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Millimeter-wave spectroscopy and modeling of 1,2-butanediol[★]

Laboratory spectrum in the 59.6–103.6 GHz region and comparison with the ALMA archived observations

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ABSTRACT

Context. The continuously enhanced sensitivity of radioastronomical observations allows the detection of increasingly complex organic molecules. These systems often exist in a large number of isomers leading to very congested spectra.

Aims. We explore the conformational space of 1,2-butanediol and provide sets of spectroscopic parameters to facilitate searches for this molecule at millimeter wavelengths.

Methods. We recorded the rotational spectrum of 1,2-butanediol in the 59.6–103.6 GHz frequency region (5.03–2.89 mm) using a free-jet millimeter-wave absorption spectrometer, and we analyzed the properties of 24 isomers with quantum chemical calculations. Selected measured transition lines were then searched on publicly available ALMA Band 3 data on IRAS 16293-2422 B.

Results. We assigned the spectra of six conformers, namely aG'Ag, gG'Aa, g'G'Ag, aG'G'g, aG'Gg, and g'GAa, to yield the rotational constants and centrifugal distortion constants up to the fourth or sixth order. The most intense signal belong to the aG'Ag species, that is the global minimum. Search for the corresponding $30_{x,30}-29_{x,29}$ transition lines toward IRAS 16293-2422 B was unsuccessful.

Conclusions. Our present data will be helpful for identifying 1,2-butanediol at millimeter wavelengths with radio telescope arrays. Among all possible conformers, first searches should be focused on the aG'Ag conformers in the 400–800 GHz frequency spectral range.

Key words. molecular data – line: identification – methods: laboratory: molecular – methods: data analysis – techniques: spectroscopic – radio lines: ISM

1. Introduction

Millimeter and submillimeter astronomy is a fundamental technique for searching complex organic molecules (COMs) in circumstellar envelopes and interstellar medium (Brünken & Schlemmer 2015). However laboratory spectra acquisition, assignment, and analysis are necessary for the correct identification of the features contained in ongoing spectral surveys.

Among all the detected molecules, diols (organic compounds with two hydroxyl groups) are an object of chemical interest because of their similarity with important biological building block molecules such as sugar alcohols. The prototype of these species, ethylene glycol, is one of the largest COMs detected in space so far, as attested by the Cologne Database for Molecular

Spectroscopy (CDMS 2001; Müller et al. 2001, 2005). Lines attributable to the most stable conformer of ethylene glycol were detected in a variety of astronomical environments, such as the Galactic center, hot cores, hot corinos, and comets (Hollis et al. 2002; Requena-Torres et al. 2008; Maury et al. 2014; Coutens et al. 2015; Jørgensen et al. 2016; Crovisier et al. 2004; Biver et al. 2014; Fuente et al. 2014; Brouillet et al. 2015), and tentatively, toward the two high-mass star-forming regions W51/e2 (Kalenskii & Johansson 2010; Lykke et al. 2015) and G34.3+0.2 (Lykke et al. 2015).

It is worth noting that recently a second conformer of ethylene glycol, lying at about 2.5 kJ mol^{-1} above the lowest energy form (Müller & Christen 2004), has also been observed both toward IRAS 16293-2422 (Jørgensen et al. 2016) and the Orion Kleinmann–Low nebula (Favre et al. 2017).

Regarding larger diols, observation searches for 1,2-propanediol (Lovas et al. 2009) and 1,3-propanediol (Plusquellic et al. 2009) toward Sgr B2 (N-LMH) were made as part of the Green Bank Telescope (GBT) Prebiotic Interstellar Molecule Survey Legacy Project (PRIMOS 2013; Remijan et al. 2013), but no transitions were detected beyond the 1σ root mean square noise limit.

* The measured and predicted transition lines for six conformers of 1,2-butanediol 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/619/A140>

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The decrease of column densities of molecules with increasing molecular weight is one of the main effects that hinder the detection of larger diols in interstellar space. Nevertheless, thanks to the enhanced sensitivity of the latest radioastronomy observations, their future detection cannot be excluded. On the other hand, the increasing size of the alkyl chain confers a molecular flexibility that is often characterized by a great conformational complexity and is not easy to solve.

In this context, we report a detailed conformational analysis and the laboratory millimeter spectrum of 1,2-butanediol (12BD, hereafter), the ethyl substituted form of ethylene glycol. The high level of flexibility of 12BD requests a nontrivial conformational study (see Sect. 2) in order to understand which are the more abundant species or which provide the most intense spectroscopic signals. The laboratory extraction and modeling of the rotational spectrum are described in Sects. 3 and 4 and the resulting prediction are described in Sect. 5. Finally in Sect. 6, the laboratory investigation has been followed by a tentative search in astronomical observation data extracted from the Atacama Large Millimeter/submillimeter Array (ALMA) Science Archive.

2. Conformational analysis

Except for 12BD, rotational spectroscopy investigations are reported for all linear diols containing up to four carbon atoms as follows: ethylene glycol or 1,2-ethanediol (12ED; Marstokk & Møllendal 1974; Walder et al. 1980; Christen et al. 1995, 2001; Christen & Müller 2003; Müller & Christen 2004), 1,2-propanediol (12PD; Caminati 1981; Lockley et al. 2002; Lovas et al. 2009; Bossa et al. 2014), 1,3-propanediol (13PD; Caminati et al. 1995; Plusquellic et al. 2009), 1,3-butanediol (13BD; Caminati & Corbelli 1982; Velino et al. 2011), 2,3-butanediol (23BD; Paul et al. 2007), and 1,4-butanediol (14BD; Evangelisti et al. 2013).

Even though diols may exist in several distinct conformations, the more stable structures are those stabilized by the intramolecular hydrogen bond between the two hydroxyl groups ($\text{OH}\cdots\text{OH}$). Actually, only this family of conformers was detected by laboratory rotational spectroscopy.

Concerning diols with hydroxyl groups at the outer carbon atoms such as 12ED, 13PD, and 14BD (Fig. 1, first row), two nonequivalent structures were observed, each of these corresponding to four equivalent conformers. The rotational spectra are complicated by a strong rotation-tunneling interaction depending on the interconversion energy barriers between the equivalent species.

In contrast, in the cases of 12PD, 12BD, and 13BD (Fig. 1, second row) the conformers are not equivalent because of the presence of an outer methyl group. Seven and five species were observed for 12PD and 13BD, respectively. It can be also pointed out that in these species the carbon atom binding four different substituents is a chiral center giving rise to the specular and isoenergetic *R* and *S* configurations. However, these configurations can be considered spectroscopically equivalent in conventional rotational experiments.

In 23BD (Fig. 1, third row), because of the presence of two chiral carbon atoms, four isomers exist; these are (*R,R*) and specular (*S,S*), (*R,S*) and specular (*S,R*). Among the possible conformations, only one species was observed for each pair, and a tunneling motion due to the interchange of the hydrogen bonded hydroxyl groups was observed for (*R,R/S,S*).

The molecular structure of 12BD is defined by the HO-CC hydroxyl dihedral angles (τ_1 and τ_4 , Fig. 1) and the OC-CO and CC-CC skeletal torsional angles (τ_2 and τ_3 , respectively). The

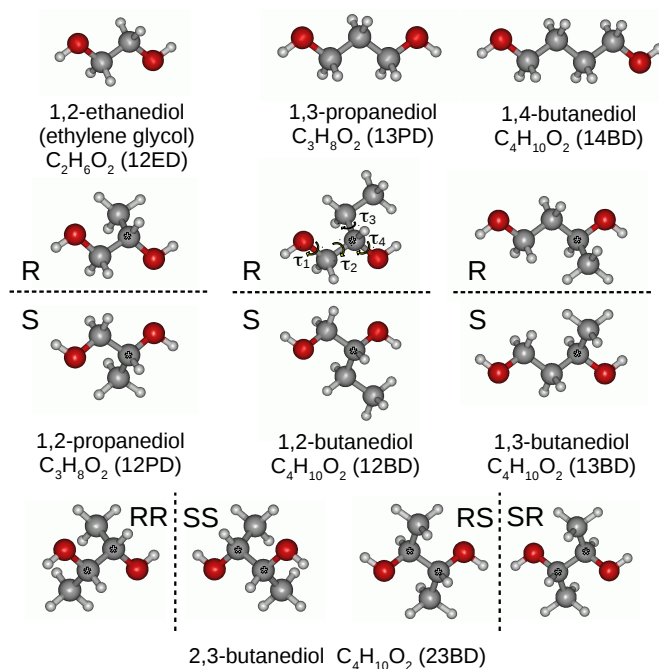


Fig. 1. Diols up to 4 carbon atoms. Gray, white, and red spheres represent the carbon (C), hydrogen (H), and oxygen (O) atoms, respectively. Asterisks indicate chiral atoms. The dashed line indicates the mirror plane between the chiral forms. The labeling of dihedral angles used in the text is shown.

methyl group internal rotation is not relevant because it leads to equivalent minima. Because of steric hindrance, for each τ_i three staggered orientations are possible, giving rise to $3^4 = 81$ possible nonequivalent conformers. The labeling chosen for the identification of these conformers reflects the τ_1 , τ_2 , τ_3 , and τ_4 values, respectively, using a sequence of the four letters xYYy (X, Y = A, G, G'; x, y = a, g, g'). These letters indicate the three staggered positions of the four dihedral angles as follows: *anti* ($\tau \sim 180^\circ$), *gauche* ($\tau \sim 60^\circ$), or *gauche'* ($\tau \sim -60^\circ$). Capital and lower case letters refer to the backbone and hydroxyl torsions, respectively.

As regards the chirality, to simplify the discussion, we refer only to the *R* configuration. The corresponding *S* configuration is achieved changing the sign of all dihedral angles.

Assuming that 12BD behaves like the already studied diols, the most stable forms are characterized by the hydrogen bond between the hydroxyl groups. This kind of interaction can take place in 24 of the 81 possible conformers as follows: aGYg', gGYg', aGYg', gGYg', g'G'Yg, aG'Yg, gG'Ya, and gG'Yg' with Y = A, G, G'. Their structures were optimized at the B3LYP-D3/aug-cc-pVTZ level of calculation using the Gaussian 09 quantum chemistry package (Frisch et al. 2013) and subsequent harmonic vibrational frequency calculations confirmed that the optimized geometries correspond to real minima in the conformational potential energy surface.

The B3LYP density functional is one of the most widely used for the quantum mechanical prediction of molecular properties. Including the empirical dispersion correction (D3; Grimme et al. 2010), and using a very extended basis set (aug-cc-pVTZ) should ensure reliable modeling of the intramolecular hydrogen bond interaction and, consequently, an accurate description of the relative energies and structures of the conformers.

The 24 hydrogen-bonded conformers, shown in Fig. 2 with their zero-point corrected relative energies, were considered in

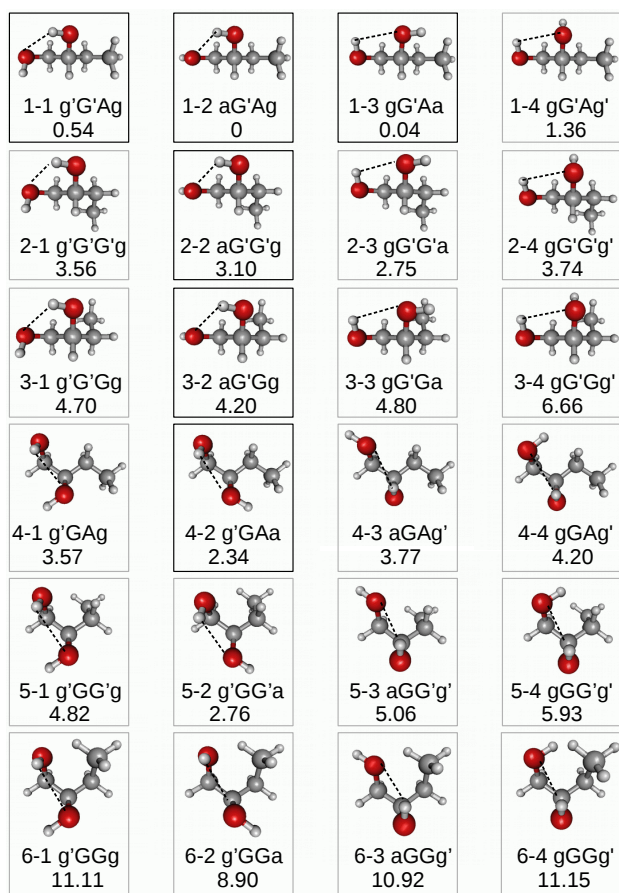


Fig. 2. Theoretical (B3LYP-D3/aug-cc-pVTZ) structures and zero-point corrected relative electronic energy values ($\Delta E_0/\text{kJ mol}^{-1}$) of the 24 hydrogen-bonded conformers of *R*-12BD. Gray, white, and red spheres represent the carbon (C), hydrogen (H) and oxygen (O) atoms, respectively. Black frames indicate the observed species.

the analysis of the rotational spectrum. For the sake of clarity, besides the label, a numerical code is also introduced. It is composed of two digits, referring to the skeletal and hydroxyl groups configurations, respectively.

The theoretical relative energy values of 20 conformers lie below 7 kJ mol^{-1} , suggesting that at room temperature the conformational population is spread on a large number of species. This behavior, common in flexible molecules, complicates both the detection and analysis of the spectra. In the present work, such a difficulty has been overcome exploiting the rotational and conformational cooling produced by a supersonic expansion of the sample seeded in monoatomic gases.

3. Experiment

12BD (purity 98%, molecular weight $90.121 \text{ g mol}^{-1}$) was purchased from Sigma-Aldrich and used without any further purification. It appears as a colorless, viscous liquid at ambient conditions. The melting point is -50°C (223 K) and the boiling point is $191\text{--}192^\circ\text{C}$ (464–465 K). Both argon and helium, purchased from SIAD (Società Italiana Acetilene e Derivati), were used as carrier gases.

The sample was heated to about 80°C (353 K) and a stream of the carrier gas ($P_0(\text{Ar}) = 20 \text{ kPa}$, $P_0(\text{He}) = 40 \text{ kPa}$) was flowed over the sample and then expanded to about $P_b = 0.5 \text{ Pa}$ through a heated 0.3 mm diameter pinhole nozzle. The data reported by

Steele et al. (1996) and Verevkin (2004) were used to extrapolate the 12BD vapour pressure at the working temperature: 0.545 kPa at 353 K . In this way the concentration of 12BD was estimated to be around 2.7% and 1.4% in argon and helium, respectively.

Rotational spectra in the millimeter-wave region ($5.03\text{--}2.89 \text{ mm}$, $59.6\text{--}103.6 \text{ GHz}$) were recorded using a Stark-modulated free-jet absorption spectrometer. The spectrometer has a resolution of about 300 kHz and an estimated accuracy of about 50 kHz . With respect to the ordinary experimental setup described in Melandri et al. (1995, 1996) and Calabrese et al. (2015), some modifications on the multipliers chain were performed to obtain frequencies in the *W* band, more precisely, the same *K*-band frequency generator equipment was driving a $\times 4$ frequency multiplier, followed by a dial attenuator. The lens limitation over 73 GHz and the over-moded driven waveguide was systematically checked and no power loss was attested up to 103 GHz .

4. Rotational spectrum of 1,2-butanediol

To acquire an overview of the 12BD rotational spectrum, two fast scans covering the $59.6\text{--}74.4 \text{ GHz}$ frequency range were run using both helium and argon as carrier gas. The spectrum recorded with helium shows a much larger number of lines with respect to the spectrum recorded in argon, suggesting that the population of the sample is constituted by several conformers and that collisional cooling favors the overcoming of the conformational potential energy barriers, giving rise to effective relaxation processes in the expansion in argon.

Predictions based on the theoretical rotational constants and electric dipole moment components reported in Table 1, allowed for the assignment of the most intense spectral lines of six different species. Then, substituting the predictions with the experimental transitions, other frequency assignments could be made until the signal-to-noise ratio was too low to measure with reasonable certainty. In this iterative process, the need for centrifugal distortion constants was checked and they were inserted as long as substantial reduction of the rms error was obtained and until their values could be determined with sufficient precision.

The assigned frequencies and corresponding rotational quantum numbers are available in the supplementary material, as outlined in the appendix. For all conformers, a distribution plot of the upper state J and K_a values of all measured transitions included in the least-squares fit is shown in Fig. 3. The CALPGM suite of programs (Pickett 1991) was used to fit the data and predict the rotational spectra, applying Watson's S reduction and the I' representation of the rotational Hamiltonian (Watson 1977).

The final sets of spectroscopic parameters (rotational and centrifugal distortion constants) are reported in Table 2 with a rough indication of the intensity ratio of the μ_b -type to μ_c -type transitions, which form a quartet of lines.

Based on the different types of rotational transitions observed for each species, theoretical relative energies, and comparison between experimental and theoretical spectroscopic constants, the assigned spectra could be uniquely correlated to the conformers aG'Ag, g'G'Ag, gG'Aa, aG'Gg, aG'G'g, and g'GAa.

The most intense spectral lines, both in helium and argon expansions, belong to the *R*-branch μ_b and μ_c degenerate transitions of conformer aG'Ag, indicating that it is the global minimum. This is in agreement both with the quantum mechanical results and experimental findings on related molecules 12ED and 12PD, for which the global minima have the same heavy atoms arrangement (aG'g).

Table 1. Theoretical results for the 24 hydrogen bonded conformers of 1,2-butanediol, obtained at the B3LYP-D3/aug-cc-pVTZ level of calculation.

Conf.		E_e	ΔE_e	E_0	ΔE_0	A_e	B_e	C_e	μ_a	μ_b	μ_c	μ
		Hartree	kJ mol^{-1}	Hartree	kJ mol^{-1}	MHz	MHz	MHz	D	D	D	D
1-1	g'G'Ag	-309.028967	0.22	-308.887333	0.54	7812.70	1921.23	1671.84	-0.21	1.68	-1.44	2.22
1-2	aG'Ag	-309.029044	0.01	-308.887539	0.00	7811.31	1937.00	1681.80	1.28	1.85	-0.44	2.29
1-3	gG'Aa	-309.029049	0.00	-308.887525	0.04	7673.24	1929.04	1666.98	-2.59	0.19	-0.36	2.62
1-4	gG'Ag'	-309.028531	1.36	-308.886956	1.53	7575.92	1928.69	1670.44	-2.00	-0.56	1.44	2.53
2-1	g'G'G'g	-309.027943	2.90	-308.886183	3.56	5462.10	2246.60	1722.97	0.72	1.87	-1.17	2.33
2-2	aG'G'g	-309.028000	2.75	-308.886357	3.10	5473.98	2266.50	1730.92	0.17	2.27	-0.71	2.38
2-3	gG'G'a	-309.028166	2.32	-308.886490	2.75	5380.29	2270.34	1723.68	-2.33	1.29	-0.46	2.71
2-4	gG'G'g'	-309.027882	3.06	-308.886114	3.74	5339.64	2275.19	1725.25	-2.17	0.55	1.38	2.63
3-1	g'G'Gg	-309.027612	3.77	-308.885747	4.70	5943.35	2179.45	1916.68	-0.31	1.24	-1.72	2.15
3-2	aG'Gg	-309.027670	3.62	-308.885939	4.20	5989.98	2190.10	1918.80	-0.75	2.16	0.20	2.30
3-3	gG'Ga	-309.027468	4.15	-308.885711	4.80	5860.98	2187.51	1905.08	-2.19	0.90	-0.86	2.52
3-4	gG'G'g'	-309.026817	5.86	-308.885003	6.66	5857.83	2172.14	1893.94	-2.50	-0.49	0.63	2.62
4-1	g'GAa	-309.027808	3.26	-308.886180	3.57	5923.82	2207.55	1929.32	-0.52	-1.19	1.71	2.15
4-2	g'GAa'	-309.028170	2.31	-308.886649	2.34	5910.35	2207.26	1926.18	-2.25	-0.48	0.83	2.44
4-3	aGAg'	-309.027642	3.69	-308.886103	3.77	5944.60	2210.23	1936.49	1.71	-1.42	-0.79	2.36
4-4	gGAg'	-309.027706	3.52	-308.885939	4.20	5968.54	2181.83	1912.71	0.32	-2.32	0.35	2.37
5-1	g'GG'g	-309.027548	3.94	-308.885701	4.82	4289.99	2766.95	2038.48	-0.10	-1.07	1.72	2.03
5-2	g'GG'a	-309.028261	2.07	-308.886489	2.76	4344.00	2735.29	2027.60	-1.61	-1.64	0.38	2.34
5-3	aGG'g'	-309.027233	4.77	-308.885610	5.06	4333.14	2736.56	2033.05	-0.18	-2.17	0.72	2.29
5-4	gGG'g'	-309.027064	5.21	-308.885278	5.93	4342.28	2696.48	2008.62	1.85	-1.70	-0.07	2.51
6-1	g'GGg	-309.025186	10.14	-308.883306	11.11	4147.98	2944.79	2376.10	-0.24	-1.89	-1.05	2.18
6-2	g'GGa	-309.025791	8.55	-308.884149	8.90	4139.47	2955.26	2368.80	1.68	-1.59	-0.73	2.43
6-3	aGGg'	-309.025068	10.45	-308.883380	10.92	4166.68	2925.26	2365.31	1.47	-1.43	-1.33	2.45
6-4	aGGg'	-309.025192	10.12	-308.883293	11.15	4181.25	2915.52	2343.73	1.07	1.68	-1.37	2.42

Notes. Columns are: electronic energy values and corresponding zero-point vibration corrected values, equilibrium rotational constants, and electric dipole moment components.

Weaker lines were then assigned to the aG'Gg, aG'G'g conformers, which differ for the orientation of the alkyl tail (τ_3) and the g'GAa conformer, which presents a different OCCO orientation (τ_2). This latter conformation can be considered the homologue of g'Ga-12ED and g'Ga-12PD. The lines belonging to these three less populated conformers appear slightly depleted in argon expansion, indicating that a partial relaxation to the global minimum takes place.

As remarked above, the rotational spectrum recorded using helium as carrier gas, showed additional lines that were assigned to the g'G'Ag and gG'Aa conformers. A clear example of this is reported in Fig. 4, where a portion of the spectra collected using helium and argon are compared. In this frequency region lie the $K_a = 6 \leftarrow 5$ Q -branch transitions of both the global (aG'Ag) and local (g'G'Ag) minima and the Q branch of the less stable conformer (g'G'Ag) appears absolutely depleted in the argon spectra. The reason for this is that, considering similar expansion velocities for both carrier gases, the higher mass of argon produces a greater kinetic energy, which entails at the ability to overcome the conformational barriers.

Since the g'G'Ag and gG'Aa conformers differ from the global minimum only for the orientation of the hydroxyl group, the depletion of their lines in argon expansion can be ascribed to a strong relaxation process along the hydroxyl internal rotation path. According to Ruoff et al. (1990) the upper limit of the corresponding barriers is 4–5 kJ mol^{-1} . It is worth noting that the hydrogen bond is maintained both in the g'G'Ag \rightarrow aG'Ag and gG'Aa \rightarrow aG'Ag isomerization processes.

Relative intensity measurements on the observed Q -branch bands match with a rotational temperature of 3 K in argon expansion and 5 K when using helium as carrier gas, confirming the greater cooling efficiency of argon with respect to helium.

The line intensity ratio of g'G'Ag to aG'Ag conformers is about 1:3. This value, weighted on the theoretical $\mu_b^2 + \mu_c^2$ values (Table 1), leads to a g'G'Ag:aG'Ag = 1:4 population ratio. Assuming that this population ratio is not modified by the expansion, and it is the ratio established at the pre-expansion equilibrium temperature, the corresponding relative energy is 3.9 kJ mol^{-1} .

Regarding the gG'Aa conformer, the intensity of the μ_a -type transition lines are similar to those of aG'Ag. Unfortunately, owing to an incomplete Stark-modulation of the spectral lines, it is not possible to determine a reliable intensity ratio. Nevertheless, considering that the μ_a ratio is about 2, we can state that the gG'Aa conformer is less populated than the aG'Ag conformer.

Finally, no internal rotation splittings due to the terminal methyl group were resolved in the spectra. The predictions based on the experimental rotational constants and theoretical geometries indicate that, in the investigated spectral region, these splittings could be appreciable mainly on some Q -type transitions, expecting a separation of 0.4 MHz between the lines. Since this value lies within the resolution limit of our spectrometer, the measured transitions appear affected in the aG'Ag and g'G'Ag fits by a high deviation.

5. Prediction of the rotational spectrum

To provide useful data for the detection of 12BD in space, the rotational spectra of the six observed conformers were predicted. The results, based on the experimental spectroscopic constants (Table 2) and theoretical electric dipole moment components (Table 1), are given in the supplementary material, as outlined in the appendix. The prediction sample comprises lines that

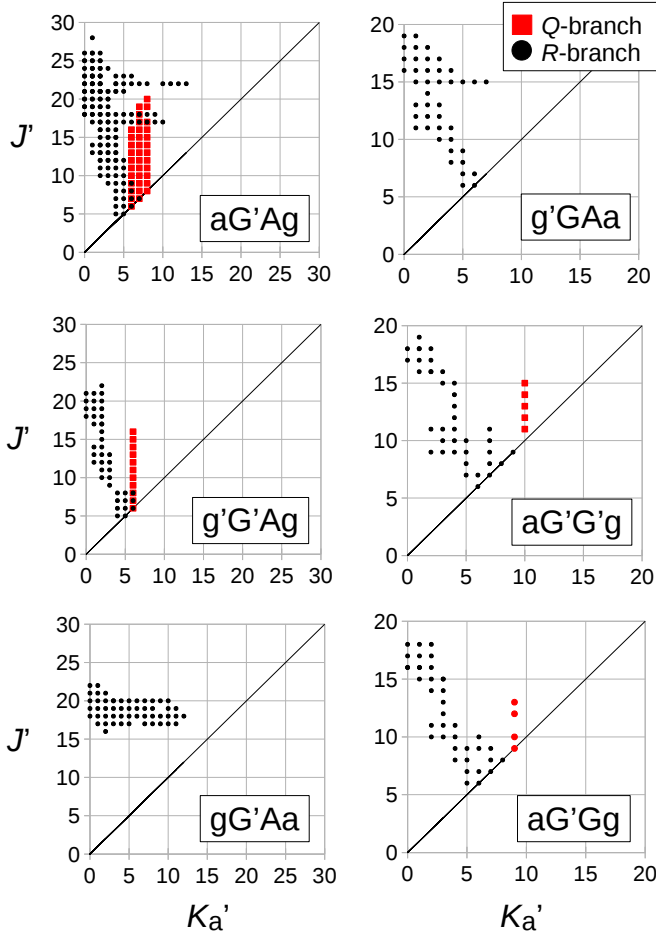


Fig. 3. Data set distribution plots of J' and K'_a of all measured transitions for the spectrum of 1,2-butanediol.

have upper state energy $E_{\text{up}} < 100$ K, and Einstein's coefficient $A > 10^{-7} \text{ s}^{-1}$ evaluated as $A = (16\pi^3\nu^3)/(3\epsilon_0hc^3)S_{ij}\mu_g^2$.

The overall shape of the predicted spontaneous emission probability of aG'Ag-12BD up to 1 THz is illustrated on the top of Fig. 5. The maximum emission is expected between 500 and 800 GHz. In Fig. 5 also the predicted absorption spectra at 5, 75, and 300 K are compared. At these temperatures, the maximum absorption intensities lie in the 50–100, 200–400, and 400–700 GHz frequency ranges, respectively. The partition function of the six observed conformers of 12BD in the 5–300 K temperature range, given in Table 3, allows for the evaluation of the absorption coefficient.

6. Search on ALMA data archives

IRAS 16293-2422 is a well-studied Class 0 protostar consisting of two main sources denoted as components A and B. This source is a prototypical hot corino and it therefore has a rich molecular line spectrum for a low-mass protostar. Previous interferometric studies (Bottinelli et al. 2004; Kuan et al. 2004) showed that the spectra toward the A component have broad lines (FWHMs up to 8 km s^{-1}), whereas the lines toward the B component are much narrower (typically less than 2 km s^{-1}), which makes it an ideal source for line identifications.

Actually, the rich spectrum of this source has made it possible to identify several COMs. Observations are reported for molecules containing up to four heavy atoms, such as ethyl

Table 2. Experimental spectroscopic parameters for the six observed conformers of 1,2-butanediol.

	aG'Ag 1–2	g'G'Ag 1–1	gG'Aa 1–3
A/MHz	7830.1015 (12) ^a	7821.8336 (50)	7694.37 (13)
B/MHz	1945.15516 (38)	1930.6686 (18)	1937.5445 (46)
C/MHz	1687.57121 (38)	1678.8135 (20)	1673.3921 (45)
D_J/kHz	0.16080 (41)	0.1632 (27)	0.1558 (21)
D_{JK}/kHz	1.9773 (25)	1.9263 (15)	1.8899 (49)
D_K/kHz	8.253 (16)	7.80 (12)	– ^b
d_1/Hz	–18.13 (15)	–17.30 (39)	–18.9(30)
d_2/Hz	–4.06 (13)	–3.26 (34)	2.72(83)
N^c	77	192	70
RMS^d/kHz	54	62	58
a/b/c ^e	y/y/y	y/y/y	y/n/n
μ_b/μ_c^f	2.1(4)	1.1(1)	–
	aG'G'g 2–3	aG'Gg 3–2	g'GAa 4–2
A/MHz	5509.3759 (53)	5966.8787 (56)	5961.4560 (44)
B/MHz	2276.8059 (36)	2207.1497 (25)	2211.2757 (20)
C/MHz	1739.5251 (40)	1936.0736 (27)	1937.8078 (21)
D_J/kHz	0.4874 (86)	0.4438 (49)	0.4597 (40)
D_{JK}/kHz	–0.508 (75)	0.386 (28)	–
D_K/kHz	9.276 (89)	7.53 (11)	7.16 (10)
d_1/Hz	–170.5 (25)	–57.60 (95)	–40.79 (87)
d_2/Hz	–16.85 (94)	6.37 (68)	3.76 (53)
H_{JK}/Hz	1.96 (24)	–	–
H_K/Hz	2.19 (45)	–3.12 (77)	–
N^c	66	57	62
RMS^d/kHz	51	52	72
a/b/c ^e	n/y/y	y/y/n	y/y/y
μ_b/μ_c^f	1.6 (2)	–	0.6 (1)

Notes. (a) Standard error in parentheses in the units of the last digit. (b) Fixed to zero in the fit. (c) Number of transitions. (d) Root mean square deviation of the fit. (e) Yes (y) or no (n) observation of μ_a -, μ_b -, and μ_c -type transitions, respectively. (f) Electric dipole moment components ratio estimated from the relative intensity of μ_b -type and μ_c -type transition quartets.

cyanide ($\text{C}_3\text{H}_5\text{N}$), methyl formate, acetic acid (Cazaux et al. 2003) and glycolaldehyde (Jørgensen et al. 2012; isomers with formula $\text{C}_2\text{H}_4\text{O}_2$), acetone and propanal (isomers with formula $\text{C}_3\text{H}_6\text{O}$), and even two conformers of ethylene glycol ($\text{C}_2\text{H}_6\text{O}_2$; Jørgensen et al. 2016).

Based on these considerations, IRAS 16293-2422 can be considered an adequate source for a search of 12BD. Several observations on this source were found in the public data of the ALMA Science Archive. Among these, project 2012.1.00712.S was selected because the frequency coverage (89.49–89.72, 92.78–93.01, 102.49–102.72, 103.18–103.41 GHz) and the spectral resolution (60 kHz channel spacing) are similar to those of our experiments.

The reduction of the data proceeded according to standard recipes exploiting the Common Astronomical Software Applications package (v4.2.1; McMullin et al. 2015). Assuming a better chance to find 12BD where the analog 12ED is present, the spectrum of the source was extracted within one $1'' \times 1''$ synthesized beam size, centered on the 102.690 GHz peak of 12ED in component B (RA = $16^{\text{h}}32^{\text{m}}22.^{\text{s}}612$; Dec = $-24^{\circ}28'32''.588$). Subsequently, each spectral window was analyzed to identify the less crowded regions of the spectrum, best suited for the search of 12BD.

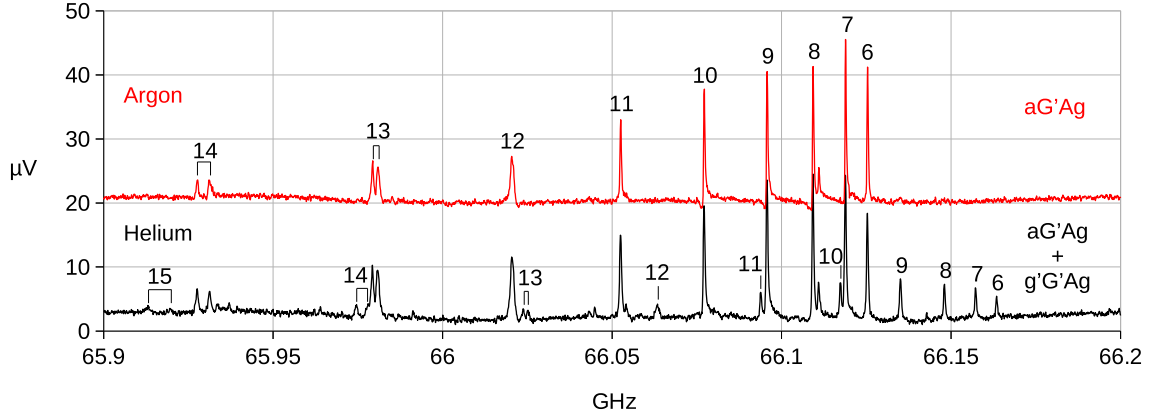


Fig. 4. Comparison between the 65.9–66.2 GHz spectra recorded using Ar (upper black red) and He (lower black trace) as carrier gas. In both spectra the aG'Ag-12BD $K_a = 6 \leftarrow 5$ Q-branch transitions can be recognized, whereas the same transitions of less stable g'G'Ag-12BD are completely depleted in the Ar spectrum. Numbers above the peaks indicate the assigned J quantum numbers.

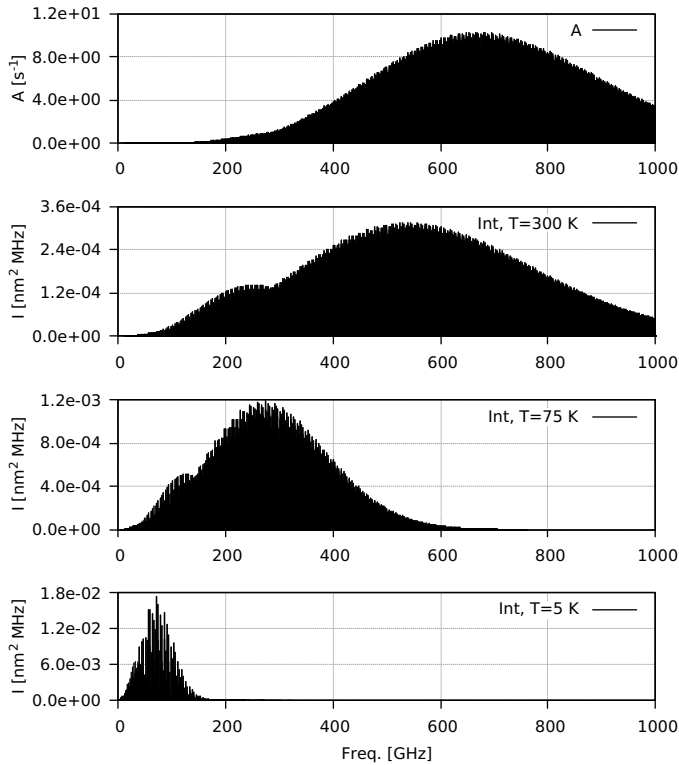


Fig. 5. Simulation of emission (top side) and absorption rotational spectra of aG'Ag-12BD. Intensities are computed at 5, 75, and 300 K.

Regarding the most stable conformer of 12BD, the predicted transition lines in the available spectral windows are listed in Table A.2. Among these, a quartet of lines constituted by the $30_{x,30} - 29_{x,29} \mu_b$ and μ_c R-type transitions appear promising to search. Indeed this quartet fulfills several requirements: (i) the value of the predicted A coefficient is high; (ii) the shape, constituted by four peaks in few MHz, characterized by a specific pattern of separation is easily recognizable; (iii) the spectrum appears relatively free of contaminating lines; and (iv) the frequency range includes that spanned by the laboratory measurements.

The predicted lines are superimposed on the cleaned spectrum in Fig. 6. Two rotational transition lines belonging to methylformate (CH_3OCHO) and methylacetylene (CH_3CCH),

Table 3. Partition function of the six observed conformers of 1,2-butanediol in the 5–300 K temperature range.

$T(\text{K})$	Q_{1-2}	Q_{1-1}	Q_{1-3}	Q_{2-3}	Q_{3-2}	Q_{4-2}
300	172 785	173 976	175 383	187 529	173 480	173 319
150	61 089	61 510	62 007	66 302	61 334	61 277
75	21 598	21 747	21 923	23 441	21 685	21 665
50	11 757	11 838	11 933	12 760	11 804	11 793
25	4157	4185	4219	4511	4173	4169
10	1052	1059	1067	1141	1056	1055
5	372	374	377	403	373	373

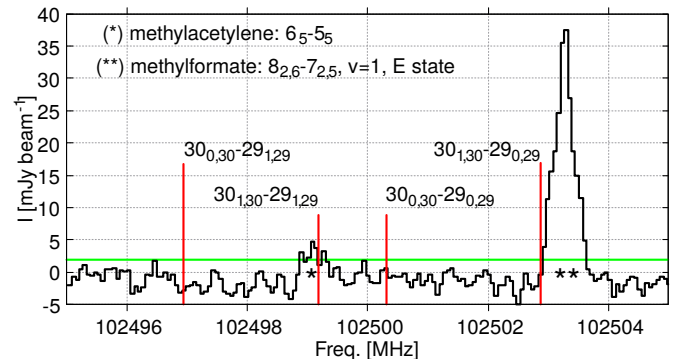


Fig. 6. Spectrum in the central beam toward the continuum peaks of IRAS 16293-2422 B. Predicted $30_{x,30} - 29_{x,29}$ transition lines of aG'Ag-12BD are overlapped as red sticks. The X-axis represents the frequencies in the rest frame of the system (i.e., corrected for the system V_{LSR} of 2.7 km s^{-1} ; Jørgensen et al. 2011). The green line is an indication of the rms level ($1.89 \text{ mJy beam}^{-1}$) represented by a spectrum extracted from an off source position.

respectively, are clearly recognizable, while a lack of detection is observed for 12BD.

We modeled the spectrum extracted from the ALMA data using MCWeeds (Giannetti et al. 2017) to obtain an upper limit for the 12BD column density. Assuming a temperature of 300 K and a line width of 1 km s^{-1} , as measured for ethylene glycol (Jørgensen et al. 2016), we estimate $N < 1 \times 10^{13} \text{ cm}^{-2}$ (95% credible interval).

However, searching of 12BD features could be successful at higher frequency spectral ranges. In fact, considering a temperature of the source around 300 K (Jørgensen et al. 2011) the most

populated rotational levels with the highest possible Einstein A coefficients lie in the 400–800 GHz frequency range (Fig. 5), corresponding to bands 8–9.

Searches for upper bands in available databases have not been performed yet, since additional frequency laboratory measurements are needed to fix the higher order centrifugal distortion constants required for reliable spectral predictions.

7. Conclusions

In this work, features of millimetre rotational spectroscopy coupled to supersonic expansion have been exploited to analyze, for the first time, the spectrum of 12BD. Actually, at room temperature, the conformational population of this highly flexible diol is distributed on a large numbers of conformations practically preventing the assignment. The spectra recorded in the 59.6–103.6 GHz frequency range have allowed the unequivocal identification of six different species, determining that the aG'Ag conformer is that with the minimum energy. Besides the rotational constants, centrifugal distortion constants were also determined: in particular, four quartic constants for gG'Aa and g'GAa, the full quartic set for aG'Ag and g'G'Ag, and additional sextic contributions for aG'Gg and aG'G'g.

The experimental spectroscopic constants and theoretical electric dipole moment components were used to predict with high precision the rotational spectrum of each of the six observed conformers up to 163 GHz based on the measurements done. Selected lines of the global minimum aG'Ag-12BD were searched toward the IRAS 16293-2422 B source in the band 3 observations of the ALMA project 2012.1.00712.S. No detection was found, but this could be justified by the fact that the maximum spectral signal of this kind of heavy compounds is predicted at higher frequencies (bands 8–9). The ALMA project, from which the data were extracted, was in fact, not specifically aimed at searching this molecule. Our predictions may allow now us to define observational ALMA campaigns dedicated to 12BD.

Hence, on the basis of our predictions, additional laboratory measurements at submillimeter wavelengths can be performed and new specific observations in COMs-reach sources could be attempted.

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Appendix A: Supplementary material

The newly recorded experimental transition frequencies for the six observed conformers of 1,2-butanediol are available at the CDS. Only the first 30 lines measured for the aG'Ag conformer of 1,2-butanediol appear here (Table A.1). The table gives the rotational quantum numbers J , K_a , and K_c for the upper state followed by those for the lower state. The observed transition frequency is given in megahertz units with the residual between observed frequency and that calculated from the final set of spectroscopic parameters and its uncertainty.

In addition also the predicted rotational spectra in the 0–163 GHz frequency range are also available at the CDS. Only a portion relative to the aG'Ag conformer is given below (Table A.2). The data listing includes the following: the predicted rest frequencies and 1σ uncertainties, the upper state degeneracy and energy, the line strength factors $S_{ij}\mu_g^2$, and the Einstein's A coefficients for spontaneous emission. The prediction sample comprises lines, which have upper state energy $E_{\text{up}} < 100$ K, and Einstein's coefficient $A > 10^{-7} \text{ s}^{-1}$.

Table A.1. Selected measured rest frequencies, residuals between observed frequency and that calculated from the final set of spectroscopic parameters and experimental uncertainties for the ground vibrational state of the aG'Ag (1-2) conformer of 1,2-butanediol.

n.	J'	K_a'	K_c'	J	K_a	K_c	Rest freq. (MHz)	O-C (MHz)	Unc. (MHz)
(1)	(2)	(3)	(4)	(5)	(6)	(7)	(8)	(9)	(10)
1	16	2	14	15	2	13	59 930.23	0.01	0.05
2	20	2	18	19	3	17	59 976.02	0.00	0.05
3	10	2	9	9	1	9	60 144.66	0.00	0.05
4	12	2	10	11	1	10	60 205.72	-0.02	0.05
5	11	3	8	11	0	11	60 207.34	0.01	0.05
6	11	3	8	11	0	11	60 207.34	0.01	0.05
7	5	4	1	4	3	2	60 261.44	0.06	0.05
8	5	4	2	4	3	2	60 261.44	0.09	0.05
9	9	2	7	8	0	8	60 506.70	-0.02	0.05
10	15	2	14	14	1	13	60 646.78	0.01	0.05
11	17	2	16	16	2	15	60 737.95	0.00	0.05
12	18	1	17	17	2	16	61 051.59	0.02	0.05
13	17	1	16	16	1	15	61 624.47	0.02	0.05
14	9	3	7	8	2	6	61 680.90	-0.01	0.05
15	18	0	18	17	1	17	61 817.80	-0.01	0.05
16	9	3	6	8	2	6	61 831.62	0.00	0.05
17	17	10	8	16	10	7	61 838.33	-0.04	0.05
18	17	10	7	16	10	6	61 838.33	-0.04	0.05
19	17	9	9	16	9	8	61 855.90	-0.03	0.05
20	17	9	8	16	9	7	61 855.90	-0.03	0.05
21	17	8	10	16	8	9	61 879.95	-0.04	0.05
22	17	8	9	16	8	8	61 879.95	-0.04	0.05
23	17	7	11	16	7	10	61 914.86	0.07	0.05
24	17	7	10	16	7	9	61 914.86	0.06	0.05
25	17	3	15	16	3	14	61 918.42	0.05	0.05
26	17	6	12	16	6	11	61 968.64	0.14	0.05
27	17	6	11	16	6	10	61 968.64	-0.26	0.05
28	18	1	18	17	1	17	61 984.63	-0.03	0.05
29	18	0	18	17	0	17	62 048.18	0.04	0.05
30	17	5	13	16	5	12	62 054.77	0.03	0.05

Notes. (1) Listing number n. (2–4) Upper state rotational quantum numbers $J'_{K_a',K_c'}$. (5–7) Lower state rotational quantum numbers J_{K_a,K_c} . (8) Observed transition frequency. (9) Residual between observed frequency and that calculated from the final set of spectroscopic parameters O-C. (10) Experimental uncertainty Unc. The full Table A.1 is available at the CDS.

Table A.2. Selected predicted rest frequencies, estimated accuracies, and lines strengths for the ground vibrational state of the aG'Ag (1-2) conformer of 1,2-butanediol.

J'	K_a'	K_c'	J	K_a	K_c	Rest freq. (MHz)	Error (MHz)	G_{up}	E_{up} (K)	$S\mu_g^2$ (D^2)	A (s^{-1})
(1)	(2)	(3)	(4)	(5)	(6)	(7)	(8)	(9)	(10)	(11)	(12)
32	4	28	32	3	30	89 492.71	0.27	65	98.520	3.889E-01	3.245E-06
17	6	12	18	3	15	89 506.84	0.02	35	37.117	1.586E-02	1.324E-07
24	8	17	24	7	17	89 533.89	0.02	49	70.881	1.991E+00	1.664E-05
24	8	16	24	7	17	89 533.95	0.02	49	70.881	4.081E+01	3.410E-04
18	5	14	18	3	15	89 534.42	0.02	37	37.118	3.602E-02	3.010E-07
24	8	17	24	7	18	89 535.92	0.02	49	70.881	4.081E+01	3.410E-04
24	8	16	24	7	18	89 535.98	0.02	49	70.881	1.991E+00	1.664E-05
21	4	17	21	2	20	89 584.59	0.04	43	45.211	3.765E-02	3.151E-07
28	7	21	29	4	26	89 589.38	0.09	57	85.174	8.523E-02	7.135E-07
23	8	16	23	7	16	89 632.77	0.02	47	66.681	1.902E+00	1.595E-05
23	8	15	23	7	16	89 632.80	0.02	47	66.681	3.863E+01	3.238E-04
23	8	16	23	7	17	89 633.89	0.02	47	66.681	3.863E+01	3.238E-04
23	8	15	23	7	17	89 633.92	0.02	47	66.681	1.902E+00	1.595E-05
22	8	15	22	7	15	89 719.41	0.02	45	62.658	1.811E+00	1.523E-05
22	8	14	22	7	15	89 719.42	0.02	45	62.658	3.645E+01	3.065E-04
22	8	15	22	7	16	89 720.01	0.02	45	62.658	3.645E+01	3.065E-04
22	8	14	22	7	16	89 720.02	0.02	45	62.658	1.811E+00	1.523E-05
18	6	12	19	3	17	92 767.31	0.03	37	40.267	1.926E-02	1.790E-07
27	7	20	28	4	25	92 974.40	0.07	55	80.261	7.105E-02	6.648E-07
30	0	30	29	1	29	102 496.62	0.03	61	77.081	9.391E+01	1.177E-03
12	4	9	12	1	12	102 496.95	0.02	25	18.248	3.639E-02	4.563E-07
30	1	30	29	1	29	102 499.18	0.03	61	77.081	4.899E+01	6.142E-04
30	0	30	29	0	29	102 500.31	0.03	61	77.080	4.899E+01	6.142E-04
30	1	30	29	0	29	102 502.87	0.03	61	77.081	9.392E+01	1.178E-03
28	6	23	27	6	22	102 591.50	0.02	57	81.506	4.376E+01	5.502E-04
28	6	22	27	6	21	102 688.25	0.02	57	81.520	4.377E+01	5.518E-04
28	5	24	27	5	23	102 719.83	0.02	57	78.429	4.440E+01	5.602E-04
18	4	15	17	3	14	103 260.20	0.01	37	34.554	1.807E+01	2.316E-04
24	3	22	24	0	24	103 267.79	0.10	49	54.948	4.944E-02	6.338E-07

Notes. (1–3) Upper state rotational quantum numbers $J'_{K_a',K_c'}$. (4–6) Lower state rotational quantum numbers J_{K_a,K_c} . (7) Rest frequency evaluated from the spectroscopic constants. (8) Estimated error of the prediction at 1σ level. (9) Upper state degeneracy. (10) Upper state energy. (11) Transition strength $S\mu_g^2$ ($g = a, b, c$ theoretical electric dipole moment component). (12) Einstein's A coefficient. The full Table A.2 is available at the CDS.