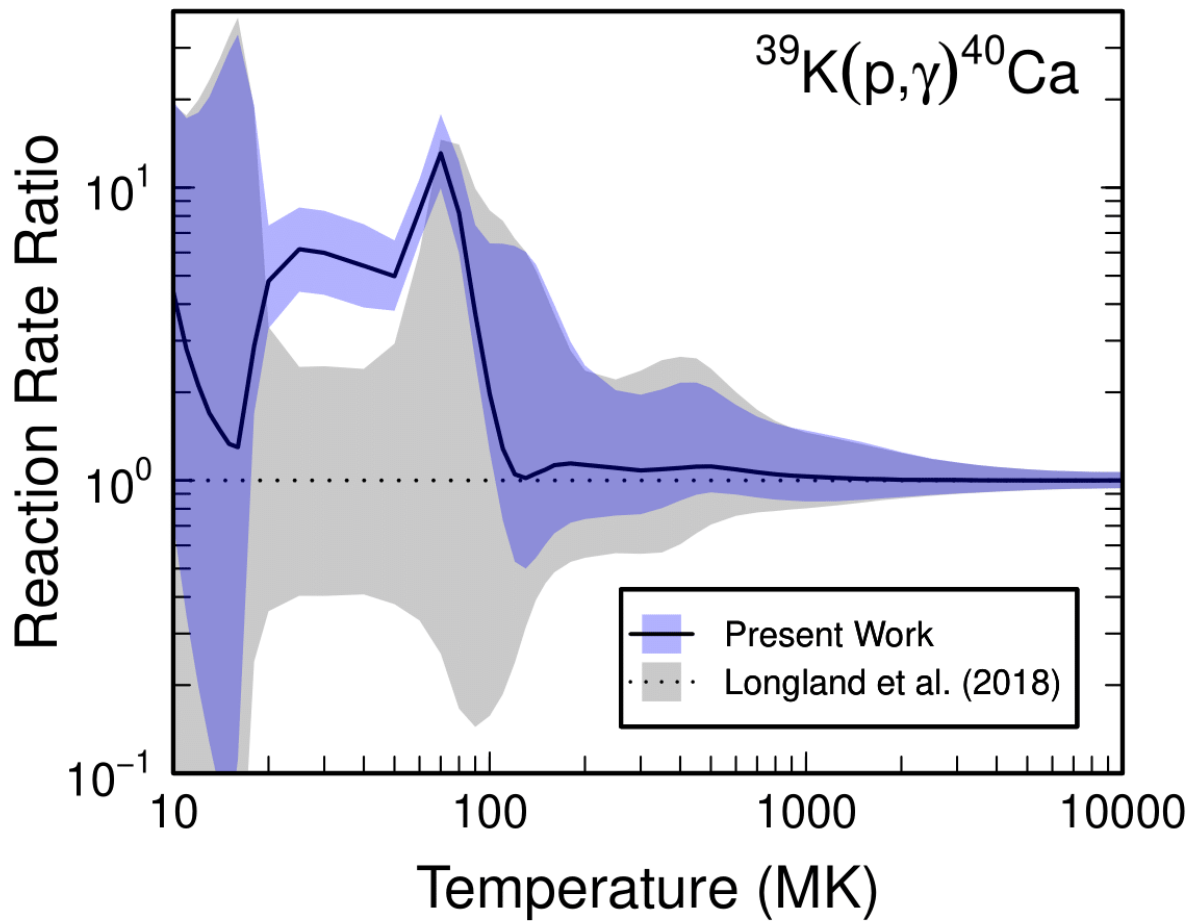


Figure 6.36: Comparison between the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate using the proton partial-widths and resonance energies of the present experiment (solid line, blue band) and the most recent evaluation of Ref. [Lon18b] (dotted line, gray band). The reaction rate ratio is taken with respect to the median, recommended rate of Ref. [Lon18b] for both calculations. The 1σ uncertainty bands are shown.

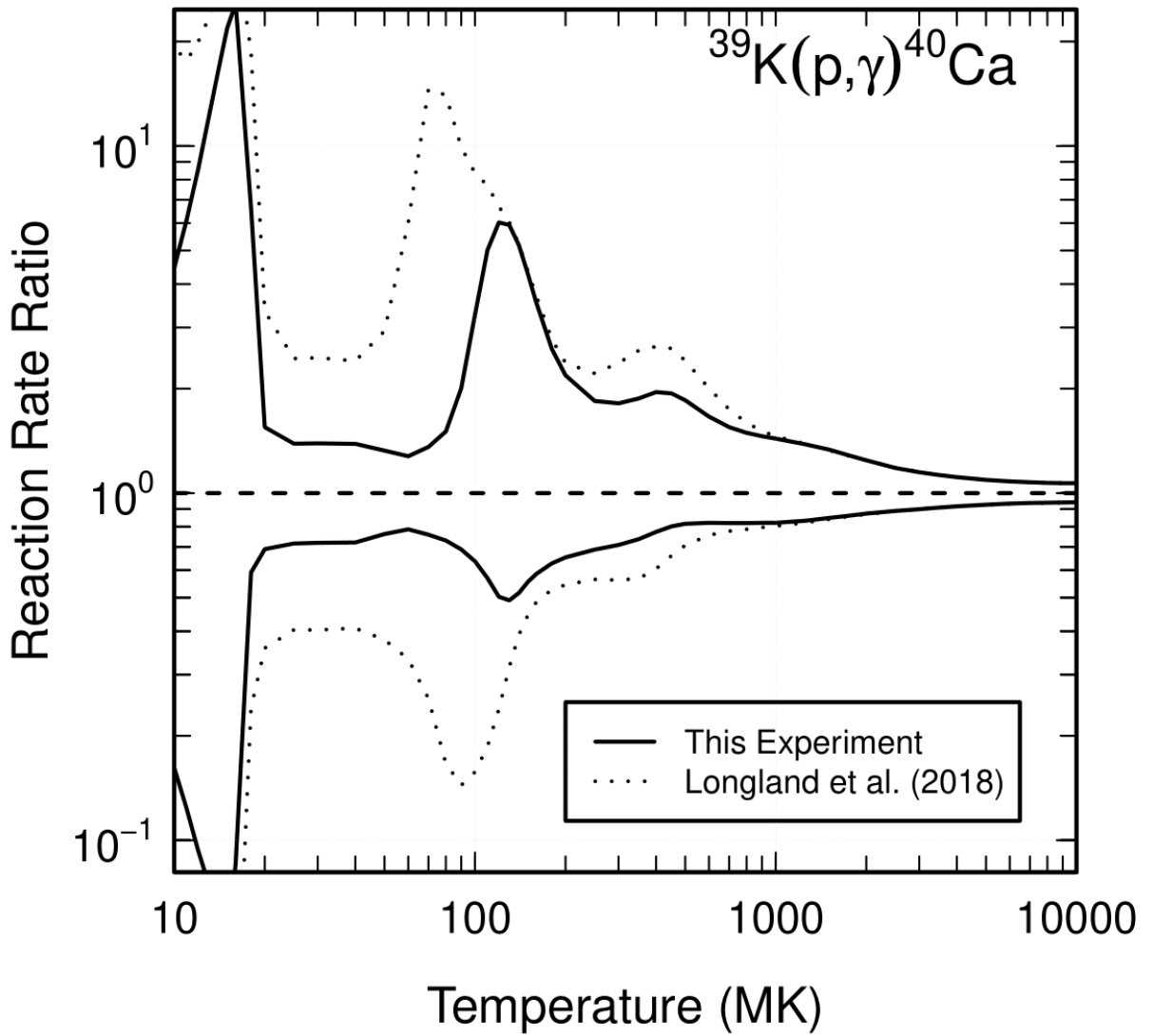


Figure 6.37: Comparison between the $^{39}\text{K}(p, \gamma)^{40}\text{Ca}$ reaction rate uncertainty using the proton partial-widths and resonance energies of the present experiment (solid band) and the most recent evaluation of Ref. [Lon18b] (dotted band). Each reaction rate ratio is taken with respect to their own median, recommended rate. The 1σ uncertainty bands are shown.

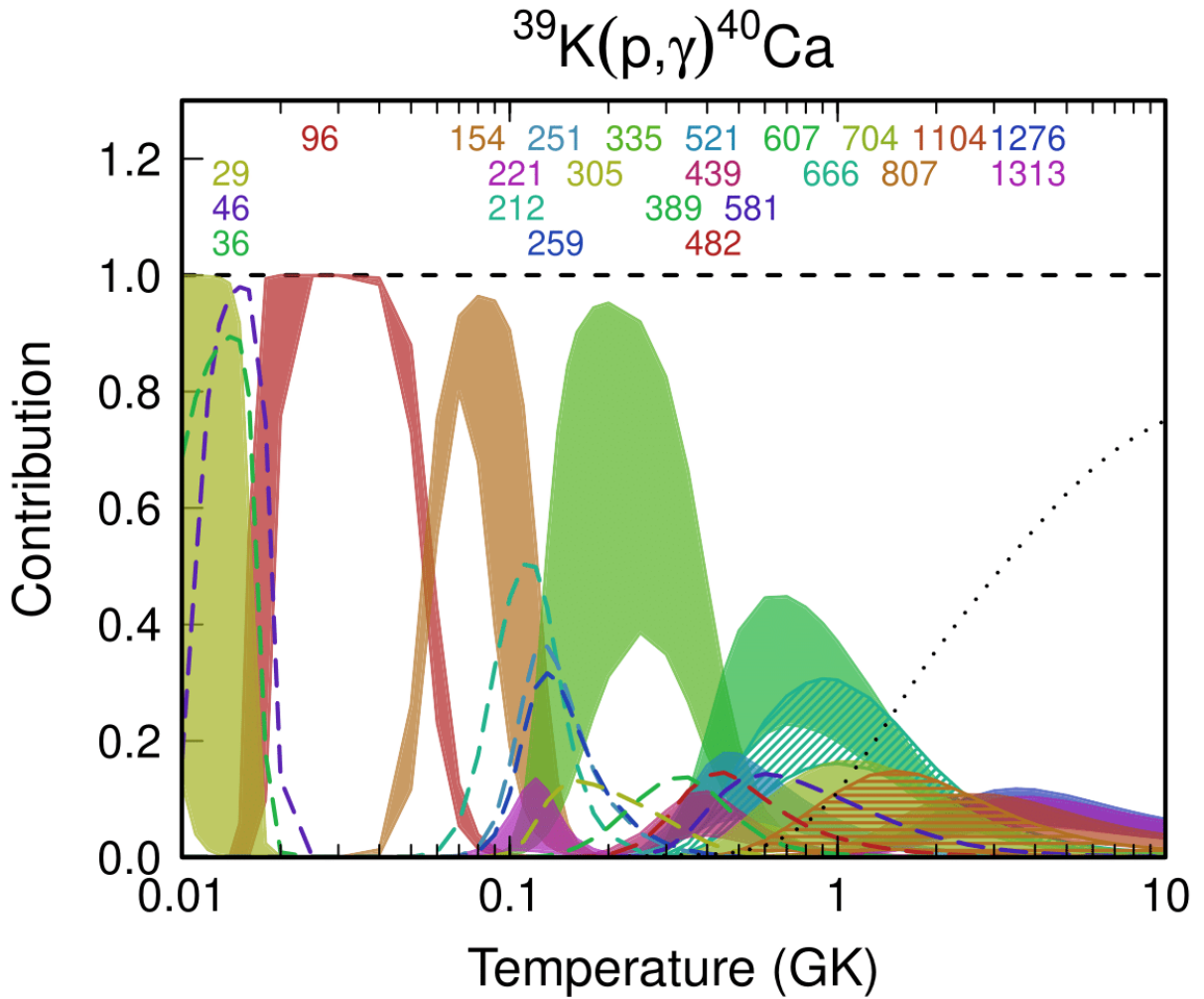


Figure 6.38: Individual resonance contributions to the $^{39}\text{K}(p,\gamma)^{40}\text{Ca}$ reaction rate, where a value of 1.0 implies that the given resonance contributes 100% to the reaction rate at that temperature. The labels correspond to center-of-mass resonance energies in keV. Resonances with shading or hatched lines have been measured and are shown with their 1σ uncertainty bands. Resonances with a single, dashed line are upper limit calculations and show their 84% 1σ value, for clarity. The resonances displayed are those that individually account for at least 10% of the total reaction rate at their corresponding temperatures. The remaining summed resonance contributions are represented by the dotted line.

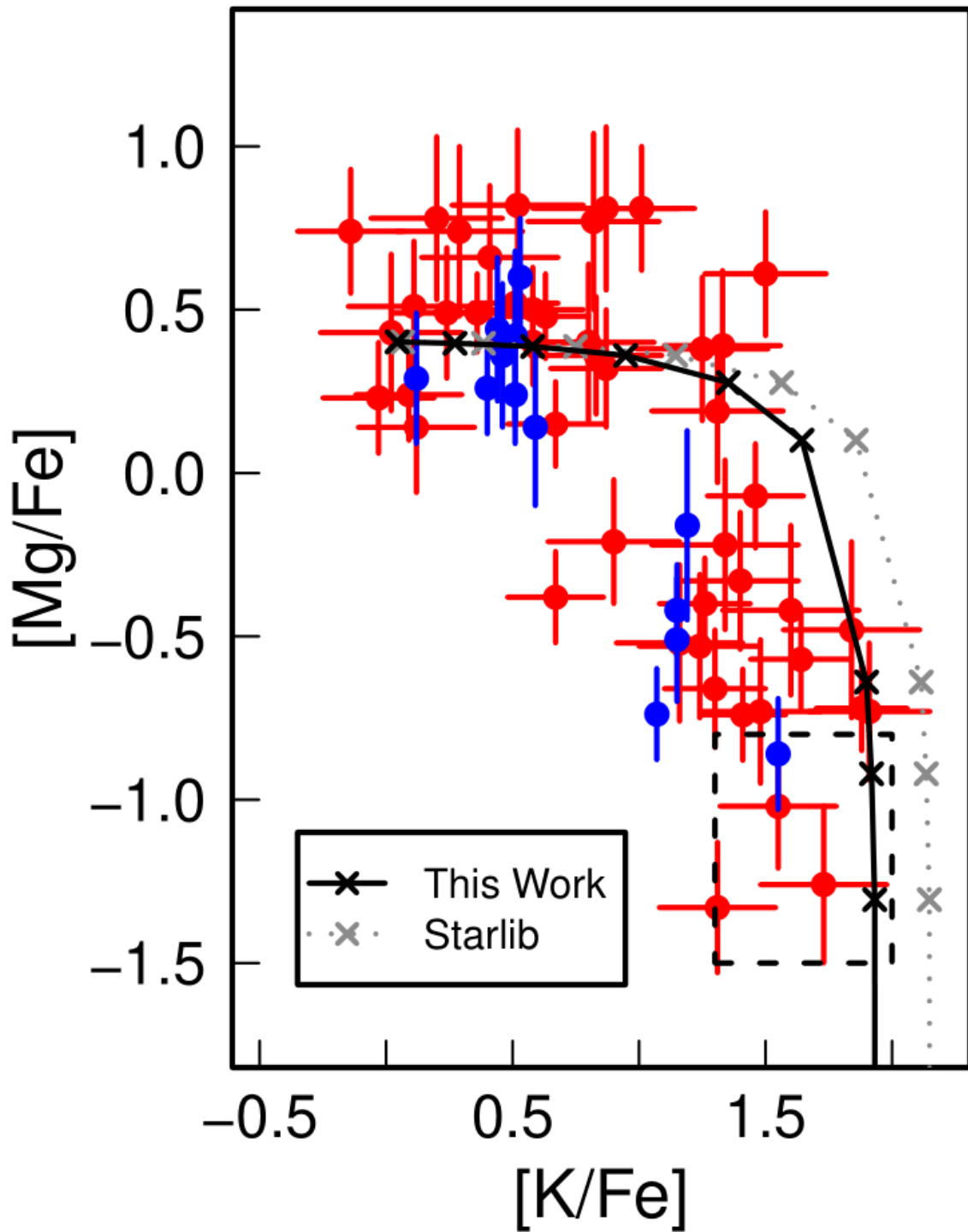


Figure 6.39: The maximum change in potassium abundance from Starlib, occurring at $T = 100$ MK, $\rho = 4 \times 10^7$ g/cm³, and $X_{3\text{He},f} = 20\%$. This corresponds to a decrease in [K/Fe] by a factor of 1.7 from Starlib. This configuration is now an acceptable configuration reproducing the Mg–K anticorrelation in NGC 2419.

CHAPTER

7

SUMMARY AND CONCLUSIONS

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APPENDIX

APPENDIX

A

^{40}Ca ENERGY CALIBRATIONS