

Laboratory and space spectroscopy of DCO⁺

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Abstract. The rotational spectra of DCO⁺ and its two isotopomers D¹³CO⁺ and DC¹⁸O⁺, produced in a negative glow discharge cell, have been recorded in the 137–792 GHz region, which includes lines from $J = 2 \leftarrow 1$ up to $J = 11 \leftarrow 10$. The determined rotational and centrifugal distortion constants allow to predict the DCO⁺ rotational spectrum up to 1000 GHz with an accuracy of 1 part in 10⁸ or better. This is important for kinematic studies of dense molecular cloud cores and for future far-infrared observations. We also report on the first detection of the hyperfine structure of the DCO⁺(1–0) line made at the IRAM 30 m antenna, toward the quiescent starless cloud core L1512, in the Taurus Molecular Cloud. We point out that this is the *first observation of the hyperfine splitting due to the deuteron*. This allowed us to quantify the effects of the hyperfine splitting on the line width determination; if the hyperfine structure is not taken into account in the line fit, the DCO⁺(1–0) line width is overestimated by a significant factor (≥ 2).

Key words. molecular data – methods: laboratory – ISM: individual objects: L1512 – ISM: molecules – radio lines: ISM

1. Introduction

In the past few years it has been realized how important is to know with high precision the frequencies of the molecular transitions used to investigate the physics and the chemistry of interstellar clouds and star forming regions (e.g. Mardones et al. 1997; Lee et al. 1999, 2001). This is because spectral line observations are unique tools to study internal motions of dense molecular cloud material, within which stars will be or are forming. In this sense, molecular line observations are crucial to test current models of cloud core and star formation and give insights on molecular cloud evolution.

In particular, before low-mass stars form, the progenitor (or starless) core is typically a cold (~ 10 K) and quiescent region, where the (optically thin) line widths are slightly (approximately a factor of 2) larger than the thermal value Δv_T (e.g. Caselli et al. 2002). This means that $\Delta v_{\text{obs}} \approx 2 \times \Delta v_T \approx 0.3 (T/10 \text{ K})^{1/2} (A_X/30 \text{ amu})^{-1/2} \text{ km s}^{-1}$, where T is the gas temperature and A_X is the molecular weight. This already shows that to determine the LSR velocity of a cold ($T = 10$ K) cloud with an accuracy comparable to the standard deviation of the corresponding Gaussian line profile ($\sigma_{V_{\text{LSR}}} = \Delta v_{\text{obs}} / \sqrt{8 \ln(2)} \lesssim 0.1 \text{ km s}^{-1}$), for a molecular species with $A_X \sim 30$ amu (such as CO, HCO⁺, and DCO⁺), the observed line frequency should have an uncertainty of $\Delta \nu (\text{kHz}) \lesssim 0.1 \times 3.3 \times \nu_0 (\text{GHz})$ (where ν_0 is the rest frequency), or about 30 kHz at millimeter wavelengths.

If observed lines are optically thick and if they are tracing inward motions, they appear broader and asymmetric, typically double peaked, with the blue peak stronger than the red peak (e.g. Leung & Brown 1977; Zhou et al. 1990). However, to quantify the infall velocity one needs to observe an optically thin tracer and compare the thin and thick profiles (e.g. Myers et al. 1996). For this purpose, one typically uses the normalized velocity difference between the thin and thick lines (δV , as defined by Mardones et al. 1997). In this case, the condition $\delta V / \sigma_{\delta V} > 3$ is only obtained if the error on V is $\sigma_V \lesssim 0.01 \text{ km s}^{-1}$ (assuming average values of $\delta V \sim 0.2$, $\sigma_{\Delta V} \sim 0.03 \text{ km s}^{-1}$, and $\Delta V \approx 0.3 \text{ km s}^{-1}$; see Mardones et al. 1997). This implies a frequency precision of $\sigma_\nu (\text{kHz}) \lesssim 0.03 \times \nu_0 (\text{GHz})$, or about 3 kHz at ~ 100 GHz. All this demonstrates the need of laboratory work to accurately measure line frequencies. Gottlieb et al. (2003) have also pointed out that it is a challenge to the spectroscopist the determination of line frequencies with a precision large enough to identify motions in molecular cloud cores from tiny velocity shifts, which typically require a precision of a few parts in 10⁸. In the case of DCO⁺ this is equivalent to 1.7% of the Doppler line width at 77 K.

DCO⁺ is a particularly suitable species to study the chemistry, in particular the deuterium fractionation (e.g. Saito et al. 2002) and the electron fraction (e.g. Guélin et al. 1977; Caselli et al. 1998; Williams et al. 1998), and the kinematics of the dense cores. The lower rotational transitions are easily detected, as it is proved by the large amount of literature on

Table 1. Observed rotational transition of DCO⁺ (MHz).

J'	J	Bogey et al. (1981)	Sastry et al. (1981)	Diff. ^a /ppb	This work	Radiastronomy ^b
2	1	144 077.342	144 077.319	160	144 077.2890	144 077.321
3	2	216 112.701	216 112.604	449	216 112.5822	
4	3	288 144.018	288 143.911	371	288 143.8583	288 143.911
5	4	360 169.980	360 169.881	275	360 169.7783	360 169.830
6	5		432 189.033		432 189.0052	

^a Difference between measurements of Bogey et al. (1981) and of Sastry et al. (1981).

^b Lovas (2004).

this molecular ion, from interstellar clouds (e.g. Wootten et al. 1982) to dense cloud cores (e.g. Butner et al. 1995), to circumstellar disks (van Dishoeck et al. 2003). DCO⁺ has been recently surveyed in a large sample of starless cores by Lee et al. (2004), with the aim of detecting infall motions. However, the previous measurements of DCO⁺ millimeter-wave frequencies, carried out in two different laboratories (Bogey et al. 1981; Sastry et al. 1981), gave results differing up to 45 part in 10⁸ for the $J = 3 \leftarrow 2$ line (see Table 1); therefore there is a need for more accurate frequency measurements of DCO⁺ lines in order to use them in the study of star-forming molecular cloud cores. In this paper, we repeat with far better precision the old measurements of millimeter-wave transitions of DCO⁺, D¹³CO⁺, and DC¹⁸O⁺ (Bogey et al. 1981), and we add new measurements of submillimeter-wave transitions: the spectroscopic constants derived from the analysis of the frequency data enable to predict spectra up to 1000 GHz with an accuracy of at least 1 part in 10⁸ for DCO⁺, 5 parts in 10⁸ for D¹³CO⁺, and of 1 part in 10⁷ for DC¹⁸O⁺; all the isotopomers have been observed in natural abundance.

2. Observations

2.1. Laboratory measurements

The laboratory spectrum was observed with a frequency-modulated millimeter-wave spectrometer (Cazzoli & Dore 1990a) equipped with a double-pass negative glow discharge cell (Dore et al. 1999) made of a Pyrex tube 3.25 m long and 5 cm in diameter. The radiation source was a frequency multiplier driven by a Gunn diode oscillator working in the region 68–76 GHz (Farran Technology Limited) for the lines up to $J = 4 \leftarrow 3$; for the high frequency transitions, a doubler in cascade with a multiplier (RPG – Radiometer Physics GmbH) was driven by Gunn oscillators working in the region 75–115 GHz (J. E. Carlstrom Co and RPG) to cover frequencies up to 792 GHz. Two phase-lock loops allow the stabilization of the Gunn oscillator with respect to a frequency synthesizer, which is driven by a 5-MHz rubidium frequency standard. The frequency modulation of the radiation is obtained by sine-wave modulating with low distortion (total harmonic distortion less than 0.01%) the reference signal of the wide-band Gunn-synchronizer. The signal, detected by a liquid-helium-cooled InSb hot electron bolometer (QMC Instr. Ltd. type QFI/2), is demodulated at 2- f by a lock-in amplifier.

DCO⁺ was produced by flowing a 1:1 mixture of CO₂ and D₂ (1 mTorr) in Ar buffer gas with a total pressure of about 5 mTorr and discharging with a current of a few mA; D¹³CO⁺ and DC¹⁸O⁺ were observed in natural abundance. The cell was cooled at 77 K by liquid nitrogen circulation, and an axial magnetic field up to about 400 G was applied throughout the length of the discharge. With this longitudinal magnetic field applied, ions are produced and observed in the negative glow (De Lucia et al. 1982), which is a nearly field free region and where they are expected to show negligible Doppler shift due to the drift velocity, occurring, instead, in the positive column (Sastry et al. 1981) where a low axial electric field is present. In addition, the double-pass arrangement would compensate for such a shift, if present.

A typical spectrum is recorded by sweeping the frequency up and down (several times if signal averaging is needed) in steps of 5 kHz at a rate of 0.8 MHz s⁻¹, with a lock-in amplifier time constant of 3 ms and a frequency modulation depth comparable to the Doppler width or larger for the rarer isotopomers. Since we have full flexibility in controlling scanning rate, number of data points and modulation depth, the values of these parameters have been adjusted to prevent any bias of the measured transition frequency, which is recovered from a line shape analysis of the spectral profile (Cazzoli & Dore 1990a; Dore 2003).

2.2. DCO⁺(1–0) toward the quiescent cloud core L1512

The DCO⁺(1–0) spectrum toward the quiescent Taurus starless core L1512 (Fig. 1) has been obtained with the IRAM–30 m antenna, located at Pico Veleta (Spain) in August 2004. The adopted coordinates were RA(1950) = 05^h00^m54.4^s, Dec(1950) = 32°39'00.0". We used the recent extension of the 3 mm tuning range below 80 GHz and carried out the observations using the frequency switching technique with a throw of 3.9 MHz. The spectral resolution is 3.3 kHz, which corresponds to 14 m/s at 72 GHz. Given that below ~75 GHz the image gain depends strongly on the tuning parameters and often varies across the bandpass of the receiver, we measured the sideband gain ratio and corrected the observed spectrum accordingly. The pointing was checked every 2 h and found to be accurate within about 10" because of anomalous

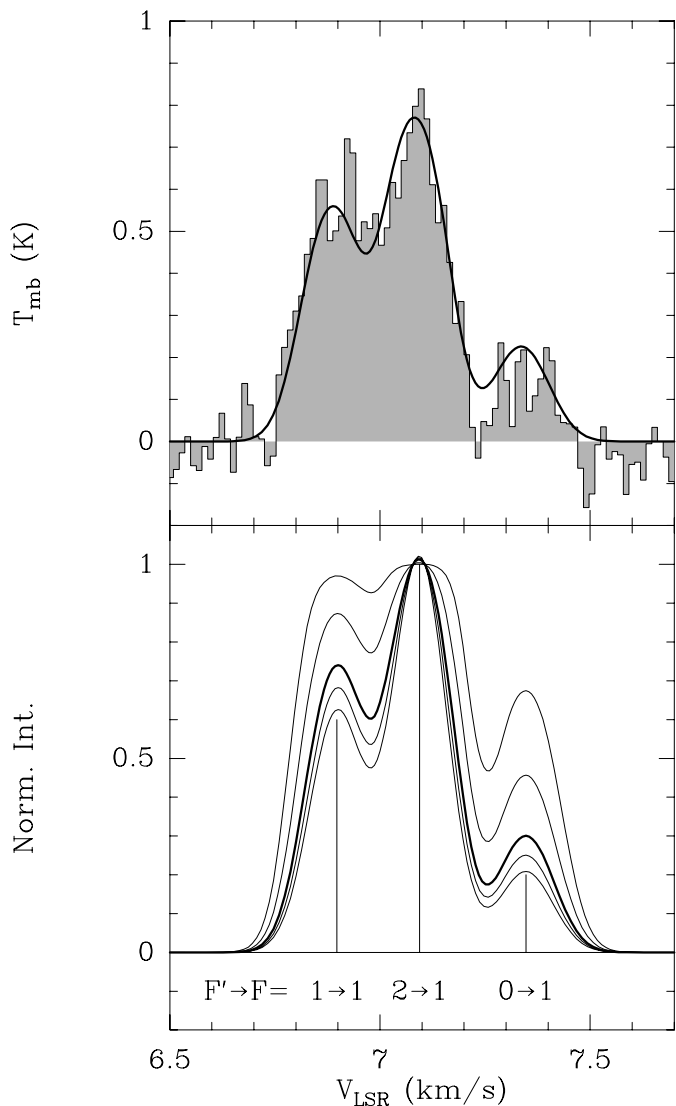


Fig. 1. *Top panel:* observed spectrum of DCO⁺(1–0) toward the quiescent starless cloud core L1512, in the Taurus Molecular Cloud. The units are in main beam brightness temperature, assuming a unity filling factor. This is the first spectrum showing the hyperfine structure of the line due to the deuteron electric quadrupole interaction. The line is significantly broadened by the hyperfine structure, and the width is a factor of 2.3 larger if the splitting is not taken into account in the fit. The black curve is the hfs fit performed in CLASS. *Bottom panel:* simulated spectrum of the DCO⁺(1–0) line assuming the same parameters of the observed line ($T_{\text{ex}} = 4$ K and intrinsic $\Delta v = 0.15$ km s^{−1}) but with different values of τ_{TOT} (from least to most intense curves: 0.1, 1, 2 (as observed), 5, and 10). Vertical bars indicate the positions of the three hyperfines with their relative intensities in the optically thin limit.

refraction problems during very good atmospheric conditions. The half power beam width (HPBW) was 33". The units are main beam brightness temperature, assuming a source filling factor of unity. The beam and forward efficiencies were 0.79 and 0.95, respectively. The rms noise of the spectrum is 0.09 K.

Figure 1 clearly shows that the DCO⁺(1–0) line profile is complex and contains structure. Indeed, the three detected features are consistent with the hyperfine splitting due to the

interaction between the molecular electric field gradient and the electric quadrupole moment of the deuteron (spin = 1), which produces the line splitting into three components. The velocity shifts predicted by theory are larger than the observed separations between the three components by about 0.1 km s^{−1}, suggesting that the quadrupole constant needs to be refined. Thus, we estimated the separations performing a Gaussian fit to the three components of the DCO⁺(1–0) spectrum. This allowed us to derive the velocity separations of -0.198 ± 0.012 km s^{−1} and 0.256 ± 0.017 km s^{−1} between the main ($F', F = 2, 1$) component and the low ($F', F = 1, 1$) and high ($F', F = 0, 1$) velocity components, respectively. With this new values of the frequency shifts and the relative intensities of the three components (3/9, 5/9, 1/9 for the (1, 1), (2, 1) and (0, 1) hyperfines, respectively) we then performed an hfs fit using the IRAM reduction package CLASS, shown in Fig. 1 (upper panel) by the thick curve. Forcing the resulting LSR velocity (V_{LSR}) to coincide with that measured with the high sensitivity N₂H⁺(1–0) spectrum reported in Caselli et al. (1995), which has been corrected to account for the new N₂H⁺(1–0) frequency value (93176.2608 MHz \pm 6 kHz for the component $F_1, F = 0, 1 \rightarrow 1, 2$) estimated by Dore et al. (2004), we obtain the frequency values reported in Table 2. These new frequencies have finally been used to refine the quadrupolar constant (see next section).

The hfs fit in CLASS gives the following parameters: $V_{\text{LSR}} = 7.094 \pm 0.004$ km s^{−1}, intrinsic (i.e. corrected for optical depth effects) line width $\Delta v = 0.149 \pm 0.008$ km s^{−1}, total optical depth (i.e. the sum of the optical depths of the three hyperfines) $\tau_{\text{TOT}} = 1.9 \pm 0.5$, excitation temperature $T_{\text{ex}} = 4 \pm 1$ K. We note here that the observed line width is only 1.25 times larger than the DCO⁺ thermal width Δv_{T} , assuming a kinetic temperature of 10 K (or $\Delta v = \Delta v_{\text{T}}$ if $T = 15$ K). If the hyperfine structure is not taken into account, the DCO⁺(1–0) line width obtained with a simple Gaussian fit is a factor of 2.3 times larger, in net contrast with the narrow (~ 0.18 km s^{−1}) N₂H⁺(1–0) line width observed in the same position (Caselli et al. 1995). Therefore, we conclude that the differences in the widths of DCO⁺(1–0) and NH₃(1, 1) lines observed in previous work (e.g. compare Butner et al. 1995; with Benson & Myers 1989) are likely due to neglecting the hyperfine splitting. The bottom panel of Fig. 1 shows how the line profile changes in case of large optical depths. For $\tau_{\text{TOT}} = 10$, the weakest ($F', F = 0, 1$) component becomes bright enough to further enlarge the line width by other ~ 0.25 km s^{−1} (so that the total linewidth will be about 0.6 km s^{−1}, if the splitting is not considered in the fit).

The effects of the hyperfine structure on the line width and profile become less important for higher J transitions. In Fig. 2, the simulated spectrum of the DCO⁺(2–1) line, with the same parameters as the DCO⁺(1–0) line in Fig. 1, is shown. The six hyperfine components are blended together, but there is still some ($\sim 20\%$) extra-broadening and a small (0.01 km s^{−1}) line center shift resulting from the hfs splitting, which should be taken into account when analysing DCO⁺(2–1) profiles. In particular, this may be important to quantify the infall velocity in the central regions of starless cores, as recently done by Lee et al. (2004). On the other hand, the components of the $J = 3 \rightarrow 2$ transition (as well as those of higher J transitions)

Table 2. Rotational transition frequencies and spectroscopic constants of DCO⁺.

$J'(F')$	$J(F)$	Observed/MHz	Obs.-calc./kHz	Uncertainties/kHz ^a
1(0)	0(1)	72 039.2413	-0.1	7.3
1(2)	0(1)	72 039.3028	-0.3	6.1
1(1)	0(1)	72 039.3504	-0.2	6.7
2	1	144 077.2890 ^b	3.9	5
3	2	216 112.5822	2.1	5
4	3	288 143.8583	0.5	5
5	4	360 169.7783	-1.2	5
6	5	432 189.0052	-1.3	5
7	6	504 200.1999	-0.5	5
8	7	576 202.0239	1.1	5
9	8	648 193.1357	0.0	5
10	9	720 172.2024	1.1	10
11	10	792 137.8811	-1.2	10
$\sigma_{\text{fit}}^c/\text{kHz}$ 1.3				
Constant ^d	This work	Previous ^e	DC ¹⁵ N	
B/MHz rotational	36 019.76765(14)	36 019.784(2)		
D_J/kHz quartic c. d. ^f	55.7960(22)	55.71(12)	54.3924(15) ^g	
H_J/Hz sextic c. d. ^f	0.054(11)		0.0633(12) ^g	
eqQ/kHz quadrupole	147.8(35)		200.6(5) ^h	
C_I/kHz spin rotation	-1.59(78)		-1.9(1) ^h	

^a Uncertainties estimated as explained in the text.

^b Not included in the fit, see text.

^c $(\sum(\text{obs.}-\text{calc.})^2/\text{degrees of freedom})^{1/2}$.

^d Standard errors are reported in parentheses in units of the last quoted digits.

^e Bogey et al. (1981).

^f Centrifugal distortion.

^g Brünken et al. (2004).

^h Cazzoli & Dore (1990b).

are heavily blended together, and the line width measurement is not affected at all by their presence.

3. Analysis and discussion

The measured transition frequencies of DCO⁺, D¹³CO⁺, and DC¹⁸O⁺ are listed in Tables 2–4; they are mean values obtained from 5 to 14 measurements. The standard errors of the mean result unrealistically small (for DCO⁺ they range from 0.1 to 1 kHz), therefore the uncertainties reported in the last column of each table were estimated from the ratio of linewidth to signal-to-noise ratio, and were roughly scaled according to the standard errors.

The experimental data were fitted, in a weighted least-squares procedure, to the standard expression for the frequency of the rotational transition $J + 1 \leftarrow J$:

$$\nu_0 = 2B(J + 1) - 4D_J(J + 1)^3 + H_J(J + 1)^3 \left[(J + 1)^3 - J^3 \right], \quad (1)$$

where B is the rotational constant and D_J and H_J are the quartic and sextic centrifugal distortion constants, respectively; the weights were the inverse-square of the uncertainties. The standard deviation σ_{fit} of the fit for each isotopomer is just a few kHz with the inclusion of H_J as fit parameter. An assessment of

the significance of the latter can be carried out by comparison with DCN centrifugal distortion constants: the values (Brünken et al. 2004) reported in Tables 2 and 3 result comparable with the values of D_J and H_J derived in this work for DCO⁺ and D¹³CO⁺, therefore these constants may be assumed correctly determined.

In the case of DCO⁺, the three hyperfine frequencies of the $J = 1 \rightarrow 0$ transition, accurately determined as reported above, could be used to derive the quadrupole coupling (eqQ) and spin rotation (C_I) constants of the D nucleus (spin quantum number $I = 1$) according to the expression:

$$\nu_{hf} = \nu_0 - eqQ[Y(J + 1, I, F') - Y(J, I, F)] + C_I[C(J + 1, I, F') - C(J, I, F)], \quad (2)$$

where F is the total angular momentum ($F = J + I$) quantum number, $C(J, I, F) = F(F + 1) - I(I + 1) - J(J + 1)$ and the Casimir function $Y(J, I, F)$ is given by:

$$Y(J, I, F) = \frac{\frac{3}{4}C(C + 1) - I(I + 1)J(J + 1)}{2(2I - 1)(2J + 3)I(2I - 1)}. \quad (3)$$

The Pickett's SPFIT fitting program (Pickett 1991) allowed us to carry out simultaneously the hfs and centrifugal analyses in

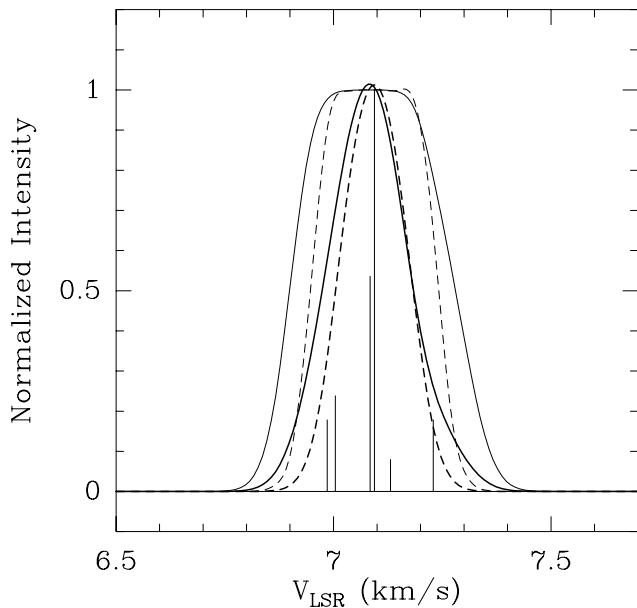


Fig. 2. Simulated spectrum of the $J = 2 \rightarrow 1$ transition of DCO⁺ in L1512, including the hyperfine splitting due to the deuteron electric quadrupole interaction (black curves). The assumed excitation temperature is 4 K and the intrinsic linewidth is 0.15 km s⁻¹, as found for the DCO⁺(1–0) line, whereas the total optical depth is 1 (thick curve) and 10 (thin curve). Dashed curves are the expected DCO⁺(2–1) lines in case of no hyperfine structure, again assuming $\tau_{\text{TOT}} = 1$ (thick dashed curve) and 10 (thin dashed curve). Vertical lines mark the velocities of the 6 hyperfine components with relative intensities given by their statistical weight. The hyperfine splitting leads to a ~ 20 and $\sim 30\%$ increase of the line width for $\tau_{\text{TOT}} = 1$ and 10, respectively.

a global fit of laboratory and radioastronomical transition frequencies, whose weights were the inverse-square of the uncertainties. For the hyperfines, their assumed uncertainties account for both the standard errors of the hfs fit and the 6 kHz uncertainty of the N₂H⁺(1–0) reference line (Dore et al. 2004).

Tables 2–4 list, for each isotopomer, the experimental frequencies with their uncertainties and the residuals of the least-squares fit carried out to determine the spectroscopic constants reported at the bottom of the table. These accurate constants (see next paragraphs for a discussion of accuracy) allow us to recommend the rest frequencies reported in Table 5: the hyperfine components of the $J = 1 \leftarrow 0$ and $J = 2 \leftarrow 1$ transitions are listed for DCO⁺, while unsplit transition frequencies from $J = 2$ up to $J = 14$ are reported for all three isotopomers. It is worth mentioning that the quoted uncertainties in Table 5 are 1σ errors derived from the least-squares variance-covariance matrix for the fitted spectroscopic parameters.

As for the accuracy of the measured transition frequencies, the sources of systematic error are of two types: those due to the experimental set-up and procedures, and those related to the physics of molecules in the experimental environment.

The sources of the first kind include: accuracy of the frequency standard, harmonic distortion of the frequency modulation, integration due to slow lock-in time constant, asymmetry in the line shape due to etalon effect in the absorption cell.

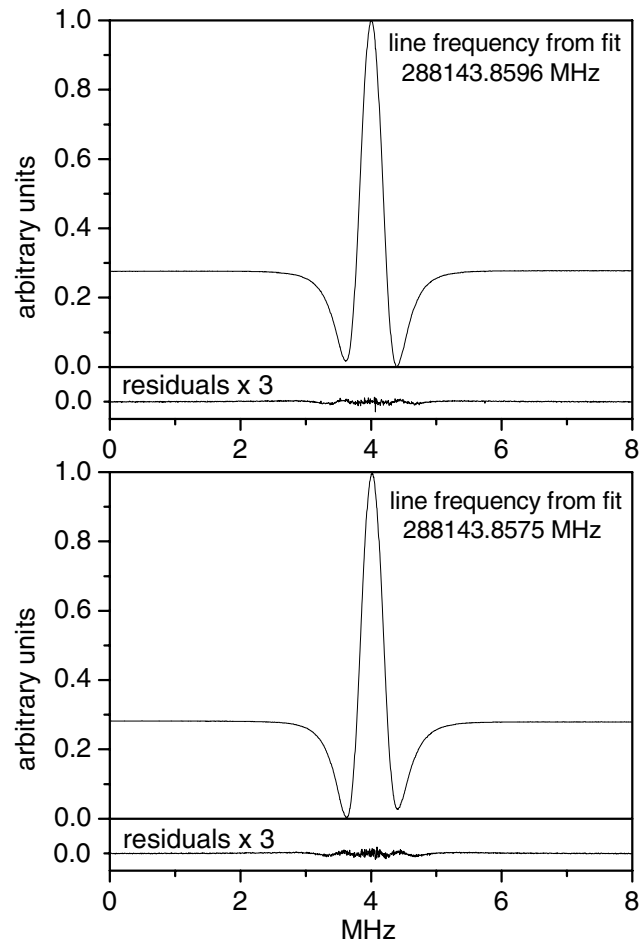


Fig. 3. Spectra of the $J = 4 \leftarrow 3$ transition of DCO⁺ with opposite line asymmetry due to different standing wave pattern in the cell: the 1601 points of each spectrum were recorded for a total time of 39 s at a rate of 0.8 MHz s⁻¹ with a lock-in time constant of 1 ms. Their profile has been fit to a model Galatry profile with the inclusion of a dispersion term to recover the line frequency.

The well tested performances of the spectrometer (Cazzoli & Dore 1990a; Puzzarini et al. 2002) and a judicious choice of the lock-in time constant (less than 1/100 of the rising time during the line recording) allowed all these error sources, except the last, to be rendered negligible: Fig. 3 shows, in fact, a pair of spectra of the $J = 5 \leftarrow 4$ transition of DCO⁺ with opposite asymmetry. The shift of the apparent center frequency, however, may be accounted for by a full profile analysis with a polynomial function describing the baseline and a dispersion term included; this procedure gives an accurate line center (Dore 2003).

The second kind of error in the line center includes: Doppler shift due to ion drift in the discharge, gas flow through the cell due to pumping, pressure shift. The first two causes of line shift should be negligible in the present experimental conditions, and are, anyway, largely suppressed by the double-pass arrangement. As for the frequency shift by Ar pressure (about 4 mTorr in this experiment), Buffa et al. (1994) have shown in the case of HCO⁺ that it is most significant for the $J = 1 \leftarrow 0$

Table 3. Rotational transition frequencies and spectroscopic constants of D¹³CO⁺.

J'	J	Observed/MHz	Obs.-calc./kHz	Uncertainties/kHz ^a
2	1	141 465.1331 ^b	3.4	5
3	2	212 194.4920	1.9	5
4	3	282 920.0055	0.3	5
5	4	353 640.3923	-1.3	5
6	5	424 354.3718	-1.8	5
7	6	495 060.6646	0.4	5
8	7	565 757.9861	2.0	5
9	8	636 445.0518	-0.8	5
10	9	707 120.5885	-0.7	15
11	10	777 783.3159	2.3	50
$\sigma_{\text{fit}}^c/\text{kHz}$ 1.8				
Constant ^d	This work	Previous ^e	D ¹³ C ¹⁵ N ^f	
B/MHz rotational	35 366.70968(21)	35 366.712(11)		
D_J/kHz quartic c. d. ^g	53.4083(40)	53.20(53)	52.26535(96)	
H_J/Hz sextic c. d. ^g	0.048(24)		0.06074(73)	

^a Uncertainties estimated as explained in the text.

^b Not included in the fit, see text.

^c $(\Sigma(\text{obs.}-\text{calc.})^2/\text{degrees of freedom})^{1/2}$.

^d Standard errors are reported in parentheses in units of the last quoted digits.

^e Bogey et al. (1981).

^f Brünken et al. (2004).

^g Centrifugal distortion.

transition; a semiclassical estimate for DCO⁺ broadened by Ar (Buffa 2004) indicates that, among the rotational lines considered here, only the $J = 2 \leftarrow 1$ center frequency should be slightly shifted (about 3 kHz); therefore it has been excluded from the fit.

In the present case, there are also the hyperfine contributions of the D and ¹³C nuclei which may affect the accuracy of the determined transition frequencies. These effects are more significant for the lower J transitions, therefore the most affected among those included in the fit should be the $J = 3 \leftarrow 2$. Its hf structure was predicted assuming the hf constants from the present work for D and from Schmid-Burgk et al. (2004) for ¹³C; then, to simulate the blended experimental profile, modulated Voigt functions (see Dore 2003) were summed at each hyperfine frequency with the relative intensity as weighting factor; finally, the center frequency of the synthetic profile was measured to check for a shift with respect to the assumed unperturbed $J = 3 \leftarrow 2$ transition frequency. From this procedure carried out for all three isotopomers, it turned out that the apparent line center is shifted at most by 0.3 kHz from the unperturbed center frequency: this indicates that the hyperfine structure is irrelevant in this case, and a fortiori for the higher J transitions, as seen in Sect. 2.2.

4. Conclusions

The present paper has shown how radioastronomical observations together with laboratory work are needed to determine

with high precision the spectroscopic parameters of molecular species. In the laboratory, it has been possible to measure the frequencies of DCO⁺, D¹³CO⁺, and DC¹⁸O⁺ lines up to 792 GHz with a fairly high accuracy. New values of the rotational, quartic, and sextic distortion constants have been determined for the three isotopomers with such a precision to allow to predict spectra up to 1000 GHz with an accuracy of at least 1 part in 10⁸ for DCO⁺, 5 parts in 10⁸ for D¹³CO⁺, and of 1 part in 10⁷ for DC¹⁸O⁺. Previous frequency estimates were much less accurate and did not allow to make accurate kinematic studies in dense molecular cloud cores.

Using the IRAM-30 m antenna, we detected for the first time the hyperfine structure of the DCO⁺ (1–0) line toward the quiescent starless core L1512, in Taurus. This is the first time that the hyperfine structure due to a deuteron has ever been observed; this gave us the possibility of obtaining the hyperfine parameters of DCO⁺. The DCO⁺(1–0) frequency has also been estimated comparing the observed line with a high sensitivity spectrum of N₂H⁺(1–0), and using the new value of the N₂H⁺(1–0) line as calculated by Dore et al. (2004). This value is however still uncertain by 6 kHz because of astronomical errors, and new sub-Doppler laboratory work on N₂H⁺ is planned to improve this precision. The estimated DCO⁺(1–0) hf frequencies were included with their uncertainties in a weighted least-squares fit of the laboratory data to derive accurate spectroscopic constants. From the fitted constants, the hyperfines have been reestimated with an accuracy of better than 2.5 parts in 10⁸.

Table 4. Rotational transition frequencies and spectroscopic constants of DC¹⁸O⁺.

J'	J	Observed/MHz	Obs.-calc./kHz	Uncertainties/kHz ^a
2	1	137 653.5239 ^b	3.1	10
3	2	206 477.2402	-0.9	5
4	3	275 297.3150	1.5	5
5	4	344 112.5213	-1.1	5
6	5	412 921.6530	0.4	5
7	6	481 723.4893	0.3	5
8	7	550 516.8156	-1.7	10
9	8	619 300.4281	4.4	35
10	9	688 073.0967	1.9	45
σ_{fit}^c /kHz 2.5				
Constant ^d	This work		Previous ^e	
B /MHz rotational	34 413.78556(18)		34 413.798(1)	
D_J /kHz quartic c. d. ^f	50.6704(44)		50.29(4)	
H_J /Hz sextic c. d. ^f	0.109(34)			

^a Uncertainties estimated as explained in the text.^b Not included in the fit, see text.^c $(\Sigma(\text{obs.}-\text{calc.})^2/\text{degrees of freedom})^{1/2}$.^d Standard errors are reported in parentheses in units of the last quoted digits.^e Bogey et al. (1981).^f Centrifugal distortion.**Table 5.** Recommended line frequencies^a (MHz) of DCO⁺, D¹³CO⁺, and DC¹⁸O⁺.

$J'(F')$	$J(F)$	DCO ⁺	D ¹³ CO ⁺	DC ¹⁸ O ⁺
1(0)	0(1)	72 039.2414(18)		
1(2)	0(1)	72 039.3031(9)		
1(1)	0(1)	72 039.3506(14)		
2(1)	1(1)	144 077.2144(18)		
2(1)	1(2)	144 077.2619(29)		
2(3)	1(2)	144 077.2804(10)		
2(2)	1(1)	144 077.2851(5)		
2(1)	1(0)	144 077.3237(15)		
2(2)	1(2)	144 077.3326(23)		
3	2	216 112.5800(6)	212 194.4901(9)	206 477.2411(7)
4	3	288 143.8577(6)	282 920.0052(9)	275 297.3135(6)
5	4	360 169.7795(6)	353 640.3936(8)	344 112.5224(6)
6	5	432 189.0066(5)	424 354.3736(7)	412 921.6526(6)
7	6	504 200.2004(6)	495 060.6642(8)	481 723.4890(8)
8	7	576 202.0228(7)	565 757.9841(9)	550 516.8173(13)
9	8	648 193.1356(8)	636 445.0526(11)	619 300.4237(31)
10	9	720 172.2013(11)	707 120.5892(28)	688 073.0947(72)
11	10	792 137.8823(22)	777 783.3135(67)	756 833.618(14)
12	11	864 088.8419(47)	848 431.946(13)	825 580.782(26)
13	12	936 023.7433(88)	919 065.206(24)	894 313.375(43)
14	13	1 007 941.251(15)	989 681.816(40)	963 030.188(68)
15	14	1 079 840.028(24)	1 060 280.497(62)	1 031 730.011(103)

^a The 1σ uncertainties are reported in parentheses in units of the last quoted digits.

The hyperfine structure of the DCO⁺(1–0) line needs to be taken into account in the analysis of astronomical data, to avoid overestimates of the line width by more than a factor of 2 and for a correct interpretation of the line profile.

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