

Submillimeter spectroscopy and astronomical searches of vinyl mercaptan, C₂H₃SH[★]

M.-A. Martin-Drumel^{1,2,★★}, K. L. K. Lee², A. Belloche³, O. Zingsheim¹, S. Thorwirth¹, H. S. P. Müller¹, F. Lewen¹, R. T. Garrod⁴, K. M. Menten³, M. C. McCarthy², and S. Schlemmer¹

¹ I. Physikalisches Institut, Universität zu Köln, Zùlpicher Str. 77, 50937 Köln, Germany
e-mail: marie-aline.martin@u-psud.fr

² Harvard-Smithsonian Center for Astrophysics, 60 Garden Street, Cambridge, MA 02138, USA

³ Max-Planck-Institut für Radioastronomie, Auf dem Hügel 69, 53121 Bonn, Germany

⁴ Departments of Astronomy & Chemistry, University of Virginia, McCormick Road, PO Box 400319, Charlottesville, VA 22904-4319, USA

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ABSTRACT

Context. New laboratory investigations of the rotational spectrum of postulated astronomical species are essential to support the assignment and analysis of current astronomical surveys. In particular, considerable interest surrounds sulfur analogs of oxygen-containing interstellar molecules and their isomers.

Aims. To enable reliable interstellar searches of vinyl mercaptan, the sulfur-containing analog to the astronomical species vinyl alcohol, we investigated its pure rotational spectrum at millimeter wavelengths.

Methods. We extended the pure rotational investigation of the two isomers *syn* and *anti* vinyl mercaptan to the millimeter domain using a frequency-multiplication spectrometer. The species were produced by a radiofrequency discharge in 1,2-ethanedithiol. Additional transitions were remeasured in the centimeter band using Fourier-transform microwave spectroscopy to better determine rest frequencies of transitions with low-*J* and low-*K_a* values. Experimental investigations were supported by quantum chemical calculations on the energetics of both the [C₂H₄S] and [C₂H₄O] isomeric families. Interstellar searches for both *syn* and *anti* vinyl mercaptan as well as vinyl alcohol were performed in the EMOCA spectral line survey carried out toward Sgr B2(N2) with ALMA.

Results. Highly accurate experimental frequencies (to better than 100 kHz accuracy) for both *syn* and *anti* isomers of vinyl mercaptan are measured up to 250 GHz; these deviate considerably from predictions based on extrapolation of previous microwave measurements. Reliable frequency predictions of the astronomically most interesting millimeter-wave lines for these two species can now be derived from the best-fit spectroscopic constants. From the energetic investigations, the four lowest singlet isomers of the [C₂H₄S] family are calculated to be nearly isoenergetic, which makes this family a fairly unique test bed for assessing possible reaction pathways. Upper limits for the column density of *syn* and *anti* vinyl mercaptan are derived toward the extremely molecule-rich star-forming region Sgr B2(N2) enabling comparison with selected complex organic molecules.

Key words. astrochemistry – molecular data – method: laboratory: molecular – ISM: individual objects: Sagittarius B2 (N) – ISM: molecules – surveys

1. Introduction

From the diatomic SH⁺ and SH radicals (Menten et al. 2011; Neufeld et al. 2012) to the large complex organic molecule ethyl mercaptan (ethanedithiol, C₂H₅SH; Kolesníková et al. 2014), nearly 20 sulfur-bearing species have been observed in the interstellar medium (ISM) and in the circumstellar envelopes of evolved stars. This total is substantial, representing roughly 10% of all the known interstellar or circumstellar compounds. While the abundance of S-containing species in the diffuse ISM is consistent with the estimated cosmic abundances of sulfur (Savage & Sembach 1996) and is well reproduced by current astrophysical models (Neufeld et al. 2015), the same set of compounds only accounts for ~0.1% of the expected sulfur abun-

dance in the cold, dense clouds and circumstellar regions around young stellar objects (Joseph et al. 1986; Tieftrunk et al. 1994). This remarkable disparity has given rise to a simple, but currently unanswered, question: Where is the missing sulfur? Several potential reservoirs have been investigated, including atomic sulfur (Anderson et al. 2013), S-bearing species trapped on icy dust grains (Jimenez-Escobar & Muñoz Caro 2011), and S-containing molecules in the gas phase (Martin-Domenech et al. 2016; Cazzoli et al. 2016), along with other possibilities. From studies of shocked gas, however, atomic sulfur can only account for at most 5–10% of the total sulfur cosmic abundance (Anderson et al. 2013), implying that additional sulfur sinks must exist. Many studies propose that the depletion of sulfur reflects accretion onto dust grains, where it could be sequestered in various forms as inorganic and organic molecules (Scappini et al. 2003), or atomic and polymeric S_n chains and rings (Wakelam et al. 2004). So far, however, only two sulfur molecules, OCS and SO₂, have been detected on grains, both in too low abundance for either species to be a major reservoir of sulfur (Palumbo et al. 1995; Boogert et al. 1997). Since H₂S

* Tables B.1 and B.2 are only available at the CDS via anonymous ftp to cdsarc.u-strasbg.fr (130.79.128.5) or via <http://cdsarc.u-strasbg.fr/viz-bin/qcat?J/A+A/623/A167>

** Present address: Institut des Sciences Moléculaires d'Orsay (ISMO), CNRS, Univ. Paris-Sud, Université Paris-Saclay, 91405 Orsay, France.

is the most abundant sulfur-bearing molecule in cometary ices (Bockelee-Morvan et al. 2000), its presence on ice grains has also received some attention, but the rather low upper limit on its abundance also appears to rule out this species as a major sink for sulfur (Smith 1991; Jimenez-Escobar & Munoz Caro 2011). Finally, photoprocessing of sulfur-containing ice products could yield gaseous S-bearing species in the dense ISM in molecular forms that are yet to be identified (Jimenez-Escobar & Munoz Caro 2011). Consequently, the identification of missing interstellar sulfur carriers requires the detection of new gaseous sulfur containing compounds, which in turn requires laboratory investigations of their rotational spectra. Transient species are particularly appealing candidates, but are often challenging to characterize in the laboratory compared to their stable, often commercially available, counterparts, especially in the millimeter- and submillimeter-wave region in which high-altitude, high-sensitivity radio facilities such as the Atacama Large Millimeter/submillimeter Array (ALMA) operate.

While Kolesníková et al. (2014) reported the detection of ethyl mercaptan in Orion, its presence there is not without controversy (Müller et al. 2016). Regardless, experimental data on related molecules that might also be present in the ISM and participate in the same chemical network, for instance unsaturated variants, may prove helpful in resolving this controversy. The isomeric family with the elemental formula $[C_2H_4S]$ – thioacetaldehyde (CH_3CHS), thiirane ($c-C_2H_4S$), and vinyl mercaptan (CH_2CHSH) – are of particular interest as all three isovalent oxygen counterparts (acetaldehyde, oxirane, and vinyl alcohol) have been detected in the ISM (Gottlieb 1973; Dickens et al. 1997; Turner & Apponi 2001). Structural isomers are an invaluable tool to probe the chemical dynamics of astrophysical environments. While a minimum energy principle was once proposed to assert molecular abundance in space (Lattelais et al. 2009), the general consensus is that chemical kinetics, rather than thermodynamic stability, is driving the formation of molecules in astronomical sources (Loomis et al. 2015). Interestingly, relatively little is known about isomers of relatively large astrophysical species (five atoms and more); this is in part because of the difficulties involved in both producing and detecting these species in the laboratory, and the $[C_2H_4S]$ family is no exception. While the pure rotational spectrum of each of these three singlet isomers has been investigated in the microwave domain (Kroto et al. 1974; Cunningham et al. 1951; Tanimoto & Saito 1977), high-resolution measurements at millimeter and shorter wavelengths have only been published for thiirane (Hirao et al. 2001; Bane et al. 2012), which is a likely reflection of the stability of this compound in the laboratory. In contrast, thioacetaldehyde polymerizes rapidly, with a lifetime on the order of ten seconds (Kroto et al. 1974), and consequently it must, as with vinyl mercaptan, be produced in situ for spectroscopic studies. To confidently detect these two transient species in the ISM, laboratory measurements at higher frequency are essential, particularly in spectral bands in which ALMA operates.

In this study we measured the pure rotational spectrum of the *syn* and *anti* isomers of vinyl mercaptan in their ground vibrational state up to 250 GHz. From the newly recorded transitions, we derived highly accurate rotational and centrifugal distortion constants for both species, ensuring that astronomical searches in the millimeter domain can now be undertaken with confidence. Using the new experimental data, we carried out searches on the spectral line survey EMOCA (Exploring Molecular Complexity with ALMA) performed toward the region of high-mass star formation Sagittarius (Sgr) B2(N) (Belloche et al. 2016). As part of this work, we also investigated the energetics of the $[C_2H_4S]$

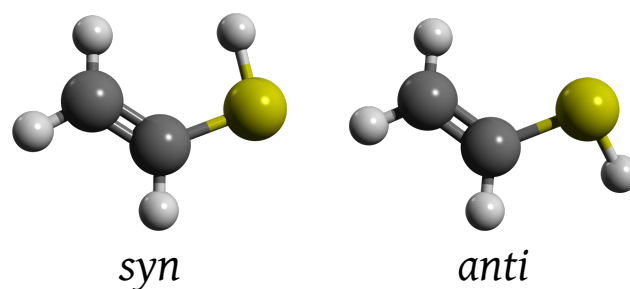


Fig. 1. Molecular representation of planar *syn* and quasi-planar *anti* vinyl mercaptan.

isomeric family and its isovalent oxygen counterparts via high-level quantum chemical calculations.

2. Methods

2.1. Spectroscopic background

Vinyl mercaptan exists in the gas phase in two distinct rotameric forms, the planar *syn* and quasi-planar *anti* isomers (Fig. 1); the former is only slightly more stable than the latter by about 0.6 kJ mol^{-1} (Almond et al. 1983, 1985; Plant et al. 1992), which yields a *syn/anti* ratio at room temperature of about 1.2. It is worth noting that while the equilibrium structure of the *anti* isomer is non-planar, with a calculated SH torsional angle of about $130\text{--}150^\circ$, the barrier to linearity (0.14 kJ mol^{-1}) is small compared to the zero-point vibration, which results in a vibrationally averaged planar structure (Tanimoto & Macdonald 1979; Almond et al. 1985). As a consequence, both isomers can be considered as belonging to the C_s point group symmetry (Almond et al. 1983).

Experimental studies on vinyl mercaptan are limited. Photoelectron (Chin et al. 1994) and low-resolution vibrational spectra (Almond et al. 1977, 1983) have been reported, while pure rotational spectra of the two isomers in the ground and several excited vibrational states have only been measured up to 50 GHz (Almond et al. 1977; Tanimoto & Saito 1977; Tanimoto et al. 1979; Tanimoto & Macdonald 1979). Isotopic investigations of the microwave spectra of the SD isotopologs¹ (Tanimoto & Saito 1977; Tanimoto et al. 1979; Tanimoto & Macdonald 1979), and extensive studies of deuterated variants of the *syn* isomer (Almond et al. 1985) that followed, enabled a determination of a partial geometry. Owing to the absence of experimental data for isotopic species involving substitution at any of the three heavy atoms, a complete structure has not been reported. Additionally, accurate dipole moments in the ground vibrational state are available from Stark measurements: $\mu_a = 0.813(1) \text{ D}$, $\mu_b = 0.376(4) \text{ D}$, $\mu_{\text{tot}} = 0.896(3) \text{ D}$ for the *syn* isomer; and $\mu_a = 0.425(10) \text{ D}$, $\mu_b = 1.033(10) \text{ D}$, $\mu_{\text{tot}} = 1.117(14) \text{ D}$ for the *anti* isomer (Tanimoto et al. 1979; Tanimoto & Macdonald 1979).

2.2. Computational details

Quantum-chemical calculations of the $[C_2H_4S]$ and $[C_2H_4O]$ isomeric families were performed at the coupled-cluster level with single, double, and perturbative triple excitations [CCSD(T)] (Raghavachari et al. 1989). All calculations were performed using the Coupled-Cluster techniques for

¹ I.e., in which the single hydrogen atom bonded to the S atom is replaced by a deuterium atom.

Table 1. Ground state molecular parameters (in MHz) of *syn* and *anti* vinyl mercaptan derived in this study compared with previously published values (Tanimoto et al. 1979; Tanimoto & Macdonald 1979) and those calculated theoretically (Calc.).

	<i>syn</i> -CH ₂ CHSH			<i>anti</i> -CH ₂ CHSH		
	This work ^a	Previous ^a	Calc. ^b	This work ^a	Previous ^a	Calc. ^b
A _e	50 304.4	49 843.5
ΔA ₀	431.2	36.4
B _e	5866.0	5912.5
ΔB ₀	38.1	31.9
C _e	5253.4	5308.9
ΔC ₀	37.3	50.6
A ₀	49 816.0376 (11)	49 815.28 (6)	49 873.2	49 423.5726 (11)	49 422.75 (5)	49 807.1
B ₀	5835.708685 (92)	5835.716 (14)	5827.9	5897.21162 (11)	5897.215 (9)	5880.6
C ₀	5222.075531 (85)	5222.081 (11)	5216.1	5279.43951 (13)	5279.436 (9)	5258.3
D _J × 10 ³	2.724203 (89)	2.85 (17)	2.716	3.09729 (17)	3.12 (11)	3.117
D _{JK} × 10 ³	-33.4874 (12)	-33.2 (21)	-35.89	-37.6145 (40)	-38.5 (17)	-40.02
D _K	0.79113 (14)	...	0.8100	0.80990 (18)	...	0.9496
d ₁ × 10 ³	-0.424456 (88)	-0.425 (35)	-0.4239	-0.47324 (19)	-0.498 (51)	-0.4143
d ₂ × 10 ³	-0.023739 (73)	...	-0.0211	-0.03110 (25)	...	-0.0411
H _{KJ} × 10 ⁶	-3.1367 (59)	-4.356 (31)
# of lines	333	37	...	205	32	...
RMS/kHz	43	49
σ ^c	0.95	1.03

Notes. ^(a)Numbers in parentheses are 1σ uncertainties in units of the last digit. ^(b)A_e, B_e, and C_e rotational constants derived from structure calculated at the ae-CCSD(T)/cc-pwCVQZ level of theory. Zero-point vibrational corrections ΔA₀, ΔB₀, and ΔC₀ and quartic centrifugal distortion constants were calculated at the fc-CCSD(T)/cc-pV(T+d)Z level. ^(c)Reduced standard deviation (unitless).

Computational Chemistry (CFOUR) program package (Stanton et al. 2017; Harding et al. 2008a) in combination with Dunning's hierarchies of correlation consistent polarized valence and polarized core-valence sets (Dunning et al. 2001; Peterson & Dunning 2002). The best estimate equilibrium structures have been calculated at the all-electron (ae) CCSD(T)/cc-pwCVQZ level of theory, which has previously been shown to yield equilibrium structures of very high quality for molecules harboring second-row elements (see, e.g., Coriani et al. 2005; Thorwirth & Harding 2009; Thorwirth et al. 2015). Centrifugal distortion constants and zero-point vibrational corrections ΔB₀ to the rotational constants were calculated in the frozen-core (fc) approximation at the fc-CCSD(T)/cc-pV(T+d)Z level and ground state rotational constants were obtained as B₀ = B_e - ΔB₀. In this work, the equilibrium rotational constants B_e are calculated from the ae-CCSD(T)/cc-pwCVQZ structures. The calculated spectroscopic parameters for both *syn* and *anti* vinyl mercaptan are given in Table 1, and the elements of their Z-matrices are given in Appendix A.

The relative energetics of the [C₂H₄O] and [C₂H₄S] isomers were calculated using a composite thermochemistry scheme similar to the High Accuracy Extrapolated Ab initio Thermochemistry (HEAT) approaches (see Bomble et al. 2006; Harding et al. 2008b). Using the CCSD(T)/cc-pwCVQZ geometries, a series of additive contributions are calculated: the extrapolated CCSD(T)/aug-cc-pCVXZ (X = D, T, Q) correlation energy; the correlation energy from perturbative quadruple excitations with fc-CCSDT(Q)/cc-pVDZ; the anharmonic zero-point contribution with CCSD(T)/cc-pV(T+d)Z; the diagonal Born-Oppenheimer correction with HF/aug-cc-pVTZ; and 1- and 2-electron scalar relativistic corrections with CCSD(T)/aug-cc-pCVTZ.

The relative stabilities, with CH₃CHO and CH₃CHS as the point of reference, are shown in Fig. 2. The energetic ordering

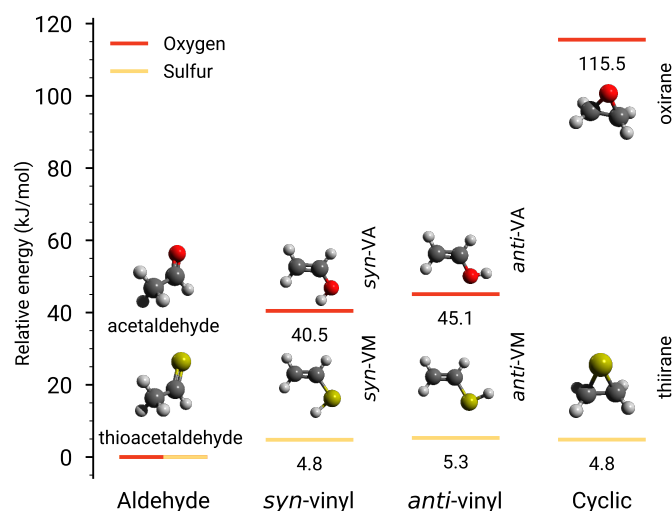


Fig. 2. Energetics of the [C₂H₄X] species (X = O or S), relative to the respective aldehydic forms, i.e., acetaldehyde and thioacetaldehyde, both set to 0 kJ mol⁻¹.

of [C₂H₄O] isomers is in agreement with previous calculations by Lattalais et al. (2009) and Karton & Talbi (2014): CH₃CHO is the most stable isomer, followed by vinyl alcohol and oxirane. In particular, the energetics computed at the W2-F12 level by Karton & Talbi (2014) for the *syn* isomer of vinyl alcohol are within ~1 kJ mol⁻¹ of the values presented in this work. The energy difference between the *syn* and *anti* isomers has been examined theoretically (Nobes et al. 1981) and experimentally; the value presented (4.6 kJ mol⁻¹) is in excellent agreement with both the microwave (4.5 ± 0.6 kJ mol⁻¹, Rodler 1985) and the far-infrared semi-experimental determination (4.0 kJ mol⁻¹,

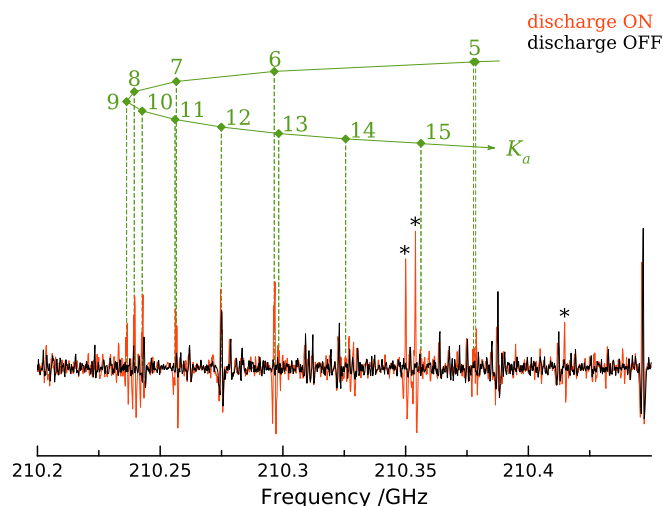


Fig. 3. Portion of the millimeter-wave spectrum recorded with RF discharge on (in red) and off (in black) showing the K_a series in the ${}^qR(18)$ ($J' \leftarrow J'' = 19 \leftarrow 18$) branch of *syn* vinyl mercaptan. For the lowest K_a value visible on the plot, the asymmetry splitting ($K_a + K_c = J$ or $J + 1$) is resolved. Transitions in black most likely arise from the precursor while transitions in red labeled with an asterisk remain unidentified (see text).

Bunn et al. 2017). The sulfur analogs [C_2H_4S] on the other hand are near-isoenergetic at the current level of treatment. With the exception of CH_3CHS , the three remaining isomers are within ~ 0.5 kJ mol $^{-1}$ of one another, where *anti*-vinyl mercaptan is the highest in energy (5.3 kJ mol $^{-1}$). Our ab initio energy difference between the vinyl mercaptan isomers (0.5 kJ mol $^{-1}$) is in quantitative agreement with the 0.6 kJ mol $^{-1}$ estimate by Almond et al. (1985).

2.3. Millimeter-wave spectroscopy

Millimeter-wave measurements on vinyl mercaptan were performed using the Cologne (sub)millimeter direct absorption spectrometer in the spectral regions 70–120 GHz and 170–250 GHz. The radiation was produced by a commercial amplifier-multiplier chain (Virginia Diodes Inc.) and optimized by in-house electronics. To reduce $1/f$ noise, frequency modulation was used with a $2f$ demodulation, yielding line shapes close to a second-derivative for all recorded transitions. The 5 m long discharge cell, equipped with a roof-top mirror on one end, allows 10 m of absorption path length. A detailed description of the apparatus can be found in Martin-Drumel et al. (2015). Vinyl mercaptan was produced by a radio-frequency (RF) discharge in a continuous flow of 1,2-ethanedithiol ($HSCH_2CH_2SH$) at a pressure of about 10 μ bar. It should be noted, however, that since the pressure was measured between the cell and pumping stage, a sizable pressure gradient may have been present in the cell. Similar to a previous study on OSSO (Martin-Drumel et al. 2015), the strongest signals of vinyl mercaptan were observed when the RF power was reduced to fairly low powers. However, to maintain stable discharge conditions in the cell, and thus to limit the noise induced by a “blinking” plasma, a discharge power of no less than 5 W was applied.

Upon ignition of the discharge, relatively strong transitions of both *syn* and *anti* isomers of vinyl mercaptan were observed, as illustrated in Fig. 3. To derive rest frequencies as accurate as possible, rotational transitions were measured individually using frequency steps of 10 and 20 kHz in the 70–120 GHz and 170–

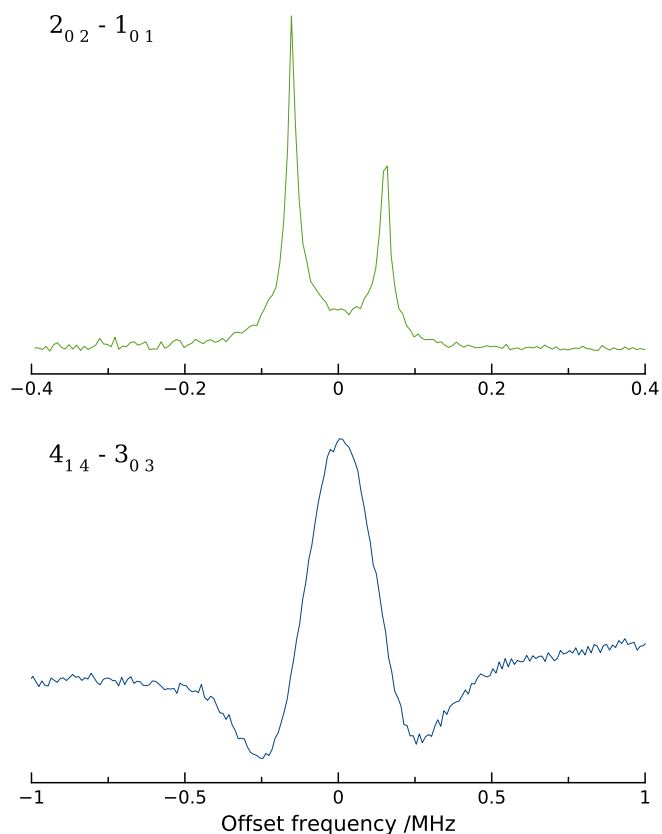


Fig. 4. Two transitions of *anti* vinyl mercaptan recorded in the microwave (top trace, in green) and millimeter-wave (bottom trace, in blue) domains. The frequency axis is given as an offset from the rest frequencies, 22 346.6731 and 85 467.985 MHz, respectively. The doublet structure of the microwave transition (top spectrum) is caused by Doppler splitting, which arises because of the coaxial arrangement of the cavity axis with respect to the axis along which the gas expands; the rest frequency is taken as the average of the two components. The second derivative line shape aspect of the millimeter transition (bottom spectrum) arises from $2f$ demodulation of the modulated frequency.

250 GHz regions, respectively, and signal averaging was used to reach a signal-to-noise ratio of at least 10, and up to 50 in some cases (Fig. 4). The accuracy on each line frequency was derived from a fit to the line profile, accounting for the full width at half maximum (FWHM), frequency step, and signal-to-noise ratio as described in Martin-Drumel et al. (2015). Using this approach, rest frequencies were derived with uncertainties as low as 10 kHz. Ultimately, nearly 300 and 150 new transitions (both *a*- and *b*-type) of *syn* and *anti* isomers of vinyl mercaptan were recorded in the millimeter domain with quantum numbers up to $J'' = 22$, $K_a'' = 17$ and $J'' = 19$, $K_a'' = 14$, respectively. We note that, besides vinyl mercaptan, other species were produced under our experimental conditions (as indicated in Fig. 3). Although some of their transitions were observed with high signal-to-noise ratios, they could not be assigned to any known species using the Cologne Database for Molecular Spectroscopy (CDMS) and NASA Jet Propulsion Laboratory (JPL) catalogs (Endres et al. 2016; Pickett et al. 1998). The carriers of these lines may be vibrational satellites of vinyl mercaptan or its precursor, 1,2-ethanedithiol, or other species produced in the discharge. Since no wide spectral surveys were conducted, however, no further line assignment attempts were made at this juncture.

2.4. Centimeter-wave spectroscopy

For each isomer of vinyl mercaptan, roughly ten transitions previously observed by Tanimoto et al. (1979) and Tanimoto & Macdonald (1979) were remeasured between 10 and 35 GHz using the Cambridge Fourier-transform microwave spectrometer, which was described elsewhere (McCarthy et al. 2000). In doing so, uncertainties on the rest frequencies of these lines were reduced by nearly two orders of magnitude, from 100 kHz to 1 kHz. In this experiment, vinyl mercaptan was generated by a 1 kV DC discharge in the throat of a small nozzle through a mixture of C₂H₂ and H₂S heavily diluted in neon and the resulting products were supersonically expanded into a large vacuum chamber. By the time the products reach the center of the Fabry-Perot cavity, where they are probed by the microwave radiation, the rotational temperature falls to 10 K or less. An example of a microwave transition of *syn*-CH₂CHSH is presented in Fig. 4.

2.5. Astronomical observations

We searched for vinyl mercaptan in the EMOCA spectral line survey performed toward the Sgr B2(N) star-forming region with ALMA. We focused our analysis on the hot molecular core Sgr B2(N2) at the position (α, δ_{J2000}) = (17^h47^m19.86^s, -28°22′13.4″). Details about the observations, data reduction, and method used to identify the detected lines and derive column densities can be found in Belloche et al. (2016). In short, the EMOCA survey covers the frequency range from 84.1 GHz to 114.4 GHz with a spectral resolution of 488.3 kHz (1.7 to 1.3 km s⁻¹). It was performed with five frequency tunings called S1 to S5. The median angular resolution is 1.6″.

3. Results and discussion

3.1. Rest frequencies and derived spectroscopic parameters

The transitions recorded in this study are summarized in Appendix B. From these measurements, and previously published data (Tanimoto et al. 1979; Tanimoto & Macdonald 1979), a fit to a Watson-*S* Hamiltonian in the *I'* representation was performed using the SPFIT/SPCAT suite of programs (Pickett 1991). The resulting best-fit spectroscopic constants are given in Table 1.

The large number of transitions recorded in this work coupled with the detection of lines that originate from much higher values of *J* and *K_a* compared to the previous studies enable centrifugal distortion (CD) parameters up to sixth order to be precisely determined. In some cases, the precision of these CD parameters has been improved by as much as four orders of magnitude. To reproduce the measured transitions to their experimental accuracy, a total of nine parameters are required, including two quartic CD terms (*D_K* and *d₂*) and one sextic term (*H_{KJ}*) that were not determined previously. The much larger dataset used in the present investigation and the relatively small number of additional parameters needed point to the rapid convergence of the Hamiltonian model and the robustness of the fit. On the basis of the best-fit parameters in Table 1, the astronomically most interesting lines either have been measured or can now be predicted throughout the entire millimeter domain to better than 50 kHz up to 250 GHz (0.06 km s⁻¹ in terms of radial velocity).

3.2. Poor predictive power of extrapolating high-frequency transitions from microwave measurements

While frequency predictions based on the spectroscopic parameters from Tanimoto et al. (1979) and Tanimoto & Macdonald

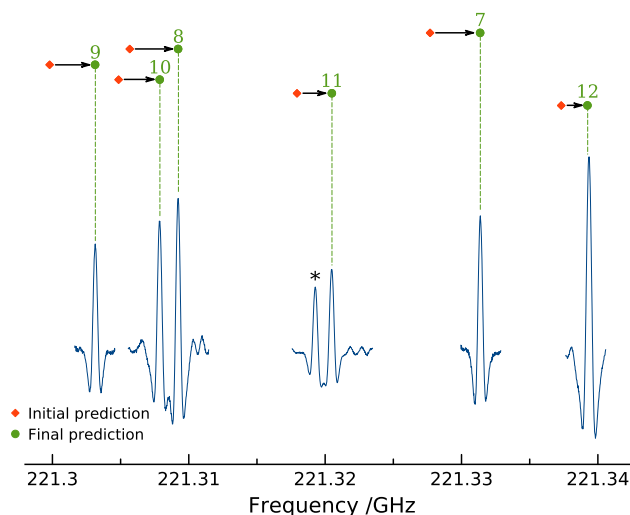


Fig. 5. Individual transitions measured in the ⁹R(19) branch of *syn* vinyl mercaptan and comparison between the initial (red diamonds) and final (green dots) predictions. The *K_a* value of each transition is indicated in green; the asymmetry splitting is not resolved for these lines. The transition labeled with an asterisk is unidentified. The line shape appears as a second derivative owing to the employed detection scheme.

(1979) have enabled many new lines to be assigned in the millimeter-wave band, this parameter set alone is inadequate to detect lines of vinyl mercaptan in astronomical sources. Many high-*J*, high-*K_a* transitions, for example, were shifted from their initial predictions by a few to up to tens of MHz; e.g., the transition 16_{3 14} – 16_{2 15} initially predicted at 226 581.2 MHz was measured at 226 532.7 MHz. In space, the most relevant transitions are the ⁹R series for the *syn* isomer ($\Delta J = 1, \Delta K_a = 0$) as these are the strongest lines at temperatures ranging from a few tens to a few hundreds of K. For these lines, the typical deviation from the initial prediction was in the range 2–5 MHz around 200 GHz (Fig. 5), corresponding to a significant fraction of the 2–10 km s⁻¹ linewidths that typify hot cores and corinos in star-forming regions. This example illustrates the pitfalls of extrapolating from the microwave to the millimeter and submillimeter domain, particularly for non-rigid molecules, and emphasizes the value of high-frequency laboratory measurements of astrophysically relevant species, even when microwave spectrum data are already available.

3.3. Isomers

The relative stability of isomers may be an important factor in predicting isomeric abundances in the ISM. With this motivation in mind, we performed high-level ab initio calculations (Fig. 2) to determine the relative stability of [C₂H₄S] isomers and the corresponding isovalent oxygen analogs; for the former, the calculations presented are – to the knowledge of the authors – the first report of the relative energetics at 0 K. Because our calculations for the [C₂H₄O] isomeric family are in excellent agreement with experiments, we can in turn be confident of the [C₂H₄S] thermochemistry, which is significantly less well constrained by measurements.

In contrast to the oxygen isomers, the four lowest single [C₂H₄S] isomers are effectively isoenergetic at our current level of theoretical sophistication. While it is unclear if a higher level of treatment (requiring sub-kJ/mol) changes the energetic ordering of these isomers, the near-degeneracy of [C₂H₄S] isomers

Table 2. Parameters of our best-fit LTE model (or upper limit) of selected complex organic molecules toward Sgr B2(N2).

Molecule	Status ^a	N_{det}^b	Size ^c ($''$)	T_{rot}^d (K)	N^e (cm^{-2})	C^f	ΔV^g (km s^{-1})	V_{off}^h (km s^{-1})	$\frac{N_{\text{ref}}}{N}^i$	Ref.
CH_3SH , $v = 0^*$	d	12	1.40	180	3.4 (17)	1.00	5.4	-0.5	1	2
<i>gauche</i> - $\text{C}_2\text{H}_5\text{SH}$	n	0	1.40	180	<1.6 (16)	1.95	5.4	-0.5	>22	2
<i>syn</i> - $\text{C}_2\text{H}_3\text{SH}$	n	0	1.40	180	<3.7 (16)	1.22	5.4	-0.5	>9	1
<i>anti</i> - $\text{C}_2\text{H}_3\text{SH}$	n	0	1.40	180	<1.6 (16)	1.63	5.4	-0.5	>21	1
CH_3CN , $v_8 = 1^*$	d	20	1.40	170	2.2 (18)	1.00	5.4	-0.5	1	3
$\text{C}_2\text{H}_5\text{CN}$, $v = 0$	d	154	1.20	150	6.2 (18)	1.38	5.0	-0.8	0.35	3
$\text{C}_2\text{H}_3\text{CN}$, $v = 0$	d	44	1.10	200	4.2 (17)	1.00	6.0	-0.6	5	3
CH_3OH , $v_t = 1^*$	d	16	1.40	160	4.0 (19)	1.00	5.4	-0.2	1	2
$\text{C}_2\text{H}_5\text{OH}$	d	168	1.25	150	2.0 (18)	1.24	5.4	0.0	20	2
<i>syn</i> - $\text{C}_2\text{H}_3\text{OH}$	n	0	1.25	150	<4.0 (16)	1.00	5.4	0.0	>1000	1
<i>anti</i> - $\text{C}_2\text{H}_3\text{OH}$	n	0	1.25	150	<1.2 (16)	1.00	5.4	0.0	>3333	1

Notes. ^(a)d: detection, n: non-detection. ^(b)Number of detected lines (conservative estimate, see Sect. 3 of Belloche et al. 2016). One line of a given species may mean a group of transitions of that species that are blended together. ^(c)Emission diameter (*FWHM*). ^(d)Rotational temperature. ^(e)Total column density of the molecule, except for the *syn* and *trans* conformers of vinyl mercaptan and vinyl alcohol, which are in each case considered as independent species. For these species, the column density represents the column density of each conformer separately. $X(Y)$ means $X \times 10^Y$. ^(f)Correction factor that was applied to the column density to account for the contribution of vibrationally or torsionally excited states or other conformers, in the cases where this contribution was not included in the partition function of the spectroscopic predictions. ^(g)Linewidth (*FWHM*). ^(h)Velocity offset with respect to the assumed systemic velocity of Sgr B2(N2), $V_{\text{lsr}} = 74 \text{ km s}^{-1}$. ⁽ⁱ⁾Column density ratio, with N_{ref} the column density of the previous reference species marked with a \star .

References. (1) This work, (2) Müller et al. (2016), (3) Belloche et al. (2016).

presents a fairly unique test bed to assess possible reaction pathways in astrochemical objects. The role of icy grains in $[\text{C}_2, \text{H}_4, \text{O}]$ formation has been suggested as the underlying factor as to why abundance ratios depart from thermodynamic equilibrium (Bennett et al. 2005). Having established that both isomers of vinyl mercaptan and thiirane are effectively isoenergetic, their relative interstellar abundance should present a more sensitive probe of local processes (e.g., shock and grain chemistry) than their oxygen analogs, which are more widely spaced in energy (Fig. 2). A systematic astronomical investigation of the $[\text{C}_2, \text{H}_4, \text{S}]$ family of isomers, however, is currently limited by the lack of reliable frequencies in the millimeter and submillimeter domains for thioacetaldehyde, which is the remaining isomer for which no data has presently been published beyond the microwave domain.

3.4. Astronomical searches

Searches for the *syn* and *anti* isomers of vinyl mercaptan were performed with the EMOCA spectrum of Sgr B2(N2) using the best-fit parameters in Table 1. Weeds (Maret et al. 2011) was used to produce synthetic spectra of both isomers under the local thermodynamic equilibrium (LTE) approximation, and the same parameters (emission size, rotational temperature, linewidth, and velocity offset with respect to the assumed systemic velocity of the source) as those derived for methyl mercaptan in Müller et al. (2016) were assumed. Hence, the only free parameters are the column densities of the *syn* and *anti* forms of vinyl mercaptan, which we consider as independent species.

No evidence was found for either isomer in the EMOCA spectrum. The upper limits of the column densities are reported in Table 2, along with the column densities or upper limits thereof derived for other molecules in our previous EMOCA studies (Müller et al. 2016; Belloche et al. 2016). In addition, no evidence was found for the *syn* and *anti* isomers of vinyl alcohol using predictions derived from the pure rotational works reported in Saito (1976) and Rodler (1985). The corresponding upper limits of their column densities are summarized in Table 2 as well. The non-detection of vinyl alco-

hol with ALMA along with its non-detection with the Institut de Radioastronomie Millimétrique (IRAM) 30 m telescope (Belloche et al. 2013) confirms that if vinyl alcohol were present in the Sgr B2(N) star-forming region at the level reported by Turner & Apponi (2001), whose observations with the Kitt Peak 12 m telescope did not resolve the multiple hot cores embedded in Sgr B2(N), then its emission must arise from the source envelope on large scales ($>1 \text{ pc}$) and not from the much more compact hot cores, as already stated by Belloche et al. (2013). Surveys like the Prebiotic Interstellar Molecular Survey (PRIMOS, Neill et al. 2012) or the survey performed at the Arizona Radio Observatory (ARO; Halfen et al. 2017) may shed more light on the presence and distribution of vinyl alcohol toward Sgr B2(N).

We find that the *syn* and *anti* isomers of vinyl mercaptan are at least 9 and 21 times less abundant than methyl mercaptan, respectively. The energy difference between the *syn* and *anti* isomers expressed in temperature units is about 60 K, which implies that the abundance ratio of the *anti* to *syn* isomers is 0.72 for a temperature of 180 K. This means that the total column density of vinyl mercaptan at this temperature is 1.72 times that of the *syn* isomer or equivalently 2.4 times that of the *anti* isomer. The most stringent constraint on the total column density of vinyl mercaptan is provided by the *anti* isomer: vinyl mercaptan is at least ~ 9 times ($21/2.4$) less abundant than methyl mercaptan.

The lower limit for the abundance ratio of vinyl to methyl mercaptan can be compared to ratios obtained for other families of molecules. It is nearly a factor of two higher than the abundance ratio measured for the cyanides ($[\text{CH}_3\text{CN}]/[\text{C}_2\text{H}_3\text{CN}] = 5$). For vinyl alcohol, the large energy difference between the *syn* and *anti* isomers (4.6 kJ mol^{-1} , i.e., 385 cm^{-1} , Fig. 2) implies that the total column density of vinyl alcohol is equal to 1.025 times that of the *syn* isomer at 150 K. We thus conclude that vinyl alcohol is at least 1000 times less abundant than methanol. This lower limit is more than two orders of magnitude higher than the ratio measured for the cyanides. Given that the abundance ratio of methanol to ethanol is also higher (by a factor ~ 60) than the abundance ratio of the corresponding cyanides, the chemistries of the cyanides and alcohols appear to differ significantly. However, the observational

constraints do not tell us whether the chemistries of the mercaptans and alcohols are similar or differ: the lower limit to the ratio of methyl to ethyl mercaptan is marginally consistent with the corresponding ratio measured for the alcohols, and the lower limits to the ratio of the methyl to vinyl species do not exclude a similar ratio in both families.

4. Conclusions

In the present study, a rotational investigation of the two isomers of vinyl mercaptan has been extended up to 250 GHz. The new set of transition frequencies and derived spectroscopic constants provide a highly reliable basis for subsequent searches for both species in the ISM. Calculations of the rotational spectra are available in the catalog section of the CDMS (Endres et al. 2016) database² while linelists, parameters, fits, and auxiliary files are available in the data section³.

The work performed in this study highlights the urgent need for high-frequency measurements of numerous species of interstellar interest that so far have only been characterized at centimeter wavelengths. Indeed, most of the current ISM observations are now performed in the millimeter and submillimeter domains, using facilities such as ALMA, NOEMA, APEX, and SOFIA, but frequency extrapolations into the millimeter domain using spectroscopic constants derived from low J and K_a observations are inherently uncertain, and make detection of new astronomical species unduly challenging. Additionally, this work provides constraints on isomeric populations in the ISM. Both *syn* and *anti* vinyl mercaptan have been the subjects of searches in the EMOCA survey recorded toward the hot core Sgr B2(N2). No transitions of either species has been detected and upper limits for their column densities have been derived, revealing that vinyl mercaptan is at least nine times less abundant than methyl mercaptan in this source.

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² <https://cdms.astro.uni-koeln.de/classic/entries/>

³ <https://cdms.astro.uni-koeln.de/classic/predictions/daten/VySH/>

Appendix A: Molecular structures of the [C₂H₄S] isomers

Internal coordinates ae-CCSD(T)/cc-pwCVQZ

syn vinyl mercaptan

H
 C 1 r1
 C 2 r2 1 a1
 S 3 r3 2 a2 1 d0
 H 4 r4 3 a3 2 d0
 H 2 r5 3 a4 1 d180
 H 3 r6 2 a5 6 d0

r1 = 1.08115
 r2 = 1.33270
 a1 = 122.05669
 r3 = 1.75221
 a2 = 127.33692
 d0 = 0.00000
 r4 = 1.33663
 a3 = 96.56932
 r5 = 1.07962
 a4 = 120.07569
 d180 = 180.00000
 r6 = 1.08206
 a5 = 120.75551

anti vinyl mercaptan

H
 C 1 r1
 C 2 r2 1 a1
 S 3 r3 2 a2 1 d1
 H 4 r4 3 a3 2 d2
 H 2 r5 3 a4 1 d3
 H 3 r6 2 a5 6 d4

r1 = 1.08105
 r2 = 1.33250
 a1 = 121.79666
 r3 = 1.75849
 a2 = 122.52864
 d1 = 3.65449
 r4 = 1.33431
 a3 = 96.59150
 d2 = -159.42502
 r5 = 1.07959
 a4 = 120.21684
 d3 = -179.52134

r6 = 1.08069
 a5 = 121.24236
 d4 = 0.62115

thiirane

S
 C 1 r1
 C 1 r1 1 a1
 H 2 r2 1 a2 3 d1
 H 2 r2 1 a2 3 md1
 H 3 r2 1 a2 2 d1
 H 3 r2 1 a2 2 md1

r1 = 1.81067
 a1 = 48.27245
 r2 = 1.07991
 a2 = 115.09725
 d1 = 110.93875
 md1 = -110.93875

thioacetaldehyde

S
 C 1 r1
 C 2 r2 1 a1
 H 3 r3 2 a2 1 d0
 H 2 r4 3 a3 1 d180
 H 3 r5 2 a4 5 d1
 H 3 r5 2 a4 5 md1

r1 = 1.61419
 r2 = 1.49172
 a1 = 125.71288
 r3 = 1.08597
 a2 = 111.24050
 d0 = 0.00000
 r4 = 1.08887
 a3 = 115.25385
 d180 = 180.00000
 r5 = 1.09181
 a4 = 109.69777
 d1 = 58.47364
 md1 = -58.47364

Appendix B: Observed experimental transitions of vinyl mercaptan

Tables B.1 and B.2 are available at the CDS.