1. Introduction

Sulfur is among the most abundant elements in the universe following hydrogen, helium, carbon, oxygen, and nitrogen. Out of the 204 molecules that have been detected in the interstellar medium so far, twenty three of them are sulfur-bearing species, nine of which have been detected in the last decade. The mercapto radical (SH) and its cation, sulfanylium, (SH+) have both been detected in various astronomical environments, but the sulfur hydrides are fairly unusual among interstellar hydrides as none of the species S, SH, S+, SH+ or H2S+ can undergo exothermic H atom abstraction with H2. This observation makes these species good tracers of turbulent dissipation regions where gas temperatures are elevated and significant ion-neutral drift is present. It was, furthermore, observed to have a particularly high relative abundance in these objects in comparison with other hydrogenated molecules. Meanwhile, SH+ can be directly formed from the reaction between vibrationally excited H3(S¹Σg⁺(ν > 1) and S+ ions or the reaction between H2S+ and atomic H. The interstellar chemistry of sulfur has recently been reviewed for dark clouds, protoplanetary disks, hot cores and corinos, and pre-stellar cores.

The spectroscopy and ultraviolet photodissociation dynamics involving the mercapto radical and the sulfanylium ion have been relatively well studied with a number of experimental techniques as well as computations. The first photoelectron spectrum (PES) of SH was recorded by Dunlavey et al. where SH was produced with the rapid reaction F + H2S → SH + HF and the products were ionized with a Helium lamp. Several products were observed on top of the SH radical but several spectroscopic features were assigned to the X2Σ−, a1Δ, b1Σ+, A1Π and c1Π electronic states of the SH+ ion. Rostas et al. reported emission spectra of SH+ which allowed the determination of equilibrium molecular constants of the X2Σ−, and A1Π states of SH+ and SD+. The first threshold photoelectron spectrum (TPES) of SH+ was recorded by Hsu et al. where SH was produced via photodissociating a H2S parent in the UV (234–240 nm) and SH was ionized via two-photon excitation to high-ν Rydberg states, followed by pulsed-field ionization. The photoionization observables of SH and S such as the anisotropy parameter, β, and relative photoionization cross-sections were further studied.

Information is much scarcer, however, when dealing with absolute photoabsorption or photoionization cross-sections, and even more so when radical species are involved. Nevertheless, these are critical data needed to model and understand the photodynamics involving these species. In astrochemistry,
these are converted to photoabsorption/photoionization rates if the photon flux is known in a given region of space. By quantitatively constraining the rates of photoprocesses, other chemical and physical parameters can be more reliably obtained, thus facilitating how observations are interpreted. In a recent seminal paper,\textsuperscript{50} photoabsorption and photoionization cross sections of interstellar species were compiled and used to model wavelength-dependent lifetimes (process rates) of interstellar molecules, using measured and calculated cross sections from several different databanks. Therein, the photoionization cross section of SH was arbitrarily simulated by a Gaussian centered at 100 nm (12.4 eV) with a maximum of 5 × 10\textsuperscript{-18} cm\textsuperscript{2} (5.0 Mb).

Recently, the photoionization cross sections of radicals have been successfully studied at the DESIRS beamline at Synchrotron SOLEIL, employing a microwave discharge-equipped flow-tube.\textsuperscript{51,52} The microwave discharge is used to create Fluorine atoms that react with Hydrogen-bearing parent molecules; abstracting H atoms to produce HF and the desired radical product.\textsuperscript{53}

In this paper we present data regarding the absolute photoionization cross section of the SH radical from threshold up to 15.0 eV, followed by the measured threshold photoelectron spectra (TPES) of H\textsubscript{2}S, the mercapto radical, SH and the S atom, and briefly discuss the astronomical implications of these measurements.

2. Methodologies

2.1 Experimental set-up

S atoms and SH radicals were produced by the following consecutive reactions inside the flow-tube:

\[
\begin{align*}
F + H_2S & \rightarrow SH + HF \quad \Delta H_{r,298K} = -1.99 \text{ eV} \\
F + SH & \rightarrow S + HF \quad \Delta H_{r,298K} = -2.22 \text{ eV}
\end{align*}
\]

These reactions are both fast, barrier-less and exothermic\textsuperscript{54,55} and hence no isotopic effects are expected.

Experiments were performed on the DESIRS VUV undulator beamline\textsuperscript{26} located at the third generation French national synchrotron facility SOLEIL (Gif-sur-Yvette, France). The beamline provided linearly polarized light with a photon flux of 10\textsuperscript{12}–10\textsuperscript{13} ph s\textsuperscript{-1} over the VUV range and we used monochromator slits corresponding to a resolution of 6 meV at 10 eV. Downstream the interaction region, a photodiode (AXUV, IRD) is used to measure the photon flux and correct energy dependent measurements.

An H\textsubscript{2}S canister was obtained commercially (air liquide, \textgreek{m} 99.5% purity) and the gas was directed into a flow-tube reactor where it mixed with Fluorine atoms that were produced by a MW discharge of a 5% F\textsubscript{2} mixture diluted in He. The flow-tube was placed inside the permanent molecular beam endstation SAPHIRS\textsuperscript{57} and has been described in detail elsewhere.\textsuperscript{53} Several experiments have been carried out using the flow-tube where the physical parameters of the hydrogen abstraction are described in detail,\textsuperscript{53} hence only the most relevant parameters are tabulated here (see Table 1).

A gas filter upstream of the beamline monochromator was filled with Argon to filter out the higher harmonics from the undulator. Calibration of the energy scale was achieved by using autoionizing resonances corresponding to transitions in neutral atomic sulfur.\textsuperscript{58} The accuracy of the calibration is better than 2 meV between 10.0 and 12.0 eV. However, above 12.0 eV the energies of autoionizing resonances are only known with 3–5 significant digits. This decreases the certainty of the calibration above 12.0 eV and hence we report an energy calibration better than 6 meV from 12.0 eV up to 15.0 eV.

The double imaging photoelectron/photoion coincidence (i\textsuperscript{2}PEPICO) spectrometer DELICIOUS\textsuperscript{359} was used to detect electrons and ions in coincidence with a velocity map imaging (VMI) setup and an imaging linear time-of-flight analyzer, respectively. The coincidence scheme yields mass-selected photoelectron images which are then converted into photoelectron spectra (PES) by Abel inversion.\textsuperscript{60} Hence, we obtained mass-tagged PES of all the flow-tube reaction products simultaneously.

Two different energy scans were performed. First, by scanning the photon energy in the 9.7–12.0 eV range in 3 meV steps, and second, in the 11.9–15.0 eV range in 10 meV steps, allowing for a scan overlap of approximately 0.1 eV. Particle acceleration was achieved with a DC field of 53 V cm\textsuperscript{-1} for the first scan while the field was increased to 176 V cm\textsuperscript{-1} for the second scan. In short, mass-selected photoelectron signals as a function of electron kinetic energy and photon energy were obtained for the flow-tube reaction products between 9.7 and 15.0 eV which included and were not limited to H\textsubscript{2}S, SH, S, and S\textsubscript{2}. Results pertaining to the production of S\textsubscript{2} will be the subject of a later publication. The data were further reduced to yield the mass-selected photoionization yields (integrating the signal over all photoelectron kinetic energies at each photon energy), and the mass-selected TPES with a 5 meV electron resolution using a method described in previous works.\textsuperscript{61,62}

2.2 Isotopic correction of the 32\textsubscript{S} SH photoionization yield

The natural abundances of the 32\textsubscript{S}, 33\textsubscript{S}, and 34\textsubscript{S} sulfur isotopes are 94.99%, 0.75%, and 4.25%, respectively. As such, the 33\textsubscript{S}, 34\textsubscript{S}, and 33\textsubscript{SH} isotopologues had to be subtracted from the m/z = 33 amu and m/z = 34 amu signals to obtain the uncontaminated 32\textsubscript{SH} and H\textsubscript{2} 32\textsubscript{S} photoionization yields.

In the 32 amu channel of the total photoionization yields (see Section 3.2 below), the appearance of metastable signals – arising from autoionization of metastable S(\textsuperscript{1}D) atoms – provided

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Experimental parameters relevant to the flow-tube through which optimal conditions were achieved to perform the photon energy scans as described in the text</th>
</tr>
</thead>
<tbody>
<tr>
<td>Experimental parameters</td>
<td>Value</td>
</tr>
<tr>
<td>F\textsubscript{2}/He mixture</td>
<td>5%/95%</td>
</tr>
<tr>
<td>Total chamber pressure</td>
<td>1 Torr</td>
</tr>
<tr>
<td>Precursor flow speed</td>
<td>10 sccm (standard cubic centimeters per minute)</td>
</tr>
<tr>
<td>Total flow speed</td>
<td>1100 sccm</td>
</tr>
<tr>
<td>F atom concentration</td>
<td>1.0·10\textsuperscript{13} atom cm\textsuperscript{-3}</td>
</tr>
<tr>
<td>H\textsubscript{2}S concentration</td>
<td>2.0·10\textsuperscript{-13} molecule cm\textsuperscript{-3}</td>
</tr>
</tbody>
</table>

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us with a benchmark to subtract the corresponding metastable \(^{33}\text{S}\) signals from the 33 amu ion yield to obtain the pure \(^{32}\text{SH}\) photoionization yield. The 32 amu spectrum arises from photoionization of \(^{32}\text{S}\) atoms and includes no other contributing sources, i.e. it is the pure photoionization spectrum of \(^{32}\text{S}\). We multiplied it by a factor which made it equal to the contribution of the ionized \(^{33}\text{S}\) metastable atoms in the 33 amu spectrum and this allowed the \(^{33}\text{S}\) contribution to be effectively subtracted from the 33 amu photoionization spectrum. The same procedure was then applied to the 34 amu photoionization spectrum with regards to the \(^{34}\text{S}\) contribution.

To obtain the pure \(^{32}\text{H}_2^{32}\text{S}\) photoionization yield, the pure \(^{32}\text{SH}\) photoionization yield (as obtained from the subtraction of \(^{33}\text{S}\) signals from the 33 amu signal) was multiplied by a factor that was optimized so that the most prominent signals coming from the \(^{33}\text{SH}\) isotopologue, e.g. the autoionizing resonances at 10.18 eV and 10.83 eV were effectively removed (see Section 3.2).

The thresholds for dissociative ionization in \(^{32}\text{H}_2\) to form \(^{32}\text{S}\) and \(^{32}\text{SH}\) are approximately 13.4 eV and 14.3 eV, respectively. \(^{32}\text{S}\) To compensate for the contributions to the \(^{32}\text{S}\) and \(^{32}\text{SH}\) ion yields from dissociative ionization of \(^{32}\text{H}_2\), a scan was performed with the MW discharge off and the ion yields of \(^{32}\text{SH}\) and \(^{32}\text{S}\) were measured in relation to \(^{32}\text{H}_2\), and subtracted from the scan where the discharge was turned on.

### 2.3 Calculations

Potential energy curves for \(^{32}\text{SH}\) were calculated by using the internally contracted multireference configuration interaction method with Davidson correction (MRCI + Q) with complete active space self-consistent field (CASSCF) wave-functions. The CASSCF and MRCI calculations were performed at full valence, namely with 16 electrons distributed in 10 orbitals with the 1s and 2s orbitals of sulfur atoms kept doubly occupied but fully optimized. All calculations were performed using the MOLPRO 2012 package. \(^{64}\)

### 3. Results and discussion

#### 3.1 Reactor composition/reaction products

The mass spectrum obtained by integrating the ion TOF over the photon energy range 11.0–12.0 eV is presented in Fig. 1. The parent compound \(^{32}\text{H}_2\) is visible with the \(^{32}\text{H}_2^{32}\text{S}, \(^{32}\text{H}_2^{33}\text{S}, \) and \(^{32}\text{H}_2^{34}\text{S}\) isotopologues discernible, as well as the radical products, \(^{32}\text{SH}\) and \(^{32}\text{S}\). As the \(^{32}\text{H}_2\) is used as a precursor, its signal is severely depleted which ultimately affected its recorded TPES (see Section 3.2).

The strongest signal, however, pertains to \(^{32}\text{S}_2\), which can form through recombination via the exothermic reactions \(^{32}\text{S} + \text{H}_2^{32}\text{S} \rightarrow \text{H}_2^{34}\text{S}\) or even \(^{32}\text{S} + \text{F}^{32}\text{S}\). The \(^{32}\text{S}^{32}\text{S}\) and \(^{32}\text{S}^{34}\text{S}\) isotopologues are visible as well in abundances of 1.5% and 8.0% compared to that of \(^{32}\text{S}_2\) or approximately twice that of the natural abundances of the \(^{33}\text{S}\) and \(^{34}\text{S}\) isotopes as expected.

Other products/impurities are observed, most notably \(^{32}\text{SO}\), but other extraneous species include \(^{32}\text{NS}, \(^{34}\text{SO}, \) \(^{32}\text{SF}, \) \(^{32}\text{HSF}\) and \(^{32}\text{S}_2\)\(^{32}\)\(^{32}\).
11.2 eV by using the method that has been applied previously to the OH, CH₃, C₂H₅, and C₆H₆ radicals. The energy for the absolute measurement was carefully chosen so that it was located in a region without autoionization resonances to minimize energy resolution effects, and for which absolute data was available for the reference species, H₂S and S. After, a series of mass spectra were recorded at different reaction times (equivalent to different injector distances), with the microwave discharge turned on and off (see Fig. S1, ESI†). With the discharge turned off, F atom production stops and thus no radicals form in the flow-tube. Under these conditions and with the photon energy set to 11.2 eV, only the parent ion, H₂S⁺, was produced and the isotopologues H₂³²S, H₂³³S, H₂³⁴S were observed in the relative ratios 94.84%, 0.87%, and 4.30%, respectively. These compare well (≤0.15%) with the natural abundances. Signals pertaining to the ³⁵S and ³⁶S isotopes were not strong enough to be deciphered from the background.

The TOF-integrated signal for the parent ion (H₂S⁺) is proportional to the number of irradiated species in the ionization volume and its photoionization cross section:

\[ S_{H_2S^+} \propto m_{H_2S} \times \sigma_{H_2S}^{ion} \]

Here, \( S_{H_2S^+} \) represents the ion signal, \( m_{H_2S} \) is the number of irradiated parent species, and \( \sigma_{H_2S}^{ion} \) is the photoionization cross section of H₂S.

Conservation of matter then assumes that the number of free radicals that are produced equals the change in the amount of H₂S that is consumed after the discharge is turned on, i.e.:

\[ \Delta n_{H_2S} = n_{H_2S}^{OFF} - n_{H_2S}^{ON} = n_{SH}^{ON} + n_{S}^{ON} \]

which allows us to write:

\[ \frac{\Delta S_{H_2S^+}}{\sigma_{H_2S}^{ion}} = \frac{S_{SH}^{ON}}{\sigma_{SH}^{ion}} + \frac{S_{S}^{ON}}{\sigma_{S}^{ion}} \]

This allows us to measure the photoionization cross section of the mercapto (SH) radical, using measured values of the photoionization cross sections of H₂S and S atoms. Barthel et al. have measured the photoionization cross section of S atoms and found 11.4 ± 2.0 Mb at 11.20 eV. Here it should be noted that Barthel et al. recorded the sum of the photoionization cross sections of both the S(3P) ground state and the S(1D) metastable state. As noted previously, we do see the contribution of S(1D) in our measured photoionization cross section, but it is significantly weaker than that of S(3P), despite the large cross-section of these autoionizing features (50 Mb), therefore the contribution from S(1D) is negligible. Watanabe and Jursa measured the photoionization cross section of H₂S to be 23.6 ± 4.7 Mb at 11.20 eV. It is also pertinent to highlight the fact that the low isotopic ³³S abundance (less than 1% of the ³²S abundance) combined with the low yield of ³²S with the discharge turned on, makes the ³³S signal from the 33 amu ion signal and the ³⁴S and ³³SH signals from the 34 amu ion signal exceedingly small in comparison with the ³²SH and H₂³⁴S signals, respectively.

Moving the flow-tube injector and altering the reaction time provides a set of 11 independent measurements to estimate the mean and standard deviation of the absolute measurement from a Monte Carlo simulation which also includes the reported errors for the H₂S and S cross-sections. Note that although the reactor conditions were set to minimize secondary reactions, for longer reaction times, S production is not negligible and needs to be taken into account as written in the above equations. The resulting distribution of values was then fitted with a Gaussian function to derive the mean value and associated error as 14.3 ± 4.8 Mb (2σ) at 11.20 eV (see Fig. S2, ESI†). Note that the majority of the error stems from the uncertainty of the previously measured cross sections of H₂S and S, as the normal distribution of our measurements showed a standard deviation smaller than 15%.

The absolute measurement was used to calibrate the photoionization yield of ³³SH on an absolute scale and the result is presented in Fig. 3, where cross sections are compared with the simulation used by Heays et al.⁵⁰

The arbitrarily simulated photoionization cross sections of SH exhibit non negligible value in the 110–90 nm (11.3–13.8 eV) range with a maximum of 5.0 Mb centered approximately 2 eV above the ionization threshold. When compared to our results this wildly underestimates the true photoionization yield that we measure. More precisely, the absolute photoionization cross section curve in the current version of the Leiden database underestimates the photoionization cross section by a factor of at least 10 at its maximum (around 12.4 eV) and furthermore does not account for any of the strong autoionizing resonances just above the ionization threshold whose maxima range between 50 and 85 Mb at our photon energy resolution. This almost equates to the maximum absolute absorption cross sections of
the SH radical which have been calculated as approaching 90 Mb at approx. 145 nm (8.55 eV).\(^{70}\)

Several structures in the cross section are still observed up until approximately 14 eV. Above 14 eV the intensity stays roughly the same within the error bar limits but one would expect a slight decrease because of the predissociation of the \(A^3 \Pi\) state in SH\(^+\) (see below).

As presented by Heays \textit{et al.}, current models estimate the photoionization rates of SH as being below \(10^{-10}\ \text{ s}^{-1}\) while the photodissociation rates of SH are estimated as being slightly above \(10^{-9}\ \text{ s}^{-1}\). Our measurements, however, suggest that the SH photoionization rates are much more competitive. Currently, SH\(^+\) formation in astrochemical models such as the Meudon PDR code is predominantly governed by the \(\text{H}_2 + \text{S} \rightarrow \text{H}_2\text{S}\) reaction but with the accompaniment of our measured photoionization cross section for SH, then photoionization might become a viable contributor of astronomical SH\(^+\).\(^{71}\)

### 3.4 Threshold photoelectron spectra (TPES) of H\(_2\)S, SH and S

The spectroscopy of the cation has been measured with the TPES technique, by scanning the photon energy while considering only electrons with little or no kinetic energy, so that signal is obtained only when the photon energy is resonant with the ionization energy. TPES of H\(_2\)S, SH and S with a 12 meV total resolution were recorded according to a method taking full advantage of our electron imaging capabilities as already described elsewhere\(^{61}\) (for a review on TPES see ref. 72 and 73). The three TPES are depicted in Fig. 4.

#### 3.4.1 Hydrogen sulfide (H\(_2\)S\(^+\)).

Hydrogen sulfide is a triatomic bent molecule of the \(C_2v\) point group. Its ground electronic state has the electron configuration \(\ldots (4a_1)^2(2b_1)^2(5a_1)^2(2b_1)^2\) and its cationic electronic ground state, H\(_2\)S\(^+\) (\(X^2B_1\)), is formed by removing an electron from the \(2b_1\) orbital of H\(_2\)S which is predominantly non-bonding. Consequently, as noted by Hochlaf \textit{et al.},\(^{74}\) the photoelectron spectrum for H\(_2\)S\(^+\) (\(X^2B_1\)) is expected to be dominated by the origin vibration band. This is indeed the case in their PFI study as in ours (see upper panel of Fig. 4).

Associated with the \((X^2B_1)\) ground state are two vibrational progressions, namely the \(\nu_2\) symmetric stretching and \(\nu_3\) bending modes. Our results are comparable to that of Hochlaf \textit{et al.} as can be seen from Table S1 in ESI\(^+\) where our results for the H\(_2\)S\(^+\) TPES are compiled and compared. Many of the peaks observed by Hochlaf \textit{et al.} could not be resolved from other peaks. The TPES peaks were fitted with Lorentzian profiles and the fitted positions are presented in Table S1 (ESI\(^+\)).

Removal of an electron from the \(5a_1\) orbital of H\(_2\)S, gives rise to the H\(_2\)S\(^+\) (\(A^2A_1\)) state. Hochlaf \textit{et al.} observed the \(\nu_2\) progression of this state between 12.7 and 14.7 eV and they were able to resolve

---

**Fig. 3** 32SH\(^+\) absolute photoionization cross section (in Mb = \(10^{-18}\ \text{cm}^2\)) as a function of the incident photon energy (blue). The red arrow corresponds to the photon energy (11.2 eV) at which the absolute cross section was normalized. The uncertainty of the photoionization yield is presented in gray and is obtained by Monte Carlo uncertainty propagation. In green the arbitrarily simulated SH photoionization cross section from the Leiden database\(^{50}\) is presented.

**Fig. 4** Experimental TPES of H\(_2\)S (top – black), SH (middle – blue), and S (bottom – red). The total energy resolution is 12 meV. The assignment marked with an asterisk in the SH spectrum corresponds to the photoionization of the \(\Omega = 1/2\) spin–orbit component of the SH ground state.
rotational structures in their spectrum. Note, however, that our mass-selected spectrum drops in intensity around 13.5 eV, due to the opening of the dissociative ionization channel that forms S(1S) + H₂(X 1Σ⁺) from 13.4 eV.

3.4.2 Sulfanylium ion (SH⁺). Neutral SH has a σ²π³ open shell electronic configuration. In the ion, the removal of a π electron from the open shell gives rise to a ground state with ³Σ⁺ symmetry and two low-lying excited singlet states, a¹Δ, and b¹Σ⁺. The removal of a σ electron leads to the cationic A¹Π state. In our recorded TPES of SH⁺ (see middle panel of Fig. 4), there are a few spectral features observed whose origins are traced to these states. The photoionizing transitions towards the two lowest excited vibrational levels of the X³Σ⁻ ground state are observed at 10.422 eV and 10.719 eV. The origin bands of the a¹Δ, b¹Σ⁺, and A¹Π states are observed at 11.643 eV, 12.728 eV, and 14.098 eV, respectively. These values are compiled and compared with the results of Dunlavey et al. in Table 2 and there are some discrepancies to be found, most notably the ground state X³Σ⁻ and the b¹Σ⁺ state.

In the experiments of Dunlavey et al., the spectrum was not mass resolved, meaning that signals pertaining to photoionization of both H₂S and S could have clouded their spectra. Furthermore, the value they cite for the ground state is the value we measure for the ionization of the Ω = 1/2 spin-orbit component of the SH ground state. It should be mentioned that contributions from a sequence band, i.e. ν⁺ = 1 ← ν⁻ = 1, would be found at very similar energies as the origin band of the same spin-orbit component. However, while the X³Π₁/₂ state is 47 meV above the X³Π₁/₂ state, the first vibrational band of the X³Π₁/₂ state is 350 meV higher in energy. The X³Π₁/₂ state is more easily populated, and this is verified by simulations (see Fig. S3, ESIF). Due to spectral congestion, lack of mass selectivity and the added spin–orbit component of the SH ground state, it is quite likely that Dunlavey et al. incorrectly assigned the origin band.

This brings about an interesting point regarding the flow-tube which is its propensity to produce species that are vibrationally hot whilst being rotationally cooled. Vibrationally hot species have already been produced in a similar manner via H-abstractions by fluorine radicals, e.g. CH₂, but the pressures within the flow-tube are high enough so that rotational excitation is quenched by collisions (e.g. NH₂, CH₂, CH₃CN). Electronic excitation with different multiplicities than the ground state are also populated, such as the metastable S(1D) atoms here. For states of the same multiplicity, the relaxation is prompt and as such the corresponding rotational temperatures would be equally small whilst the vibrational temperature is relatively high.

As for the b¹Σ⁺ state, it was found to exhibit broader peaks as compared with the rest of the bands in their PES study. The accuracy in the determination of the ionization onset of the state can be called into question due to the aforementioned spectral congestion. As stated previously, our values are calibrated with the appearance of S atomic lines from the NIST database. As such, we present our results with a confidence interval that is 2 meV below 12 eV and 6 meV between 12 eV and 15 eV.

Dunlavey et al. observed and assigned more vibrational bands of the A¹Π state and we do indeed observe these bands as well. However, they are observed in the S⁺ channel of the TPES (Fig. 5) as the A¹Π state interacts with a repulsive state (³Σ⁻) and thus predissociates to S⁺(1S) + H⁺(S) for ν⁺ ≥ 1, as seen in the potential energy curves shown in Fig. 6. This observation is in line with prior studies, both experimental18,19,40 and theoretical.19,43,46 This confirms that the ν⁺ > 0 vibrational levels are purely dissociative within the timeframe that ions spend inside the acceleration region of DELICIOUS3, i.e., a few μs.

According to the calculations of Brites et al.46 the predissociative lifetime of the A¹Π decreases significantly above ν⁺ = 0. Namely, the A¹Π (ν⁺ = 1, 2, 3) states were calculated to exhibit a predissociative lifetime of 10.26 ns, 44.89 ps and 1.54 ps, respectively. In DELICIOUS3, under the current experimental conditions, ions traverse the acceleration region in a matter of a few μs. Thus, we would not expect to observe any parent species

| Table 2 | Energies of the TPES peaks for the electronic states of SH⁺ observed in the middle panel of Fig. 4 |
|-----------------|-----------------|-----------------|
| Electronic state | Energy (eV) – our values | Energy (eV) – Dunlavey et al.17,79 |
| X³Σ⁻ (ν⁺ = 0)⁷⁸ | 10.375 | 10.375 |
| X³Σ⁻ (ν⁺ = 0) | 10.422 | 10.37 |
| X³Σ⁻ (ν⁺ = 1) | 10.719 | — |
| a¹S (ν⁺ = 0) | 11.643 | 11.65 |
| b¹Σ⁺ (ν⁺ = 0) | 12.728 | 12.76 |
| A¹Π (ν⁺ = 0) | 14.098 | 14.11 |
| A¹Π (ν⁺ = 1) | 14.296 | 14.30b |
| A¹Π (ν⁺ = 2) | 14.477 | 14.49b |
| A¹Π (ν⁺ = 3) | 14.652 | 14.66b |

⁷⁸ Excitation from the Ω = 1/2 spin–orbit component of the SH ground state & contribution from the ν⁺ = 1 ← ν⁻ = 1 hot band.⁸ Estimated values from the ωₓ and ωₓCₓ values presented by Dunlavey et al.
with such short predissociative lifetimes. Conversely, the \( \text{A}^2\Pi (\nu' = 0) \) state was reported to have a predissociative lifetime of 38.68 \( \mu s \), meaning that it would predissociate only after exiting the acceleration region of DELICIOUS3 and therefore we do not detect it in the \( S^+ \) channel.

### 3.4.3 \( S^+ \) atoms

An exhaustive study of the photoionization of neutral \( S \) atoms in the ground state and metastable states has already been carried out by Barthel et al.\(^{60}\) by using tunable VUV radiation in the energy range 9-30 eV. Hence, only the most prominent features in our TPES will be briefly discussed here below.

Our TPES of the \( S \) atoms reveal several weak autoionizations from Rydberg series as well as direct ionizations, where the excited/ejected electron is originated from the ground state of sulfur (3s\(^2\)3p\(^3\)1P\(_{\text{b},1,0}\)) as well as from the metastable 3s\(^2\)3p\(^4\)1D\(_2\) state. The 3P\(_{1,0}\), 3P\(_0\), and 4D\(_2\) states all lie 396.055 cm\(^{-1}\) (1.1454 eV), 573.640 cm\(^{-1}\) (0.0711 eV), and 9238.609 cm\(^{-1}\) (1.1454 eV), respectively, above the 3P\(_2\) state. Valence shell photoionization of neutral \( S(3p_2) \) atoms produces the ionic continuum states \( S'(3s), S'(3d), \) and \( S'(3p) \) which have ionization potentials of 10.36, 12.20, and 13.40 eV, respectively.\(^{58}\) Our results are compiled in Table S2 (ESI).

Direct ionizations from the 3P\(_{2,1,0}\) ground state to the \( S'(3s) \) 3s\(^2\)3p\(^3\)1S\(_{1/2}\) ground state are observed at 10.360 eV, 10.309 eV, and 10.288 eV, respectively. Similarly, direct ionizations to the \( S'(3p) \) 3s\(^2\)3p\(^3\)2P\(_{1/2}\) state from the 3P\(_{2,1}\) ground state are observed at 12.199 eV and 12.149 eV. A very weak signal at 11.051 eV is attributed to direct ionization of metastable \( S(4D_2) \) atoms to the \( S'(2D_{1/2}) \) ionic state.

Three weak peaks are observed around 9.5 eV, approximately 0.87 eV below the direct ionization threshold of the 3P\(_2\) ground state. These signals correspond to excitations from \( S(4D_2) \) to a Rydberg series above the ionization threshold which promptly autoionize. These transitions have been well characterized but the peak we observe at 9.413 eV was observed but only tentatively assigned as a 3s\(^3\)3p\(^1\)1D\(_2\) 3d state but without a corresponding term symbol.\(^{60}\) Unassigned lines correspond to autoionization due to imperfect hot electron subtraction.

### 4. Summary and conclusions

Here we present the first measurement of the absolute photoionization cross section of the \( 32\text{SH} \) radical over a large photon energy range, calibrated at the photon energy of 11.2 eV using the known values for \( \text{H}_2\text{S} \)\(^{65}\) and \( S \).\(^{69}\) The uncertainty we present with the measured cross section mainly stems from these reference values, but within our error bars the cross section we measured is significantly different from the arbitrarily estimated photoionization cross section that is currently used in the Leiden cross section database.\(^{50}\)

The mass-selectivity of our \( \text{i}^2\text{PEPICO} \) spectra allowed us to experimentally verify the predissociative inclination of the \( \text{A}^2\Pi (\nu' > 0) \) states, as a vibrational progression was observed in the \( S^+ \) portion of the TPES and only the \( \text{A}^2\Pi (\nu' = 0) \) state was observed in the TPES of \( \text{SH} \). To estimate any effects these results would have on the overall sulfur chemistry of the interstellar medium, further calculations would be required that are outside the scope of this work.

### Conflicts of interest

There are no conflicts to declare.

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### References

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