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Perturbations in the A¹ Π , v = 0 state of ¹²C¹⁸O investigated via complementary spectroscopic techniques

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ABSTRACT

The A¹ $\Pi(v = 0)$ level of ¹²C¹⁸O has been reinvestigated using three different high-resolution spectroscopic methods: (1) 2 + 1' resonance-enhanced multiphoton ionisation of the A¹ $\Pi - X^{1}\Sigma^{+}(0, 0)$ band using narrowband lasers in a Doppler-free geometry; (2) Fourier-transform emission spectroscopy in the visible range probing the B¹ $\Sigma^{+} - A^{1}\Pi(0, 0)$ band in a discharge; (3) Fourier-transform absorption spectroscopy in the vacuum-ultraviolet range measuring the A¹ $\Pi - X^{1}\Sigma^{+}(0, 0)$ and B¹ $\Sigma^{+} - X^{1}\Sigma^{+}(0, 0)$ bands at multiple temperatures ranging from 90 to 900 K. An effective-Hamiltonian analysis of A¹ Π , v = 0 levels was performed up to J = 44 which quantitatively addresses perturbations by the e³ $\Sigma^{-}(v = 1)$, d³ $\Delta(v = 4)$, a'³ $\Sigma^{+}(v = 9)$, D¹ $\Delta(v = 0)$, and I¹ $\Sigma^{-}(v = 0, 1)$ levels.



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1. Introduction

Spectroscopy of the carbon monoxide molecule and, in particular, its pronounced 4th-positive or $A^{1}\Pi - X^{1}\Sigma^{+}$ system has been investigated repeatedly since the 1920s, beginning with Birge [1] and Herzberg [2]. It is a benchmark system for analysing perturbations in diatomic molecules. A comprehensive theoretical framework for such analysis of perturbations in the A¹\Pi state was developed by Field *et al.* [3,4]. The A¹\Pi state of the specific ¹²C¹⁸O isotopomer has been investigated via emission spectroscopy of the B¹\Sigma^{+} \rightarrow A¹Π Ångström bands [5,6], the C¹\Sigma^{+} \rightarrow A¹Π Herzberg bands [7], and the E¹Π \rightarrow A¹Π Kępa–Rytel system [8,9], and in absorption via the $A^1\Pi \leftarrow X^1\Sigma^+$ system using vacuumultraviolet (VUV) absorption [10–12]. These experiments reveal multiple non-Born–Oppenheimer interactions of the $A^1\Pi$ state with several other states, which do not themselves readily appear in experimental spectra: $d^3\Delta$, $e^3\Sigma^-$, $a'^3\Sigma^+$, $D^1\Delta$, and $I^1\Sigma^-$. Previously, Beaty *et al.* [12] used their high-resolution $A^1\Pi \rightarrow X^1\Sigma^+$ emission spectra to model the interaction of the $A^1\Pi(v = 0$ - 9) levels with these states and deduced deperturbed molecular constants. VUV-laser excitation studies on the $A^1\Pi \rightarrow X^1\Sigma^+$ system for ${}^{12}C^{18}O$ have been performed as well [13,14].

Other spectroscopic studies involving ${}^{12}C^{18}O$ have focused on the $B^{1}\Sigma^{+} - X^{1}\Sigma^{+}$ system [15], on transition

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oscillator strengths [16,17], and on the measurement of lifetimes [18,19]. The spectroscopy of the ${}^{12}C{}^{18}O$ ground state has been studied to high accuracy in microwave [e.g. 20,21] and far infrared (FIR) experiments [e.g. 22]. A comprehensive analysis was made by Coxon and Haji-georgiou [23] leading to a very accurate model for the ground-state in CO isotopologues expressed in terms of molecular parameters.

The present investigation is one of a series analysing perturbations in low- ν levels of the A¹ Π state in detail for various isotopologues. These studies have provided a highly accurate description of the level energies and the corresponding perturbations of the A¹ Π state in the ¹²C¹⁶O main isotopologue [24–27], and in the ¹³C¹⁶O [28], ¹²C¹⁷O [29] and ¹³C¹⁷O [30] isotopologues. One goal of this work is to quantify the electronic isotope shifts in the A¹ Π state and determine whether it shows anomalous behaviour in common with observations of the a ³ Π state [31,32]. The new spectroscopic measurements of A¹ Π ($\nu = 0$) state presented here are necessary to obtain the precision required to probe this effect.

The general methodology of this work is similar to our recent studies of other CO isotopologues [28–30]. Three spectroscopic techniques are employed: Doppler-free two-photon laser spectroscopy of the $A^1\Pi - X^1\Sigma^+(0, 0)$ band, Fourier-transform emission spectroscopy in the visible range on the $B^1\Sigma^+ - A^1\Pi(0, 0)$ band, and Fourier-transform absorption spectroscopy in the VUV range on the $A^1\Pi - X^1\Sigma^+(0, 0)$ and $B^1\Sigma^+ - X^1\Sigma^+(0, 0)$ bands. The first technique provides transition frequencies with ultrahigh accuracy and the latter two provide high-resolution broadband measurements. These independent measurements of the level energy separations for all combinations of $X^1\Sigma^+(\nu=0)$, $A^1\Pi(\nu=0)$, and $B^1\Sigma^+(\nu=0)$ levels provide a check on the absolute frequency uncertainties estimated for each experiment.

Based on this variety of complementary spectroscopic experiments, accurate level energies are determined for the rotational levels of $A^1\Pi(v=0)$ and $B^1\Sigma^+(v=0)$ and some levels in the perturbing $d^3\Delta(v=4)$, $e^3\Sigma^-(v=1)$ and $a'^3\Sigma^+(v=9)$ states. None of the latter states support allowed transitions in our spectra due to spin and orbital angular momentum selection rules, but appear due to their spin–orbit and rotational–electronic (*L*uncoupling) interactions with $A^1\Pi(v=0)$. An effective Hamiltonian is used to model the level structures and to determine deperturbed molecular constants for each of the states. Mutual interactions, as well as interactions with some more remote levels are included in the Hamiltonian matrix, leading to an accurate reproduction of the measured level energies to within experimental accuracy.

Hereafter, we abbreviate vibrational levels and transitions, for example, $A^1\Pi(v = 0)$ and $B^1\Sigma^+ - X^1\Sigma^+(v' =$ 0, $\nu'' = 0$) to A(0) and B-A(0, 0). Rotational branches for transitions involving triplet states are labelled in the tables, presented in this paper, as, e.g. Q_{21fe} , indicating a $\Delta J = 0$ transition with an upper-state *f*-symmetry F_2 level and lower-state *e*-symmetry F_1 level. The important cases are the d³ Δ and e³ Σ^- states for which the F_1 , F_2 , and F_3 rotational stacks correspond to Hund's case (a) levels with J = N - 1, N, and N + 1, respectively. There exist *e*- and *f*-symmetry versions of all F_i levels of d³ Δ whereas e³ $\Sigma^$ is restricted to the pairs (F_1 , e), (F_2 , f) and (F_3 , e).

2. 2 + 1' REMPI laser spectroscopy

2.1. Experimental details

A narrowband tunable pulsed dye amplifier laser system at LaserLaB Amsterdam [33] was used in a molecular beam study. Doppler-free spectra of the A-X(0,0)band of ¹²C¹⁸O were recorded using counter-propagating laser beams of ultraviolet (UV) radiation perpendicularly crossing a molecular beam of ¹⁸O isotopically enriched carbon monoxide (Sigma Aldrich, 99.9% ¹²C and 95% ¹⁸O). The excited molecules were ionised by a second pulse-time-delayed laser at 202 nm, forming a 2 + 1' resonance enhanced multiphoton ionisation (REMPI) process. Signal detection was established by monitoring 30 amu ions in a time-of-flight mass spectrometer. An accurate frequency calibration was performed by simultaneously measuring a saturated absorption spectrum of molecular iodine with reference to pre-calibrated I2 resonances [34] and a stabilised etalon, as was done in previous experiments with other CO isotopologues [26,27]. The measured transition frequencies of the two-photon resonances were corrected for frequency chirp effects in the pulsed dye amplifier [26,33,35] and extrapolated to zero intensity to correct for AC Stark effects [25]. These procedures yielded two-photon transition frequencies with an accuracy of 0.001 cm^{-1} , which corresponds to 30 MHz. This error estimate is composed of contributions from frequency chirp, absolute frequency calibration, and the AC-Stark extrapolation, in order of importance.

2.2. Results

Figure 1 shows a recording of the R(2) two-photon transition in the A–X(0, 0) band of ¹²C¹⁸O. The I₂ saturation spectrum used for absolute calibration and the transmission intensity from the stabilised etalon are also shown. The latter exhibits a low finesse at the wavelength of the fundamental, $\lambda = 617$ nm, but after fitting a sine function through the markers this does not contribute significantly to the calibration uncertainty. In the ¹²C¹⁸O molecular



Figure 1. Recording of the R(2) two-photon transition in the A¹ Π – X¹ Σ ⁺(0, 0) band of ¹²C¹⁸O measured by 2 + 1' REMPI (red points and fitted black curve). The lower blue and black lines represent etalon markers and the saturated absorption spectrum of I₂ used for frequency interpolation and calibration. The asterisk indicates the a13 (7,7) hyperfine component of the B-X (10,3) R(81) iodine line at 16 189.695 59 cm⁻¹ [34] that was used for calibration.

beam expansion only the lowest *J*-values can be probed due to rotational cooling. Accurate transition frequencies were measured for 10 lines in the two-photon spectrum and their results are listed in Table 1.

After the laser spectroscopic measurements on ${}^{12}C^{18}O$, a separate measurement was performed to correct for the chirp phenomenon [26]. Typical values for the chirp-induced shift were -18(5) MHz at the fundamental frequency. Even after quantitative assessment of this contribution to the systematic frequency shift it remains the major source of uncertainty in the error budget for frequencies listed in Table 1.

Table 1. Rotational transition frequencies,¹ ν_{obs} , the deviations between observed values and calculated values from a fit to the entire data set involving all laser, FT-emission and FT-absorption lines $\Delta_{obs-calc.}$, and AC-Stark slope coefficient², C_{AC} , of $A^{1}\Pi - X^{1}\Sigma^{+}(0, 0)$ in ¹²C¹⁸O as measured in the 2 + 1' REMPI laser experiment.

Line	$\nu_{\sf obs}$	$\Delta_{\rm obscalc.}$	C _{AC}
P(2)	64,743.6807 (10)	-0.0022	0.45
P(3)	64,738.7324 (10)	0.0001	0.33
P(4)	64,733.1271 (10)	-0.0007	0.44
Q(1)	64,750.9905 (20)	0.0006	
Q(2)	64,749.6591 (10)	- 0.0021	0.37
Q(3)	64,747.6367 (20)	- 0.0016	
R(1)	64,757.0383 (10)	- 0.0031	0.34
R(2)	64,758.7614 (10)	0.0019	0.43
S(0)	64,760.6457 (10)	- 0.0011	0.40
S(1)	64,765.9457 (10)	- 0.0016	0.43

 1 Units of cm $^{-1}$ and 1σ uncertainties given in parentheses in units of the least-significant digit)

²Units of MHz/(MW/cm²).



Figure 2. AC-Stark-plots for four transitions measured by two-photon Doppler-free spectroscopy in the $A^1\Pi - X^1\Sigma^+(0, 0)$ band of $^{12}C^{18}O$.

The transition frequencies of all lines were measured under various intensities of the laser field, providing information for making AC-Stark plots. These are shown in Figure 2 for four lines plotted against an absolute scale of laser intensity in the focal region. The uncertainty of the absolute intensity value may be as large as 25%, the relative intensities are accurate within 10%, allowing for an extrapolation to zero intensity to obtain an AC-Starkfree transition frequency. The values listed in Table 1 include such an extrapolation. An AC-Stark slope parameter CAC was also determined for most lines, representing the proportionality between the laser intensity in units of MW/cm² and the AC-Stark induced line shift. For the Q(1) and Q(3) lines only indicative values for the slopes were measured. On average a value of $C_{AC} = 0.41(10)$ is found for the two-photon lines in the A-X(0, 0) band of ¹²C¹⁸O. This value falls within a factor of two of that found in an early high-power study of the AC-Stark effect in CO [36]. Results on dynamic polarisabilities may provide information on the structure and internal dynamics in molecules, but will not be discussed further in this paper, which focuses on high resolution spectra and perturbations in the excited state.

3. Fourier-transform emission in the visible range (VIS-FT)

3.1. Experimental details

Visible (VIS) wavelength emission spectra of ${}^{12}C^{18}O$ were recorded at the University of Rzeszów using an air-cooled, carbon hollow-cathode (HC) lamp operated at 780 V, 54 mA dc. The lamp was filled with a static mixture of ${}^{18}O_2$ and ${}^{16}O_2$ (Sigma-Aldrich, 98.1% ${}^{18}O_2$) at a pressure of 3 mbar. During the discharge process, the O_2 molecules react with the ¹²C atoms ejected from the carbon filler placed inside the cathode, thus forming ¹²C¹⁸O and ¹²C¹⁶O molecules in the gas phase, in sufficient amounts to achieve a signal-to-noise ratio (SNR) of 70: 1 for the ¹²C¹⁸O spectrum. For the current experiment, we used an improved version of the HC lamp with respect to previous investigations [37,38]. A higher temperature dc-plasma (up to 900 K) was obtained at the centre of the cathode and allowed for observations of B–A(0, 0) rotational transitions with *J* as high as 40. The associated line-broadening increased by only 0.02 cm⁻¹ relative to previous studies [39,40], in which a dc-plasma temperature of about 650 K was used.

The discharge glow was focused by a plano-convex quartz lens onto the entrance aperture (1.0 mm diameter) of a 1.71 m Fourier-transform (FT) spectrometer [28,30]. A recent modification of this spectrometer significantly reduced the reflection of the HeNe laser beam, used for calibration and control of the position of a movable retroreflecting mirror, from optical elements in the interferometer and detector compartments, and noticeably improved its sensitivity in the VIS and UV regions. A visible-wavelength quartz beamsplitter and a photomultiplier tube running in integration mode were used to record the ${}^{12}C^{18}O B - A(0, 0)$ spectrum between 22,100 and 22,900 cm⁻¹. The spectrometer was operated under vacuum conditions (pressure below 0.003 mbar). Emission spectra were accumulated over 128 scans to obtain the desired SNR with a spectral resolution of 0.018 cm^{-1} . Under these settings the observed widths of ¹²C¹⁸O lines did not depend on the spectrometer resolution, but were limited by the Doppler broadening, yielding linewidths of 0.1 cm⁻¹. The molecular gas composition used to obtain the spectrum was ${}^{12}C^{18}O: {}^{12}C^{16}O = 1:0.1$.

The spectrum was frequency calibrated using the 633 nm line of an internal frequency-stabilised HeNe laser. The central positions of emission lines were measured by fitting Voigt lineshape functions to the experimental spectrum using a least-squares fitting procedure. The estimated absolute calibration uncertainty (1σ) is 0.003 cm⁻¹. The fitting uncertainty of the line frequency measurements is estimated to be 0.003 cm⁻¹ for single medium-strength lines and 0.01–0.02 cm⁻¹ for weak and/or blended ones.

3.2. Results

An overview of the wavelength region studied in VISemission (437–452 nm), together with rotational assignments and a simulated spectrum, is shown in Figure 3. In this spectrum some ${}^{12}C^{18}OB-A(0, 0)$ lines are blended

Table 2. Transition frequencies of the $^{12}C^{18}O~B^1\Sigma^+ - A^1\Pi$ (0, 0) band obtained in the VIS-FTS experiment. 1

J″	R	Q	Р
1	22,173.18 ^{wb}	22,165.76 ^{wb}	22,162.06 ^b
2	22,178.31 ^{wb}	22,167.13 ^b	22,159.76 ^b
3	22,184.191	22,169.21 ^b	22,158.220
4	22,190.86 ^{wb}	22,172.02 ^b	22,157.48 ^b
5	22,198.473	22,175.56 ^b	22,157.66 ^b
6	22,207.358	22,179.91 ^b	22,159.13 ^b
7	22,218.343	22,185.14 ^b	22,162.70 ^b
8	22,217.920	22,191.41 ^b	22,154.869
9	22,228.927	22,199.06 ^b	22,158.466
10	22,240.030	22,191.563	22,162.16 ^b
11	22,251.629 *	22,201.330	22,166.36 ^b
12	22,264.02 ^b	22,210.788 *	22,171.34 ^b
13	22,277.546 *	22,220.305	22,177.458 *
14	22,293.024	22,230.131 *	22,185.559
15	22,297.486	22,240.411	22,182.639 *
16	22,313.068	22,251.238	22,190.84 ^b
17	22,328.66 ^b	22,262.659	22,199.06 ^b
18	22,344.626	22,274.700	22,207.634
19	22,361.130	22,287.389	22,216.770
20	22,378.243	22,300.745	22,226.521
21	22,396.035	22,314.806	22,236.953
22	22,414.657	22,329.714	22,248.22 ^b
23	22,431.519 *	22,342.853	22,257.736
24	22,452.843	22,360.530 *	22,271.729
25	22,473.905	22,377.918	22,285.467 *
26	22,496.898	22,397.234	22,301.132
27	22,510.07 ^b	22,406.756	22,307.006
28	22,535.354 *	22,428.37 ^b	22,324.962
29	22,559.85 ^{wb}	22,449.219 *	22,342.145
30	22,586.31 ^{wb}	22,472.041	22,361.33 ^{b*}
31	22,599.42 ^{wb}	22,481.500	22,367.17 ^{wb}
32	22,627.33 ^{wb}	22,505.824	22,387.81 ^{wb}
33	22,653.81 ^{wb*}	22,528.868	22,407.04 ^b
34	22,680.43 ^{wb}	22,551.528 *	22,426.36 ^w
35	22,707.45 ^{wb}	22,574.959	22,446.14 ^{wb}
36	22,734.54 ^{wb}	22,598.91 ^{wb}	22,465.98 ^{wb}
37	22,762.99 ^{wb}	22,623.47 ^{wb}	22,487.27 ^{wb}
38	22,791.78 ^{wb}	22,648.67 ^{wb}	22,508.86 ^{bs}
39	22,821.20 ^{wb}	22,674.20 ^{wb}	22,531.06 ^{wb}
40		22,700.89 ^{wb}	22,553.92 ^{wb}

¹In units of cm⁻¹. Lines marked with *w* and/or *b* are weak and/or blended. For lines marked by an asterisk (^{*}) a small perturbation in the B¹ Σ^+ (*v* = 0) excited state was found. The instrumental resolution was 0.018 cm⁻¹. The estimated absolute calibration uncertainty (1 σ) is 0.003 cm⁻¹. The absolute accuracy of the line frequency measurements are estimated to be 0.003 – 0.02 cm⁻¹ depending on line strength and blending.

with ¹²C¹⁶O B–A (0, 0) as well as with lines from B–A (1, 1) bands of ¹²C¹⁸O and ¹²C¹⁶O. This contamination was taken into consideration during the analysis of the spectral lines in the ¹²C¹⁸O B–A (0, 0) band. In total, we assigned and analysed 168 molecular emission lines of ¹²C¹⁸O, among which 119 belong to B–A (0, 0) band, and 49 to the so-called 'extra lines" associated with the B¹ Σ^+ – $e^3\Sigma^-(0, 1)$, B¹ Σ^+ – $d^3\Delta(0, 4)$ and B¹ Σ^+ – $a'^3\Sigma^+(0, 9)$ transitions terminating on perturber states and gaining intensity from mixing with the A¹ Π state. Wavenumbers of the ¹²C¹⁸O B–A (0, 0) band are listed in Table 2 and



Figure 3. High resolution emission spectrum with rotational assignments of the ${}^{12}C^{18}O B^{1}\Sigma^{+} - A^{1}\Pi (0, 0), B^{1}\Sigma^{+} - e^{3}\Sigma^{-} (0, 1), B^{1}\Sigma^{+} - d^{3}\Delta (0, 4), and B^{1}\Sigma^{+} - a'^{3}\Sigma^{+} (0, 9)$ bands recorded with the VIS-FT spectrometer. Upper trace: experimental spectrum; Lower trace: simulation of ${}^{12}C^{18}O B^{1}\Sigma^{+} - A^{1}\Pi (0, 0)$ obtained with the Pgopher software [41]. Lines indicated by red labels represent *J*-values with the strongest perturbations.

transition frequencies of the extra lines are presented in Table 3.

4. Fourier-transform absorption in the vacuum ultraviolet (VUV-FT)

The Fourier-transform (FT) vacuum ultraviolet (VUV) spectrometer on the DESIRS beamline at the SOLEIL synchrotron was used to record spectra of the A-X(0, 0) and B-X(0, 0) bands of ${}^{12}C^{18}O$. Its operation and performance have already been described in detail [42,43] as

well as its application for the measurement of CO spectra [24,27,44].

4.1. $A^{1}\Pi \leftarrow X^{1}\Sigma^{+}(0, 0)$

Figure 4 shows three absorption spectra of A-X(0, 0) recorded with successively greater sample temperatures and column densities. The quasi-static CO gas was outflowing from a windowless cell under three different temperature regimes: liquid-nitrogen cooled (90 K), room temperature (300 K) and a heated

Table 3. Spin-forbidden lines appearing in the VIS-FT absorption spectra.¹

<i>J''</i>	^s R _{11ee}	^r Q _{11ef}	^q P _{11ee}	rR _{12ee}	^q Q _{12ef}	PP _{12ee}	^q R _{13ee}	^p Q _{13ef}	°P _{13ee}
3 4 5 6 7 8 9 10 11 12 13 14 15 16 17	22,180.76 ^{wb} 22,193.56 ^{wb} 22,206.250 22,232.36 ^{wb}		22,131.60 ^{wb} 22,135.25 ^{wb} 22,139.93 ^{wb} 22,145.33 ^{wb} 22,150.611 22,169.29 ^b 22,178.58 ^{wb}	Β	¹ Σ ⁺ - e ³ Σ ⁻ (0,1) 22,134.34 ^{wb} 22,139.43 ^{wb} 22,145.73 ^{wb} 22,153.20 ^{wb} 22,161.824 22,171.34 ^b 22,181.422 22,208.669 22,220.60 ^b 22,234.816 [*] 22,250.99 ^w		22,246.59 ^{wb} 22,263.93 ^{wb*} 22,281.187 22,311.44 ^{wb} 22,332.51 ^{wb} 22,355.57 ^{wb}		22,129.86 ^{wb} 22,136.72 ^{wb} 22,144.82 ^{wb} 22,153.94 ^{wb} 22,163.83 ^{b*} 22,173.717 22,196.57 ^{b*} 22,210.27 ^b
18				В	${}^{1}\Sigma^{+} - \mathbf{d}^{3}\Delta (0,4)$				
23	22,437.37 ^{wb*}	22,348.657	22,263.57 ^{wb}						
 27				22,526.95 ^{wb}	22,423.60	22,323.88 ^{wb}			
 30 31				fa	5 + 2/35 + (0.0)	22,565.14 ^{wb} 22,619.97 ^{wb}	22,450.816 22,502.03 ^w	22,387.69 ^b
32 33		22,473.76 ^{wb*} 22,523.68 ^{wb}		Β.	2 - a 2 (0,9	,			

¹All transition frequencies are in cm⁻¹. Lines marked with *w* and/or *b* are weak and/or blended. Lines marked with an asterisk (*) indicate a small perturbation in the B¹ $\Sigma^+(v = 0)$ excited state. The branch-label subscripts *e* and *f* indicate the upper-state/lower-state symmetry and superscripts *o*, *p*, *q*, *r* and *s* denote change in the total angular momentum excluding spin.

cell (900 K) [45]. Doppler-broadening of the observed spectral lines increases at higher temperatures and limits the accuracy of line-center measurements. The advantage of high-temperature spectra is that more rotational states can be probed, in the present study up to J = 41. Analysis of the combined spectra permitted precise frequency measurements for strong A–X(0, 0) lines and much-weaker forbidden transitions. Lines arising from the A–X(1, 1) hot band are also evident in Figure 4(c) between 63,800 and 64,130 cm⁻¹.

The integrated cross sections of all lines were optimised to best fit the 900 K spectrum after application of the Beer–Lambert law to their Doppler broadened summation. This required consideration of the sinc-shaped instrumental broadening of the VUV-FT instrument, with a full-width half-maximum of 0.27 cm⁻¹ (a higher spectral resolution, 0.07 cm⁻¹, was adopted while recording 90 and 300 K spectra). To improve the fitting of blended lines, the spectra were analysed while assuming combination differences of *P*- and *R*-branch transitions fixed to known ground state term values [23]. Also, no perturbers of A(0) have dipole-allowed transitions to the ground state and we assumed no intensity-interference effects afflicting A–X(0, 0) *P*- and *R*-branch lines with common *J'*, which might otherwise arise from the mixing of parallel and perpendicular transition moments [46]. Under this assumption, the relative strengths of P(J' + 1) and R(J' - 1) lines are then given by the ratio of their Hönl–London factors and a Boltzmann distribution of ground state J'' level populations [47,48].

Linearity of the FT frequency scale is intrinsically reliable, being dictated by the macroscopic interferometrically controlled stepping of a mirror in the wave-front-division interferometer. An absolute frequency calibration of the A-X(0, 0) spectrum was obtained by comparison with our two-photon absorption laser-based measurements of low-J $A^{1}\Pi - X^{1}\Sigma^{+}$ lines, described in Section 2. This required consideration of the parity of excited levels probed in one- and two-photon absorption because of significant Λ -doubling of the $A^1\Pi$ state. The estimated accuracy of this absolute calibration is 0.0015 cm⁻¹. The final list includes 126 rotational transitions of A-X(0, 0), and 73 spin-forbidden lines associated with the $e^{3}\Sigma^{-} - X^{1}\Sigma^{+}(1,0)$ and $d^{3}\Delta$ - $X^{1}\Sigma^{+}(4,0)$ bands. All measured frequencies are listed in Tables 4 and 5.

In the zero-order model, only the A–X(0, 0) band provides any oscillator strength and has a J'/J''independent band *f*-value when dividing by the Hönl– London factors for a ${}^{1}\Pi - {}^{1}\Sigma^{+}$ transition [47,48]. After



Figure 4. Spectra of the A–X(0, 0) band, and perturbing $e^{3}\Sigma^{-} - X^{1}\Sigma^{+}(1,0)$ and $d^{3}\Delta - X^{1}\Sigma^{+}(4,0)$ bands, recorded with the VUV-FT spectrometer at the SOLEIL synchrotron for three different sample temperatures, T = 90 K, T = 300 K and T = 900 K, as indicated.

level mixing is considered, some transitions to d(4) and e(1) levels borrow intensity by mixing with nearby A(0) levels.

The integrated cross sections of A-X(0, 0) lines were converted into band *f*-values by factorising their ground state populations and Hönl-London transition factors and are shown in Figure 5. Band f-values will be J' independent in the absence of perturbations, whereas A-X(0, 0) shows locally reduced f-values due to intensity borrowing by levels of $d^3\Delta - X^1\Sigma^+(4,0)$ and $e^3\Sigma^-$ - $X^{1}\Sigma^{+}(1,0)$. The integrated cross sections of extra lines of $d^3\Delta - X^1\Sigma^+(4,0)$ and $e^3\Sigma^- - X^1\Sigma^+(1,0)$ were converted into band f-values assuming Hönl-London factors for a ${}^{1}\Pi - {}^{1}\Sigma^{+}$ transition and are also shown in Figure 5. The summation of transitions to all interacting levels is I' independent and represents the unperturbed A-X(0, 0) band *f*-value. For *P*/*R* branch transitions connected to *e*-symmetry excited levels J' = 7 and 14 the admixture of A(0) into the F_3 component of e(1) is nearly

50%, indicated by the equal sharing of intensity. The *f*-symmetry J' = 10 levels of A(0) and e(1) are similarly affected.

4.2. $B^1\Sigma^+ \leftarrow X^1\Sigma^+(0,0)$

A spectrum of the B-X(0, 0) absorption band near 115 nm is shown in Figure 6, recorded with the FT-VUV instrument set at a spectral resolution of 0.11 cm⁻¹. A list of experimental transition energies is given in Table 6. Transition energies measured from this spectrum were combined with the VIS-FT spectrum of A-X(0, 0) to arrive at absolute term values for A(0), which can be compared with those derived from the REMPI laser experiment. Additionally, the direct measurement of small perturbations in the B(0) energy levels, indicated with (*) in Table 6, prevented these from being misinterpreted in B-A(0, 0) spectra as arising from A(0). The nature of levels interacting with B(0) interactions is beyond

Table 4. Transition frequencies of $A^{1}\Pi-X^{1}\Sigma^{+}(0,\,0)$ from the VUV-FT experiment. 1

J″	R	Q	Р
0	64,754.654(5)	_	_
1	64,756.983(4)	64,751.007(4)	-
2	64,758.614(4)	64,749.716(3)	64,743.668(5)
3	64,759.504(3)	64,747.779(3)	64,738.674(4)
4	64,759.522(3)	64,745.156(2)	64,732.982(4)
5	64,758.297(3)	64,741.852(2)	64,726.550(3)
6	64,755.022(3)	64,737.805(3)	64,719.248(3)
7	64,763.192(2)	64,732.927(2)	64,710.704(3)
8	64,759.989(2)	64,727.040(2)	64,700.111(3)
9	64,756.728(2)	64,719.811(2)	64,700.966(2)
10	64,753.016(2)	64,727.812(2)	64,690.450(2)
11	64,748.569(2)	64,718.578(2)	64,679.879(2)
12	64,743.025(2)	64,709.693(2)	64,668.859(2)
13	64,735.565(2)	64,700.814(3)	64,657.107(2)
14	64,739.169(2)	64,691.676(2)	64,644.262(2)
15	64,731.703(2)	64,682.119(2)	64,629.504(2)
16	64,724.267(2)	64,672.061(2)	64,625.814(2)
17	64,716.488(2)	64,661.463(2)	64,611.058(2)
18	64,708.217(2)	64,650.289(2)	64,596.337(2)
19	64,699.376(2)	64,638.514(2)	64,581.277(2)
20	64,689.910(2)	64,626.117(2)	64,565.730(2)
21	64,679.649(2)	64,613.065(2)	64,549.619(2)
22	64,671.189(3)	64,599.202(2)	64,532.887(2)
23	64,658.293(3)	64,587.172(2)	64,515.367(2)
24	64,645.698(3)	64,5/0.634(2)	64,499.653(3)
25	64,631.218(3)	64,554.432(2)	64,4/9.510(3)
26	64,626.582(4)	64,536.348(3)	64,459.674(3)
2/	64,609.894(4)	64,528.103(3)	64,437.960(3)
28	64,594.030(4)	64,507.805(3)	64,426.097(4)
29	04,570.222(5)	04,488.330(9)	64,402.190(4)
30 21	04,5/1./98(0)	04,400.910(4)	64,3/9.114(4)
31	04,332.392(0)	04,438.893(5)	64,354.101(5)
32 33	04,334.030(7)	04,430.009(3)	04,542.401(0)
20	04,317.007(0) 64.409.93(1)	04,414.333(0) 64,202,452(6)	04,310.000(0) 64.301.173(7)
24 25	04,490.02(1) 64,490.61(1)	64,393.433(0) 64,371,654(7)	04,291.1/2(/) 64.266.214(9)
26	64,460.01(1)	64,371.034(7)	04,200.214(0) 64.240.91(1)
20 27	64,401.00(1) 64,441.12(2)	04,349.330(9) 64 236 51(1)	04,240.01(1) 64 215 44(1)
32	64 420 67(2)	64 303 04(1)	64,213.44(1) 64,188,69(1)
30	64 300 58(3)	64 279 27(2)	64 161 69(2)
40	64 377 84(5)	64 254 42(2)	64 134 11(2)
41	(כ)דט. <i>ד</i> וכ _ו דט –	64 229 12(2)	(2)וו.דכו,דט –
42	64 332 51(6)	64 203 22(4)	_
43	-	64 176 64(4)	_
44	_	64 149 47(4)	_
-		57,172,72(7)	

¹In units of cm⁻¹ and with 1σ fitting uncertainties given in parentheses in units of the least-significant digit that are additional to a 0.0015 cm⁻¹ systematic uncertainty.

the scope of the current work and will be subsequently studied.

An absolute frequency calibration of the spectrum containing B-X(0, 0) was made with respect to NIST data for several lines of atomic Kr, Xe and H₂ contaminating the spectrum. These lines were induced by absorption upstream in the synchrotron beam line at the location of the gas filter used for discriminating the harmonics produced by the undulator [43]. Since these lines are Doppler broadened under static gas conditions there should be no shift of their centre frequencies. Lines of ${}^{12}C^{18}O$ itself, previously measured in several high-resolution laser-based experiments [18,19,49] concerning B-X(0, 0) and other well-known transitions $E^1\Pi$ –



Figure 5. Intensity sharing between $A^1\Pi - X^1\Sigma^+(0, 0)$ and overlapping bands. Band *f*-values for all transitions were calculated assuming the Hönl-London factors for a ${}^1\Pi - {}^1\Sigma^+$ transition. Points / error bars: experimental data (circles: *P*/*Q* branches, triangles: *R* branch). Lines: Results of the deperturbation model.

 $X^{1}\Sigma^{+}(1,0)$ and $C^{1}\Sigma^{+} - X^{1}\Sigma^{+}(1,0)$ were used for calibration as well. The estimated accuracy of this absolute calibration is 0.02 cm⁻¹.

5. Perturbation analysis

The CO $A^1\Pi$ state, especially its v = 0 level, has one of the more complicated rotational structures among lowlying states of diatomic molecules due to the multiple rotational-electronic (*L*-uncoupling) and spin–orbit perturbations that affect it. The spin–orbit interactions of A(0) with the $e^3\Sigma^-(v=1)$, $d^3\Delta(v=4)$ and $a'^3\Sigma^+(v=9)$ levels, as well as electronic– rotational interactions with $D^1\Delta(v=0)$ and $I^1\Sigma^-(v=0, 1)$ are responsible for these irregularities.

The rotational term values of ¹²C¹⁸O A¹ $\Pi(\nu = 0)$ are plotted in Figure 7 in reduced form. The crossing term series of e³ $\Sigma^-(\nu = 1)$, d³ $\Delta(\nu = 4)$, a'³ $\Sigma^+(\nu = 9)$, and D¹ $\Delta(\nu = 0)$ are also indicated and are associated with shifts of the A(0) energy levels. The strongest perturbation of A(0) rotational structure is due to spin–orbit interaction with the e³ $\Sigma^-(\nu = 1)$ at low *J* and is associated

Table 5. Spin-forbidden lines arising in the VUV-FT absorption spectra.¹

<i>J''</i>	^s R _{11ee}	^r Q _{11fe}	^q P _{11ee}	^r R _{12ee}	^q Q _{12fe}	^p P _{12ee}	^q R _{13ee}	^p Q _{13fe}	°P _{13ee}	
	$\mathbf{e}^{3}\boldsymbol{\Sigma}^{-}-\boldsymbol{X}^{1}\boldsymbol{\Sigma}^{+}\left(1,0\right)$									
1	64,787.53(9)				64,788.8(1)		64,799.0(2)			
2	64,785.26(7)				64,786.29(3)		64,801.30(6)			
3	64,781.72(2)				64,782.65(2)		64,802.2(2)			
4	64,777.24(1)				64,777.76(1)		64,802.0(3)			
5	64,772.109(5)		64,748.79(1)		64,771.70(1)		64,800.49(3)			
6	64,767.109(3)		64,736.97(1)		64,764.491(7)		64,797.76(2)			
7	64,748.755(6)		64,724.516(5)		64,756.242(6)		64,793.83(2)		64,752.90(3)	
8	64,739.87(1)		64,712.199(3)		64,747.134(5)		64,788.60(3)		64,742.85(2)	
9	64,729.11(2)		64,686.529(6)		64,737.477(4)		64,782.16(2)		64,731.60(2)	
10			64,670.33(1)		64,710.706(3)		64,774.57(1)		64,719.06(3)	
11			64,652.26(2)		64,699.299(6)		64,765.974(6)		64,705.31(2)	
12					64,685.667(5)		64,756.651(6)		64,690.41(1)	
13					64,670.15(2)		64,747.407(3)		64,674.512(6)	
14					64,653.01(1)		64,725.223(4)		64,657.888(6)	
15					64,634.40(2)		64,712.24(1)		64,641.346(3)	
16					64,614.45(1)		64,697.36(2)		64,611.868(4)	
17					64,593.08(1)				64,591.59(1)	
18									64,569.43(2)	
				($\mathbf{d}^{3} \mathbf{\Delta} - \mathbf{X}^{1} \mathbf{\Sigma}^{\mathbf{+}} (4,0)$					
22	64,665.341(5)									
23		64,581.366(7)								
24			64,493.805(5)	64,678.48(4)						
25			64,458.25(1)	64,649.95(1)						
26				64,609.73(1)	64,555.177(8)					
27					64,511.257(7)	64,456.70(1)				
28					64,477.29(3)	64,409.24(1)				
29							64,597.40(1)	64,522.68(3)		
30							64,551.25(2)	64,488.13(3)	(1 275 20/5)	
31								64,438.39(1)	64,3/5.28(1)	
32								64,401.58(1)	64,321.94(2)	





Figure 6. Photoabsorption spectrum of $B^1\Sigma^+ - X^1\Sigma^+(0, 0)$ band with a sample temperature of 1000 \pm 100 K. The B-X(1, 1) band also appears due to the higher temperature. The overall intensity slope is due to the continuum profile of the synchrotron beam. Also shown is the residual difference of the experimental spectrum and its modelled profile. Residuals are plotted on the same scale as the spectrum.

Table 6. Transition frequencies of $B^1\Sigma^+ - X^1\Sigma^+(0,\,0)$ in $^{12}C^{18}O$ from the VUV-FT experiment.^1

0 86,920.396(2) - 1 86,924.156(2) 86,913.022(4) 2 86,927.966(2) 86,909.847(2) 4 86,935.730(2) 86,902.334(2) 5 86,939.880(2) 86,898.872(2) 6 86,943.681(2) 86,895.456(2) 7 86,947.730(2) 86,892.087(2) 8 86,951.826(2) 86,882.77(2) 9 86,955.967(2) 86,882.287(2) 10 86,964.382(2)* 86,882.287(2) 11 86,964.382(2)* 86,872.920(2)* 12 86,964.382(2)* 86,872.920(2)* 13 86,973.010(2)* 86,865.99111(2) 15 86,981.795(2) 86,866.949(2)* 16 86,986.259(2) 86,864.024(2) 17 86,995.319(2) 86,855.359(2) 20 87,004.557(2) 86,855.358(2) 21 87,004.557(2) 86,854.332(2) 22 87,013.966(2) 86,844.5314(2) 23 87,013.392(2) 86,833.5015(2) 24	J''	R	Р
1 86,924,156(2) 86,913,022(4) 2 86,927,966(2) 86,909,411(2) 3 86,935,730(2) 86,902,334(2) 4 86,935,730(2) 86,902,334(2) 5 86,939,880(2) 86,888,872(2) 6 86,943,681(2) 86,892,887(2) 8 86,951,826(2) 86,882,771(2) 9 86,955,967(2) 86,882,287(2) 10 86,966,874(2) 86,882,287(2) 11 86,964,332(2)* 86,882,287(2) 12 86,966,874(2) 86,879,917(2) 13 86,973,370(0)* 86,872,920(2)* 14 86,977,379(2) 86,866,949(2)* 16 86,986,259(2) 86,864,024(2) 17 86,990,770(2) 86,851,350(2) 18 86,995,319(2) 86,853,329(2) 19 86,999,916(2) 86,851,384(2) 20 87,004,557(2) 86,852,833(2) 21 87,003,28(2) 86,847,534(2) 22 87,013,266(2) 86,841,954(2) 23	0	86,920.396(2)	-
2 86,927.966(2) 86,909.411(2) 3 86,931.826(2) 86,909.847(2) 4 86,931.826(2) 86,902.334(2) 5 86,939.680(2) 86,898.872(2) 6 86,943.681(2) 86,898.872(2) 7 86,947.730(2) 86,885.504(2) 8 86,951.826(2) 86,888.771(2) 9 86,955.967(2) 86,885.504(2) 10 86,964.674(2) 86,879.117(2) 12 86,968.674(2) 86,879.117(2) 13 86,973.010(2)* 86,872.920(2)* 14 86,977.379(2) 86,864.999.91(2) 15 86,981.795(2) 86,864.999.91(2) 16 86,980.599.916(2) 86,855.559(2) 20 87,004.557(2) 86,855.559(2) 20 87,004.557(2) 86,852.833(2) 21 87,003.287(2) 86,841.956(2)* 24 87,023.542(2) 86,844.956(2)* 25 87,023.932(2) 86,835.516(2)* 26 87,033.287(2) 86,835.164(2)*	1	86,924.156(2)	86,913.022(4)
3 86,931,826(2) 86,905,847(2) 4 86,935,730(2) 86,902,334(2) 5 86,939,680(2) 86,898,872(2) 6 86,943,681(2) 86,895,455(2) 7 86,947,730(2) 86,895,455(2) 8 86,951,826(2) 86,885,771(2) 9 86,955,957(2) 86,885,771(2) 10 86,960,155(2) 86,882,287(2) 11 86,964,382(2)* 86,872,920(2)* 12 86,968,674(2) 86,872,920(2)* 13 86,973,010(2)* 86,862,991(1) 15 86,981,795(2) 86,864,024(2)* 16 86,995,319(2) 86,864,024(2)* 16 86,995,319(2) 86,855,559(2) 20 87,004,557(2) 86,855,559(2) 21 87,004,557(2) 86,850,158(2) 22 87,013,96(2) 86,851,519(2) 23 87,013,92(3)* 86,844,956(2) 24 87,023,542(2) 86,837,519(2) 25 87,023,837(2) 86,837,519(2) 26 <td>2</td> <td>86,927.966(2)</td> <td>86,909.411(2)</td>	2	86,927.966(2)	86,909.411(2)
4 86,935.730(2) 86,902.334(2) 5 86,939.680(2) 86,898.872(2) 6 86,943.681(2) 86,892.872(2) 7 86,947.730(2) 86,885.504(2) 9 86,955.967(2) 86,885.504(2) 10 86,964.382(2)* 86,882.87(2) 11 86,964.382(2)* 86,887.517(2) 12 86,968.674(2) 86,875.998(2) 13 86,973.010(2)* 86,867.991(12) 15 86,981.795(2) 86,866.949(2)* 16 86,982.59(2) 86,864.024(2) 17 86,990.770(2) 86,861.150(2) 18 86,995.319(2) 86,855.559(2) 20 87,004.557(2) 86,852.833(2) 21 87,003.287(2) 86,850.158(2) 22 87,013.966(2) 86,847.534(2) 23 87,013.966(2) 86,847.534(2) 24 87,023.542(2) 86,837.519(2) 25 87,023.3287(2) 86,837.519(2) 26 87,033.287(2) 86,837.519(2) 27 </td <td>3</td> <td>86,931.826(2)</td> <td>86,905.847(2)</td>	3	86,931.826(2)	86,905.847(2)
5 86,939,680(2) 86,898,872(2) 6 86,934,3681(2) 86,895,456(2) 7 86,951,826(2) 86,885,504(2) 8 86,951,826(2) 86,885,504(2) 10 86,960,155(2) 86,882,287(2) 11 86,966,8674(2) 86,875,5988(2) 13 86,973,010(2)* 86,872,920(2)* 14 86,977,379(2) 86,869,911(2) 15 86,981,795(2) 86,866,949(2)* 16 86,986,259(2) 86,866,949(2)* 16 86,999,319(2) 86,858,329(2) 17 86,990,770(2) 86,851,559(2) 20 87,004,557(2) 86,853,559(2) 20 87,004,557(2) 86,853,232(2) 21 87,003,282(2) 86,850,158(2) 22 87,013,966(2) 86,847,534(2) 23 87,013,966(2) 86,847,534(2) 24 87,023,542(2) 86,833,519(2) 25 87,023,3287(2) 86,832,803(2) 26 87,033,287(2) 86,832,803(2) 27 87,038,218(2) 86,832,803(2) 28 <t< td=""><td>4</td><td>86,935.730(2)</td><td>86,902.334(2)</td></t<>	4	86,935.730(2)	86,902.334(2)
6 86,943,681(2) 86,895,456(2) 7 86,947,730(2) 86,892,087(2) 8 86,951,826(2) 86,888,771(2) 9 86,955,967(2) 86,885,504(2) 10 86,960,155(2) 86,882,287(2) 11 86,964,382(2)* 86,875,998(2) 12 86,968,674(2) 86,875,998(2) 13 86,977,379(2) 86,866,991(12) 15 86,981,795(2) 86,866,949(2)* 16 86,990,770(2) 86,864,024(2) 17 86,990,770(2) 86,851,55(2) 20 87,004,557(2) 86,852,833(2) 21 87,009,238(2) 86,852,833(2) 22 87,013,966(2) 86,854,454(2) 23 87,018,739(2)* 86,837,519(2) 24 87,023,542(2) 86,837,519(2) 25 87,023,542(2) 86,837,519(2) 26 87,033,287(2) 86,837,519(2) 27 87,033,287(2) 86,837,519(2) 26 87,033,287(2) 86,831,56(2) 27 <td>5</td> <td>86,939.680(2)</td> <td>86,898.872(2)</td>	5	86,939.680(2)	86,898.872(2)
7 86,947,730(2) 86,892,087(2) 8 86,955,1826(2) 86,888,771(2) 9 86,955,967(2) 86,885,504(2) 10 86,960,155(2) 86,885,504(2) 11 86,964,382(2)* 86,875,988(2) 13 86,973,010(2)* 86,875,998(2) 13 86,973,010(2)* 86,867,99117(2) 14 86,973,379(2) 86,866,949(2)* 16 86,986,259(2) 86,866,949(2)* 16 86,990,770(2) 86,861,150(2) 17 86,990,770(2) 86,850,158(2) 18 86,995,319(2) 86,852,832(2) 20 87,004,557(2) 86,852,833(2) 21 87,009,238(2) 86,850,158(2) 22 87,013,966(2) 86,844,4956(2) 24 87,023,542(2) 86,842,431(2) 25 87,023,287(2) 86,837,519(2) 26 87,033,287(2) 86,837,519(2) 27 87,038,218(2) 86,830,514(2) 30 87,053,257(2) 86,822,828(2) <	6	86,943.681(2)	86,895.456(2)
8 86,951,826(2) 86,888,771(2) 9 86,955.967(2) 86,885.504(2) 10 86,960.155(2) 86,882.287(2) 11 86,964.332(2)* 86,875.998(2) 13 86,973.010(2)* 86,875.998(2) 14 86,977.379(2) 86,869.911(2) 15 86,981.795(2) 86,866.949(2)* 16 86,986.259(2) 86,864.024(2) 17 86,990.770(2) 86,861.150(2) 18 86,995.319(2) 86,855.559(2) 20 87,004.557(2) 86,855.559(2) 20 87,004.557(2) 86,851.58(2) 21 87,003.238(2) 86,851.58(2) 22 87,013.966(2) 86,844.956(2) 24 87,023.542(2) 86,833.956(2)* 25 87,028.393(2) 86,835.158(2) 25 87,043.198(2)* 86,833.956(2)* 26 87,033.287(2) 86,835.136(2) 28 87,043.198(2)* 86,832.803(2) 29 87,043.398(2) 86,832.803(2)	7	86,947.730(2)	86,892.087(2)
9 86,955.967(2) 86,885.504(2) 10 86,960.155(2) 86,885.297(2) 11 86,964.382(2)* 86,875.998(2) 12 86,968.674(2) 86,875.998(2) 13 86,973.010(2)* 86,875.998(2) 14 86,977.379(2) 86,866.949(2)* 14 86,977.379(2) 86,866.949(2)* 16 86,986.259(2) 86,864.024(2) 17 86,990.770(2) 86,861.150(2) 18 86,995.319(2) 86,858.329(2) 19 86,999.916(2) 86,855.559(2) 20 87,004.557(2) 86,854.7534(2) 21 87,003.280(2) 86,847.534(2) 22 87,013.966(2) 86,847.534(2) 23 87,013.96(2) 86,837.519(2) 24 87,023.542(2) 86,837.519(2) 25 87,028.393(2) 86,830.514(2) 26 87,033.287(2) 86,830.514(2) 27 87,038.218(2) 86,830.514(2) 28 87,043.198(2)* 86,821.2803(2) <t< td=""><td>8</td><td>86,951.826(2)</td><td>86,888.771(2)</td></t<>	8	86,951.826(2)	86,888.771(2)
10 86,960.155(2) 86,882.287(2) 11 86,964.382(2)* 86,879.117(2) 12 86,968.674(2) 86,875.998(2)* 13 86,973.010(2)* 86,867.990.720(2)* 14 86,977.379(2) 86,866.9911(2)* 15 86,980.259(2) 86,866.949(2)* 16 86,990.770(2) 86,864.024(2) 17 86,990.770(2) 86,855.559(2) 18 86,995.319(2) 86,858.329(2) 18 86,999.319(2) 86,858.329(2) 20 87,004.557(2) 86,855.559(2) 20 87,013.966(2) 86,851.558(2) 21 87,013.966(2) 86,854.451(2) 22 87,013.966(2) 86,834.956(2)* 24 87,023.542(2) 86,834.956(2)* 25 87,028.393(2) 86,837.519(2) 26 87,033.287(2) 86,837.519(2) 27 87,033.287(2) 86,832.803(2) 28 87,043.198(2)* 86,832.803(2) 29 87,044.308(3) 86,832.803(2)	9	86,955.967(2)	86,885.504(2)
11 86,964.382(2)* 86,879.117(2) 12 86,968.674(2) 86,875.998(2) 13 86,973.010(2)* 86,869.911(2) 14 86,977.379(2) 86,866.949(2)* 16 86,986.259(2) 86,866.949(2)* 16 86,990.770(2) 86,866.949(2)* 16 86,990.770(2) 86,865.559(2) 17 86,990.916(2) 86,855.559(2) 20 87,004.557(2) 86,852.833(2) 21 87,009.238(2) 86,850.158(2) 22 87,013.966(2) 86,844.956(2) 24 87,023.542(2) 86,837.519(2) 25 87,023.542(2) 86,837.519(2) 26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,832.803(2) 29 87,048.207(2) 86,832.835(2) 31 87,058.339(2) 86,824.82(2)* 31 87,058.339(2) 86,821.835(2) 32 87,079.060(2) 86,817.746(2)* 36	10	86,960.155(2)	86,882.287(2)
12 86,968.674(2) 86,875.998(2) 13 86,973.010(2)* 86,872.920(2)* 14 86,977.379(2) 86,866.949(2)* 15 86,981.795(2) 86,866.4024(2) 16 86,996.259(2) 86,864.024(2) 17 86,990.770(2) 86,864.024(2) 18 86,999.916(2) 86,865.2832(2) 20 87,004.557(2) 86,855.559(2) 20 87,004.557(2) 86,855.158(2) 21 87,003.281(2) 86,851.58(2) 22 87,013.966(2) 86,851.58(2) 23 87,018.739(2)* 86,837.519(2) 24 87,023.542(2) 86,837.519(2) 25 87,028.393(2) 86,837.519(2) 26 87,033.287(2) 86,837.519(2) 27 87,033.287(2) 86,832.803(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,832.803(2) 31 87,058.339(2) 86,828.282(2)* 31 87,058.600(2)* 86,821.852(2)	11	86,964.382(2)*	86,879.117(2)
13 86,973.010(2)* 86,872.920(2)* 14 86,977.379(2) 86,869.911(2) 15 86,981.795(2) 86,866.949(2)* 16 86,986.259(2) 86,864.024(2) 17 86,990.770(2) 86,861.150(2) 18 86,995.319(2) 86,858.329(2) 19 86,999.916(2) 86,855.559(2) 20 87,004.557(2) 86,850.158(2) 22 87,013.966(2) 86,847.534(2) 23 87,018.739(2)* 86,844.956(2) 24 87,023.542(2) 86,837.519(2) 25 87,028.393(2) 86,837.519(2) 26 87,033.287(2) 86,833.5136(2) 27 87,038.218(2) 86,833.5136(2) 28 87,043.198(2)* 86,832.803(2) 29 87,043.297(2) 86,830.514(2) 30 87,053.252(2) 86,822.822(2)* 31 87,058.339(2) 86,812.835(2) 32 87,064.600(2)* 86,812.835(2) 33 87,068.600(2)* 86,812.835(2)	12	86,968.674(2)	86,875.998(2)
14 86,977.379(2) 86,869.911(2) 15 86,981.795(2) 86,866.949(2)* 16 86,986.259(2) 86,861.150(2) 17 86,990.770(2) 86,851.329(2) 18 86,999.916(2) 86,855.559(2) 20 87,004.557(2) 86,852.833(2) 21 87,009.238(2) 86,850.158(2) 22 87,013.966(2) 86,844.4956(2) 24 87,023.542(2) 86,837.519(2) 25 87,023.542(2) 86,837.519(2) 26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,832.803(2) 29 87,043.198(2)* 86,822.82(2)* 31 87,058.339(2) 86,821.835(2) 32 87,043.490(2) 86,817.746(2)* 33 87,068.600(2)* 86,817.746(2)* 36 87,073.829(2) 86,817.746(2)* 36 87,073.829(2) 86,818.774(2)* 36 87,094.978(4) 86,812.029(3) 39	13	86,973.010(2)*	86,872.920(2)*
15 86,981.795(2) 86,866.949(2)* 16 86,986.259(2) 86,864.024(2) 17 86,990.770(2) 86,851.359(2) 18 86,995.319(2) 86,855.559(2) 19 86,999.916(2) 86,855.559(2) 20 87,004.557(2) 86,855.559(2) 20 87,004.557(2) 86,851.538(2) 21 87,009.238(2) 86,850.158(2) 22 87,013.966(2) 86,844.431(2) 23 87,018.739(2)* 86,844.431(2) 24 87,023.542(2) 86,837.519(2) 25 87,028.393(2) 86,837.519(2) 26 87,033.287(2) 86,835.136(2) 27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,830.514(2) 30 87,053.252(2) 86,828.282(2)* 31 87,058.339(2) 86,817.746(2)* 32 87,063.460(2) 86,817.746(2)* 33 87,064.600(2)* 86,817.746(2)* 34 87,073.829(2) 86,818.1834(2) 35	14	86,977.379(2)	86,869.911(2)
16 86,986.259(2) 86,864.024(2) 17 86,990.770(2) 86,861.150(2) 18 86,999.916(2) 86,858.329(2) 19 86,999.916(2) 86,852.833(2) 20 87,004.557(2) 86,852.833(2) 21 87,009.238(2) 86,850.158(2) 22 87,013.966(2) 86,844.956(2) 24 87,023.542(2) 86,839.956(2)* 26 87,033.287(2) 86,839.956(2)* 26 87,033.287(2) 86,835.136(2) 27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,835.136(2) 29 87,043.207(2) 86,830.514(2) 30 87,053.329(2) 86,826.08(2) 31 87,058.339(2) 86,826.08(62) 32 87,063.460(2) 86,817.746(2)* 36 87,079.060(2) 86,817.746(2)* 36 87,084.338(3) 86,815.814(2) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) <td< td=""><td>15</td><td>86,981.795(2)</td><td>86,866.949(2)*</td></td<>	15	86,981.795(2)	86,866.949(2)*
17 86,990.770(2) 86,861.150(2) 18 86,995.319(2) 86,858.329(2) 19 86,999.916(2) 86,855.559(2) 20 87,004.557(2) 86,855.559(2) 21 87,009.238(2) 86,850.158(2) 22 87,018.739(2)* 86,844.956(2) 23 87,018.739(2)* 86,844.956(2) 24 87,023.542(2) 86,839.956(2)* 26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,832.803(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,823.803(2) 29 87,063.460(2) 86,823.803(2) 30 87,053.252(2) 86,823.803(2) 31 87,058.339(2) 86,826.086(2) 32 87,063.460(2) 86,821.835(2) 34 87,079.060(2) 86,817.746(2)* 35 87,079.060(2) 86,813.894(2) 38 87,089.643(3) 86,813.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,804.560(2) 39 <	16	86,986.259(2)	86,864.024(2)
18 86,995,319(2) 86,858,329(2) 19 86,999,916(2) 86,855,559(2) 20 87,004,557(2) 86,852,833(2) 21 87,009,238(2) 86,850,158(2) 22 87,013,966(2) 86,844,956(2) 23 87,018,739(2)* 86,844,431(2) 25 87,023,542(2) 86,839,956(2)* 26 87,033,287(2) 86,837,519(2) 27 87,038,218(2) 86,835,136(2) 28 87,043,198(2)* 86,832,803(2) 29 87,048,207(2) 86,832,803(2) 29 87,053,252(2) 86,828,282(2)* 31 87,053,460(2) 86,823,936(2) 32 87,063,460(2) 86,821,835(2) 33 87,068,600(2)* 86,813,894(2) 34 87,073,829(2) 86,813,894(2) 35 87,079,060(2) 86,813,894(2) 36 87,084,338(3) 86,813,894(2) 37 87,089,643(3) 86,813,894(2) 38 87,074,976(5) 86,808,416(4) <t< td=""><td>17</td><td>86,990.770(2)</td><td>86,861.150(2)</td></t<>	17	86,990.770(2)	86,861.150(2)
19 86,999.916(2) 86,855.559(2) 20 87,004.557(2) 86,852.833(2) 21 87,009.238(2) 86,852.833(2) 22 87,013.966(2) 86,847.158(2) 23 87,013.966(2) 86,844.956(2) 24 87,023.542(2) 86,842.431(2) 25 87,028.393(2) 86,839.956(2)* 26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,823.803(2) 29 87,053.252(2) 86,824.82(2)* 30 87,053.460(2) 86,821.835(2) 31 87,053.460(2) 86,821.835(2) 33 87,068.600(2)* 86,821.835(2) 34 87,073.829(2) 86,817.746(2)* 36 87,084.338(3) 86,813.894(2) 37 87,089.643(3) 86,813.894(2) 38 87,094.978(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) <td< td=""><td>18</td><td>86,995.319(2)</td><td>86,858.329(2)</td></td<>	18	86,995.319(2)	86,858.329(2)
20 87,004.557(2) 86,852.833(2) 21 87,009.238(2) 86,850.158(2) 22 87,013.966(2) 86,847.534(2) 23 87,018.739(2)* 86,844.956(2) 24 87,023.542(2) 86,834.956(2)* 26 87,023.542(2) 86,837.919(2)* 26 87,033.287(2) 86,837.919(2) 27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,835.136(2) 29 87,048.207(2) 86,830.514(2) 30 87,053.252(2) 86,828.282(2)* 31 87,058.339(2) 86,826.086(2) 32 87,063.460(2)* 86,821.835(2) 33 87,068.600(2)* 86,819.776(2) 35 87,079.060(2) 86,817.746(2)* 36 87,094.978(4) 86,812.029(3) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,804.970(5) 43 87,122.135(9) 86,803.322(6)	19	86,999.916(2)	86,855.559(2)
21 87,009.238(2) 86,850.158(2) 22 87,013.966(2) 86,847.534(2) 23 87,018.739(2)* 86,844.956(2) 24 87,023.542(2) 86,844.956(2)* 25 87,023.393(2) 86,837.519(2)* 26 87,033.287(2) 86,837.519(2)* 26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,830.514(2) 30 87,053.252(2) 86,828.282(2)* 31 87,058.339(2) 86,826.086(2) 32 87,063.460(2)* 86,821.835(2) 33 87,068.600(2)* 86,817.746(2)* 36 87,079.060(2) 86,817.746(2)* 36 87,094.43(3) 86,813.894(2) 37 87,089.643(3) 86,813.894(2) 38 87,094.978(4) 86,812.029(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44	20	87,004.557(2)	86,852.833(2)
22 87,013.966(2) 86,847.534(2) 23 87,018.739(2)* 86,844.956(2) 24 87,023.542(2) 86,842.431(2) 25 87,028.393(2) 86,839.956(2)* 26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,835.5136(2) 28 87,043.198(2)* 86,835.0514(2) 30 87,053.252(2) 86,828.282(2)* 31 87,058.339(2) 86,826.086(2) 32 87,063.460(2) 86,821.835(2) 34 87,073.829(2) 86,817.746(2)* 35 87,079.060(2) 86,817.746(2)* 36 87,084.338(3) 86,813.814(2) 37 87,089.643(3) 86,813.894(2) 38 87,094.978(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,803.322(6) 42 87,113.20(1) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 <t< td=""><td>21</td><td>87,009.238(2)</td><td>86,850.158(2)</td></t<>	21	87,009.238(2)	86,850.158(2)
23 87,018.739(2)* 86,844.956(2) 24 87,023.542(2) 86,839.956(2)* 25 87,028.393(2) 86,837.519(2) 26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,837.519(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,828.282(2)* 31 87,053.252(2) 86,828.282(2)* 31 87,058.339(2) 86,821.835(2) 33 87,068.600(2)* 86,821.835(2) 34 87,073.829(2) 86,817.746(2)* 35 87,079.060(2) 86,813.814(2) 36 87,084.338(3) 86,813.894(2) 37 87,089.643(3) 86,813.894(2) 38 87,019.745(5) 86,804.672(4) 40 87,105.745(5) 86,804.4970(5) 41 87,111.187(6) 86,804.672(4) 42 87,116.669(7) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,803.322(6)	22	87,013.966(2)	86,847.534(2)
24 87,023.542(2) 86,842.431(2) 25 87,028.393(2) 86,839.956(2)* 26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,837.519(2) 28 87,043.198(2)* 86,832.80(2) 29 87,048.207(2) 86,830.514(2) 30 87,053.252(2) 86,826.8262(2)* 31 87,058.339(2) 86,826.086(2) 32 87,063.460(2) 86,821.835(2) 34 87,073.829(2) 86,817.746(2)* 35 87,079.060(2) 86,817.746(2)* 36 87,084.338(3) 86,815.814(2) 37 87,089.643(3) 86,815.814(2) 38 87,094.978(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,801.726(7) 46 87,133.20(1) 86,800.125(9) 47 87,144.44(2) 86,797.69(1) 48 87	23	87,018.739(2) [*]	86,844.956(2)
25 87,028.393(2) 86,839.956(2)* 26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,832.803(2) 30 87,053.252(2) 86,828.282(2)* 31 87,053.359(2) 86,823.936(2) 32 87,063.460(2) 86,823.936(2) 33 87,068.600(2)* 86,813.936(2) 34 87,073.829(2) 86,819.776(2) 35 87,079.060(2) 86,817.746(2)* 36 87,084.338(3) 86,813.894(2) 37 87,089.643(3) 86,813.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,801.726(7) 46 87,133.20(1) 86,803.322(6) 44 87,150.03(3) 86,795.65(2) 45 87,133.20(1) 86,800.125(9) 46 87,	24	87,023.542(2)	86,842.431(2)
26 87,033.287(2) 86,837.519(2) 27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,835.136(2) 29 87,048.207(2) 86,830.514(2) 30 87,053.252(2) 86,828.282(2)* 31 87,058.339(2) 86,826.086(2) 32 87,063.460(2) 86,823.936(2) 33 87,068.600(2)* 86,821.835(2) 34 87,073.829(2) 86,817.746(2)* 36 87,084.338(3) 86,817.746(2)* 36 87,094.978(4) 86,813.894(2) 37 87,089.643(3) 86,813.894(2) 38 87,010.346(4) 86,810.202(3) 40 87,110.346(4) 86,810.202(3) 40 87,111.187(6) 86,804.416(4) 41 87,111.187(6) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,803.322(6) 44 87,150.03(3) 86,795.65(2) 45	25	87,028.393(2)	86,839.956(2)*
27 87,038.218(2) 86,835.136(2) 28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,830.514(2) 30 87,053.252(2) 86,828.282(2)* 31 87,058.339(2) 86,826.086(2) 32 87,063.460(2) 86,821.936(2) 33 87,068.600(2)* 86,819.376(2) 34 87,073.829(2) 86,819.376(2) 35 87,079.060(2) 86,817.746(2)* 36 87,084.338(3) 86,815.814(2) 37 87,089.643(3) 86,813.894(2) 38 87,094.978(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,803.322(6) 42 87,116.669(7) 86,803.322(6) 44 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.126(7) 45 87,133.20(1) 86,803.322(6) 44 87,127.67(1) 86,797.09(1) 45 87,133.20(1) 86,798.61(1) 47 87,138.79(2) 86,798.61(1) 48 87,150	26	87,033.287(2)	86,837.519(2)
28 87,043.198(2)* 86,832.803(2) 29 87,048.207(2) 86,830.514(2) 30 87,053.252(2) 86,828.282(2)* 31 87,058.339(2) 86,826.086(2) 32 87,063.460(2) 86,823.936(2) 33 87,068.600(2)* 86,821.835(2) 34 87,073.829(2) 86,817.776(2) 35 87,079.060(2) 86,817.776(2)* 36 87,089.643(3) 86,815.814(2) 37 87,089.643(3) 86,813.894(2) 38 87,094.978(4) 86,812.029(3) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,803.322(6) 42 87,116.669(7) 86,803.322(6) 44 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,138.79(2) 86,798.61(1) 48 87,150.03(3) 86,795.65(2) 49 <td>27</td> <td>87,038.218(2)</td> <td>86,835.136(2)</td>	27	87,038.218(2)	86,835.136(2)
29 87,048.207(2) 86,830.514(2) 30 87,053.252(2) 86,828.282(2)* 31 87,058.339(2) 86,826.086(2) 32 87,063.460(2) 86,823.936(2) 33 87,068.600(2)* 86,821.835(2) 34 87,073.829(2) 86,819.776(2)* 35 87,079.060(2) 86,817.814(2)* 36 87,089.643(3) 86,815.814(2)* 37 87,089.643(3) 86,813.894(2) 38 87,094.978(4) 86,812.029(3) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,800.125(9) 45 87,133.20(1) 86,800.125(9) 46 87,133.20(1) 86,797.09(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 <td< td=""><td>28</td><td>87,043.198(2)*</td><td>86,832.803(2)</td></td<>	28	87,043.198(2)*	86,832.803(2)
30 87,053.252(2) 86,828.282(2)* 31 87,053.339(2) 86,826.086(2) 32 87,063.460(2) 86,823.936(2) 33 87,068.600(2)* 86,821.835(2) 34 87,073.829(2) 86,819.776(2)* 35 87,094.060(2) 86,817.746(2)* 36 87,094.0643(3) 86,815.814(2) 37 87,094.978(4) 86,812.029(3) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.25(9) 45 87,133.20(1) 86,800.125(9) 46 87,133.20(1) 86,797.09(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,797.69(2) 49 87,155.66(3) - 50 87,161.39(4) -	29	87,048.207(2)	86,830.514(2)
31 87,058.339(2) 86,826.086(2) 32 87,063.460(2) 86,823.936(2) 33 87,068.600(2)* 86,821.835(2) 34 87,073.829(2) 86,819.776(2)* 35 87,079.060(2) 86,817.746(2)* 36 87,084.338(3) 86,813.814(2) 37 87,089.643(3) 86,813.814(2) 38 87,094.978(4) 86,812.029(3) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,807.92(9) 46 87,133.20(1) 86,797.69(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	30	87,053.252(2)	86,828.282(2)*
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33 87,068.600(2)* 86,821.835(2) 34 87,073.829(2) 86,819.776(2) 35 87,079.060(2) 86,817.746(2)* 36 87,084.338(3) 86,815.814(2) 37 87,089.643(3) 86,815.814(2) 38 87,094.978(4) 86,812.029(3) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,804.672(4) 42 87,116.669(7) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,800.125(9) 46 87,133.20(1) 86,799.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	32	87,063.460(2)	86,823.936(2)
34 87,073,829(2) 86,819,776(2) 35 87,079,060(2) 86,817,746(2)* 36 87,084,338(3) 86,815,814(2) 37 87,089,643(3) 86,813,894(2) 38 87,094,978(4) 86,812,029(3) 39 87,100,346(4) 86,810,202(3) 40 87,105,745(5) 86,808,416(4) 41 87,111,187(6) 86,808,416(4) 42 87,116,669(7) 86,804,970(5) 43 87,122,135(9) 86,803,322(6) 44 87,127,67(1) 86,801,726(7) 45 87,133,20(1) 86,801,726(7) 46 87,133,20(1) 86,798,61(1) 47 87,144,44(2) 86,797,09(1) 48 87,150,03(3) 86,795,65(2) 49 87,155,66(3) - 50 87,161,39(4) -	33	87,068.600(2)*	86,821.835(2)
35 87,079.060(2) 86,817.746(2)* 36 87,084.338(3) 86,815.814(2) 37 87,089.643(3) 86,813.894(2) 38 87,094.978(4) 86,812.029(3) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,806.672(4) 42 87,116.669(7) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,800.125(9) 46 87,133.20(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	34	87,073.829(2)	86,819.776(2)
36 87,084.338(3) 86,815.814(2) 37 87,089,643(3) 86,813.894(2) 38 87,094.978(4) 86,812.029(3) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,806.672(4) 42 87,116.669(7) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,800.125(7) 45 87,133.20(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	35	87.079.060(2)	86,817,746(2)*
37 87,089,643(3) 86,813,894(2) 38 87,094,978(4) 86,812,029(3) 39 87,100,346(4) 86,810,202(3) 40 87,105,745(5) 86,808,416(4) 41 87,111,187(6) 86,806,672(4) 42 87,116,669(7) 86,804,970(5) 43 87,122,135(9) 86,803,322(6) 44 87,127,67(1) 86,800,125(7) 45 87,133,20(1) 86,800,125(9) 46 87,138,79(2) 86,798,61(1) 47 87,144,44(2) 86,795,65(2) 49 87,155,66(3) - 50 87,161,39(4) -	36	87,084.338(3)	86,815.814(2)
38 87,094.978(4) 86,812.029(3) 39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,806.672(4) 42 87,116.669(7) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.125(7) 45 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	37	87,089.643(3)	86,813.894(2)
39 87,100.346(4) 86,810.202(3) 40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,806.672(4) 42 87,116.669(7) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	38	87,094.978(4)	86,812.029(3)
40 87,105.745(5) 86,808.416(4) 41 87,111.187(6) 86,806.672(4) 42 87,116.669(7) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,797.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	39	87,100.346(4)	86,810.202(3)
41 87,111.187(6) 86,806.672(4) 42 87,116.669(7) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	40	87,105.745(5)	86,808.416(4)
42 87,116.669(7) 86,804.970(5) 43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	41	87,111.187(6)	86,806.672(4)
43 87,122.135(9) 86,803.322(6) 44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	42	87,116.669(7)	86,804.970(5)
44 87,127.67(1) 86,801.726(7) 45 87,133.20(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	43	87,122.135(9)	86,803.322(6)
45 87,133.20(1) 86,800.125(9) 46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	44	87,127.67(1)	86,801.726(7)
46 87,138.79(2) 86,798.61(1) 47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	45	87,133.20(1)	86,800.125(9)
47 87,144.44(2) 86,797.09(1) 48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	46	87,138.79(2)	86,798.61(1)
48 87,150.03(3) 86,795.65(2) 49 87,155.66(3) - 50 87,161.39(4) -	47	87,144.44(2)	86,797.09(1)
49 87,155.66(3) - 50 87,161.39(4) -	48	87,150.03(3)	86,795.65(2)
50 87,161.39(4) -	49	87,155.66(3)	-
	50	87,161.39(4)	-

¹In units of cm⁻¹ and with 1σ fitting uncertainties given in parentheses in units of the least-significant digit that are additional to a 0.02 cm⁻¹ systematic uncertainty. Lines undergoing small perturbations in the B¹\Sigma⁺ state are indicated with (*).

with a large splitting of *e*- and *f*-symmetry levels. An accurate treatment of this interaction is essential for determining a deperturbed value for the A(0) term origin and its mass dependence. All three spin substates of the $d^3\Delta(v = 4)$ level strongly perturb A(0) at mid-*J*. The *F*₃ substate induces the largest shifts because of the Hund's case (a) selection rule limiting the spin–orbit interaction of ${}^1\Pi$ and ${}^3\Delta$ states to their $\Omega = 1$ components, in this case



Figure 7. Reduced term values (in cm⁻¹) of the ${}^{12}C^{18}O A^{1}\Pi$ (v = 0) level and the perturbing rovibronic levels.

the F_3 level of the inverted $d^3\Delta$. Relaxation of this rule by spin-rotational mixing then permits the further interactions with F_1 and F_2 levels. The A(0)–a'(9) interaction is weaker, with associated level shifts of ~0.5 cm⁻¹ occurring between J = 33 and 39.

When comparing the perturbations evident in Figure 7 with similar data and figures addressing other isotopologues (e.g. ${}^{12}C^{16}O$ [25,50], ${}^{13}C^{16}O$ [28,51], and ${}^{13}C^{17}O$ [30]) the crossings of A(0) with higher-v e(1), d(4), a'(9) levels all occur at decreasing *J* with increasing reduced-mass, μ , as is expected from the $-\sqrt{\mu}\omega_e(v + \frac{1}{2})$ proportionality of first-order isotope shifts. Conversely, the level crossings of A(0) and D(0) occur at higher *J* for higher mass because of the smaller vibrational constant, ω_e , of D ${}^{1}\Delta$.

To quantify the interaction energies and deperturbed molecular constants of all directly and indirectly observed levels we employ the same methodology as Niu et al. [25] and Hakalla [30] applied to other isotopologues. We used 476 transition frequencies obtained from the laser, VIS-FT and VUV-FT studies of A-X(0, 0), B-A(0, 0)0) and B-X(0,0) transitions and their satellite extra lines. We computed level energies and transition frequencies in a matrix diagonalisation of all interacting levels. The effective Hamiltonian used has matrix elements described in Niu et al. (see Table 6 in [25]) and the deperturbation was performed using the Pgopher software [41]. The interpretation of the interaction parameters ξ and η was also discussed in [25] and in [29]. The rotational structure of the ground state was fixed to the ¹²C¹⁸O constants of Coxon and Hajigeorgiou [23] throughout the analysis in the present paper and are listed in a footnote to Table 7.

The different sets of experimental data have different accuracies so we use relative weights to integrate them into a single molecular-parameter fit. A relative weight of 5 is assigned to data obtained from 2 + 1' REMPI

Constant	$A^{1}\Pi(v=0)$ (this work)	$A^{1}\Pi(v=0)$ Ref. [12] ^e	$B^1\Sigma^+(\nu=0)$ (this work)	$D^1 \Delta(v=0)$ (this work)	$I^1 \Sigma^- (\nu = 0)$ (this work)	$1^{1}\Sigma^{-}(v=1)$ (this work)
T _v	64755.71358(61)	64755.7020(92)	86916.6923(15)	65435.784 ^f	64558.877 ^h	65604.206 ^h
B	1.527662(10)	1.527624(29)	1.8554295(50)	1.18878 ^f	1.20084 ^h	1.184498 ^h
$D imes 10^6$	6.610(19)	6.700(16)	6.1022(28)	6.38 ^f	6.21 ^g	6.23 ^g
$H \times 10^{12}$	— 36.6(85)			- 0.26 ^g	2.59 ^g	2.59 ^g
ξ				0.0229(48)		0.0944(24)
ξ_{theoret}				0.0247	- 0.04104	0.06942
δ_{ξ}^{a}				7.1		36.0
Constant	$e^{3}\Sigma^{-}(v=1)$	$e^{3}\Sigma^{-}(v=1)$	$d^3 \Delta (v = 4)$	$d^3 \Delta (v = 4)$	$a'^{3}\Sigma^{+}(v=9)$	$a'^{3}\Sigma^{+}(v=9)$
	(this work)	Ref. [12] ^e	(this work)	Ref. [12] ^e	(this work)	Ref. [12] ^e
$T_{\nu}(F_1)$	64787.6689(13)	64787.651(10)		64982.44(13)	65281.120(11)	65281.14(63)
$T_v(F_2)$			65015.4993(38)	65014.71(19)		
$T_v(F_3)$				65048.25(12)	:	
В	1.197568(17)		1.1759189(60)		1.1287 ^j	
A	0 5011/07)	0 5 () (17)	- 16.5441(41)		1 15	
λ	0.5311(27)	0.562(17)	1.257(17) oi		- 1.15	
$\gamma \times 10^{\circ}$	- 2.11(25)		- 9 ⁰		E 01İ	
$D \times 10^{12}$	0.10 1.72 ⁰		5.96 ⁹		0.0P	
$\Lambda \times 10^4$	- 1./5		- 0.09 ⁹ - 0.1j		- 0.55	
$n_D \wedge 10$	14 7564(21)	14 780(14) ⁱ	- 0.1° - 21 9065(53)	_ 21 273(36) ⁱ	_ 2 538(22)	$-254(14)^{i}$
", c	14 6659	14 6659	- 21.9003(33) - 21.7426	- 21.273(30) - 21.7426	- 2.550(22) - 2.554	— 2.54(14) — 2.554
$\delta^{\prime\prime}$ theoret	0.6	0.8	0.8	21,720	0.6	0.5
γ_{η}	0.0	0.0	0.0	<i></i>	0.0	0.5

Table 7. Deperturbed molecular parameters of ¹²C¹⁸O for the A(0) state and its perturbers, and B(0).^{*a*, *b*} Perturbation parameters as discussed in Refs. [25,29].

^aAll values are in cm⁻¹ except for relative errors which are given as percentages.

^bMolecular constants fitted during the model optimisation have uncertainties indicated in parentheses (1 σ , in units of the least significant digit). All other parameters were fixed during the fitting procedure. The ¹²C¹⁸O fundamental ground state level, X($\nu = 0$), was fixed to the following constants determined by Coxon and Hajigeorgiou [23]: $G_{\nu} = 1055.7172740$, $B_{\nu} = 1.830980706$, $D_{\nu} = 5.55078179 \times 10^{-6}$, $H_{\nu} = 5.01843 \times 10^{-12}$, $L_{\nu} = -3.0008 \times 10^{-17}$, $M_{\nu} = -3.80 \times 10^{-23}$, $N_{\nu} = -6.0 \times 10^{-28}$, and $O_{\nu} = -3.0 \times 10^{-33}$ cm⁻¹.

^cTheoretical rotation and spin-orbit interaction parameters calculated on the basis of isotopically-invariant scaling and calculation of vibrational overlap integrals for ¹²C¹⁸O.

^dDifference between theoretical value, based on isotopic scaling, and the experimentally determined value in terms of a percentage: $\delta_{\eta} = (\eta_{\text{theoret}} - \eta)/\eta_{\text{theoret}} \times 100.$

^eWe only list the free parameters fitted in [12].

^fObtained by isotopic scaling values taken from [52].

^gObtained by isotopic scaling values taken from [50].

^hObtained by isotopic scaling values taken from [53].

^{*i*}Recalculated to the current definition of the spin-orbit parameter, η , using the relationship $\eta = \alpha \times \sqrt{3}$.

^jObtained by isotopic scaling values taken from [12].

laser spectroscopy; 2 to isolated and strong lines in the VIS-FTS spectrum, and 1–0.5 for weak and/or blended transitions and 1 to isolated and strong lines in the the VUV-FTS spectrum and 0.5–0.1 to weak and/or blended transitions.

Fitted deperturbed molecular constants and interaction parameters are given in Table 7 with the same parameter definitions as in Niu *et al.* [25]. The root-meansquare error (RMSE) of unweighted transition-frequency residuals is 0.014 cm⁻¹ after finalising the deperturbation fit. This measure of the overall quality-of-fit of the model to the experimental data is dominated by uncertainty in the least accurate $B^1\Sigma^+ - X^1\Sigma^+(0, 0)$ lines. This demonstrates an overall high level of agreement between the experimental and model frequencies. A correlation matrix was calculated during the least-squares fitting procedure and checked to ensure satisfactorily low correlation coefficients between the model parameters.

Figure 8 shows the residual error of the deperturbation model when compared with experimental frequencies. Histograms for each data-set contributing to the deperturbation fit have a width representative of the random uncertainties for each experiment. As expected, the laser-absorption REMPI frequencies are least scattered; all values are within 0.003 cm^{-1} (see Table 1). The greater scatter of VUV-FTS data in the A-X(0, 0)region relative to B-X(0, 0) is due to a combination of the somewhat better instrumental resolution adopted for measurements of the latter and the greater number of weak and overlapped lines present in the former. A small +0.005 cm⁻¹ bias of the VUV-FTS B-X(0,0) model residuals indicates a slight disagreement of the absolute calibration of this experiment relative to the others with a superior calibration to the REMPI laser experiment. Actually, under optimal conditions the VUV-FTS experiment is limited to an accuracy of 0.01 cm⁻¹ [24], which



Figure 8. Residual error of modelled line frequencies after deperturbation, shown separately for each experimental measurement. The histograms consist of residuals for the main bands and all observed extra lines.

is significantly larger than the systematic offset found here.

Modelled energies for all bands and excited-state term values are given as an online data archive. All bands and levels are extrapolated to J = 60 but increasing errors are expected for levels which are not well constrained by the observed lines and crossings. There are also archived experimental spectra and the Pgopher input file used to perform our deperturbation.

The level mixing deduced from deperturbing the interaction of A(0) and its perturbers also provides a means to calculate the linestrength borrowed by extra lines that appear in the experimental $B \rightarrow A$ and $A \leftarrow X$ spectra. Modelled band *f*-values are compared with $A \leftarrow X(0, 0)$ absorption *f*-values in Figure 5 and show good agreement, further validating the deperturbation model deduced from level energies.

The careful absolute calibration of our laser-based measurements with respect to the ground state energy means that the value for the deperturbed A(0) T_{ν} constant can be determined at high accuracy. Least-squares fitting of the deperturbation model returns a statistical uncertainty of 0.0006 cm⁻¹ for this parameter. However, this statistical uncertainty is smaller than the 0.001 cm⁻¹ absolute frequency uncertainty of our data (tied to the

REMPI-laser experiment) and we conclude that the estimated uncertainty of T_0 is 0.001 cm⁻¹. We list all residual differences of laser-measured and modelled frequencies in Table 1 and these show agreement between 0.0001 and 0.003 cm⁻¹.

A deperturbation analysis of ¹²C¹⁸O A(0) and perturbers was previously performed by Beaty et al. [12] based on comprehensive emission spectra of A-X(0, 0)and extra lines from perturbing states, with their deperturbed constants also listed in Table 7. A similar scale of experimental information was used in that previous work but with frequency uncertainties of 0.05 cm⁻¹ or more. Our new measurements, with uncertainties between about 0.001 and 0.02 cm⁻¹, enable the determination of more constants defining the A(0) and perturber levels than previously, and with better precision. All comparable parameters show consistency between this work and that of Beaty et al. [12], given their estimated uncertainties, apart from the A(0) and d(4) term origins and the $\eta_{A(0) \sim d(4)}$ interaction parameter. The T_{ν} discrepancies likely arise from correlation of this parameter with high-order rotational constants and our inclusion of an additional constant, H, relative to Beaty et al. [12].

Some parameters in Table 7 are experimentally determined for the first time, for example, the ξ parameters that control the L-uncoupling matrix elements $\langle A(0)|J_+L_-|D(0)\rangle$ and $\langle A(0)|J_+L_-|I(1)\rangle$. We do not experimentally observe spectral lines associated with D(0)nor I(1) and their avoided crossings with A(0) occur above our highest-observed J level. Fixed molecular constants representing the D(0) and I(1) levels are nevertheless included in our perturbation analysis by isotopically rescaling their electronic-state equilibrium constants found in the literature [52-55]. The D(0)-A(0) and I(1)-A(0) interaction parameters then exhibit quantitatively well-defined values $\xi_{A(0) \sim D(0)} = 0.0229(48) \text{ cm}^{-1}$ and $\xi_{A(0) \sim I(1)} = 0.0944(24)$ cm⁻¹ when included in the deperturbation optimisation. The $\xi_{A(0)} \sim D(0)$ parameter is consistent within its uncertainty with a value calculated from the isotopically invariant rotationalinteraction parameter determined by Lefloch et al. [50] (parameter b in their Table IV). A similar mass-scaling calculation is described by Hakalla [30]. The newly determined $\xi_{A(0)} \sim I(1)$ does not show such good agreement, differing by 36% from its rescaled value. This may be due to a poorer determination of this parameter in our deperturbation than suggested by its estimated uncertainty. The crossing point of A(0) and I(1) rotational series in ${}^{12}C^{18}O$ occurs near J = 49 while the maximum J of our experimentally constrained term values is significantly lower J = 44, suggesting the possibility of some uncertainty.

The isotopologue-independent $A^1\Pi \sim a'^3\Sigma^+$ perturbation parameter has a typical magnitude for spin–orbit interactions in CO [30], 84.02(10) cm⁻¹, and the weakness of the fitted A(0)– a'(9) interaction energy is then due to a relatively small overlap of vibrational wavefunctions. We calculate a normalised value for this matrix element, $\langle v_{A(0)} | v_{a'(9)} \rangle = 0.071$ using the method of Refs. [29,30] and find a value similar to other CO isotopologues, e.g. 0.070 and 0.072 with respect to ${}^{12}C{}^{17}O$ and ${}^{13}C{}^{17}O$.

6. Conclusion

Three different state-of-the-art techniques were employed to gather transition frequencies of the $A^{1}\Pi - X^{1}\Sigma^{+}(0, 0)$, $B^{1}\Sigma^{+} - X^{1}\Sigma^{+}(0, 0)$, and $B^{1}\Sigma^{+} - A^{1}\Pi(0, 0)$ bands in ${}^{12}C^{18}O$ with the highest resolution and accuracy obtained to date. The spectra show many extra lines probing optically forbidden transitions to the $d^{3}\Delta(4)$, $e^{3}\Sigma^{-}(1)$, and $a'^{3}\Sigma^{+}(9)$ levels, due to their interaction with $A^{1}\Pi(0)$ by spin-orbit and rotational-electronic *L*-uncoupling. This spectroscopic data was included into an effective Hamiltonian model of all observed levels and their interactions.

Lines probing the $d^{3}\Delta(4)$, $e^{3}\Sigma^{-}(1)$, and $a'^{3}\Sigma^{+}(9)$ triplet state perturbers were observed directly, while interactions with the further remote perturber states $D^{1}\Delta(0)$ and $I^{1}\Sigma^{-}(1)$ were found to measurably affect the A(0) term values. As expected, the extents and strengths of perturbations arising in $A^{1}\Pi(0)$ in ${}^{12}C^{18}O$ level are similar to the cases of ${}^{12}C^{16}O$ [25], ${}^{13}C^{16}O$ [28], and ${}^{13}C^{17}O$ [30].

The deperturbed term origin of $A^1\Pi(\nu = 0)$ is found with a precision of 0.001 cm⁻¹. This term origin will be compared with our past and future measurements in other CO isotopologues to determine the precise magnitude of the $A^1\Pi - X^1\Sigma^+(0, 0)$ electronic isotopic shift.

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Disclosure statement

No potential conflict of interest was reported by the authors.

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References

- [1] R.T. Birge, Phys. Rev. 28, 1157-1181 (1926).
- [2] G. Herzberg, Zeitschr. f. Physik 52, 815–845 (1929).
- [3] R.W. Field, S.G. Tilford, R.A. Howard, and J.D. Simmons, J. Mol. Spectr. 44(2), 347–382 (1972).
- [4] R.W. Field, B.G. Wicke, J.D. Simmons, and S.G. Tilford, J. Mol. Spectr. 44(2), 383–399 (1972).
- [5] M. Rytel, Acta Phys. Polon. A 38, 299 (1970).
- [6] J. Janjić, J. Danielak, R. Kepa, and M. Rytel, Acta Phys. Polon. A 41, 757 (1972).
- [7] J. Janjić, L.U. Cŏnkić, D.S. Pesiš, R. Kepa, and M. Rytel, J. Mol. Spectr. 72(2), 297–300 (1978).
- [8] R. Kepa, M. Rytel, and Z. Rzeszut, Acta Phys. Polon. A 54, 355–361 (1978).
- [9] R. Kepa, Acta Phys. Hung. 60, 227-237 (1986).
- [10] C. Haridass, S.P. Reddy, and A.C. Le Floch, J. Mol. Spectr. 168(2), 429–441 (1994).
- [11] C. Haridass, S.P. Reddy, and A.C. Le Floch, J. Mol. Spectr. 167(2), 334–352 (1994).
- [12] L.M. Beaty, V.D. Braun, K.P. Huber, and A.C. Le Floch, Ap. J. Suppl. Ser. **109**(1), 269 (1997).
- [13] C.M. Steinmann, E.G. Rohwer, and H. Stafast, Astrophys. J. Lett. **590**(2), L123 (2003).
- [14] A. du Plessis, E.G. Rohwer, and C.M. Steenkamp, Astrophys. J. Suppl. Ser. 165(1), 432 (2006).
- [15] M. Eidelsberg, J.Y. Roncin, A.L. Floch, F. Launay, C. Letzelter, and J. Rostas, J. Mol. Spectr. **121**(2), 309–336 (1987).
- [16] G. Stark, A.N. Heays, J.R. Lyons, P.L. Smith, M. Eidelsberg, S.R. Federman, J.L. Lemaire, L. Gavilan, N. de Oliveira, D. Joyeux, and L. Nahon, Astroph. J. 788(1), 67 (2014).
- [17] M. Eidelsberg, J.L. Lemaire, S.R. Federman, G. Stark, A.N. Heays, L. Gavilan, J.R. Lyons, P.L. Smith, N. de Oliveira, and D. Joyeux, Astron. Astroph. 566, A96 (2014).
- [18] P. Cacciani, W. Ubachs, P.C. Hinnen, C. Lyngå, A. L'Huillier, and C.G. Wahlström, Astroph. J. Lett. 499(2), L223 (1998).

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- [19] W. Ubachs, I. Velchev, and P. Cacciani, J. Chem. Phys. 113(2), 547 (2000).
- [20] G. Klapper, F. Lewen, R. Gendriesch, S.P. Belov, and G. Winnewisser, Zeitschr. f
 ür Naturf. A 56(3-4), 329-332 (2014).
- [21] G. Cazzoli, C. Puzzarini, and A.V. Lapinov, Astrophys. J. Lett. **592**(2), L95 (2003).
- [22] P. de Natale, M. Inguscio, C.R. Orza, and L.R. Zink, Astroph. J. Lett. **370**, L53–L55 (1991).
- [23] J.A. Coxon and P.G. Hajigeorgiou, J. Chem. Phys. 121(7), 2992–3008 (2004).
- [24] E.J. Salumbides, M.L. Niu, J. Bagdonaite, N. de Oliveira, D. Joyeux, L. Nahon, and W. Ubachs, Phys. Rev. A 86, 022510 (2012).
- [25] M.L. Niu, E.J. Salumbides, D. Zhao, N. de Oliveira, D. Joyeux, L. Nahon, R.W. Field, and W. Ubachs, Mol. Phys. 111, 2163–2174 (2013).
- [26] M.L. Niu, F. Ramirez, E.J. Salumbides, and W. Ubachs, J. Chem. Phys. 142, 044302 (2015).
- [27] M.L. Niu, E.J. Salumbides, A.N. Heays, N. de Oliveira, R.W. Field, and W. Ubachs, Mol. Phys. 114(5), 627–636 (2016).
- [28] M.L. Niu, R. Hakalla, T.M. Trivikram, A.N. Heays, N. de Oliveira, E.J. Salumbides, and W. Ubachs, Mol. Phys. 114(19), 2857–2867 (2016).
- [29] R. Hakalla, M.L. Niu, R.W. Field, E.J. Salumbides, A.N. Heays, G. Stark, J.R. Lyons, M. Eidelsberg, J.L. Lemaire, S.R. Federman, M. Zachwieja, W. Szajna, P. Kolek, I. Piotrowska, M. Ostrowska-Kopec, R. Kepa, N. de Oliveira, and W. Ubachs, Roy. Soc. Chem. Adv. 6, 31588– 31606 (2016).
- [30] R. Hakalla, M.L. Niu, R.W. Field, A.N. Heays, E.J. Salumbides, G. Stark, J.R. Lyons, M. Eidelsberg, J.L. Lemaire, S.R. Federman, N. de Oliveira, and W. Ubachs, J. Quant. Spectr. Rad. Transfer 189, 312–328 (2017).
- [31] A.J. de Nijs, E.J. Salumbides, K.S.E. Eikema, W. Ubachs, and H.L. Bethlem, Phys. Rev. A 84, 052509 (2011).
- [32] A.J. de Nijs, D. Zhao, H.L. Bethlem, H. Linnartz, and W. Ubachs, J. Mol. Spectr. 292, 20–22 (2013).
- [33] W. Ubachs, K.S.E. Eikema, W. Hogervorst, and P.C. Cacciani, J. Opt. Soc. Am. B 14(10), 2469–2476 (1997).
- [34] S. Xu, R. van Dierendonck, W. Hogervorst, and W. Ubachs, J. Mol. Spectr. 201, 256–266 (2000).
- [35] E.E. Eyler, A. Yiannopoulou, S. Gangopadhyay, and N. Melikechi, Opt. Lett. 22, 49–51 (1997).

- [36] B. Girard, N. Billy, J. Vigue, and J.C. Lehmann, Chem. Phys. Lett. **102**(2–3), 168–173 (1983).
- [37] R. Hakalla, W. Szajna, and M. Zachwieja, J. Phys. B 45(21), 215102 (2012).
- [38] R. Hakalla, M. Zachwieja, and W. Szajna, J. Phys. Chem. A 117(47), 12299–12312 (2013).
- [39] R. Hakalla, Roy. Soc. Chem. Adv. 4, 44394-44407 (2014).
- [40] R. Hakalla, M. Zachwieja, and W. Szajna, J. Quant. Spectr. Rad. Trans. 140, 7–17 (2014).
- [41] C.M. Western, Pgopher: a program for simulating rotational structure (University of Bristol, Bristol, 2017).
- [42] N. de Oliveira, M. Roudjane, D. Joyeux, D. Phalippou, J.C. Rodier, and L. Nahon, Nat. Photon. 5(3), 149–153 (2011).
- [43] N. de Oliveira, D. Joyeux, M. Roudjane, J.F. Gil, B. Pilette, L. Archer, K. Ito, and L. Nahon, J. Synch. Rad. 23, 887– 900 (2016).
- [44] J.L. Lemaire, M. Eidelsberg, A.N. Heays, L. Gavilan, S.R. Federman, G. Stark, J.R. Lyons, N. de Oliveira, and D. Joyeux, J. Phys. B 49(15), 154001 (2016).
- [45] M.L. Niu, A.N. Heays, S. Jones, E.J. Salumbides, E.F. van Dishoeck, N. de Oliveira, L. Nahon, and W. Ubachs, J. Mol. Spectr. 315, 137–146 (2015).
- [46] H. Lefebvre-Brion and R.W. Field, *The Spectra and Dynamics of Diatomic Molecules* (Elsevier, Amsterdam, 2004).
- [47] M. Larsson, Astron. Astrophys. 128, 291-298 (1983).
- [48] G. Herzberg, Molecular Spectra and Molecular Structure I: Spectra of Diatomic Molecules, 2nd ed. (Krieger Publishing Company, New York, NY, 1989).
- [49] M. Drabbels, J. Heinze, J.J. Ter Meulen, and W.L. Meerts, J. Chem. Phys. 99(8), 5701–5711 (1993).
- [50] A.C. Le Floch, F. Launay, J. Rostas, R.W. Field, C.M. Brown, and K. Yoshino, J. Mol. Spectr. **121**(2), 337–379 (1987).
- [51] C. Haridass and K.P. Huber, Astroph. J. 420, 433–438 (1994).
- [52] C. Kittrell and B.A. Garetz, Spectrochim. Acta 1(45), 31– 40 (1989).
- [53] R.W. Field, Ph. D. thesis, Harvard University, 1971.
- [54] S.G. Tilford and J.D. Simmons, J. Phys. Chem. Ref. Data 1, 147–188 (1972).
- [55] K.P. Huber and G. Herzberg, Constants of diatomic molecules. (Van Nostrand Reinhold, New York, NY, 1979).