Cross section measurement of the astrophysically important $^{17}\text{O}(p,\gamma)^{18}\text{F}$ reaction in a wide energy range

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**Background:** The $^{17}\text{O}(p,\gamma)^{18}\text{F}$ reaction plays an important role in hydrogen burning processes in several different stages of stellar evolution. The rate of this reaction must therefore be known with high accuracy at the relevant temperatures in order to provide the necessary input for astrophysical models.

**Purpose:** The cross section of $^{17}\text{O}(p,\gamma)^{18}\text{F}$ is characterized by a complicated resonance structure at low energies which needs to be reproduced by theoretical models if a reliable extrapolation to astrophysical energies is required. Experimental data, however, are scarce in a wide energy range, which increases the uncertainty of the extrapolations. The purpose of the present work is therefore to provide consistent and precise cross section values in a wide energy range for the $^{17}\text{O}(p,\gamma)^{18}\text{F}$ reaction.

**Method:** The cross section is measured using the activation method. This method provides directly the total cross section which can be compared with model calculations. With this technique some typical systematic uncertainties encountered in in-beam $\gamma$-spectroscopy experiments can be avoided.

**Results:** The cross section was measured between 500 keV and 1.8 MeV proton energies with a total uncertainty of typically 10%. The results are compared with earlier measurements and it is found that the gross features of the $^{17}\text{O}(p,\gamma)^{18}\text{F}$ excitation function are relatively well reproduced by the present data. Deviation of roughly a factor of 1.5 is found in the case of the total cross section when compared with the only high energy dataset. At the lowest measured energy our result is in agreement with two recent datasets within one standard deviation and deviates by roughly two standard deviations from a third one. An $R$-matrix analysis of the present and previous data strengthens the reliability of the extrapolated zero energy astrophysical 5 factor.

**Conclusions:** Using an independent experimental technique, the literature cross section data of $^{17}\text{O}(p,\gamma)^{18}\text{F}$ is confirmed in the energy region of the resonances while a lower direct capture cross section is recommended at higher energies. The present dataset provides a constraint for the theoretical cross sections.

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I. INTRODUCTION

Hydrogen burning, the conversion of four protons into an $\alpha$ particle in the interior of stars, is the most important energy source in the universe and it is also responsible for the existence of several chemical elements. Besides the $pp$ chains powering, e.g., our sun, catalytic reactions cycles, like the various CNO cycles, play the major role in hydrogen burning [1,2]. Depending on the temperature and chemical composition of the stellar plasma, different CNO cycles can take place involving various isotopes of carbon, nitrogen, oxygen, and fluorine.

The $^{17}\text{O}(p,\gamma)^{18}\text{F}$ reaction, which competes with the $\alpha$ decay in $^{17}\text{O}(p,\alpha)^{14}\text{N}$ [3], is the starting point of the third CNO cycle. This cycle is activated in various stellar conditions such as red giant and asymptotic giant stars and classical novae. The abundances of fluorine and the heavy oxygen isotopes are strongly related to the operation of this cycle and therefore the rates of the participating reactions must be known.

Below 1.5 MeV the $^{17}\text{O}(p,\gamma)^{18}\text{F}$ reaction is characterized by many broad and narrow resonances. Therefore, the temperature dependence of the $^{17}\text{O}(p,\gamma)^{18}\text{F}$ thermonuclear reaction rate shows a complicated picture. The rate depends on the direct capture component as well as on the narrow low energy resonances and the tails of the higher energy broad resonances (see, e.g., Fig. 10 in Ref. [4] for the contribution of the different components to the reaction rate). An $R$-matrix fit to the experimental data is therefore necessary to provide reaction rates at various temperatures for stellar models.

The first cross section measurement of $^{17}\text{O}(p,\gamma)^{18}\text{F}$ was carried out several decades ago by Rolfs in a wide energy range between 300 keV and 1.9 MeV [5]. After the turn of the century, several experimental studies were carried out, mostly concentrating on the low energy region below about 500 keV [4,6–14]. (The only exception was the work of Kontos et al. [11] which extended up to 1.6 MeV.) The comparison of the new precise data with the results of [5] revealed some discrepancies both in the absolute scale and the energy dependence of the cross section at the lowest energies studied by [5].

In most of the previous experiments, the cross section of $^{17}\text{O}(p,\gamma)^{18}\text{F}$ was measured with in-beam $\gamma$ spectroscopy: the prompt $\gamma$ radiation from the formed $^{18}\text{F}$ nucleus was detected. The complicated level scheme of $^{18}\text{F}$ (see, e.g., Fig. 1 in Ref. [14]) implies that the detection of many primary and secondary transitions is necessary for the cross section determination. This represents a source of uncertainty in the experiments. In order to provide the astrophysically relevant total cross section, all the transitions must be measured and
care must be taken to measure even the weakest $\gamma$ lines. The angular distributions of all the $\gamma$ emissions must also be known. Moreover, in order to measure low cross sections, close target-detector geometries are typically used, leading to strong true coincidence summing effects.

All these experimental difficulties can be avoided by the application of the activation method, which was used by only two experiments before at energies below 400 keV $[4,8,9,13]$. The reaction product of $^{17}\text{O}(p,\gamma)^{18}\text{F}$ is radioactive, decaying by positron emission$^1$ with a half-life of 109.77 ± 0.05 min $[15]$. The decay is entirely to the ground state of $^{18}\text{O}$; no $\gamma$ radiation follows the decay. The emission of the 511 keV $\gamma$ radiation following the positron annihilation, on the other hand, allows the measurement of the decay by $\gamma$ detection. By measuring the $^{18}\text{F}$ activity, the number of reaction products and therefore the total reaction cross section can be determined directly. The activation measurement of $^{17}\text{O}(p,\gamma)^{18}\text{F}$ provides therefore an independent means of cross section determination which can be used to check earlier experimental data and provide a constraint for $R$-matrix calculations regarding the total reaction cross section.

The aim of the present work is therefore to measure the $^{17}\text{O}(p,\gamma)^{18}\text{F}$ cross section with the activation method in a wide energy range. The next section provides detailed information about the experimental technique, the results are presented in Sec. III, while Sec. VI provides the summary and conclusions.

II. EXPERIMENTAL PROCEDURE

A. Target preparation and characterization

Solid state oxygen targets were produced by anodic oxidation of tantalum disks in water enriched in $^{17}\text{O}$. With this technique Ta$_2$O$_5$ layers can be produced with well defined Ta:O ratio, and the targets have high stability under beam bombardment. The anodization setup was the same as that used recently by the LUNA Collaboration for low energy $^{16}\text{O}(p,\gamma)^{17}\text{F}$ cross section measurements $[4,13]$. Full details of the anodization device and the preparation procedure have been published by the LUNA Collaboration $[16]$, here only the most important features and the differences are summarized.

Two water samples were used for the target preparations. The isotopic abundances of the $^{16}\text{O}$, $^{17}\text{O}$, and $^{18}\text{O}$ isotopes, respectively, were the following: $(15.5 \pm 0.6)\%$, $(77.8 \pm 0.6)\%$, and $(6.7 \pm 0.2)\%$ (sample 1) and $(39.5 \pm 0.6)\%$, $(27.4 \pm 0.6)\%$, and $(33.1 \pm 0.6)\%$ (sample 2). These values are quoted by the supplier.

Applying two different anodization voltages (24 and 50 V), targets with different thicknesses were produced. Altogether seven targets were prepared from the two water samples and with the two thicknesses. Intercomparison of the different targets was done by carrying out activation at the same proton energy on targets with different isotopic composition and/or thickness.

As the determination of the number of target atoms is crucial for the precise cross section measurements, different experimental techniques were used to determine this quantity. First, the Ta:O stoichiometry ratio and the thickness of the oxide layer were measured with Rutherford backscattering spectrometry (RBS). The first set of RBS measurements were carried out before the start of the activation experiments at the microbeam setup installed at the 5 MV Van de Graaff accelerator of Atomki $[17]$. A 1.6 MeV $\alpha$ beam bombarded the Ta$_2$O$_5$ targets, and the scattered particles were detected by two ion implanted Si detectors positioned at 135 and 165 degrees with respect to the beam direction. Exploiting the high lateral resolution of the microbeam setup, spectra were recorded at several different positions on the target surfaces. This test proved that the thickness and stoichiometry of the targets are uniform along the whole surface of the targets. The spectra were analyzed using the SIMNRA code $[18]$, which provided the areal density of the O atoms as well as the Ta:O ratio.

A second set of RBS measurement was carried out after the activation experiments using a completely independent setup, namely the activation chamber itself (i.e., similar beam size and position with respect to the proton beam used for the activations; see below). At the Tandetron accelerator a 10 MeV $^{16}\text{O}^{+}$ beam bombarded the Ta$_2$O$_5$ targets and a Si detector built into the activation chamber detected the backscattered ions.

Figure 1 shows typical spectra of the two RBS measurements. The measured data as well as the fits using the SIMNRA code are shown. The results of the $^{18}\text{O}$ RBS measurements were in good agreement with the ones obtained with $\alpha$-RBS (see below). The ratio of the Ta:O atoms was found to be $0.411 \pm 0.015$, in agreement with the stoichiometric value of 0.4.

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$^1$A weak electron capture decay branching also exists with 3% probability.
FIG. 2. Measured profile on the $E_p = 1098$ keV resonance in $^{17}$O($p, \gamma$)$^{18}$F. The target thickness was obtained from the width of the fitted resonance profile.

If the Ta:O ratio is known, a totally independent target thickness value can be obtained by the measurement of the resonance profile on a suitable nuclear resonance. We have investigated the target thicknesses also by this method using both $^{17}$O and $^{18}$O isotope content of the targets. The $E_p = 1098$ keV and $E_p = 1925$ keV resonances in the $^{17}$O($p, \gamma$)$^{18}$F and $^{18}$O($p, \gamma$)$^{19}$F reactions, respectively, were used to measure the target profiles. A 100% relative efficiency high-purity germanium (HPGe) detector was placed next to the activation chamber at zero degrees with its front face about 1 cm distant from the target. The yield of the strongest transition was used for the measurement of the profiles, which was the 937 keV transition of the first excited state to the ground state in the case of $^{18}$F$^{17}$O($p, \gamma$)$^{18}$F reaction and the 197 keV transition of the second excited state to the ground state in the case of $^{19}$F$^{18}$O($p, \gamma$)$^{19}$F reaction. The number of target atoms was obtained from the width of the target profiles using the Ta:O ratio given by the RBS measurements.

Figure 2 shows a typical resonance profile measured with the $^{17}$O($p, \gamma$)$^{18}$F reaction. The target thickness obtained from the resonance profile measurement using the two reactions gave consistent results. The comparison with the RBS results, however, revealed a roughly 9% systematic difference. The RBS measurements resulted in systematically higher thickness values. Table I summarizes the thickness results of a given target (prepared with 50 V anodization voltage) obtained with the four measurements.

The uncertainties quoted in the table are statistical only, stemming from the fit of the RBS spectra and the resonance profiles. Taking into account only these errors, the two methods are in contradiction. If, however, one includes the uncertainty of the stopping power, the results can be considered to be consistent. It is difficult to quantify the uncertainty of stopping power in our experiment as the stopping of protons, $\alpha$ particles, and $^{16}$O isotopes should be considered in O and Ta, and the related information [19] in the widely used SRIM code indicates uncertainties from about 2% up to 6%. Most likely the deviation of the thickness values has its origin in the uncertainty of the stopping power. Therefore, we have adopted the average of the two methods and assigned a conservative 6% uncertainty to the number of target atoms.

B. Activations

The activations were carried out at the new Tandetron laboratory of Atomki where a 2 MV Tandetron accelerator manufactured by High Voltage Engineering Europa B.V. was installed in 2015. The energy calibration of the accelerator was carried out by measuring resonances in the $^{27}$Al($p, \gamma$)$^{28}$Si reaction and the neutron thresholds in $^{7}$Li($p, n$)$^{7}$Be and $^{12}$C($p, n$)$^{13}$O reactions [20].

The Tandetron provided proton beams in the energy range between 500 keV and 1.8 MeV and the beam current was limited to about 5 $\mu$A in order to avoid target degradation. The lifetime of the targets was also increased by using an off-axis target chamber where the beam spot was shifted from the target center by 6 mm. By rotating the target between the consecutive activations, fresh or not heavily bombarded target spots could be selected. The target chamber was insulated from the rest of the beam line and served as a Faraday cup in order to determine the number of projectiles by charge integration. A secondary electron suppression voltage of $\sim 300$ V was applied behind the 4 mm diameter entrance aperture of the chamber.

Depending on the cross section, the length of the irradiations varied between 15 minutes and 5 hours. Although the beam intensity during the irradiations were typically very stable, in order to follow the possible fluctuations, the beam current was recorded in multichannel scaling mode with one minute time basis. The recorded time dependence of the beam current was then used in the analysis.

C. Measurement of the $^{18}$F decay

After the irradiation the target was removed from the chamber and transported to the counting laboratory where a 100% relative efficiency HPGe detector equipped with full 4π lead shielding was used to measure the annihilation $\gamma$ radiation of the targets. The $\gamma$ countings started typically 15 minutes after the end of the irradiation and the spectra were recorded every 10 minutes in order to follow the $^{18}$F decay.

Since the 511 keV annihilation radiation is present also in the laboratory background and can come from many possible sources, it is crucial to determine the background. The length of the countings was therefore typically 16 hours. Towards the
end of this counting period, the activity of $^{18}$F decayed to a negligible level and therefore the 511 keV background level could be estimated. This was always found to be consistent with the laboratory background measured without target, indicating that no long-lived positron emitter was created in the targets.

In some cases excess in the 511 keV activity was observed at the beginning of the counting period, indicating the production of some short-lived positron emitter. From its decay rate it was identified as $^{13}$N produced by the $^{12}$C($p,\gamma$)$^{13}$N reaction on carbon impurity of the target. This identification was also supported by the fact that such a deviation from the pure $^{18}$F decay was observed mostly around 500 keV proton energy where the $^{13}$C($p,\gamma$)$^{13}$N reaction has high cross section due to a broad resonance at about 420 keV [21]. In such cases roughly the first one hour of the counting was omitted from the analysis.

The decay of the 511 keV activity could always be fitted well using the literature half-life of $^{18}$F. As an example, Fig. 3 shows the decay curve measured after the irradiation at 520 keV. The figure indicates the above discussed $^{13}$N contribution, the $^{18}$F decay fitted with the literature half-life and the laboratory background level of the 511 keV line.

In order to maximize the detection efficiency, the targets were placed in close geometry onto the detector, directly on top of the detector end cap. Since the two annihilation photons are emitted from the source at opposite directions and no other x-ray or $\gamma$ radiation follows the $^{18}$F decay, true coincidence summing effect was not present in this measurement in spite of the close source-to-detector geometry.\footnote{The true coincidence of the two annihilation $\gamma$'s through a Compton scattering process was observed to cause less than 0.5% loss of counts from the 511 keV peak, and was therefore neglected.} The summing effect, on the other hand, is significant in the case of any multiline calibration source which could be used for the measurement of the absolute detection efficiency. The absolute efficiency was therefore measured in the counting geometry only with single line calibration sources. Calibrated $^{7}$Be, $^{65}$Zn, $^{85}$Sr, and $^{137}$Cs sources were used to obtain the efficiency curve of the detector. The $^{85}$Sr source was especially useful for the efficiency determination as it has a single $\gamma$ line at 514 keV, very close to the relevant 511 keV. The absolute efficiency was measured with 3% uncertainty, which includes also the beam spot size and target positioning effect.

\section{III. Experimental Results}

The cross section of $^{17}$O($p,\gamma$)$^{18}$F was measured between proton energies of 500 keV and 1.8 MeV. The selection of the actual proton energies was based on the structure of the $^{17}$O($p,\gamma$)$^{18}$F excitation function. The two broad resonances at 590 and 717 keV proton energies were measured with fine energy steps. The low energy tails of these resonances have significant contributions to the astrophysical reaction rate of $^{17}$O($p,\gamma$)$^{18}$F, especially at higher temperatures.

The energy regions where there are no resonances were covered with fewer data points. The aim of these measurements was to fix the absolute value of the direct capture part of the cross section, which again has an important contribution to the reaction rate.

In the studied energy range there are several narrow resonances at proton energies of 517, 673, 741, 826, 926, 1098, 1240, 1270, and 1345 keV. Activation runs at these energies were also carried out with the aim of confirming their existence in the ($p,\gamma$) channel and checking their resonance energies. The widths of these resonances, on the other hand, are often comparable with the target thicknesses used in the present work; the determination of the strengths of these resonances was therefore not a goal.

The obtained cross section results are listed in Table II. The first column shows the energy of the proton beam provided by the Tandetron accelerator. Based on the accelerator calibration, this value is known with a total uncertainty of less than 0.5 keV. The energy loss of the beam in the target layer is given in the second column. Especially near the resonances, the cross section changes significantly in the energy range covered by the target thickness. An effective proton energy was therefore calculated. For this calculation it was supposed that the cross section has a linear energy dependence in the energy range of the target. The slope of the cross section was estimated based on the adjacent experimental data points and on the shape of the excitation function as measured by previous works. The effective energy was then given by the median of the yield curve within the target thickness [22]. The uncertainty of the effective energy as listed in the table was estimated based on the slope of the cross section function. Higher energy uncertainties were assigned to the data points near the narrow resonances, where the cross section changes strongly within the target thickness.

In the table only the statistical uncertainty of the cross section values is quoted. This is obtained simply from the peak integration of the 511 keV $\gamma$ peak and the background subtraction. Typically the statistical uncertainties are between 0.5% and 5%. Higher statistical uncertainties can be found in the case of the lowest cross sections and for those points...
where, based on the literature data, higher cross sections were expected at a resonance but the actual resonance was found at slightly shifted energy (see below).

In order to obtain the total uncertainty of the cross section values, 7.6% systematic uncertainty must be added quadratically to the relative statistical uncertainties. This systematic uncertainty is the quadratic sum of the following components: number of oxygen atoms in the target (6%), \( \gamma \)-detection efficiency (3%), number of protons hitting the target (3% from charge integration), \(^{17}\text{O} \) enrichment (2%). Uncertainties well below 1%, such as the uncertainty of \(^{18}\text{F} \) decay parameters or the measurement of irradiation and counting times, were neglected.

In order to increase the reliability of our experiments, repeated activations were carried out at a few different proton energies using different targets. The results were always in agreement within the statistical uncertainties of the measurements. In Table II either the weighted average of these points are shown or—if targets with different energies were used—the more precise value was adopted.

### IV. Discussion

As most of the previous experiments yielded partial cross sections for the various transitions in \(^{18}\text{F} \) measured at a
FIG. 4. Experimental cross section of the $^{17}$O($p,\gamma$)$^{18}$F reaction in the upper part of the studied energy range. Besides the present work, the total and partial cross sections of Rolfs [5] are also shown as discussed in the text. The lines through the points are only to guide the eye.

given angle, it is rather difficult to compare the results of the present work with previous experiments. Above 500 keV proton energy the only total cross section (in the form of an astrophysical $S$-factor figure) is provided by Rolfs [5]. Although Rolfs studied the reaction in a wide energy range, total $S$ factor is only provided in the energy regions far from the resonances, i.e., below 450 keV (outside the energy range of the present work) and above 900 keV.

Figure 4 shows the total cross section determined in the present work and that of Rolfs in this high energy range. The points of Rolfs are taken from the EXFOR [23] database where they were obtained by scanning Fig. 17 of [5]. As one can see, the data of Rolfs are on average a factor of 1.5 higher than the present data, although the agreement becomes somewhat better at the highest energies.

In addition to the total cross section data, the partial cross section involving the first excited state to ground state transition in $^{18}$F measured by Rolfs is also included in the figure taken from EXFOR. Obviously, this partial cross section is lower than the total one, but it is included in the figure in order to compare the observed resonances in the high energy region. The existence of the narrow resonances observed by Rolfs is confirmed by the present work. There is an apparent energy shift between the two datasets: the resonances in the present work are observed at slightly higher energies than in Ref. [5] as given in EXFOR. However, the numerical values of the resonance energies, as given in Ref. [24], are in reasonably good agreement with the present work. The apparent discrepancy as can be seen in the figure can therefore most likely be attributed to the digitization uncertainty of the low resolution Fig. 11 of [5].

Between 500 and 900 keV proton energies, no total cross section values are available in the literature, making the comparison of our data with the existing database even more difficult. Figure 5 shows the measured cross section in this energy range. In order to compare at least the energy dependence of the cross section, besides the present data, partial cross sections measured by Rolfs [5] and Kontos et al. [11] are also included in the figure. As in Fig. 4, the data of Rolfs [5] are taken from its Fig. 11 as compiled in EXFOR. In the case of Kontos et al., capture to the first excited state of $^{18}$F measured at 135 degrees is arbitrarily chosen. The energy dependence of the cross section is very similar in the three datasets. The partial cross section of Rolfs exceeds the total cross section measured in the present work. This is similar to the observation on the direct capture cross section at higher energies.

A further comparison with literature data can be made at the lowest studied energy of the present work at $(E)_p = 500$ keV. Several recent low energy datasets extend up to this energy and some of them quote total cross section (or $S$ factor) which can be compared with the present work. Table III lists the experimental (or quasieperimental, see below) cross section values at 500 keV proton energy. The following literature data were considered: Hager et al. [12] measured the total cross section with the DRAGON recoil separator at $(E)_{c.m.} = 470$ keV corresponding to 497.9 keV proton energy, which matches exactly our lowest energy. The value is taken...
FIG. 5. Experimental cross section of the $^{17}\text{O}(p,\gamma)^{18}\text{F}$ reaction in the lower part of the studied energy range. Besides the present work, the partial cross sections of Rolfs [5] and Kontos et al. [11] are also shown as discussed in the text. The lines through the points are only to guide the eye.

from Table VI of [12]. In-beam $\gamma$-spectroscopy measurement of Newton et al. [10] provided a total cross section at 500 keV proton energy which again coincides with our data point, taking into account the energy uncertainties. The value is taken from Table I of [10]. Kontos et al. [11] do not provide total cross section data directly at $(E_p) = 500$ keV. A measured partial cross section around this energy region, however, is available and, based on these data, the authors provide total $S$-factor values in tabular form in their Table V. An interpolated value for $(E_{c.m.}) = 470$ keV [corresponding to $(E_p) = 500$ keV] is put into Table III, keeping the 12% relative experimental uncertainty.

The result of the present work at $(E_p) = 500$ keV is in good agreement with Hager et al. [12] and Kontos et al. [11]. The cross section of Newton et al. [10], on the other hand, is almost 20% lower than—and therefore barely consistent with—the other three values.

### V. R-MATRIX ANALYSIS

AZURE2, a multichannel and multilevel $R$-matrix code [25], was used to simultaneously fit the total cross section, measured by Newton et al. [10], Hager et al. [12], Di Leva et al. [4], Buckner et al. [14], and by the present work, as well as the primary transitions, measured by Kontos et al. [11], of the $^{17}\text{O}(p,\gamma)^{18}\text{F}$ reaction. The fit using these data will be referred to as “our” fit in the following.

First, we have compared the total $S$ factor obtained from an $R$-matrix fit made by Kontos et al. [11] with our experimental data. One can see in Fig. 6 that there is a good agreement between our data and the calculated one of Kontos, although some narrow resonances are omitted from their plot. Values of Kontos are obtained from Fig. 9 of [11] by figure digitization using the software PLOTDIGITIZER 2.6.8 [26].

In our $R$-matrix fit for the determination of direct capture, because of the nice agreement with the results of Kontos, we used same asymptotic normalization coefficient (ANC) values and high energy background poles that they used. Table IV lists these fixed ANC values. In addition, 15 MeV as the excitation energy of the background poles was selected. There are no proton scattering data to provide restrictions for the proton partial widths of the poles, so they were fixed at $\Gamma_p = 6$ MeV, close to the Wigner limit. The $R$-matrix radius was taken as

### TABLE III. Experimental cross section of $^{17}\text{O}(p,\gamma)^{18}\text{F}$ at $E_p = 500$ keV from the present and previous works.

| Reference          | Cross section at $E_p = 500$ keV (nb) |
|--------------------|-------------------------------------|
| U. Hager et al. [12]| $585 \pm 8_{\text{stat.}} \pm 75_{\text{syst.}}$ |
| J. R. Newton et al. [10]| $488 \pm 49$ |
| A. Kontos et al. [11]| $588 \pm 71^a$ |
| Present work       | $592 \pm 21_{\text{stat.}} \pm 45_{\text{syst.}}$ |

$^a$Not purely experimental value. See text.
zero energy is \(\sim\) transitions to the total \(S\) factor.

The uncertainty of the extrapolation of the total change of the above parameters around their fixed values.

These values are estimated from the manual manual.

The estimated dependence of the \(R\)-matrix extrapolation on the choice of the channel radius, the position of the background poles, and \(ANC\) values are \(\sim 4\%, \sim 7\%\) and \(\sim 15\%\), respectively. These values are estimated from the manual manual.

The uncertainty of the extrapolation of the total \(S\) factor to zero energy is \(\sim 20\%\).

Table V lists the calculated contributions of all the measured transitions to the total \(S\) factor at zero energy. The second column is calculated by Kontos et al. [11], the third one by Di Leva et al. [4], and the last one comes from our fit.

The error of our data is \(\sim 15\%\) because of the uncertainty of the choice of \(ANC\). The uncertainties are statistical only. The total \(S(0)\) value of Kontos and Di Leva are 5.4 \(\pm (th.)1.0\) \(\pm (exp.)0.6\) keV b and 5.0 \(\pm 0.3\) keV b, respectively. Our total \(S\) factor value at zero energy is 4.7 \(\pm 1.0\) keV b, where the error is statistical only.

Figure 7 shows the total \(S\) factor obtained from our \(R\)-matrix fit (continuous line) as well as experimental datasets of Newton et al. [10], Hager et al. [12], Di Leva et al. [4], Buckner et al. [14], and the present work. Narrow resonances are also included. The \(\chi^2\) value of our dataset is 7.3 without any normalization of datasets. The contribution of the direct capture to the total \(S\) factor at zero energy in our fit is \(S_{DC} = 4.3 \pm 1.0\) keV b, where the uncertainty is statistical only. Figure 8 shows the low energy total \(S\) factor obtained from our \(R\)-matrix fit with the above experimental datasets.

| Energy (keV) | \(\ell\) | ANC (fm\(^{-1/2}\)) |
|-------------|--------|----------------|
| 937         | 0      | 6.1            |
| 937         | 2      | 1.2            |
| 1121        | 2      | 2.7            |
| 2523        | 0      | 1.4            |
| 3062        | 0      | 4.5            |
| 3062        | 2      | 1.0            |
| 3839        | 0      | 4.6            |
| 3839        | 2      | 0.6            |
| 4115        | 0      | 2.5            |
| 4115        | 2      | 1.0            |
| 4652        | 2      | 1.3            |
| 4964        | 0      | 3.2            |
| 4964        | 2      | 0.7            |

\(r_c = r_0 \times (A_1^{1/3} + A_p^{1/3}) = 4.46\) fm, with \(r_0 = 1.25\) fm. More details about the selected values are in Ref. [11].

Our \(R\)-matrix analysis used the data set of Table II. As the quoted effective energies were used, no target effect was taken into account. No normalization of datasets was applied and for physical parameters the Brune parametrization [27] was used. The full parameter list of our \(R\)-matrix fit is provided as Supplemental Material [28]. It contains the used datasets and the AZURE2 input file with all parameters.

Table V lists the calculated contributions of all the measured transitions to the total \(S\) factor at zero energy. The second column is calculated by Kontos et al. [11], the third one by Di Leva et al. [4], and the last one comes from our fit.

| Transition (keV) | \(S(0)_{\text{Kontos}}\) [11] | \(S(0)_{\text{Di Leva}}\) [4] | \(S(0)_{\text{Present}}\) |
|-----------------|-------------------------------|-------------------------------|--------------------------|
| \(R/DC \to 937\) | 1.7 \(\pm 0.3\)              | 1.48 \(\pm 0.08\)             | 1.73 \(\pm 0.26\)        |
| \(R/DC \to 1121\) | 0.66 \(\pm 0.13\)           | 0.47 \(\pm 0.05\)             | 0.65 \(\pm 0.10\)        |
| \(R/DC \to 1700\) | 0.013 \(\pm 0.002\)         | 0.12 \(\pm 0.03\)             | 0.15 \(\pm 0.02\)        |
| \(R/DC \to 2523\) | 0.17 \(\pm 0.03\)           | 0.59 \(\pm 0.03\)             | 0.45 \(\pm 0.07\)        |
| \(R/DC \to 3062\) | 0.66 \(\pm 0.11\)           | 0.20 \(\pm 0.05\)             | 0.03 \(\pm 0.005\)       |
| \(R/DC \to 3839\) | 0.93 \(\pm 0.14\)           | 0.92 \(\pm 0.04\)             | 0.66 \(\pm 0.10\)        |
| \(R/DC \to 4115\) | 0.55 \(\pm 0.08\)           | 0.50 \(\pm 0.03\)             | 0.51 \(\pm 0.08\)        |
| \(R/DC \to 4652\) | 0.21 \(\pm 0.03\)           | 0.10 \(\pm 0.03\)             | 0.19 \(\pm 0.03\)        |
| \(R/DC \to 4964\) | 0.49 \(\pm 0.07\)           | 0.43 \(\pm 0.03\)             | 0.35 \(\pm 0.05\)        |

\(^a\)The \(S(0)\) of the present work was obtained by using for the fit simultaneously the partial cross sections from the literature (see text) and the total cross section presented in this paper.

![FIG. 6. Total \(S\) factor obtained from an \(R\)-matrix fit made by Kontos et al. [11] is compared with our experimental data of the \(^{17}\text{O}(p,\gamma)^{18}\text{F}\) reaction.](image1)

![FIG. 7. Total \(S\) factor obtained from our \(R\)-matrix fit (continuous line) is compared with experimental data of the \(^{17}\text{O}(p,\gamma)^{18}\text{F}\) reaction. The dashed line is the contribution of the direct capture to the total \(S\) factor (background poles included).](image2)
VI. SUMMARY AND CONCLUSIONS

In the present work the total cross section of the $^{17}$O($p, \gamma$)$^{18}$F reaction was measured with the activation method in a wide energy range for the first time with a total uncertainty of about 10%. Since this method provides cross sections which are in several aspects independent of the ones obtained with in-beam $\gamma$-spectroscopy and some systematic errors are not present, our results can be used to check the validity of previous data.

In general, our data are in good qualitative agreement with the structure of the excitation function of previous works. The possibility of the comparison of total cross sections is limited owing to the scarcity of total cross section data in the literature in the studied energy range. At energies above 900 keV, our results are on average a factor of 1.5 lower than those of Rolfs [5]. Similar deviation is found at lower energies where the partial cross sections of Rolfs exceed substantially our total cross section. Too high values of Rolfs were also pointed out earlier by Kontos et al. [11] in the case of individual transitions. This observation is confirmed by the present work.

A direct comparison of our cross section data with the literature was carried out also at a single proton energy of 500 keV. It is found that our value agrees well with that of Hager [12] and Kontos et al. [11], while the result of Newton et al. [10] is about 20% (two standard deviations) lower.

An $R$-matrix analysis with the AZURE2 code was performed to check the conformity of our measured total cross section dataset and to extrapolate the astrophysical $S$ factor to lower energies. In this analysis all primary transitions observed from $^{18}$F were simultaneously fitted with some total cross section datasets, including the present one. The resulting total $S$ factor is in good agreement with previous measurements and calculations within the experimental uncertainties.

Our total cross section data can be used to constrain any future theoretical description of the $^{17}$O($p, \gamma$)$^{18}$F reaction.

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