Millimeter Wave Rotational Spectrum and Centrifugal Distortion of Thioketene, H₂C=C=S

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a-type rotational transitions of molecules in the vibrational ground state of thioketene, H₂C=C=S, have been measured in the millimeter wavelength region. The measurements yielded improved rotational constants:

\[ A = 286.655(82) \text{ MHz}, \]
\[ B = 5.659.47596(72) \text{ MHz}, \]
\[ C = 5.544.51260(72) \text{ MHz}. \]

A detailed centrifugal distortion analysis by means of Watson’s S-reduced Hamiltonian led to the determination of four quartic, two sextic and two higher order distortion constants:

\[ D_{JK} = 1.08569(4) \text{ kHz}, \quad H_{JK} = 0.716(20) \text{ Hz}, \]
\[ D_{JK} = 168.269(77) \text{ kHz}, \quad H_{KJ} = -408.7(73) \text{ Hz}, \]
\[ d_1 = -25.46(88) \text{ Hz}, \quad L_{KJ} = 0.65(24) \text{ Hz}, \]
\[ d_2 = -5.21(35) \text{ Hz}, \quad S_{KJ} = -0.0533(24) \text{ Hz}. \]

Effective rotational and centrifugal distortion constants using planarity conditions were calculated. The electric dipole moment of thioketene was determined to be \( \mu = 1.013(3) \text{ D.} \)

I. Introduction

Intensive studies have been carried out on the rotational spectra of various isotopic species of ketene, H₂C=C=O, since 1950 [1—6]. The centrifugal distortion analysis of ketene [3, 4], together with information from additional infrared data [3, 5], enabled Mallinson and Nemes [6] to reevaluate the molecular force field for ketene. Recently, microwave investigations on the higher homologues thioketene, H₂C=C=S [7, 8], and selenoketene, H₂C=C=Se [9, 10], have been reported.

The first interstellar microwave detection of interstellar ketene was reported in 1977 by Turner [11]. The detection in the source Sgr B2 was based on three transitions: 5_14 — 4_13, 5_15 — 4_14, and 4_13 — 3_12 at the frequencies 101.98139, 100.09451 and 81.586.19 MHz, respectively. Further millimeter wave transitions were tentatively detected and assigned. Recently, Thaddeus and coworkers have identified isothiocyanic acid, HNCS [12], and methyl mercaptan, CH₃SH [13], in Sgr B2 from observations of a number of millimeter wave rotational transitions. The abundances of both of these sulfur-containing molecules, relative to their oxygen analogues which had previously been detected in interstellar dust clouds, imply a sulfur/oxygen ratio not very different from the terrestrial value. Therefore, the detection of the five-atomic molecule H₂C=C=S in interstellar dust clouds might become possible with more sensitive instrumentation. The predicted rotational transition frequencies based on the measured millimeter wave spectrum, together with calculated standard deviations, provide the basis for a possible search for interstellar thioketene.

Moreover, thioketene has raised some interest due to the fact that it is a more stable isomer of thiirene, H — C = C — H, according to SCF-calculations [14, 15]. These calculations also reveal the fact that ethynylmercaptane, H — C ≡ C — SH, should have the same stability as thioketene. As a matter of fact, Kranz and Laurenzi [16] detected both ethynylmercaptane and thioketene in the photolysis of 1,2,3-thiadiazole. However, no ethynylmercaptane was found in the products of pyrolysis reactions [14]. This was confirmed by our microwave experiments.

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Several methods of preparing thioketene are known [17]. All reports confirm its instability at room temperature, so that only pyrolysis experiments in flow-systems can be taken into consideration for spectroscopic experiments. In such an experiment Georgiou, Kroto and Landsberg measured four rotational transitions and could determine the rotational constants $B$ and $C$ and the centrifugal distortion constant $D_{JK}$ of thioketene [7].

In this paper a-type R-branch ground state transitions are reported which have been studied in the millimeter-wave region from 60 up to 230 GHz. 125 transitions have been measured and fitted with the S-reduced Hamiltonian according to Watson [18].
II. Experimental Procedures

Thioketene was produced in two different ways (Scheme I).

\[ \text{Me-S-Me} \rightarrow 1300 \text{K} \]
\[ \text{Me-C=C-S-Me} \rightarrow 1100 \text{K} \]

Scheme I.

As in the report on the earlier microwave measurements of thioketene [8], hexamethyltrithiane (I) was pyrolysed at 1300 K to give thioketene, which was directly pumped through the 2 m glass cell of the millimeter wave spectrometer as shown in Figure 1. Infrared analysis of the gas mixture after pyrolysis showed that a large fraction of the products consisted of ethylene, propylene and allene, thus indicating that only a rather poor yield of thioketene is obtainable by this method. A recently reported procedure [14] of preparing thioketene by pyrolysing dithioacetic acid (Scheme I) was found to yield a considerably higher intensity of the absorption lines, thus indicating higher column densities of H$_2$CCS.

Attempts to isolate thioketene have failed so far. It could be trapped (spectrum disappears) at temperatures below about 150 K, but polymerised upon warming. Even in the gas phase, at sample pressures of 40 µbar, decay took place with a halflife of about five minutes, independent of the way of preparation. The use of argon as carrier gas in order to entrain the thioketene did not improve the stability.

Microwave measurements were performed near 22 GHz in order to determine the dipole moment using a Hewlett-Packard spectrometer, model 8460 MRR. The millimeter-wave measurements were carried out with a spectrometer described by Winnewisser et al. [19, 20]. A block diagram of the newly improved version of the spectrometer is shown in Fig. 1 and a detailed description of recent improvements is given by Helms et al. [21]. The microwave power was generated by reflex klystrons in the range from 28 - 40 GHz and multiplied by means of harmonic generation. The multiplier diodes used were phosphorus diffused p-type Si/W point contact diodes and were prepared and mounted in our laboratory using a slightly simplified variant of the method of Burrus [22, 17]. Used together with a recently installed liquid helium cooled InSb detector [21] they proved to generate sufficient power for measuring transitions up to 475 GHz. All measurements were carried out with the help of a digital data acquisition system [20] to improve the signal-to-noise ratio and to measure automatically the transition frequencies. The accuracy of the measurements is better than 20 kHz at a klystron fundamental frequency of 40 GHz. Sample pressure in the cell was in the range 20 - 60 µbar.

III. Assignment of the Measured Rotational Transitions

The available data on thioketene from the earlier measurements [7] enabled us to predict the entire ground-state rotational spectrum in the millimeter wave region with some certainty. The most intense absorption lines in the observed frequency range could easily be assigned to thioketene ground state transitions. The spectrum shows the characteristic pattern of a-type 4R-branch rotational transitions. It differs only slightly from the ketene spectrum, having slightly less inertial asymmetry:

\[ \alpha(CH_2CCS) = -0.99918, \alpha(H_2CCO) = -0.99722. \]

Figure 2 shows an oscilloscope display of the most intense lines of the \( J = 10 \rightarrow 9 \) transition together with the assignment of the ground state spectrum. The splitting of the \( K_a = 1 \) lines is 1.15 GHz, which places them outside the represented frequency range. Their intensity is comparable to the \( K_a = 5 \) line intensity. The intensities of the transitions follow basically an asymmetric top intensity distribution modulated by the spin statistics due to the two protons; because of the molecular C$_2$V symmetry, transitions arising from odd \( K_a \) levels have three times the statistical weight of those arising from even \( K_a \) levels. Ground state transitions with \( K_a > 7 \) have intensities comparable to excited vibrational state transitions. Therefore the measurement and assignment of these weaker lines require precise predictions, which however could not be provided by the centrifugal distortion analysis (see discussion).

Table 1 gives the line frequencies of the observed and calculated transitions. It was impossible to resolve \( K_a \) doublets for \( K_a > 2, K_a > 3 \) and \( K_a = 3 \). \( J > 11 \) line frequencies were assigned and fitted to just one of the doublet transitions. For \( J > 10 \) the observed \( K_a = 3 \) lines were fitted to both components which were weighted equally. For \( J > 17 \) these lines...
had to be rejected. Several other measured transitions were also rejected from the fit, indicating an error in measurement or overlap with excited state lines. The standard deviation of the least squares analysis was obtained to be 16.5 kHz, which corresponds to the assumed reproducibility of the frequency measurements.

IV. Centrifugal Distortion Analysis

The centrifugal distortion analysis of the observed rotational spectrum was performed using Watson’s reduced Hamiltonian in the \( I_1 \) axis representation [18]. As in previous papers on ketene [3, 4], the A-reduced form of the Hamiltonian operator with \( R_6 = 0 \) was used first to calculate the rotational energy levels. Further refinement of the distortion constants could not be achieved with this form. The higher inertial symmetry of thioketene made it necessary to use the S-reduced Hamiltonian according to Watson, in which \( R_5 \) is zero. The complete form of this operator with the constants used is given below:

\[
\begin{align*}
\hat{\mathcal{H}} &= \frac{1}{2} (B + C) \hat{P}_z^2 + \frac{(A - (B + C)/2)}{2} \hat{P}_z^2 \\
&+ \frac{(B-C)/4}{2} \hat{P}_z^2 \left( \hat{P}_+^2 + \hat{P}_-^2 \right) \\
&+ D \hat{P}_z^4 - D_{JK} \hat{P}_z^2 \hat{P}_x^2 - D_{KK} \hat{P}_z^4 + H_{JJ} \hat{P}_z^6 + H_{JK} \hat{P}_z^4 \hat{P}_x^2 + H_{KJ} \hat{P}_z^4 \hat{P}_y^2 \\
&+ L_{KJ} \hat{P}_z^6 + S_{KJ} \hat{P}_z^2 \hat{P}_x^2 \hat{P}_y^4
\end{align*}
\]

with \( \hat{P}_z = \hat{P}_z + i \hat{P}_y \).

\( \hat{P}, \hat{P}_x, \hat{P}_y, \) and \( \hat{P}_z \) are the operators for the total angular momentum and its components. The constants \( A, B \) and \( C \) are Watson’s reduced rotational constants in the \( I_1 \) axis representation. Only two sextic constants have been used in our calculations. In addition to the sextic terms we found it necessary to add two higher order terms, \( L_{KJ} \) and \( S_{KJ} \), in order to fit the observed transitions with \( K_a \) values of 6 and 7. The reduced spectroscopic constants in the \( I_1 \) axis representation are given in Table 2.

As was pointed out by Nemes and Winnewisser [4], a-type rotational spectra of nearly symmetric prolate rotors contain information about the difference \( (A - D_K) \), but not about \( A \) and \( D_K \) separately. In order to obtain the best possible \( A \) constant the value of \( D_K \) was taken from ketene [3] and was held fixed in the least squares analysis. The uncertainty of \( D_K \) might cause a larger error of \( A \) than that calculated from the least squares analysis and given in Table 2. The remaining rotational constants and the quartic and sextic constants are well determined. The octic constant \( L_{KJ} \) is poorly
determined and the dentic constant $S_{KJ}$ shows that the Hamiltonian is not converging properly. These two constants therefore have to be considered only as fitting parameters to allow the measured frequencies of the $K_a=6$ and $K_a=7$ lines to be included in the calculations. In order to complete the distortion analysis, perpendicular rovibrational infrared transitions would have to be included.

V. Dipole Moment

The electric dipole moment for thioketene was determined from Stark effect measurements of the $2_{02} - 1_{01}$ transition to be 1.01(3) D. This value places thioketene between the homologues ketene, with $\mu = 1.41$ D [3], and selenoketene with $\mu = 0.90$ D [9].

VI. Discussion

The reduced spectroscopic constants given in Table 2 can be transformed to the so-called determinable constants, $\mathfrak{A}$, $\mathfrak{B}$, $\mathfrak{C}$, $T_{aa}$, $T_{bb}$, $T_{cc}$, $T_1$ and $T_2$, following the notation of Watson [18] and Yamada and Winnewisser [23]. These constants are invariant to a unitary transformation of the Hamiltonian and they are listed together with the derived inertial moments and the $\tau$-defect $\Delta T_{cc}$ in Table 3. The errors of these parameters are estimated from the standard deviations obtained in the least squares analysis. The small value for the $\tau$-defect $\Delta T_{cc}$ permits the application of any of the various planarity relations [23] in order to obtain the complete set of derived quartic distortion constants as was done in the case of ketene [4].

In the discussion of the previous section it was shown that the addition of higher order terms in the Hamiltonian was necessary to fit the $K_a=6$ and $K_a=7$ transitions. However, the inclusion of these terms did not prove useful in predicting the line positions of the $K_a=8$ lines, which could not be found at the expected positions. Though their intensities are rather low, we believe that the Hamiltonian used is insufficient to allow good frequency predictions outside the observed range of $K_a$ states. No predictions for $K_a > 7$ lines were therefore included in Table 1.

In a recent communication Yamada [24] has explained and analyzed the observed anomaly found in the $^1R_5$ and $^1R_6$ branches of the far infrared spectrum of HNCO. An interaction between the $K_a=6$ and 7 levels in the ground vibrational state and the $K_a=5$ and 6 levels in the lowest excited vibrational state is caused in HNCO by a centrifugal distortion correction term in the Hamiltonian. Although the $K_a$ energy levels in the ground vibrational state of $\text{H}_2\text{C}=\text{C}=\text{S}$ are lower in energy ($A \sim 30$ cm$^{-1}$ for HNCO, 9.5 cm$^{-1}$ for $\text{H}_2\text{CCS}$), the ratio of the rotational constant $A$ to the energy of the lowest bending mode is probably similar for HNCO and $\text{H}_2\text{CCS}$. Thus the lowest bending mode of $\text{H}_2\text{CCS}$ can be expected to be involved in this type.

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**Table 2. Spectroscopic parameters of thioketene for Watson’s reduced Hamiltonian in the $I^c$-axis representation.**

<table>
<thead>
<tr>
<th>$\mathfrak{A}$/MHz</th>
<th>286 655 (82)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mathfrak{B}$/MHz</td>
<td>5 659.47596 (72)</td>
</tr>
<tr>
<td>$\mathfrak{C}$/MHz</td>
<td>5 544.51269 (75)</td>
</tr>
<tr>
<td>$D_{11}$/kHz</td>
<td>1.08569 (4)</td>
</tr>
<tr>
<td>$D_{22}$/kHz</td>
<td>168.269 (77)</td>
</tr>
<tr>
<td>$D_{33}$/MHz</td>
<td>23.5 (fixed)</td>
</tr>
<tr>
<td>$d_1$/Hz</td>
<td>-25.46 (68)</td>
</tr>
<tr>
<td>$d_2$/Hz</td>
<td>-5.21 (35)</td>
</tr>
<tr>
<td>$H_{JK}$/Hz</td>
<td>0.716 (20)</td>
</tr>
<tr>
<td>$H_{KJ}$/Hz</td>
<td>-408.7 (73)</td>
</tr>
<tr>
<td>$L_{KJ}$/Hz</td>
<td>0.65 (24)</td>
</tr>
<tr>
<td>$S_{KJ}$/Hz</td>
<td>-0.0533 (24)</td>
</tr>
</tbody>
</table>

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**Table 3. Watson’s determinable constants and moments of inertia for thioketene.**

<table>
<thead>
<tr>
<th>$\mathfrak{A}$/MHz</th>
<th>286 655 (82)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mathfrak{B}$/MHz</td>
<td>5 659.64633 (80)</td>
</tr>
<tr>
<td>$\mathfrak{C}$/MHz</td>
<td>5 544.68316 (83)</td>
</tr>
<tr>
<td>$I_{aa}/\mu\text{A}^2$</td>
<td>1.76301 (50)</td>
</tr>
<tr>
<td>$I_{bb}/\mu\text{A}^2$</td>
<td>89.294625 (12)</td>
</tr>
<tr>
<td>$I_{cc}/\mu\text{A}^2$</td>
<td>91.146056 (13)</td>
</tr>
<tr>
<td>$A/\mu\text{A}^2$</td>
<td>0.08842 (53)</td>
</tr>
<tr>
<td>$T_{aa}$/MHz</td>
<td>-23.66935 (8)</td>
</tr>
<tr>
<td>$T_{bb}$/kHz</td>
<td>-1.1470 (25)</td>
</tr>
<tr>
<td>$T_{cc}$/kHz</td>
<td>-1.0452 (25)</td>
</tr>
<tr>
<td>$T_1$/kHz</td>
<td>-171.495 (79)</td>
</tr>
<tr>
<td>$T_2^x$/kHz</td>
<td>-4.2204 (34)</td>
</tr>
<tr>
<td>$\Delta T_{cc}$/Hz</td>
<td>2.3 (73) undefined</td>
</tr>
</tbody>
</table>

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\[ a \] The standard deviation of the fit is 16.5 kHz for 125 equally weighted lines. Numbers in parentheses are standard errors.
of resonance for some value of $K\alpha$. The symmetry selection rules are analogous in the two cases. The network of Coriolis interactions in HNCO is similar to that in H$_2$CCO [5, 6] and therefore a corresponding set of interactions can be expected in H$_2$CCS, favoring an interaction with the ground state.

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