The high resolution spectrum of methyltrioxorhenium reinvestigated with new infrared and millimeter-wave measurements

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Following our first paper about high resolution spectroscopy of methyltrioxorhenium (MTO) [Stoeffler et al. PCCP, **13**, 854, (2011)], the present study reports a deeper investigation of the ground state, and Re=O antisymmetric (v_{as}) and symmetric (v_s) stretching excited states of both CH_3^{187} ReO₃ and CH_3^{185} ReO₃ isotopologues, thanks to new devices implemented within our consortium. We carry out high resolution millimeter-wave (MMW) and infrared (IR) spectroscopy in room temperature absorption cells, in a pulsed supersonic jet and in a cryogenic buffer gas cell. This collection of sensitive spectrometers enables us to probe both levels of a vibrational transition in low and room temperature gaseous environments. We thus report a new series of measurements providing particularly accurate rotational and rovibrational data for such a large and heavy organometallic molecule that is solid at room temperature.

The combination of the new MMW and IR data leads to an improvement of the rovibrational model of MTO: (i) rotational spectra are extended to the 150-300 GHz range and are analyzed to characterize the ground state rotational and hyperfine structure up to $J =$ 43 and $K = 41$, resulting in the significant refinement of rotational, quartic and hyperfine parameters, as well as the determination of sextic parameters and a centrifugal distortion correction to the quadrupolar hyperfine constant,

(ii) rovibrational data recorded at very different temperatures (between 6 and 300 K) in the 970-1015 cm^{-1} range, at resolutions down to 8 MHz and frequency accuracies as high as 30 MHz (using the buffer gas technique), are used in a first step to extract more precise excited-state rotational, Coriolis and quartic parameters as well as the ground-state centrifugal distortion parameter D_K of the ¹⁸⁷Re isotopologue, which isn't accessible from rotational spectroscopy alone. In a second step, taking the hyperfine structure into account, the upper state rhenium atom quadrupole coupling constant *eQq'* is determined from the assignment of partially hyperfine-resolved CO² laser absorption spectra of a skimmed molecular beam.

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I- Introduction

There is an increasing demand for precision spectroscopy of polyatomic molecules. For example, precise knowledge of molecular constants is required for modeling atmospheres, whether it be our own^{1,2} or those found in exo-planets³ and in the interstellar medium⁴, for studying collision physics^{5,6}, and as crucial groundwork for performing fundamental physics tests7,8,9,10,11,12,13,14,15,16,17 .

One such test is the measurement of the parity-violating (PV) energy difference between opposite enantiomers of a chiral molecule, induced by the weak interaction. ¹⁸ The chiral derivatives of methyltrioxorhenium (MTO) are excellent candidate species for this measurement, because calculations predict relatively large PV energy differences, and because it is possible to synthesize these molecules and bring them into the gas phase. ^{[8,](#page-1-0)[17,1](#page-1-1)9,20}

In the present work, we obtain much new spectroscopic information about MTO. A few years ago, a first set of measurements studied the ground and first excited states of the v_{as} antisymmetric Re=O stretching mode of MTO. In that work^{[19](#page-1-2)} we reported microwave spectra obtained using a Fourier transform microwave (FTMW) spectrometer, Fourier transform infrared (FTIR) and laser absorption experiments both in cells and in supersonic beams. By combining these data, we constructed a model for MTO. Here, we report new experiments with largely improved sensitivity, resolution and spectral coverage leading to definite assignments of both Re=O stretching rovibrational spectra and to a substantial refinement of the model. New and better resolved data allow us to improve the accuracy of some spectroscopic constants and to determine new higher-order interaction terms. Importantly, we establish a working procedure for extracting the most precise molecular parameters from a large amount of spectroscopic data spanning centimeter to infrared wavelengths, taken at different resolutions from a variety of different experiments.

The paper is organized as follows. Section II describes new apparatus used for this work. These comprise the incorporation of an external-cavity quantum cascade laser (EC-QCL) into a pulsed supersonic jet setup, a distributed-feedback quantum cascade laser (DFB-QCL) and its use for acquiring spectra from laser-ablated molecules in a cryogenic cell and a new millimeter wave (MMW) spectrometer. Section III describes new spectra recorded in the 150- 300 GHz range, which contribute to an improved understanding of the vibrational ground state with parameters derived up to the sextic centrifugal distortion contributions. Section IV reports full analyses of new vibrational spectra of the v_{as} antisymmetric and v_{s} symmetric Re=O stretches recorded in the mid-infrared. Those were obtained *via* laser spectroscopy of molecules cooled in a pulsed jet, a skimmed continuous molecular beam and a cryogenic helium buffer gas cell, or *via* FTIR spectroscopy in a long path absorption cell at room temperature. Section IV also discusses the resulting refinements made to the model and to the analysis of the hyperfine structure (HFS) of the upper vibrational state of MTO

.**II- Experimental details**

MTO powder (from Strem Chemicals Inc., 98% purity) has been used without purification in all the experiments described in this paper.

II-1 *Millimetre-wave experiments:* **room temperature millimetre-wave spectroscopy in a cell**

The spectrum of MTO was recorded in the 150-300 GHz range on the Lille millimeter-wave spectrometer²¹ at room temperature (294 K), with a vapour pressure of about 20 bar. The frequency source is a 20-GHz synthesizer (Agilent, model E8257D), stabilized on the Global Positioning System. The output frequency (12.5-18.4 GHz) is multiplied and amplified by an active sextupler AMC-10 (Virginia Diodes Inc.) providing an output power of +14 dBm in the W-band ranging from 75 to 110 GHz. Passive Schottky multipliers (\times 2, \times 3, \times 5, \times 3 \times 2, \times 3 \times 3, Virginia Diodes Inc.) are used in the last stage of the frequency multiplication chain in order to provide a useful signal in the 150-990 GHz spectral range. The absorption cell is a stainless-steel tube (6-cm diameter, 220-cm path length). In order to improve the signal sensitivity, the source is frequency modulated at 10 kHz and second harmonic detection in a lock-in amplifier is carried out. Absorption signals are detected with an InSb liquid He-cooled bolometer (QMC Instruments Ltd.). The absolute accuracy of the line-center's frequency is estimated to be better than 30 kHz for isolated lines, but worsen to 100 kHz for blended or very weak lines.

II-2 *IR experiments*

II-2-1 The extended cavity quantum cascade laser spectrometer coupled to a pulsed jet

The v_{as} band of MTO has been recorded at high resolution using an extended cavity quantum cascade laser (EC-QCL) based spectrometer that includes a multipass absorption device recently developed at MONARIS. The experimental setup coupling this mid-infrared spectrometer to a pulsed supersonic jet is shown in Fig. 1. The apparatus consists in three main parts: 1) The laser and optical system, 2) The vacuum system and supersonic jet, 3) The data acquisition electronics and software.

II-2-1-a The laser and the optical system

The light source is a continuous-wave room-temperature mode-hop-free EC-QCL which covers the 930-990 cm⁻¹ range (Daylight Solutions). The QCL chip and a diffraction grating mounted on a piezoelectric transducer (PT) form an external cavity, and high-resolution measurements are obtained by scanning the length of this cavity. This is done by applying a sine wave with an amplitude of up to 80 V to the PT at frequencies up to 100Hz via an external controller (MDT693B, Thorlabs).

About 7.5% of the light is sent through an etalon consisting of a 0.025 cm^{-1} freespectral-range confocal Fabry–Perot cavity, to provide a relative frequency scale. A linear interpolation of the positions of the etalon maxima establishes the relationship between the voltage applied to the PT and the relative frequency. Absolute laser frequencies are obtained by sending 7.5% of the light through a reference cell containing a known reference gas. A toroidal mirror (500-mm radius of curvature) mode matches the 85 % of the remaining laser beam to a multipass near-concentric Perry cavity²² (made of two spherical mirrors with a 125mm radius of curvature) placed inside the vacuum chamber. The laser light crosses the

supersonic expansion 20 times and focuses midway between the two mirrors at each path. Three liquid-nitrogen-cooled HgCdTe detectors (Fermionics Corporation PV 12-1 and Judson J15D12) are used to measure the powers transmitted through the multipass cavity, the etalon and the reference gas cell.

II-2-1-b The vacuum system and supersonic jet

The 400 mm diameter jet chamber (Kurt J. Lesker Company) is evacuated by a combination of a 2000 l/s oil diffusion pump (Edwards, Diffstak 250/2000), a roots blower (Sogev AR350) and a rotary pump (Edwards E2M40). The diffusion pump is equipped with water-cooled baffles in order to minimize back streaming of the oil vapour towards the multipass optics. The molecular jet is produced using a pulsed circular (1-mm diameter) nozzle from General Valve Series 9 model, controlled by a valve driver (Iota One, Parker Hannifin). MTO is seeded in the supersonic jet using a brass reservoir heated to 370 K (after a design from NIST²³ and PhLAM²⁴). The reservoir is filled with 0.5 g of MTO powder, and the sublimated vapour is swept away by argon at a backing pressure of 1.2 bar. Pulsed expansions are produced at a repetition rate of typically 1 Hz. The distance between the nozzle and the waist collectively formed by the reflected infrared beams inside the multipass cavity is between 3 and 7 mm.

II-2-1-c The data acquisition electronics and software

Our rapid scan scheme is similar to previous designs developed for high resolution molecular spectroscopy.^{25,26} The experiment is controlled by a computer running a Labview code. Spectra are recorded by driving the PT of the EC-QCL diffraction grating with a 100 Hz sine wave. This allows the QCL frequency to be scanned once per period over 0.8 $cm⁻¹$ in 5 ms. During this time window, the intensities transmitted through the supersonic jet, the etalon and the reference gas cell are digitized (12-bit acquisition card NI-PCI-6110, 5 MS/s) and recorded. A baseline-free transmittance through the multipass cavity is obtained by taking the ratio of signals recorded in presence and absence of the jet (by opening or not the nozzle).

The data acquisition sequence is illustrated in Fig. 2: the experiment starts with a TTL pulse generated from the timer card (NI-PCI-6602) that triggers both the subsequent laser sweeps (*Diffraction grating PT voltage* in Fig. 2) and 5-ms duration data acquisitions (*Acquisition triggers* and *Data acquisition* in Fig.2). It also sends a *Nozzle driver pulse* with delay d_1 to the Iota One controller to open the pulsed valve. For obtaining a good coincidence between the gas pulse, a laser sweep and the data acquisition, d_1 has to be chosen by taking into account d₂, the molecules' travelling times from the nozzle to the laser beams. A sequence typically consists of four background scans (in the absence of molecular jet), two taken before (scans 1 and 2 in Fig. 2) and two after (scans 4 and 5 in Fig. 2) an absorption scan (scan 3 in Fig. 2, in presence of the jet). The recorded scans are then transferred from the acquisition card to the computer to be processed and saved. This sequence is repeated at a user-defined rate of typically 1 Hz.

II-2-1-d Performances of the EC-QCL/pulsed jet spectrometer

The laser rapid-scan data show a remarkable improvement in signal-to-noise ratio (by a factor between 5 and 10) over the jet FTIR spectrum of our previous work.[19](#page-1-2)

The resolution (full-width-at-half-maximum, FWHM, of isolated rovibrational lines) of the jet-cooled QCL spectrum reported in this work (see Fig. 6 and 7 later) is about 120 MHz, limited by the Doppler broadening that results from the circular nozzle expansion geometry.[†] This is an improvement on the continuous jet-FTIR v_{as} spectrum of our previous work^{[19](#page-1-2)}, where the resolution was 160 MHz, resulting from a combination of a similar Doppler broadening (due to the use of the same geometry) and the instrumental resolution of 100 MHz. Furthermore, compared to continuous expansions, the higher backing pressures achievable with pulsed expansions allows us to obtain better rovibrational cooling. From the analysis of section IV-2-1, we estimate the MTO jet rotational temperature to be 6 ± 1 K, substantially lower than in the continuous supersonic expansion of our previous work $(\sim 10K)$. In our conditions, a good signal-to-noise ratio was obtained after 5000 gas pulses of 5 ms duration. This corresponds to a consumption of 200 mg of MTO, which is about 75 times less than for the continuous jet.

II-3 Room temperature FTIR spectroscopy in a cell

We recorded both the v_{as} and v_s bands of the two isotopologues of CH₃ReO₃ (centered at about 976 and 1004 cm^{-1} , respectively) using a room temperature absorption cell with a path length of 150 m that has already been described elsewhere.²⁷ The cell was coupled to a high resolution FTIR spectrometer (IFS 125 HR, Bruker) with a maximal resolution of 0.00102 cm⁻¹. For those experiments a globar source, a KBr beam splitter and an HgCdTe detector were used. Because of the large difference in intensity of the two Re=O stretching bands – v_{as} is calculated to be about 14 times more intense than v_s^{28} – the antisymmetric band was recorded after injecting 0.005 mbar of MTO for an acquisition time of about 36 hours while the symmetric one was recorded by injecting ~0.014 mbar for an acquisition time of about 49 hours.

Aside from focusing on these two stretching modes, we also recorded a wide infrared vibrational spectrum from 50 to 5000 cm^{-1} , which will be presented elsewhere.

II-4 The DFB QCL spectrometer coupled to a 6 K buffer-gas cell

Details of this buffer-gas cell and associated laser setup are to be found elsewhere,²⁹ and are thus only briefly reviewed here. A copper cell is filled with $\sim 10^{-2}$ mbar of helium and is cooled to 6 K using a cryocooler (Sumitomo RDK-415D). A target containing MTO is ablated with 30 mJ pulses of 8 ns from an Nd:YAG laser at 1064 nm. The resulting plume is rapidly cooled by collisions to the helium temperature, permitting spectroscopy at low temperatures for a few milliseconds, the typical diffusion time of molecules to the cold cell walls, where they freeze and are lost from the experiment. Light from a free-running Peltiercooled (to typically -10°C) DFB QCL laser emitting at ~976 cm⁻¹ was sent through the cell at right angles with the plume, and absorption spectra were recorded. In order to reconstruct a

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[†] FWHM as narrow as 50 MHz have recently been obtained on the same set-up equipped with a pulsed slit nozzle for a SF₆ expansion seeded in argon.

frequency axis, some of the light was also sent through reference cells containing ethene or methanol, and some through a temperature-stabilized germanium reference etalon. These signals allowed for an *a posteriori* reconstruction of a frequency scale with an absolute accuracy of about 30 MHz, and a relative uncertainty of a few megahertz. 29 29 29

This setup was used to investigate several regions of the v_{as} mode of CH₃ReO₃. We obtain a resolution of ~ 8 MHz (FWHM), limited by a combination of laser frequency fluctuations, collisional and Doppler broadenings.^{[29](#page-4-0)} We estimate the temperature of the molecules to be 6 ± 3 K^{[29](#page-4-0)} (see Fig. 8 later).

II-5 CO² laser linear absorption spectroscopy of a supersonic beam

Again, details of the setup can be found elsewhere. ^{[9,](#page-1-3)[19](#page-1-2)} A mixture of MTO and He is prepared in an oven at \sim 370 K. This mixture is sent to a 100 μ m nozzle, through which it is expanded into vacuum to form a continuous supersonic beam. Downstream of a skimmer, a CO² laser crosses this beam at right angles for linear absorption spectroscopy. In order to enhance the signal to noise ratio, the light crosses the beam 18 times using a multi pass cell. A rotating slotted disk was used to chop the molecular beam, resulting in a ∼1 kHz modulation of the absorption, detected with a lock-in amplifier. The $CO₂$ laser is locked via a phase lock loop to a second CO_2 laser, which is in turn locked to a saturated absorption signal of $OsO₄$ in the vicinity of the $R(20)$ CO₂ emission line. Using this locking scheme, we know the absolute frequency of the laser to within less than 100 Hz and obtain a stability at 1 s and a line width of respectively 1 Hz and 10 Hz.^{30,31} With this apparatus, we probed a region of the v_{as} transition of MTO at around 976 cm^{-1} . From the analysis in section IV-2-3, we estimate the MTO beam rotational temperature to be about 15 K (see Fig. 9 later).

III- Millimetre-wave rotational spectroscopy: Results and analysis

Compared to our previous work, we extend here the rotational analysis of the MTO spectrum in the millimeter wave region. The rotational and hyperfine structure of both CH₃¹⁸⁷ReO₃ and CH₃¹⁸⁵ReO₃ is characterized up to $J = 43$ and $K = 41$ in the ground state (with *J* and *K*, the quantum numbers associated with the total orbital angular momentum and its projection on the molecular symmetry axis respectively). An example spectrum is presented in Fig. 3. The ground state of both isotopologues is characterized using a standard Hamiltonian that takes into account the rotational structure (B) , a few quartic (D_J, D_{JK}) and sextic (*H_J*, *H_{JJK}*) centrifugal distortion parameters, the spin-rotation constants (C_{aa} , C_{bb}), and the rhenium atom hyperfine quadrupole coupling constant (eQq) , and its centrifugal distortion correction $(eQqJ)$. The SPFIT/SPCAT suite of programs of H. Pickett is used to determine molecular parameters from the frequencies of assigned transitions.³² 966 and 530 lines have been assigned for the ¹⁸⁷Re and ¹⁸⁵Re isotopologues, respectively. Of these, 46 and 19 lines for which the difference between experimental and calculated frequency is bigger than 3 times the standard deviation have been rejected in the analysis. The corresponding line lists (with the rejected lines removed) are available as ESI (ESI1). A new set of parameters has been obtained and is presented in Table 1. The value of A was fixed to the value obtained in the structural determination of Wikrent et al.³³ The standard deviation of the difference between experimental and calculated frequencies is about 40 kHz, close but higher than the experimental accuracy because of the presence of several overlapped lines. Note that 21 rotational lines assigned to each isotopologue in the microwave (μW) study reported in our previous work^{[19](#page-1-2)} have been included in the present analysis.[‡]

IV- Mid-infrared rovibrational spectroscopy: Results and analysis IV-1 Experimental results

We have recorded new high resolution spectra of both the symmetric v_s and antisymmetric v_{as} Re=O stretching bands of MTO.

Both the v_s and v_{as} modes have been probed at a resolution of 0.00102 cm⁻¹ using the room-temperature cell-FTIR setup described in Section II-3. Fig. 4 shows the full v_s band while Fig. 5 shows a zoom of five rotational progressions of the *R*-branch. v_s being a parallel band, it exhibits evident rotational progressions in *J* which makes it possible to directly interpret the room temperature cell data and assign transitions to it. On the other hand, a direct interpretation of the room temperature v_{as} perpendicular band spectrum is out of reach. MTO has four vibrational modes below 600 cm^{-1} and, being heavy, possesses small rotational constants. Its room temperature v_{as} spectrum is therefore very dense given the large number of occupied rotational states and the presence of several hot bands. ³⁴ Typical rotational progressions in *K* are thus blurred. We therefore resort to the supersonic jet technique in which the rovibrational cooling simplifies the spectrum. Fig. 6 shows the jet-cooled EC-QCL spectrum of the full v_{as} band recorded with the setup described in Section II-2-1 and Fig. 7 is a zoom of rotational progressions of the *P*-branch.

Moving to higher resolution, linear absorption spectra of laser ablated MTO buffergas-cooled to ~6 K have been recorded as described in Section II-4. Fig. 8 shows an example spectrum obtained by tuning the free-running DFB QCL frequency to the *P*-branch of the v_{as} mode, where isolated rovibrational lines can be observed. The resolution is about 8 MHz. As illustrated in Fig. 8, this allows neighbouring rovibrational lines of the two isotopologues to be resolved, an improvement on the two spectroscopic studies of supersonic jet-cooled MTO mentioned in the previous paragraph. The hyperfine structure is also partially resolved, and will be discussed below. The inset of Fig. 8 shows spectroscopy of the *Q*-branch of the ¹⁸⁷Re MTO v_{as} mode (here, the splitting in *J* is far smaller and is thus unresolved).

Finally, Fig. 9 shows examples of $CO₂$ laser linear absorption spectra of a continuous helium supersonic beam seeded with a few percent of MTO as described in Section II-5. The structures on the low frequency side correspond to the ${}^R O(J,2)$ band, for *J* between 2 and 10. At these low *J* values, the expected regular rovibrational structure is strongly modified by the hyperfine interaction between the rhenium nuclear quadrupole moment and the gradient of the electric field, resulting in a dense and complex spectrum. On the high frequency side, high *J* transitions of the R *Q*(*J*,1) band exhibit reduced hyperfine splitting, and the amplitudes are weaker in accordance with the Boltzmann distribution at 15 K. As will be shown later, these

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[‡]Amongst those, only 16 lines for the ¹⁸⁷Re isotopologue and 17 lines for the ¹⁸⁵Re one have been kept for the present analysis (see ESI1).

data will be used to refine our model and estimate *eQq'*, the quadrupole coupling constant in the excited vibrational state.

IV-2 Rovibrational analysis of the Re=O stretching bands

MTO is a prolate symmetric top with C_{3v} symmetry and 18 vibrational modes³⁵. In the 1000 cm⁻¹ region two stretching modes are infrared active: the $v_{as}(E)$ and $v_s(A_1)$ bands measured at 959 and 998 cm^{-1} respectively in the solid phase³⁶, the rovibrational analysis of which is detailed in the following sections.

Experimental spectra of MTO are analyzed with the prolate symmetric rovibrator model, including the rotational constants A and B, the centrifugal distortion constants D_J , D_{JK} and D_K and sextic constants H_I and H_{IJK} . Ground state rotational energy levels are given by:

$$
E(J,K) = BJ(J+1) + (A-B)K^2 - D_J J^2 (J+1)^2 - D_{JK} J(J+1)K^2
$$
 (1)
-D_KK⁴ - H_JJ³(J+1)³ - H_{JJK}J⁴(K+1)²

For the v_{as} perpendicular band, rovibrational energy levels are calculated from the following expression:

$$
E(v_{as}, J, K) = v_{as} + B'J(J+1) + (A'-B')K^2 - 2\xi A'Kl
$$

-D'_{J}J^2(J+1)^2 - D'_{JK}J(J+1)K^2 - D'_{K}K^4 - H'_{J}J^3(J+1)^3 - H'_{JJK}J^4(K+1)^2 (2)

where ξ is the first-order Coriolis constant, characteristic of a degenerate vibration. Transitions follow the selection rules $\Delta J = 0, \pm 1$; $\Delta K = \pm 1$; $\Delta l = \pm 1$ and $\Delta |K - l| = 0$ (with *l*, the quantum number associated to the vibrational angular momentum). The v_{as} perpendicular band exhibits different kinds of transitions designated by $P, R, P \left(J, K \right)$ $P, R \left(Q \left(J, K \right) \right)$ and ^{P,R}R(*J*, *K*) where the superscripts P, R correspond to $\Delta K = -1, +1$, the main capital letter P, Q, R to $\Delta I = -1.0,1$, respectively³⁷, and where *J* and *K* are the ground state rotational quantum numbers.

For the v_s parallel band, rovibrational energy levels are calculated from the following expression:

$$
E(vs, J, K) = vs + B'J(J + 1) + (A' - B')K2 - D'_{J}J2(J + 1)2 - D'_{JK}J(J + 1)K2
$$

-D'_KK⁴ - H'_JJ³(J + 1)³ - H'_{JJK}J⁴(K + 1)² (3)

Transitions follow the selection rules $\Delta J = 0, \pm 1$; $\Delta K = 0$. Rotational lines for the v_s parallel band are designated by ${}^{Q}P(J,K)$, ${}^{Q}Q(J,K)$ and ${}^{Q}R(J,K)$ where the superscript Q means $\Delta K = 0$.

The lower-state statistical weights is affected by the $2J + 1$ rotational degeneracy factor and the rules of nuclear spin statistics for a C_{3v} molecule possessing three identical H atoms of nuclear spin $I_H = 1/2$ outside the axis of symmetry. This results in transitions originating from rotational levels (in the ground vibrational state) with $K = 3n$ which are twice as strong as those with $K = 3n \pm 1$ (n is an integer).

As for the MMW rotational analysis of Section III, the SPFIT/SPCAT suite of programs is used^{[30](#page-5-0)} in the following analyses, while simulated spectra are plotted using the PGOPHER program.³⁸ Note also that the ground state value of \vec{A} is again fixed to the value obtained in the structural determination of Wikrent et al^{[33](#page-5-1)}

IV-2-1 The as spectrum

The rovibrational structure of the jet-cooled v_{as} spectrum (Fig. 6) which extends between 973.8 and 978.5 $cm⁻¹$ is composed of two similar rovibrational band contours centered at 976.0 and 976.6 cm⁻¹, respectively, with a 10.5:6 intensity ratio estimated from the integrated intensities of *Q* branches. This corresponds to the isotopic abundance of the rhenium atom (62.93% for 187 Re and 37.07% for 185 Re), allowing us to assign each isotopologue a band contour. Those perpendicular bands consist of two types of rotational structures. The central part exhibits a series of strong $P, RQ(J, K)$ branches for which K is fixed (see also inset of Fig. 8). Each of these *K* branches is composed of a number of unresolved lines, each corresponding to a different value of *J*, with $J \geq K$. Further from the band center, there are several progressions composed of single isolated rovibrational P, R, P, V and $P, R \mid R(J, K)$ lines.

In these *P* and *R* regions, the QCL-jet v_{as} spectrum displays characteristic clusters of lines equally spaced in frequency by about 0.2 cm⁻¹ which are likely to correspond to $P_P(f, K)$ or ${}^R R(J, K)$ transitions for which *J* is fixed. We start by assigning the most intense lines of these clusters to the stronger $P_P(f, 3n)$ and $R(f, 3n)$ transitions. We note that the lines' frequency is an increasing function of the *K* quantum number in ${}^{P}P(J,K)$ clusters, while it is a decreasing function in ${}^R R(J,K)$ clusters. Equations (1) and (2) show that this happens when the K-dependent term $(-2\zeta A')$ is much larger than the K^2 -dependent term $(A' - A) - (B' - B)$. The near zero value of the Coriolis coupling constant ξ of our previous work^{[19](#page-1-2)} (which had been determined from the simulation of the Q -branch alone) contradicts our observations and is now refuted. Theoretical predictions using $\xi = 0.20$ derived from a DFT (B3LYP) calculation nicely confirm our observations.

All intense patterns of the jet-cooled spectrum are then assigned to the ¹⁸⁷Re isotopologue. Most of unassigned lines are likely to correspond to the less abundant ¹⁸⁵Re isotopologue. As shown in Fig. 7, an almost perfect overlap is observed between the $P_{P}(J,K)$ transitions of the ¹⁸⁵Re isotopologue and the $P, R_{P}(J - 3, K - 3)$ lines of the ¹⁸⁷Re one.

The rovibrational lines assigned up to $I = 10$ at that stage provide a first set of new constants including one ground-state parameter (D_K) , which was not accessible from the µW/MMW experiments) and 4 excited-state parameters (v_{as} , A', B' and ξ) for each isotopologue. In these preliminary analyses, ground-state rotational and both ground- and excited-state centrifugal distortion parameters are fixed to the μ W/MMW values of Table 1.

We now return to the room temperature spectrum, for which the model is now good enough to assign higher *J* lines. 240 (37) lines for CH_3^{187} ReO₃ (CH_3^{185} ReO₃) have been identified, reaching quantum numbers of up to $J=33$ (17) for P transitions and 25 (12) for

 ${}^{R}R$ transitions. Note that 15 lines for CH₃¹⁸⁷ReO₃ (5 for CH₃¹⁸⁵ReO₃) were observed in the *R*branch of both the jet and cell/FTIR spectra. In the final line list (see ESI2), for these lines, the more precise frequencies extracted from the cell/FTIR data (30 MHz resolution) have been preferred.

Having assigned these higher *J* lines, we now focus on *P*-branch transitions where 185 Re and 187 Re isotopologues transitions are nearly overlapped in both supersonic jet and cell/FTIR spectra (see Fig. 7). Although the absolute accuracy using the buffer-gas setup is comparable to the other methods, the relative uncertainty of the frequency scale is only a few megahertz. These lines are thus resolved in the buffer-gas-cooled spectra, (see Fig. 8). The positions of 6 jet-cooled ^PP lines of ¹⁸⁷Re MTO, and 7 jet-cooled ^PP and ^RP lines of ¹⁸⁵Re MTO have thus been refined and replaced by frequencies extracted from buffer-gas-cooled spectra (those are labeled in the final line lists given in ESI2). We point out that 5 amongst the 7 CH₃¹⁸⁵ReO₃ lines are now fully resolved and assigned a position different (shifted by about 100 MHz) from the neighbouring ¹⁸⁷Re isotopologue line frequency.

Finally, a global fit to a total of 323 line positions was performed for the ¹⁸⁷Re isotopologue. The assignment of higher *lines allow the extraction of yet 3 additional* parameters, namely excited-state D'_J , D'_{JK} , D'_K centrifugal distortion parameters. The set of parameters obtained at that stage is given in set 1 of Table 2.

A similar fit to a total of 96 lines was realized for $CH₃¹⁸⁵ReO₃$, and led to set 3 in Table 2. Here centrifugal distortion parameters of both the ground and excited state were constrained to the μ W/MMW ground-state values. Only D_K ['] could be floated while D_K was fixed to the ¹⁸⁷Re MTO ground-state value of the previous analysis. Note that including the buffer-gas lines allows us to better reproduce the relative line positions between isotopologues (see Fig. 8).

Fig. 7 shows the good agreement between a simulated spectrum of the v_{as} mode of both isotopologues with a rotational temperature of 6 ± 1 K and jet-cooled experimental data for rotational progressions in the $974.8-975.6$ cm⁻¹ range. Fig. 6 also shows our best agreement between the data and a simulated stick spectrum over the entire band at the same temperature.

IV-2-2 The ^s spectrum

The v_s band is observed in the cell-FTIR spectrum recorded at the maximal resolution of 0.00102 cm⁻¹ (see Sections II-3 and IV-1). The whole transition spans 40 cm⁻¹ from 985 to 1025 cm⁻¹ but only the 995-1015 cm⁻¹ range shown in Fig. 4 has been considered in the analysis. The ${}^{Q}P(J,K)$ branch is strongly perturbed by red shifted hot bands and the nearby v_{as} band centered at 976 cm⁻¹ but the characteristic *PQR* structure of the v_{s} parallel band of a symmetric top molecule is still visible with resolved K substructures in the $P-$ and R branches. 848 (296) lines have been assigned to the ¹⁸⁷Re (¹⁸⁵Re) isotopologue, with highest *J* values equal to 51 (30) for ^QR transitions and 40 (30) for ^QP transitions. Five rotational parameters (v_s, A', B', D'_j and D'_{jK}) in the v_s = 1 state have been adjusted to reproduce the assigned transition frequencies of both isotopologues.

As for the v_{as} band of CH₃¹⁸⁵ReO₃, only D_K ^{*'*} was adjusted independently while D_K was fixed to the ¹⁸⁷Re MTO ground-state value of the previous analysis. The results of those two least squares optimizations are reported in set 1 and set 3 of Table 3, for the 187 Re and 185 Re isotopologues respectively. The final line lists for the v_s band is reported in ESI3. Fig. 5 shows the good agreement between the data and a simulation of the v_s modes of both isotopologues for five rotational progressions of the *R*-branch with resolved *K* lines. Fig. 4 also shows good agreement between the data and a simulated stick spectrum on a broader span.

IV-2-3 Including the hyperfine structure

The HFS of MTO has not been taken into account in the above analyses of the Re=O vibrational modes. Microwave^{[33,3](#page-5-1)9} and millimeter-wave experiments show that the rhenium quadrupole coupling constant $\epsilon Qq > 700$ MHz is relatively close in magnitude to the rotational constants, resulting in the break-up of microwave and millimeter-wave transitions into hyperfine components. The HFS must be included in the analysis of the most resolved data.

In particular, to refine our understanding of the HFS in the excited vibrational state of the v_{as} perpendicular band of CH₃¹⁸⁷ReO₃, we proceed to the assignment of the structures observed in the continuous supersonic beam $CO₂$ laser absorption spectra displayed in Fig. 9. As mentioned above, these structures are strongly modified by the hyperfine interaction. In a parallel work^{[29](#page-4-0)}, an analytical hyperfine Hamiltonian which included the rhenium electric quadrupole interaction and spin-rotation magnetic interactions was fitted to four *^PP* lines from the buffer-gas setup (including the $PP(6,3)$ line in Fig. 8). The quadrupole coupling constant in the upper state, *eQq*', was found to be 716(3) MHz, to be compared to *eQq* = 716.5725(26) in the ground state (see Table 1), and so we conclude that there is little variation of the quadrupole coupling constant with vibration. To assign hyperfine components to the structures observed in the supersonic beam linear absorption spectra of Fig. 9, we thus compare the data to a simulation that uses both the rovibrational parameters of set 1 in Table 2 as well as the HFS parameters obtained from µW/MMW spectroscopy (Table 1), using the latter for both the ground and excited vibrational states. From this comparison, a total of 36 hyperfine components of the ${}^R Q(J,2)$ lines with $2 \le J \le 10$ and of the ${}^R Q(16,1)$ line have been assigned, with frequency uncertainties ranging from 1×10^{-4} cm⁻¹ to 3×10^{-4} cm⁻¹, depending on the width of the assigned structure. These have been added to the previous 187 Re MTO v_{as} band line list (see ESI2, where they are labeled).

For both the v_s and v_{as} bands of ¹⁸⁷Re and ¹⁸⁵Re isotopologues of MTO, we now fit and simulate the transition frequencies with a Hamiltonian containing the rhenium quadrupole coupling constants eQq and its centrifugal distortion correction eQq , and the spin-rotation interaction terms C_{aa} and C_{bb} in both the ground and excited states. For the ¹⁸⁷Re (and ¹⁸⁵Re) isotopologue, we have to fit line lists including: (i) the 16(17) assigned rotational lines from the microwave study reported in our previous work^{[19](#page-1-2)} (ii), the 920(511) assigned rotational lines at higher values of \bar{I} in the MMW study of section III, and (iii), the most intense hyperfine components (those with $\Delta F = \Delta I$) of all the rovibrational transitions assigned in section IV-2. Note that hyperfine components of a given rovibrational line are unresolved and

thus are all assigned the same observed frequency. For the particular case of the ¹⁸⁷Re MTO v_{as} mode, the 36 partially resolved hyperfine components assigned in the spectrum of Fig. 9 are added to the line list. A total of 2891 (respectively 6024) P^P , P^R and P^R lines are assigned in *J*, *K* and *F* to the v_{as} (respectively v_s) band of the ¹⁸⁷Re isotopologue, and 1101 (respectively 2304) lines are assigned to the v_{as} (respectively v_s) band of the ¹⁸⁵Re isotopologue. The complete lists of assigned lines are available as ESI for the two modes of each isotopologue.

For each line list, we perform a fit in which we float all parameters but A (D_K is also fixed in 3 amongst the 4 cases). However, with the exception of the upper-state quadrupole coupling constant of the v_{as} mode of 187 Re MTO, all hyperfine parameters and sextic centrifugal distortion constants were constrained to the same values for both ground and excited states. Note that, for the v_{as} band of CH_3^{185} ReO₃ *D'_J* and *D'_{JK}* were also constrained to be equal in both states due to the limited number of lines. The four corresponding sets of molecular parameters are gathered in sets 2 and 4 of Table 2 and 3. For the v_{as} mode of 187 Re MTO, it was also possible to float eOq' , after the assignment of hyperfine components to the structures in Fig. 9. We obtained $eOq' = 705(44)$ MHz, which is in good agreement with our analysis of the buffer-gas-cooled spectra detailed in Ref. [29,](#page-4-0) confirming the very weak variation of the quadrupole coupling constant with vibration.

Fig. 7, 8 and 9 compare data and simulations that include HFS. Although this makes little difference for the EC-QCL spectra in Fig 7 where the resolution is similar to the HFS splitting $\left(\sim\right)100$ MHz), it is indispensable for obtaining agreement for the CO₂ laser (Fig. 9) and buffer gas spectra (Fig. 8), at a rotational temperature of 6 K and 15 K, respectively.

V- Conclusion

Three new MMW and IR high resolution spectroscopic devices coupled either to a room temperature cell, a cryogenic cell or a pulsed supersonic expansion, were used to fully reinvestigate the spectrum of MTO in the ground state and Re=O stretching v_s and v_{as} excited states, as well as to refine the rovibrational model of MTO.

A full set of rotational and hyperfine parameters of the v_s and v_{as} modes of both isotopologues 187 Re and 185 Re of MTO, including centrifugal distortion contributions, has been extracted from the combined analysis of different μ W, MMW and IR spectra. The assignment of more than thirty partially resolved hyperfine transitions in $CO₂$ laser absorption spectra of a skimmed molecular beam led to a determination of the quadrupole coupling constant in the excited state.

Studying the MTO molecule provides insight into how to study its chiral derivatives, which have a pronounced dissymmetry around the stereogenic rhenium center and may thus exhibit measurable parity violation effects. A full characterization of such organometallic molecules requires the use of several sensitive spectroscopic techniques, much like the experiments reported here. We thus expect to perform this sort of analysis on the chiral species in order to identify the right transition with which to observe parity violation.

We also establish a procedure for extracting very accurate molecular parameters of complex species, both in the ground and excited state, from a large amount of spectroscopic

data spanning various spectral regions, taken at different resolutions from a variety of experiments. We believe that this sort of analysis will become increasingly necessary for any precise measurement on polyatomic molecules, whether it be for atmospheric, planetary or interstellar studies, chemistry, biology or other fundamental physics tests.

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Experiment control

Figure 1: Experimental setup combining a pulsed supersonic jet originating from a circular nozzle and an EC-QCL based spectrometer that includes a near concentric multipass optics. EC-QCL: external-cavity quantum cascade laser.

Pulse sequence

Figure 2: Data acquisition sequence for the jet-cooled EC-QCL experiment. The TTL pulse timer card triggers three channels: 1) the *Diffraction grating PT voltage* sine wave , 2) the *Acquisition triggers* sequence which defines the starting time of each 5-ms acquisition period, 3) the *Nozzle driver pulse* after an adjustable delay d₁. After each TTL pulse the signal of only 5 (amongst 100 at a 1-Hz repetition rate) acquisition windows are kept: 4 bkg (background signal in the absence of molecular jet) and 1 jet (in the presence of the molecular jet). The delay d_2 is the molecules' travelling times from the nozzle to the laser beams. PT: piezoelectric transducer.

Figure 3: Small portion of the millimeter-wave spectrum illustrating the hyperfine structure observed on the rotational spectrum of CH_3^{187} ReO₃. The assignments of the $\Delta F=+1$ hyperfine components for the ${}^{Q}R(J=21,K=18,F)$ line are presented in green, as well as the *K* structure of the $\Delta F=+1$ ^Q*R*(*J*=21,*K*,*F*=24) lines in red. *J*, *K* and *F* are the ground state rotational quantum numbers.

Figure 4: Comparison between the observed room temperature cell-FTIR spectrum of the v_s band of MTO and the calculated stick spectrum (using sets of parameters 1 and 3 of Table 3, for the ¹⁸⁷Re and ¹⁸⁵Re isotopologues respectively). The experimental ^QQ branch is strongly broadened by the many hot bands red shifted with respect to the band center. Note also that below 995 cm⁻¹, high *J* lines of v_s *P*-branch overlap with the v_{as} *R*-branch.

Figure 5: Comparison between observed room temperature cell-FTIR spectrum and our best calculated spectrum (using sets of parameters 1 and 3 of Table 3), with assigned rotational progressions of both isotopologues, located in the R -branch of the v_s band. Those K progressions are composed of ${}^{Q}R(J,K)$ lines for which *J* is fixed, each corresponding to different *K* values with $K \leq J$. Three progressions of the ¹⁸⁷Re isotopologue for $J = 21-23$ and two progressions of the ¹⁸⁵Re isotopologue for $J = 21$ and 22 are visible. The features observed between the assigned progressions of the fundamental of v_s are attributed to hot bands.

Figure 6: Comparison between the observed jet-cooled EC-QCL spectrum of the v_{as} band and a simulated stick spectrum of the v_{as} modes of both isotopologues without the contribution of the hyperfine structure (using set of parameters 1 and 3 of Table 2), at a rotational temperature of 6 K.

Figure 7: Comparison between the observed jet-cooled EC-QCL spectrum and two calculated spectra at 6 K, without the HFS (using set of parameters 1 and 3 of Table 2) and with the HFS (using set of parameters 2 and 4 of Table 2), for four assigned rotational progressions of both isotopologues, located in the P -branch of the v_{as} band. Those K progressions are composed of $P(P(J,K)$ and $P(P(J,K))$ lines for which *J* is fixed, each corresponding to a different value of *K*, with $K \leq J$. The four progressions of the ¹⁸⁷Re and ¹⁸⁵Re MTO are those for which $I = 2$ -5 and $I = 5$ -8 respectively. Calculated spectra are obtained by convolving stick spectra with a Lorentzian profile of 100 MHz full-width-at-halfmaximum, corresponding to the expected resolution. Including the HFS does not significantly modify the simulations because the HFS splitting is of the same order as the resolution.

Figure 8: Upper black curve: Linear absorption spectrum of the ablation plume of MTO buffer-gas-cooled to ~6 K recorded using a free-running DFB QCL. It exhibits two isolated rovibrational lines, the ^{*P*} $P(9,6)$ and ^{*P*} $P(6,3)$ lines of the ¹⁸⁵Re and ¹⁸⁷Re MTO isotopologues respectively, both showing a partially resolved hyperfine structure. Lower dashed blue curve: simulated $CH₃¹⁸⁷ReO₃$ spectrum at a rotational temperature of 6 K, using set 2 of Table 2. Lower solid red curve: simulated $CH₃¹⁸⁵ReO₃$ spectrum at a rotational temperature of 6 K, using set 4 of Table 2. Stick spectra (blue for CH_3^{187} ReO₃ and red for CH_3^{185} ReO₃) are convolved by a Lorentzian profile of 8 MHz full-width-at-half-maximum, corresponding to the expected resolution.^{[29](#page-4-0)} The small offset between the experimental and simulated spectra is consistent with the 30 MHz absolute accuracy of the frequency scale^{[29](#page-4-0)} (see section II-4). Insert: Buffer-gas-cooled spectrum showing the rotational contour of the *Q* branch of the antisymmetric Re=O stretching mode of the 187 Re isotopologue at ~976 cm⁻¹ from which we infer a rotational temperature $T_{\text{rot}} = 6 \pm 3 \text{ K.}^{29}$ $T_{\text{rot}} = 6 \pm 3 \text{ K.}^{29}$ $T_{\text{rot}} = 6 \pm 3 \text{ K.}^{29}$

Figure 9: Upper black curve: linear absorption spectroscopy of an MTO-seeded continuous skimmed supersonic beam, in the vicinity of the $R(20)$ CO₂ laser line frequency (located at \sim 975.93044 cm⁻¹); experimental condition[s9:](#page-1-3) 100 µm nozzle diameter, 750 µm skimmer diameter, helium backing pressure ∼ 1 bar, oven temperature ∼ 90°C, translational longitudinal temperature ∼ 1 K, 9 mm nozzle-to-skimmer distance, 1 point recorded every 100 kHz, ~ 1 s of integration time per point, 1.5×10^{-5} cm⁻¹ binning, ~ 0.5 μW of laser power in each of the 18 passes on the multi-pass cell (see text), ∼ 5 mm beam waist. Dashed blue curve: simulated $CH₃¹⁸⁷ReO₃$ spectrum at a rotational temperature of 15 K, using set of parameters 2 of Table 2. Solid red curve: simulated $CH₃¹⁸⁵ReO₃$ spectrum at a rotational temperature of 15 K, using set of parameters 4 of Table 2. Stick spectra (blue for $CH₃¹⁸⁷ReO₃$ and red for $CH₃¹⁸⁵ReO₃$ are convolved by a Lorentzian profile of full-width-at-halfmaximum 2.5×10^{-4} cm⁻¹, corresponding to the expected Doppler broadening resulting from the divergence of the beam.^{[9](#page-1-3)}

Table 1: Molecular parameters (in MHz) for the ground states of CH_3^{187} ReO₃ and CH_3^{185} ReO₃ obtained from µW and MMW data. Uncertainties corresponding to one standard deviation are indicated into parentheses. St. dev.: standard deviation (in MHz) of the difference between experimental and calculated frequencies.

Table 2: Ground- and excited-state parameters of the v_{as} band for both MTO isotopologues. All microwave, millimeter-wave and mid-infrared experiments described in this work were used. The uncertainties in parentheses correspond to one standard deviation.

Table 3: Ground- and excited-state parameters of the v_s band for both MTO isotopologues. Microwave and millimeter-wave (section III), and mid-infrared data from the roomtemperature cell-FTIR experiments (section IV-2-2) were used. The uncertainties in parentheses correspond to one standard deviation.

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