# Synchrotron-based High Resolution Far-infrared Spectroscopy of *trans*-butadiene

Marie-Aline Martin-Drumel<sup>1</sup>, Jessica P. Porterfield<sup>2</sup>, Manuel Goubet<sup>3</sup>, Pierre Asselin<sup>4</sup>, Robert Georges<sup>5</sup>, Pascale Soulard<sup>4</sup>, Matthew Nava<sup>6</sup>, P. Bryan Changala<sup>7</sup>, Brant Billinghurst<sup>8</sup>, Olivier Pirali<sup>1</sup>, Michael C. McCarthy<sup>2</sup>, and Joshua H. Baraban<sup>9</sup>

<sup>1</sup> Université Paris-Saclay, CNRS, Institut des Sciences Moléculaires d'Orsay, 91405 Orsay, France
 <sup>2</sup> Center for Astrophysics, Harvard & Smithsonian, Cambridge, Massachusetts 02138, United States
 <sup>3</sup> Université Lille, CNRS, UMR 8523-PhLAM-Physique des Lasers, Atomes et Molécules, F-59000 Lille, France
 <sup>4</sup> Sorbonne Universités, UPMC Univ Paris 06, UMR 8233, MONARIS, F-75005 Paris, France & CNRS, UMR 8233, MONARIS, F-75005 Paris, France
 <sup>5</sup> Institut de Physique de Rennes, CNRS, UMR 6251, Université de Rennes 1, F-35042 Rennes, France
 <sup>6</sup> Department of Chemistry, Harvard University, Cambridge, MA 02138, United States
 <sup>7</sup> JILA, University of Colorado, Boulder, CO 80309, United States
 <sup>8</sup> Canadian Light Source, University of Saskatchewan, Canada
 <sup>9</sup> Department of Chemistry, Ben-Gurion University of the Negev Beer-Sheva 8410501, Israel

Contact corresponding author(s): marie-aline.martin@universite-paris-saclay.fr

The high resolution far-infrared spectrum of trans-butadiene has been re-investigated by Fourier-transform spectroscopy at two synchrotron radiation facilities, SOLEIL and the Canadian Light Source, at temperatures ranging from 50 to 340 K. Beyond the well-studied bands, two new fundamental bands lying below 1100 cm<sup>-1</sup>,  $v_{10}$  and  $v_{24}$ , have been assigned using a combination of cross-correlation (ASAP software) and Loomis-Wood type (LWWa software) diagrams. While the  $v_{24}$  analysis was rather straightforward,  $v_{10}$  exhibits obvious signs of a strong perturbation, presumably owing to interaction with the dark  $v_9 + v_{12}$  state. Effective rotational constants have been derived for both the  $v_{10} = 1$  and  $v_{24} = 1$  states. Since only one weak, infrared active fundamental band ( $v_{23}$ ) of trans-butadiene remains to be observed at high resolution in the far-infrared, searches for the elusive gauche conformer can now be undertaken with considerably greater confidence in the dense ro-vibrational spectrum of the trans form.

#### 1. Introduction

Much attention has surrounded 1,3-butadiene ( $C_4H_6$ ) because this conjugated diene is an ideal candidate to observe the effects of  $\pi$ -electron delocalization [1, 2], and because it is of central importance in a wide range of chemistry applications, from its archetypal role in the Diels-Alder reaction [3] to industrial rubber production [4]. Characterizing the structural conformers and isomerization dynamics of this diene, however, has proven surprisingly challenging. 1,3-butadiene exists in two stable forms, the more stable planar *trans*, and the long-elusive *gauche*, lying 12 kJ/mol higher in energy [5]. The latter was only conclusively detected in the gas phase recently by several authors of the present work, an effort which also yielded a nearly complete semi-experimental equilibrium structure ( $r_{\rm e}^{\rm SE}$ ) from a combination of pure rotational

measurements of various isotopic species and high-level quantum chemical calculations [5].

While electron diffraction studies provided insight into the structure of *trans*-butadiene as early as the late 1930's [6, 7], an accurate determination of its equilibrium structure was only achieved in 2006 [1]. This centro-symmetric species possesses no permanent dipole moment, and consequently no pure rotational spectrum. Determination of the ground state constants thus relied on the assignments of rotational transitions in its dense vibrational bands, a challenging feat overcome in 2004 with the high-resolution analysis of three infrared bands — namely  $\nu_{17}$  (3100 cm<sup>-1</sup>) [8],  $\nu_{11}$  (908 cm<sup>-1</sup>) [9], and  $\nu_{20}$  (1596 cm<sup>-1</sup>) [9]. From extensive isotopic investigations, Craig and co-workers [1] soon afterwards derived a full semi-experimental structure from the ground state rotational

Accepted article 1/10

constants of eight isotopologues: the normal [9], 1,1-d<sub>2</sub> [10], 2,3-d<sub>2</sub> [9], 1,4-trans,trans-d<sub>2</sub> [11], 1,4-cis,cis-d<sub>2</sub> [11], 1,4-cis,trans-d<sub>2</sub> [11],  $1^{-13}C_1$  [12], and 2,3- $^{13}C_2$  [13] species. With the notable exception of butadiene-1,1-d<sub>2</sub> which is slightly polar and was thus investigated using microwave spectroscopy by Caminati et al. [10], the rotational constants of all other isotopologues were derived from high-resolution ro-vibrational measurements.

trans-butadiene belongs to the C<sub>2h</sub> point group and possesses 12 IR-active modes, 4 of  $a_u$  and 8 of  $b_u$  symmetries. Among them, 6 modes lie below  $1100 \,\mathrm{cm}^{-1}$ , *i.e.* in the far-IR domain:  $v_{13}$  (162.42 cm<sup>-1</sup>,  $a_u$ ) [14],  $v_{24}$  (~299.1 cm<sup>-1</sup>,  $b_u$ )[15],  $v_{12}$  (524.57 cm<sup>-1</sup>,  $a_u$ ) [14],  $v_{11}$  (908.07 cm<sup>-1</sup>,  $a_u$ ) [9],  $v_{23}$  ( $\sim$  990.3 cm<sup>-1</sup>,  $b_u$ ) [15], and  $v_{10}$  ( $\sim$  1013.8 cm<sup>-1</sup>,  $a_u$ ) [15]. Of these, only three —  $\nu_{10}$ ,  $\nu_{23}$ , and  $\nu_{24}$  — have not been observed at high resolution prior to this work, and thus their band centers (listed above) are probably accurate to at most a few wavenumbers based on low resolution gas phase investigation [15]. In this article, we report the first ro-vibrational analysis of two of these,  $v_{10}$  (outof-plane C-H wagging[16]) and  $v_{24}$  (in-plane C=C-C deformation[16]). The spectra were recorded at two synchrotron radiation facilities, SOLEIL and the Canadian Light Source (CLS), during a search for the ro-vibrational spectrum of the gauche-butadiene.

#### 2. Methods

#### 2.1. Quantum chemical calculations

Vibrational frequencies and rotational constants were computed by second-order vibrational perturbation theory (VPT2)[17]. The force field constants were obtained using a hybrid approach, wherein a coupled-cluster treatment of electron correlation with singles, doubles, and perturbative triples and the frozen-core approximation [fc-CCSD(T)] was employed with the ANO1 basis set for the harmonic frequencies and the ANO0 basis set for the anharmonic constants[18]. Calculations were performed using the CFOUR suite of electronic structure programs [19]. The expected far-IR spectrum of butadiene from literature experimental values, when available, or these calculations, otherwise, is simulated in Fig 1. Tables S1 and S2 report the calculated band centers and intensities, and the calculated rotational constants, respectively.

A particularly noteworthy result of the VPT2 calculations is the predicted resonance between  $v_{10}=1$  ( $a_u$  symmetry) and  $v_9=v_{12}=1$  ( $a_u$ ), whose zero-order energies lie less than 1 cm<sup>-1</sup> apart, around 1034 cm<sup>-1</sup>, according to the ANO1 harmonic frequencies. Using the GUINEA module of CFOUR with hybrid ANO1/ANO0 constants, the effective Hamiltonian between the  $v_{10}=1$  and  $v_9=v_{12}=1$  states has been constructed and diagonalized. Once anharmonicity is taken into account, the states are found to be 22 cm<sup>-1</sup> apart (with  $v_{10}=1$  lower in energy, lying at 1012 cm<sup>-1</sup> and  $v_9=v_{12}=1$  lying at 1034 cm<sup>-1</sup>, see Table S1) with an overall (pure vibrational) coupling matrix element of  $0.8 \, \mathrm{cm}^{-1}$ .

#### 2.2. Butadiene sample preparation

In all of the experiments performed here, 1,3-butadiene was synthesized from thermal decomposition of sulfolene ( $C_4H_6SO_2$ ) at 85–100 °C according to the reversible reaction:

$$C_4H_6SO_2(s) \stackrel{\Delta}{\rightleftharpoons} C_4H_6(g) + SO_2(g).$$
 (1)

Theoretical calculations of this reaction [20] predict gaseous butadiene is initially produced in the *gauche* form through a *cis* transition state via a retro-Diels-Alder mechanism.

To prevent the reverse Diels-Alder reaction, and to obtain a high purity sample of butadiene, an SO<sub>2</sub> trap consisting of KOH pellets was used:

$$SO_2(g) + KOH(s) \longrightarrow K_2SO_3(s) + H_2O(l)$$
, (2)

where water released during reaction Eq. (2) was condensed in a cold trap maintained at about 0 °C. Butadiene passes through this trap, but was subsequently collected either by condensation in a vessel cooled with liquid nitrogen or was directly injected into the measurement cell. This purification procedure precludes the obtainment of nonthermal rotamer populations of butadiene (gauche:trans ratio) due to numerous collisions with the vessel walls.

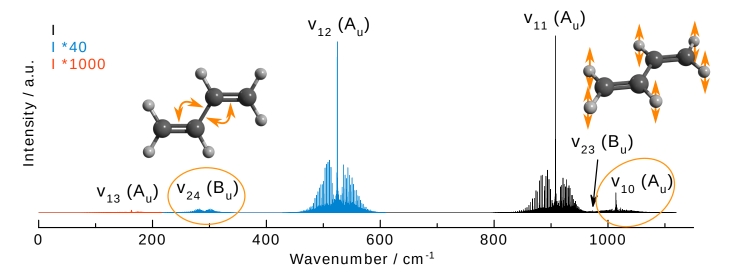
#### 2.3. Synchrotron-based FT-IR spectroscopy

The high resolution ro-vibrational spectrum of butadiene has been investigated using three complementary set-ups installed at two synchrotron facilities, SOLEIL and the Canadian Light Source. These set-ups allowed us to probe the sample at three different temperatures, namely 50, 273, and 343 K. In all cases, butadiene spectra were recorded using Fourier-transform spectroscopy (in the far- and mid-IR) using a Bruker IFS125HR spectrometer at its ultimate resolution of about  $1\times 10^{-3}~{\rm cm}^{-1}$ . When used as the light source, the synchrotron radiation was focused onto the entrance iris of the interferometer. In all cases, the interferometer was continuously evacuated to  $10^{-5}$  mbar to minimize absorption by atmospheric water. Each experiment is described in greater details in the following paragraphs.

#### 2.3.1. Warm cell experiment

A synchrotron-based ro-vibrational spectrum of butadiene was recorded in the 400–1300 cm $^{-1}$  region at the far-IR beamline of the CLS facility [21]. About 40 µbar of sample was injected into a 2-m base-length multipass White-type cell aligned for a 72 m optical path length and maintained at a 343 K (70  $^{\circ}$ C) temperature [22]. The cell was separated from the interferometer using KBr windows. 280 scans were collected at 0.00096 cm $^{-1}$  resolution (according to Bruker definition) using a 80kHz scanner velocity, a 1.15 mm entrance iris aperture, a KBr beamsplitter, and a Ge:Cu detector installed inside a QMC pulse tube cooled cryostat, and equipped with a KRS5 window.

Accepted article 2/10



**Fig. 1.** Simulation of the high resolution (at  $0.001~\rm cm^{-1}$  resolution and assuming a Doppler profile) far-IR spectrum of *trans*-butadiene (fundamental bands only) performed with the PGOPHER software using the experimental band centers and rotational constants when available, or those derived from the quantum chemical calculations (Table S1-S2). Computed intensities are plotted in arbitrary units. For the sake of clarity, the weak bands have been magnified by a factor  $40~(\nu_{12}, \nu_{24}, \text{ in blue})$  or  $1000~(\nu_{13}, \text{ in red})$ . The two new bands investigated in this work ( $\nu_{10}$  and  $\nu_{24}$ ) have been highlighted in orange and the deformation associated with the two corresponding vibrations is represented by orange arrows on the molecular pictures.

The spectrum was calibrated using residual lines from HDO (around  $1200\,\mathrm{cm^{-1}}$ ), HCN (around  $715\,\mathrm{cm^{-1}}$ ), and CO<sub>2</sub> (around  $670\,\mathrm{cm^{-1}}$ ) whose rest wavenumbers were taken from the HITRAN database [23] and includes experimental data pertinent to this work [24–26]. Residual lines of water, saturated in our experimental spectrum, were not considered in the calibration procedure. From the difference between the calibrated wavenumbers and the reference ones, accuracy of line positions for the entire spectrum should be better than  $\pm 2 \times 10^{-4}\,\mathrm{cm^{-1}}$  (Fig. S1). Around  $1100\,\mathrm{cm^{-1}}$ , where the  $\nu_{10}$  band of butadiene is predicted to lie, the Doppler full width at half maximum of the lines at 343 K is of about  $2 \times 10^{-3}\,\mathrm{cm^{-1}}$ ; thus, a wavenumber accuracy of about a tenth that value appears appropriate.

#### 2.3.2. Room temperature experiment

A synchrotron-based ro-vibrational spectrum of butadiene was recorded in the  $50\text{--}650~\text{cm}^{-1}$  region at the AILES beamline of the SOLEIL synchrotron facility [27]. A spectrum consisting of 220 co-added interferograms was recorded at a resolution of  $0.00102~\text{cm}^{-1}$  (Bruker definition) using a 2.5-m base-length room temperature multipass White-type cell aligned for about 150~m optical path length [28] and filled with  $180~\text{\mu}$ bar of sample. A 2 mm iris, a 6 µm thick mylar-silicon composite beamsplitter, and a liquid-helium Si bolometer were used for the measurements. The cell was isolated from the interferometer by two  $50~\text{\mu}$ m thickness polypropylene windows.

Calibration was performed using residual H<sub>2</sub>O and

 $CO_2$  lines whose reference wavenumbers were taken from Refs. 29, 30, which in turn allowed line frequencies to be determined to better than  $\pm 1 \times 10^{-4}$  cm<sup>-1</sup> below 500 cm<sup>-1</sup> and  $\pm 2 \times 10^{-4}$  cm<sup>-1</sup> above 500 cm<sup>-1</sup> (Fig. S1).

#### 2.3.3. Supersonic-jet experiment

Finally, a spectrum of butadiene in a supersonic jet expansion was recorded in the 850–1050 cm<sup>-1</sup> region using the Jet-AILES apparatus at SOLEIL[31, 32]. A slit nozzle of 60 mm length and 130 µm width was used to expand the sample seeded in helium buffer gas, with respective flows of 1 and 20 standard liters per minutes (slm), into a large chamber that was continuously evacuated by a set of Roots pumps. The reservoir and chamber pressures were maintained at 260 and 0.8 mbar, respectively, and the chamber was separated from the interferometer and the detection compartment by two KBr windows. The high resolution spectrum is the result of the co-addition of 130 individual scans recorded at 0.00102 cm<sup>-1</sup> resolution using a globar source, a 1.13 mm iris aperture, a KBr beamsplitter, and photovoltaic MCT detector. A low-pass optical filter was placed in front of the detector thereby limiting the upper wavenumber value of the measurements to  $1050 \,\mathrm{cm}^{-1}$ . Under our experimental conditions, the use of the synchrotron radiation instead of the globar source resulted in a lower signal-to-noise ratio.

Additional spectra were recorded at lower resolution  $(0.01 \, \text{cm}^{-1})$  for different sample to buffer gas ratios  $(2 \, \text{slm})$  of butadiene and  $x \, \text{slm}$  of He, where x = 0, 5, 10, 20, 30) but otherwise identical experimental conditions. Since

Accepted article 3/10

accurate rotational constants of the  $\nu_{11}$  band of *trans*-butadiene were available from prior work[9], simulations of the band profile (using the PGOPHER software [33]) allowed us to infer the rotational temperature of the spectra with different mixing ratios. A temperature of 30 K well reproduced the high resolution Jet-AILES spectrum, while temperatures of 150, 100, 70, 50, and 40 K were estimated to reproduce the lower resolution spectra with 0, 5, 10, 20, and 30 slm of He, respectively (Fig. S2).

In the relatively narrow range covered by the Jet-AILES spectrum, only two bands of *trans*-butadiene,  $v_{10}$  and  $v_{11}$ , are visible, which precludes a conventional frequency calibration using standard molecules. Instead, calibration was performed by comparing strong, isolated lines in both the well-calibrated warm spectrum recorded at CLS with those in this jet spectrum. Several drawbacks to this procedure should be noted: i) the wavenumber accuracy of the CLS transitions is at best  $2 \times 10^{-4}$  cm<sup>-1</sup> whereas accuracy as low as  $1 \times 10^{-4}$  cm<sup>-1</sup> should be possible in the cold Jet-AILES spectrum owing to the narrower lines and sparser spectrum that arises at colder rotational temperatures; and ii) because of the temperature difference between the two spectra, lines that are strong in the Jet-AILES spectrum may not be intense in the CLS spectrum, and vice versa. To increase our calibration reliability, we have selected as many lines as possible on the Jet-AILES spectrum that correspond to lines having a transmittance higher than 0.2 in the warm CLS spectrum. A conservative wavenumber accuracy of  $5 \times 10^{-4}$  cm<sup>-1</sup> is expected from this procedure (Fig. S1).

#### 2.4. Analysis procedure

Since several ro-vibrational bands of *trans*-butadiene have previously been assigned at high resolution, it is reasonable to assume that the vibrational ground state is very well-described, even though no pure rotational measurements of the main isotopologue are available. As a consequence, analysis of the newly-recorded bands appears well-suited for the Automated Spectral Analysis Approach (ASAP) developed by C. P. Endres and several of the authors of this paper [34, 35]. Briefly, this procedure facilitates fast analysis of a ro-vibrational band when one of the two vibrational levels involved, the so-called reference state, is well-characterized. The second, or target state is thus the one from which spectroscopic information needs to be extracted. The method exploits the power of the Loomis-Wood approach for ro-vibrational assignments (see, e.g., Refs. 36, 37) and the redundancy of the spectroscopic information in a ro-vibrational band with multiple selection rules (typically P-, Q-, R-branches and asymmetric splitting), i.e. when transitions originating from different rotational energy levels of the reference state reach the same rotational energy level of the target state. Since the energy levels of the reference state are well-determined, the error in the predicted ro-vibrational frequencies between the reference and target state are dominated by the error in the prediction of the target state. Since each

set of transitions reaches the same rotational level, the frequencies of the sought-after experimental transitions all share the same offset relative to their prediction, with the value of that offset being the error in the predicted energy of the particular target rotational level. Assigning ro-vibrational transitions in such a set thus entails identifying lines lying at the same offset value from the prediction in each portion of the spectrum. ASAP simplifies this step by cross-correlating (multiplying) these portions of the spectrum. For spectra plotted in absorbance, the signal approaches zero between transitions, provided the spectrum is well-resolved. Thus, most frequency points will be multiplied by zero at least once except if a line is always present on each plot. This method allows for significant spectral simplification, as only a few points will stand out from the baseline, and by plotting successive *J* values in a Loomis-Wood fashion, one can assign series of lines. One consequence of the cross-correlation is that the actual rovibrational frequencies are no longer assigned; rather the energy of the sought after target state is directly extracted (predicted energy + assigned offset). Since it is no longer necessary to assign each transition for each selection rule, but only the target energy levels involved in the band, this procedure greatly speeds-up analysis time. The energy levels together with other available data are input to the SPFIT/SPCAT suite of programs[38] to derive the rotational constants and band center of the target state.

The ASAP method has proven extremely helpful in assigning several bands of S<sub>2</sub>O, for which the rotational ground state was accurately known from extensive pure rotational investigations [34, 35]. In the present case, however, since no such pure rotational data exists and since the ro-vibrational bands assigned in the literature were recorded at similar or lower resolution compared to the present work, once the target state rotational constants were derived from the ASAP analysis, a conventional assignment of each ro-vibrational transition was also undertaken. To do so, the LWWa software developed by W. Lodyga was used [37].

#### 3. Results and discussion

## 3.1. Initial fit of the literature data

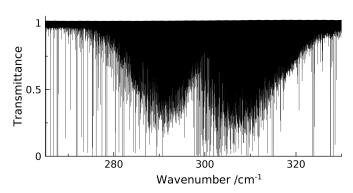
To obtain ground state constants that are as accurate as possible, we have refitted all of the ro-vibrational data previously published, *i.e.* in the  $v_{11}$  [9],  $v_{12}$  [14],  $v_{13}$  [14],  $v_{17}$  [8], and  $v_{20}$  [9] bands. A Watson *S*-reduced Hamiltonian in the  $I^r$  representation and the Pickett SPFIT program were used. Since  $A >> B \sim C$ , we have chosen *S*-reduction although literature data were fitted using the *A*-reduction. Data from  $v_{12}$  and  $v_{13}$  [14] were weighted according to their experimental uncertainty (0.0002 cm<sup>-1</sup>), while those in  $v_{11}$  and  $v_{20}$  [9], for which the estimated line accuracy was not stated, were weighted using the standard deviation obtained in the fit reported by the authors (0.0002 and 0.007 cm<sup>-1</sup>, respectively). In the case of  $v_{17}$ , the authors [8] noted that the expected wavenumber accuracy

Accepted article 4/10

on each transition should be 10 times better than the standard deviation of their fit  $(0.003 \,\mathrm{cm}^{-1})$  but perturbations, most likely from a rovibrational dark state, limited the fit convergence. Since in our study no attempt was made toward treating this perturbation, the value of this standard deviation was retained as the line uncertainty. In total, 7015 transitions (6929 different frequencies) are included in our fit with J'' = 0 - 73 and  $K_a'' = 0 - 18$ . All quartic centrifugal distortion (CD) terms were varied in the ground state while those unfitted in the excited states were constrained to the ground state value. By using a  $5\sigma$  rejection criterion (i.e. transitions were discarded if the observed—calculated frequency was five or more times the postulated uncertainty ), 16 lines from Refs. 9, 14 were rejected from the fit. In these instances, it appears that the inability to reproduce the literature data to within the experimental accuracy is due to perturbations rather than misassignments. We have therefore decided to keep these lines in the fit but to reduce their wavenumber accuracy by a factor 10 (see Table S4 for the list of transitions). The reduced standard deviation (unitless) of the combined fit is 1.16, ranging from 0.73 to 1.54 for individual bands. To our knowledge, only ground state combination differences and some individual ro-vibrational bands were fitted together prior to this study. Since a combined effective fit has been performed here, the ground state rotational constants should be quite reliable.

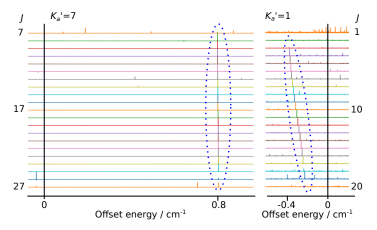
#### 3.2. The $v_{24}$ vibrational band

The  $v_{24}$  band of *trans*-butadiene (a/b hybrid) is observed in the room temperature spectrum recorded at SOLEIL (Fig. 2).



**Fig. 2.** Overview of the high-resolution spectrum of the  $v_{24}$  band of *trans*-butadiene observed at room temperature. Isolated lines arise from residual water vapor, and most of these are saturated. A portion of the Q-branch is also saturated.

**ASAP assignments** Using the ground state as the reference state and  $v_{24} = 1$  as the target state, ASAP searches were performed. The initial rotational constants for  $v_{24} = 1$  were obtained by scaling the VPT2 results by the experimentally-derived versus calculated constants in the ground state (Table S1-S2). The initial frequency for



**Fig. 3.** Initial ASAP views of the  $v_{24}$  band (*left panel*, for one of the two  $K'_a = 7$  asymmetric branches) and the  $v_{10}$  band (*right panel* for a  $K'_a = 1$  branch). Predicted energies are represented by the black vertical line (0 cm<sup>-1</sup> offset) and visible series of lines are highlighted in blue.

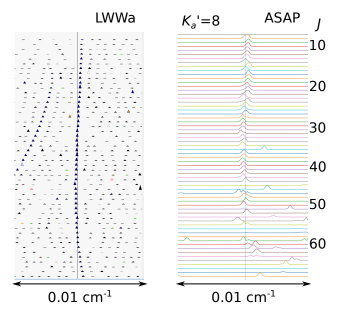
the band center was taken from the low-resolution gas phase assignment of Ref. 15.

Because several series of lines are readily identifiable on the ASAP plots (Fig. 3), a preliminary assignment of the band could be performed almost immediately; it consists of 290 energy levels in  $v_{24} = 1$  with J' = 4 - 44,  $K'_a =$ 4-14, measured to  $0.0001 \, \mathrm{cm}^{-1}$  accuracy. These energy levels were then added to the list of literature assignments, as defined in the previous section, and constants fit using the Pickett software. The resulting parameters are reported in Table 1. Even though we suspected that inclusion of higher-order CD parameters would help to reduce the standard deviation, at this juncture of the analysis the number of fitted parameters was limited because it appeared that the energies derived in the ASAP procedure were not as accurate as expected given our experimental resolution. Ostensibly this situation arises because of residual error in the ground (reference) state energies owing to the absence of extremely accurate pure rotational data. Indeed, after fitting, the ASAP series are still not uniformly aligned, and the distribution of residuals is larger than expected for a spectrum recorded at a resolution of 0.001 cm<sup>-1</sup>. This may come from the fact that the data defining the ground state has at best an uncertainty twice what we expect from the  $v_{24}$  measurements (0.0002) vs.  $0.0001 \,\mathrm{cm}^{-1}$ ). Additionally, some Q-branch lines are saturated, which induces broadening of the ASAP lines. Because it is not possible to omit specific lines in the crosscorrelation procedure using the current version of the software, this effect may skew the ASAP plots. For both of these reasons, a subsequent line-by-line assignment of the  $v_{24}$  band has been performed using the LWWa software.

**LWWa assignments** Using the preliminary constants derived for  $v_{24} = 1$ , assignments using the LWWa software are straightforward. Comparison of the LWWa diagrams with those from ASAP (Fig. 4) indicates that lines ap-

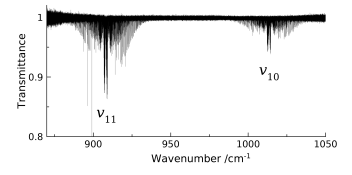
Accepted article 5/10

pear better aligned (i.e. have a smaller dispersion) compared to the former plots. It should be noted, however, that the LWWa software represents a line as a triangle placed at its center frequency, with no consideration for the width of the actual line, while the ASAP plot displays a cross-correlated line profile. Nevertheless, the LWWa plots allow assignment over a wider range of quantum numbers without ambiguity. In total, 2840 transitions with J'=4-87 and  $K_a'=2-18$  have been assigned with a frequency uncertainty of 0.0001 cm<sup>-1</sup> using this approach.



**Fig. 4.** Initial LWWa view of the  $v_{24}$  band for one of the two  $K'_a = 8$  branches in comparison with the corresponding final ASAP plot (i.e., following the initial fit).

#### 3.3. The $v_{10}$ vibrational band



**Fig. 5.** Overview of the high resolution, rotationally cold spectrum recorded using the Jet-AILES apparatus: both the  $v_{10}$  and  $v_{11}$  bands of *trans*-butadiene are clearly visible.

The c-type  $v_{10}$  vibrational band of butadiene was observed both in the cold supersonic jet spectrum recorded at SOLEIL (Fig. 5) and in the warm one recorded at CLS. During the measurements with the supersonic jet, we were also able to observe the effect of rotational cooling on

the band profile: from a standard *c*-type band with a sharp Q-branch at moderate temperature, the band evolves to a completely different profile with two Q-branch lobes at low temperature (Fig. 6).

**ASAP assignments** Following the same procedure as for the  $v_{24}$  band, assignments were first performed using the ASAP software. The ground state was used as the reference state, the initial constants of the target state (here  $v_{10} = 1$ ) were obtained by scaling the VPT2 calculation (Table S2), and the initial frequency of the band center was taken from the low-resolution gas-phase assignment of Ref. 15.

As with the  $v_{24}$  analysis, several series of lines, slightly displaced from the prediction, were readily observed and assigned (Fig. 3): over 100 energy levels in total, with J' = 1-24 and  $K'_a = 0-6$ . As before, the reduced standard deviation of the fit, assuming a  $0.0005 \, \mathrm{cm}^{-1}$  accuracy for the energy levels, is high ( $\sigma = 16$  when fitting a single quartic CD term in addition to the rotational constants and band center; see Table 1). Owing to the limited range of the quantum numbers that are accessible at this temperature, no additional CD terms were varied at this stage of the analysis.

**LWWa assignments** Using the preliminary set of rotational constants and the LWWa software we were able to assign 927 lines with 682 different frequencies, including 131 lines with  $\Delta K_c = \pm 2$ , in the Jet-AILES spectrum. These transitions involve J' = 1 - 29 and  $K'_a = 0 - 7$ .

Additionally, 1176 transitions (964 different frequencies) were also assigned in the warm CLS spectrum with J' = 2 - 49 and  $K'_a = 0 - 7$ . More lines, likely involving higher J and  $K_a$  values, are present in the warm spectrum, but it was not possible to assign them with confidence using our best model (see the Combined fit section).

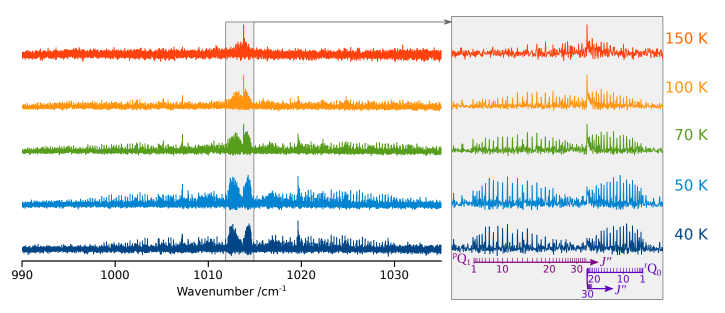
#### 3.4. Other bands

In the FIR spectra we recorded, limited sensitivity prevented detection of any combination bands. Searches for  $v_{23}$  (lying around 977 cm<sup>-1</sup>) were also unsuccessful. This band is expected to be extremely weak (see Fig. 1 and Table S1) and increasing the pressure in hopes of detecting it resulted in full optical saturation of  $v_{10}$  and  $v_{11}$  in the room temperature and warm spectra.

#### 3.5. Combined effective fit

A fit has been performed using the Pickett software for all the literature data and the newly assigned transitions. Energy levels assigned at the ASAP step were not included, nor were the assignments of the  $\nu_{10}$  band from the warm CLS spectrum (only the cold Jet-AILES data were considered for this band). The best-fit constants and relevant fit parameters for the  $\nu_{10}$  and  $\nu_{24}$  bands are summarized in Table 1, while the complete set of constants (all states in the fit) is given in Table S3. For  $\nu_{24}$ , inclusion of two higher-order (sextic and octic:  $H_K$  and  $L_K$ ) CD terms, not needed for the ground state, are required to reproduce

Accepted article 6/10



**Fig. 6.** Evolution of  $\nu_{10}$  band profile as a function of rotational temperature showing an expanded view of the Q-branch (gray insert). Spectra were all recorded at medium resolution (0.01 cm<sup>-1</sup>) but with different sample:He ratios. Rotational temperatures determined from simulations of the  $\nu_{11}$  band profile are indicated on the far right (see text in the Methods section and Fig. S2 for further details).

all assigned transitions to their experimental accuracy (resulting in a reduced standard deviation of 1.1). Taken together with the larger value of  $D_K$  in  $v_{24}=1$  compared to that in v=0, this behavior likely indicates  $v_{24}=1$  is somewhat less "rigid" than the ground state.

The situation for the  $v_{10}$  band is significantly more complicated; the band shows obvious signs of a (strong) perturbation. To reproduce as many transitions from the cold spectrum as possible, three sextic ( $H_K$ ,  $H_{IK}$ , and  $H_{KI}$ ) CD terms are required. These parameters, however, as well as the quartic CD terms, have magnitudes that are much larger than expected (e.g.,  $D_{IK}$  is 38 times larger than the ground state value and  $H_K$  is 50 times larger than the corresponding value in  $v_{24} = 1$ ). Despite their inclusion, 163 transitions (120 different frequencies) were still rejected from the fit using a  $5\sigma$  criterion. These transitions are almost exclusively high I values in each  $K_a$  branch, meaning that the highest values of *J* included in the fit decrease with  $K_a$ . For completeness these transitions are reported in the Supporting Information. In light of the difficulties encountered in the cold Jet-AILES spectrum, which samples a limited range of rotational excitation, no attempt was made to fit the warm CLS spectrum. Based on the available data it appears that a strong perturbation occurs for  $K'_a \ge 4$ , likely from the nearly degenerate dark state  $v_9 + v_{12}$ , if the VPT2 calculations can be trusted.

Treatment of the perturbation in  $v_{10}$  has been attempted. From our quantum chemical calculations, the  $v_9 = v_{12} = 1$  level is predicted to lie at  $1034\,\mathrm{cm}^{-1}$ , and the interaction between this state and  $v_{10} = 1$  is predicted to be Fermi and/or c-type Coriolis, the former with a coupling value of order  $0.8\,\mathrm{cm}^{-1}$ . The same calculation also predicts  $v_{10} = 1$  at  $1012.2\,\mathrm{cm}^{-1}$  (about  $1\,\mathrm{cm}^{-1}$  away from the

experimental value) and scaled rotational constants that are consistently very close to the experimental values for the other bands investigated in this work, i.e. about 0.2 % for *B* and *C*, and 5% for the *A* constant (see Tables 1 and S2). Nevertheless, no satisfactory fit was obtained for the combined fit, regardless of the magnitude of the coupling parameter. At present, we believe this fitting failure is no accident, but rather due to a coincidence between  $K_a = 4$ of  $v_{10} = 1$  and the first  $K_a$  values in  $v_9 = v_{12} = 1$ , based on effective Hamiltonian predictions (Fig. 7). This is consistent with the large residuals associated with the  $K'_a = 4$ value in the  $v_{10}$  band. The main obstacle is the lack of accurate data on the dark  $v_9 = v_{12} = 1$  state: no pure rotational data are available and there is no evidence of the  $v_9 + v_{12}$  band in the experimental spectra, presumably because its calculated intensity is more than two orders of magnitude lower than that of  $v_{10}$ .

While unsatisfactory in some respects, the effective fit reported here improves our understanding of the rovibrational structure of *trans*-butadiene. Concerning the band centers, for instance, the value determined in this work for the  $v_{24}$  band is  $2\,\mathrm{cm}^{-1}$  lower than the previous value, derived from low-resolution spectroscopy reported in Ref. 15. A much closer agreement is obtained for the  $v_{10}$  band  $(0.5\,\mathrm{cm}^{-1})$  as a consequence of this band being a c-type, which is characterized by a sharp Q-branch at room temperature. The rotational constants in the ground state are also extremely reliable, since they have now been derived from measurements of roughly 10,000 lines.

Accepted article 7/10

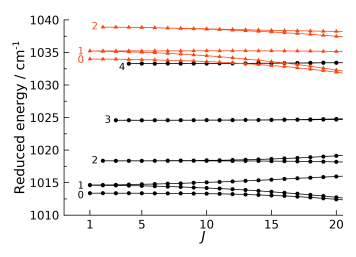
the ground state values. comparison with predicted values. Numbers in parenthesis are one standard deviation in units of the last quoted digit. Brackets indicate parameters fixed to **Table 1.** Rotational constants of butadiene in the ground state,  $v_{24} = 1$ , and  $v_{10} = 1$  after initial assignments using the ASAP software and the final fit in

$N(t)$ $I_{ma}$ $rms$ $\sigma^d$			H	d $a$	D	D	D	C	В	A	E	C	
$N(n)^b$ $J_{\max}, K_{a\max}^c c$ $rms/cm^{-1}$ $\sigma^d$		$H_K \times 10^9$ $L_K \times 10^{12}$	$H_{JK} \times 10^9$ $H_{KJ} \times 10^6$	$d_1 \times 10^9$ $d_2 \times 10^9$	$D_K \times 10^6$	$D_{JK} \times 10^6$	$D_J \times 10^6$					Constant	
10663 (9068) 88, 18 0.00061 1.29	Total			-3.5988(53) -0.1883(24)	7.2929(11)	-0.23874(16)	0.0287444(89)	0.133694116(62)	0.147885664(59)	1.39038285(40)		Final	v = 0
				$\Xi\Xi$	]	<u> </u>	$\equiv$	_		1.3455	299.1	$\mathrm{Pred}^{a}$	
290 44, 14 0.00085 8.52	$v_{24} = 1$			$\overline{\mathbb{L}}$	19.4396(32)	-0.22862(60)	0.027890(37)	0.13364383(222)	0.14793757(207)	1.43140808(57)	297.0970719(258)	Ini. (ASAP)	$v_{24} = 1$
2840 (1953) 87, 18 0.00011 1.08	$v_{24}$	8.485(30) -4.010(49)		-3.5927(71) $-0.5070(27)$	21.9587(59)	-0.24369(17)	0.0278283(90)	0.133616011(65)	0.147960763(63)	1.43163076(55)	297.0923451(62)	Final	
				$\overline{z}$	]	$\Box$	$\equiv$	0.1337	0.1469	1.3950	1013.8	$\mathrm{Pred}.^a$	
134 24,6 0.00803 16.05	$v_{10} = 1$			$\Box$	. <u>.</u>	$-16.39\overline{23}(253)$	$\equiv$	0.13381633(72)	0.14945258(72)	1.3861088(66)	1013.371993(93)	Ini. (ASAP)	$v_{10} = 1$
764 (562) 26, 7 0.00075 1.50	$ u_{10}$	434.(13)	$-6.985(70) \\ 0.3054(15)$	$\Xi$	53.51(82)	-9.191(57)	0.0470(12)	0.13382711(68)	0.14952571(74)	1.386966(15)	1013.368967(66)	Final	

Accepted article 8/10

<sup>&</sup>lt;sup>a</sup> Experimental band centers determined from low-resolution spectroscopy (Ref. 15) and scaled ab initio rotational constants (see text and Tab. S2)
<sup>b</sup> Number of energy levels in ASAP or transitions in SPFIT (N) and number of different frequencies (n) included in the fit (this number may differ from the number of assigned

 $<sup>^</sup>c$  Upper state values except in the column "Total" where the ground state values are reported  $^d$  Reduced standard deviation (unitless)



**Fig. 7.** Rotational energy level diagram in  $v_{10} = 1$  (best fit, black dots) and  $v_9 = v_{12} = 1$  (best prediction, red triangles). The term 1/2(B+C)J(J+1) is subtracted from the energy.  $K_a$  values are noted at the head of each series.

#### 4. Conclusions

While attempting to record the ro-vibrational spectrum of gauche-butadiene, we have re-investigated the far-infrared spectrum of its more stable trans conformer. Two new bands,  $v_{24}$  and  $v_{10}$ , have been observed and assigned, meaning that only one IR active band below  $1100 \text{ cm}^{-1}$ namely  $v_{23}$ , has evaded detection at high resolution. The ASAP software was used to initially assign the spectra of both bands in a few hours. While the assignment was straightforward, the fit has proven more challenging since the  $v_{10}$  band appears strongly perturbed, most likely by the dark  $v_9 = v_{12} = 1$  state. In absence of any experimental information for this state (rotational constants or band center), no reasonable treatment of the perturbation could be achieved. Further investigation of the species under rotationally cooled conditions may prove useful to determine the  $v_9 + v_{12}$  band centers, and ideally rotational constants, assuming a sufficient signal-to-noise ratio can be reached. Such measurements may also enable the detection of the  $\nu_{23}$  band. Nevertheless, now that the far-IR ro-vibrational spectrum of the most abundant conformer of butadiene is nearly complete, renewed attempts to assign the gauche spectrum can be undertaken with greater confidence.

**Supporting Information.** Calibration plots (Figure S1); Experimental Jet-AILES spectra and comparison with simulations (Figure S2); Calculated band centers and intensities (Tables S1); Predicted rotational constants (Table S2); Full list of derived rotational constants (Table S3); Literature transitions for which the estimated uncertainty was increased in the present fit (Table S4). Complete list of observed transitions and fit files (in text format).

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Accepted article 10/10

# Synchrotron-based High Resolution Far-infrared Spectroscopy of *trans*-butadiene

Marie-Aline Martin-Drumel<sup>1</sup>, Jessica P. Porterfield<sup>2</sup>, Manuel Goubet<sup>3</sup>, Pierre Asselin<sup>4</sup>, Robert Georges<sup>5</sup>, Pascale Soulard<sup>4</sup>, Matthew Nava<sup>6</sup>, P. Bryan Changala<sup>7</sup>, Brant Billinghurst<sup>8</sup>, Olivier Pirali<sup>1</sup>, Michael C. McCarthy<sup>2</sup>, and Joshua Baraban<sup>9</sup>

Contact corresponding author(s): marie-aline.martin@universite-paris-saclay.fr

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#### ELECTRONIC SUPPORTING INFORMATION

C	ONIE	N 1 5			LIST	OF FIGURES	
1	Supple	emen	ntary figures	<b>S2</b>	S1 S2	Calibration plots	
2	Supple	emen	ntary tables	<b>S4</b>		son with PGOPHER simulations	\$3
3	Lists o	f ass	igned transitions	<b>S7</b>	LIST (	OF TABLES	
		<sub>24</sub> ba 1.1	Assigned and fitted transitions on the room temperature spectrum		S1 S2	Calculated band centers and intensities Predicted rotational constants in the vibrational states of interest for this study and comparison	S4
	_	<sub>'10</sub> ba 3.1	Assigned and fitted transitions from the		<b>S</b> 3	with experimental values	
	В	3.2	Jet-AILES spectrum  Additional lines assigned in the Jet- AILES spectrum (unfitted)		S4	diene after the global fit performed in this work Transitions from the literature for which the estimated uncertainty was increased	
	В	3.3	Assigned transitions in the warm CLS	S9		estimated uncertainty was increased	30

<sup>&</sup>lt;sup>1</sup> Institut des Sciences Moléculaires d'Orsay (ISMO), CNRS, Univ. Paris-Sud, Université Paris-Saclay, F-91405 Orsay, France

<sup>&</sup>lt;sup>2</sup> Harvard-Smithsonian Center for Astrophysics, Cambridge, Massachusetts 02138, United States

<sup>&</sup>lt;sup>3</sup> Université Lille, CNRS, UMR 8523-PhLAM-Physique des Lasers, Atomes et Molécules, F-59000 Lille, France

<sup>&</sup>lt;sup>4</sup> Sorbonne Universités, UPMC Univ Paris 06, UMR 8233, MONARIS, F-75005 Paris, France & CNRS, UMR 8233, MONARIS, F-75005 Paris, France

<sup>&</sup>lt;sup>5</sup> Institut de Physique de Rennes, CNRS, UMR 6251, Université de Rennes 1, F-35042 Rennes, France

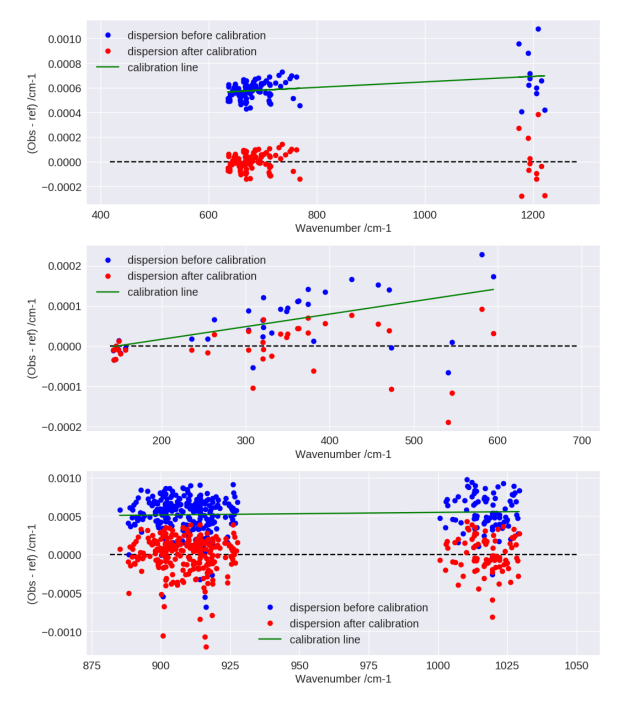
<sup>&</sup>lt;sup>6</sup>Department of Chemistry, Harvard University, Cambridge, MA 02138, United States

<sup>&</sup>lt;sup>7</sup>JILA, University of Colorado, Boulder, CO 80309, United States

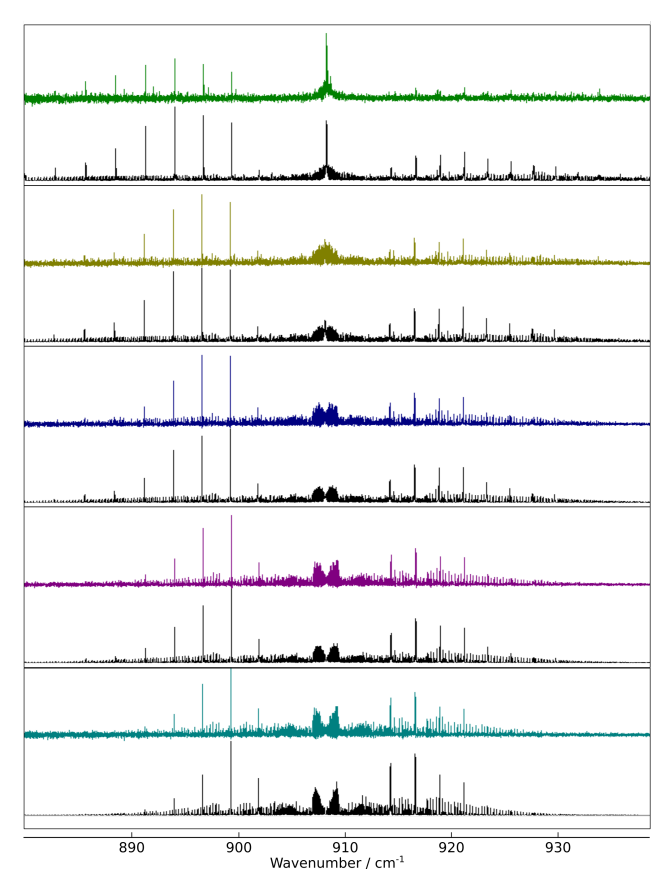
<sup>&</sup>lt;sup>8</sup> Canadian Light Source, University of Saskatchewan, Canada

<sup>&</sup>lt;sup>9</sup> Department of Chemistry, Ben-Gurion University of the Negev Beer-Sheva 8410501, Israel

## 1. SUPPLEMENTARY FIGURES



**Fig. S1.** Calibration plots for the warm spectrum recorded at CLS (top trace), the room temperature one recorded at SOLEIL (middle trace), and the cold spectrum recorded using the Jet-AILES apparatus at SOLEIL (bottom trace). Each dot represent the value of  $\tilde{nu}_{exp} - \tilde{nu}_{ref}$ , where exp refers to the experimental wavenumber and ref the accurate wavenumber used for calibration (see main text for information on reference molecules), before (in blue) and after (in red) calibration. The green lines are the calibration lines whose parameters were used to calibrate the experimental spectra. The residuals after calibration is an indication of the wavenumber accuracy one can expect from the experimental data.



**Fig. S2.** Experimental Jet-AILES spectra recorded with 0, 5, 10, 20, and 30 slm of He seeded in 2 slm of butadiene (from top to bottom, colored traced) and comparison with PGOPHER [1] simulations at temperatures of 150, 100, 70, 50, and 40 K, respectively (black traces)

# 2. SUPPLEMENTARY TABLES

Calculated band centers determined at the CCSD(T)/ANO1 ("Harmonic"), hybrid harmonic/anharmonic Table S1. CCSD(T)/ANO1//CCSD(T)/ANO0 ("Anharmonic"), and VPT2+K ("GUINEA") levels of theory [2]. Intensities calculated at the harmonic CCSD(T)/ANO1 level.

v	Symmetry	F	requency / cm <sup>-1</sup>		IR intensity	/ km⋅mol <sup>-1</sup>
		Harmonic	Anharmonic	GUINEA	Harmonic	GUINEA
13	$a_u$	164.7543	158.1349		0.47	
24	$b_u$	289.6453	290.9936		2.16	
9	$a_g$	509.4167	504.6210		0.00	
12	$a_u$	533.9530	516.9960		11.76	
16	$b_g$	766.0941	738.7697		0.00	
8	$a_g$	895.6995	878.9920		0.00	
11	$a_u$	923.2770	884.8401		72.09	
15	$b_g$	925.1164	889.9208		0.00	
14	$b_g$	988.8452	956.5181		0.00	
23	$b_u$	994.7242	977.3987		1.24	
10	$a_u$	1042.5074	1014.1572	1012.2453	36.11	28.79
7	$a_g$	1222.3291	1194.5204		0.00	
6	$a_g$	1307.8309	1285.3375		0.00	
22	$b_u$	1310.7191	1284.7354		2.09	
21	$b_u$	1408.7609	1369.7341		4.00	
5	$a_g$	1473.3658	1428.5660		0.00	
20	$b_u$	1633.5237	1586.8279		14.39	
4	$a_g$	1695.4519	1651.6894		0.00	
3	$a_g$	3144.9188	2941.0555		0.00	
19	$b_u$	3147.1063	3038.2044		13.43	
2	$a_g$	3156.4329	3041.4562		0.00	
18	$b_u$	3162.1996	3030.8058		10.50	
1	$a_g$	3241.3186	3105.6929		0.00	
17	$b_u$	3241.5700	3106.5926		19.48	
9+12	$a_u$			1034.4503		0.22

**Table S2.** Predicted rotational constants (in  $cm^{-1}$ ) in the vibrational states of interest for this study and comparison with experimental values. Predicted values for the  $v_i = 1$  state have been obtained using the scaling formula  $B_{i \text{calc.}} = B_{i \text{calc.}} \times B_{0 \text{exp.}} / B_{0 \text{calc.}}$ 

$v_i = 1$		Ab initio <sup>a</sup>			Scaled			Е	xperiment	al			$\delta^b$	
i	A	В	C	$\overline{A}$	В	C	-	A	В	<i>C</i>	_	$\delta_{\!A}$	$\delta_B$	$\delta_C$
0	1.3674	0.1445	0.1307	_	_	_		1.3904	0.1479	0.1337				
10	1.3720	0.1435	0.1307	1.3950	0.1469	0.1337		1.3870	0.1495	0.1338		0.58	-1.75	-0.11
24	1.3233	0.1443	0.1307	1.3455	0.1478	0.1338		1.4316	0.1480	0.1336		-6.02	-0.13	0.11
9 + 12	1.3685	0.1446	0.1308	1.3915	0.1480	0.1338								

<sup>&</sup>lt;sup>a</sup> CCSD(T)/ANO0 VPT2 (harmonic) <sup>b</sup>  $\delta = \frac{B_{i \exp } - B_{i scaled}}{B_{i} \text{ scaled}} * 100, \text{ in } \%$ 

**Table S3.** Rotational constants and band centers (in cm<sup>-1</sup>) of butadiene after the global fit performed in this work: refit in the S reduction of all the available literature data (V<sub>13</sub>, V<sub>12</sub>, V<sub>11</sub>, V<sub>20</sub>, and V<sub>17</sub> bands) and new assignments in V<sub>10</sub> and V<sub>24</sub>. Brackets indicate parameters fixed to the GS values. Relevant parameters to the fit are also reported. Numbers in parenthesis are one standard deviation in units of the last quoted digit.

Constant	$\nu = 0$	$v_{13} = 1$	$v_{24} = 1$	$v_{12} = 1$	$v_{11} = 1$	$\nu_{10} = 1$	$v_{20} = 1$	$v_{17}=1$
E		162.419463(12)	297.0923451(62) 524.5738512(91) 908.071968(16)	524.5738512(91)		1013.368967(66)	1596.44603(88) 3100.63227(29)	100.63227(29)
A	1.39038236(43)			1.38902369(45)	1.36826885(95)	1.386966(15)	1.39260(12)	1.387928(28)
C C	0.147885699(62) 0.133694207(64)	0.14/961986(7) 0.134118551(80)	0.14/960/63(63) 0.133616011(65)	0.147747042(66) $0.133708028(68)$	0.147/38901(80) $0.133725433(79)$	0.14952571(74) $0.13382711(68)$	0.14/3896(45) $0.1333877(32)$	0.14/6913(82) $0.1335834(50)$
$D_J{ imes}10^6$	0.0287509(91)	0.030249(11)	0.0278283(90)	0.0288470(96)	0.029224(11)	0.0470(12)	0.0284(13)	
$D_{JK}{ imes}10^6$	-0.23859(17)	-0.24946(27)	-0.24369(17)	-0.25072(18)	-0.20877(29)	-9.191(57)	-4.59(29)	$\equiv$
$D_K \times 10^6$	7.2913(12)	-5.4398(71)	21.9587(59)	7.1452(13)	2.1983(98)	53.51(82)	$\equiv$	$\equiv$
$d_1 \times 10^9$	-3.5889(65)	-3.459(13)	-3.5927(71)	-3.5555(80)	-3.3252(96)		$\equiv$	$\equiv$
$d_2{ imes}10^9$	-0.1888(25)	0.1844(60)	-0.5070(27)		· 	:Ξ	:Ξ	:Ξ
$H_{JK}{ imes}10^9$						-6.985(70)		
$H_{KJ} \times 10^6$						0.3054(15)		
$H_K \times 10^9$			8.485(30)			434.(13)		
$L_K \times 10^{12}$			-4.010(49)					
	Total	<b>V</b> <sub>13</sub>	<i>V</i> 24	V <sub>12</sub>	<i>V</i> <sub>11</sub>	V <sub>10</sub>	<b>V</b> 20	<b>V</b> 17
$N(n)^a$	10619 (9444)	1510 (1468)	2840 (1953)	2838 (2806)	2081(2076)	764 (562)	311 (304)	275 (275)
$J_{\max}$ , $K_{a\max}$	88, 18	70, 10	87, 18	72, 17	72, 10	26, 7	51,4	17, 6
$\rm rms \ / cm^{-1}$	0.00140	0.00033	0.00011	0.00021	0.00032	0.00075	0.00070	0.00302
$\mathbf{Q}_{c}$	1.16	1.54	1.08	1.02	1.04	1.50	0.73	1.01

<sup>a</sup> Number of lines (N) and number of different frequencies (n) considered in the fit (this number may differ from the number of assigned transitions, see text)

<sup>&</sup>lt;sup>b</sup> Upper state values except in the column "Total" where the ground state values are reported <sup>c</sup> Reduced standard deviation (unitless)

**Table S4.** Transitions from the literature for which the estimated uncertainty was increased. The value used in this work is reported in the "Unc." column. Reference to the experimental measurement is also indicated.

Band	J'	$K'_a$	$K'_c$	J''	$K_a^{\prime\prime}$	$K_c''$	Freq.	Unc.	Ref.
$v_{13}$	48	6	42	49	7	42	131.502667	0.002	[3]
$v_{13}$	35	3	32	35	2	34	171.462708	0.002	[3]
$v_{13}$	7	6	1	6	5	1	176.513082	0.002	[3]
$v_{13}$	29	5	25	28	4	25	180.853477	0.002	[3]
$v_{13}$	30	5	26	29	4	26	181.143133	0.002	[3]
$v_{13}$	63	9	54	62	8	54	198.520900	0.002	[3]
$v_{12}$	30	0	30	31	1	30	509.759518	0.002	[3]
$v_{12}$	33	3	31	32	2	31	541.217248	0.002	[3]
$v_{12}$	58	5	53	57	4	53	549.323537	0.002	[3]
$v_{12}$	56	12	44	55	11	44	568.615349	0.002	[3]
$v_{11}$	65	4	62	66	5	62	876.337884	0.003	[4]
$v_{11}$	48	0	48	47	1	46	909.847553	0.003	[4]
$v_{11}$	44	0	44	43	1	42	909.920503	0.003	[4]
$v_{20}$	37	3	35	36	3	34	1606.419508	0.07	[4]
$v_{20}$	38	3	36	37	3	35	1606.679400	0.07	[4]
$v_{20}$	39	3	37	38	3	36	1606.939468	0.07	[4]

# 3. LISTS OF ASSIGNED TRANSITIONS

All lists are reproduced in the following format:  $J' K'_a K'_c J'' K''_a K''_c$  wavenumber uncertainty (wavenumber and uncertainty in cm<sup>-1</sup>)

Below are reported the first 100 lines of each list (by frequency order), the full lists are reported as electronic files.

#### A. $v_{24}$ band

# A.1. Assigned and fitted transitions on the room temperature spectrum

The full list can be found in the file  $jp0c00623\_si\_003.txt$  where  $v_{24} = 1$  is coded at v = 5.

87 1 86	88 1 87	273.381097	0.0001
87 3 84	88 3 85	273.587850	0.0001
86 1 85	87 1 86	273.649241	0.0001
86 3 83	87 3 84	273.840097	0.0001
85 1 84	86 1 85	273.917131	0.0001
85 3 82	86 3 83	274.091812	0.0001
84 1 83	85 1 84	274.185065	0.0001
84 3 81	85 3 82	274.343286	0.0001
83 1 82	84 1 83	274.453095	0.0001
		274.433033	
83 3 80			0.0001
82 1 81	83 1 82	274.720822	0.0001
82 2 80	83 2 81	274.836181	0.0001
82 3 79	83 3 80	274.844949	0.0001
81 1 80	82 1 81	274.988684	0.0001
81 3 78	82 3 79	275.095212	0.0001
81 2 79	82 2 80	275.096833	0.0001
80 1 79	81 1 80	275.256329	0.0001
80 3 77	81 3 78	275.344857	0.0001
80 2 78	81 2 79	275.357385	0.0001
79 1 78	80 1 79	275.524066	0.0001
79 3 76	80 3 77	275.593907	0.0001
79 2 77	80 2 78	275.618045	0.0001
78 1 77	79 1 78	275.791637	0.0001
78 3 75	79 3 76	275.842199	0.0001
78 2 76	79 2 77	275.878435	0.0001
77 1 76	78 1 77	276.059155	0.0001
77 3 74	78 3 75	276.090261	0.0001
77 2 75	78 2 76	276.138788	0.0001
76 1 75	77 1 76	276.326538	0.0001
76 3 73	77 3 74	276.337707	0.0001
76 2 74	77 2 75	276.398775	0.0001
75 3 72	76 3 73	276.584181	0.0001
75 2 73	76 2 74	276.658563	0.0001
74 3 71	75 3 72	276.830003	0.0001
74 2 72	75 2 73	276.918388	0.0001
	74 3 71	277.075368	0.0001
73 2 71	74 2 72	277.177864	0.0001
72 3 69	73 3 70	277.319876	0.0001
72 2 70	73 2 71	277.437073	0.0001
71 3 68	72 3 69	277.564034	0.0001
71 1 70	72 1 71	277.662227	0.0001
71 2 69	72 2 70	277.696161	0.0001
70 3 67	71 3 68	277.807638	0.0001
84 11 73	85 11 74	277.922049	0.0001
70 1 69	71 1 70	277.929348	0.0001
70 2 68	71 2 69	277.954923	0.0001
69 3 66	70 3 67	278.050611	0.0001
69 1 68	70 1 69	278.196057	0.0001
83 11 72	84 11 73	278.204331	0.0001
69 2 67	70 2 68	278.213303	0.0001
68 3 65	69 3 66	278.293448	0.0001
68 1 67	69 1 68	278.462838	0.0001
68 2 66	69 2 67	278.471504	0.0001
82 11 71	83 11 72	278.486327	0.0001
67 3 64	68 3 65	278.536039	0.0001
67 1 66	68 1 67	278.729378	0.0001
67 2 65	68 2 66	278.729378	0.0001
81 11 70	82 11 71	278.768557	0.0001
66 3 63	67 3 64	278.778484	0.0001
66 2 64	67 2 65	278.986606	0.0001
66 1 65	67 1 66	278.995958	0.0001
65 3 62	66 3 63	279.021110	0.0001
80 11 69	81 11 70	279.021110	0.0001
65 2 63		279.030733	0.0001
00 2 03	66 2 64	2/9.243308	0.0001

65	1	64	66	1	65	279.262310	0.0001
79	11	68	80	11	69	279.332789	0.0001
64	2	62	65	2	63	279.500145	0.0001
63	3	60	64	3	61	279.508204	0.0001
64	1	63	65	1	64	279.528605	0.0001
78	11	67	79	11	68	279.614955	0.0001
62	3	59	63	3	60	279.752810	0.0001
63	2	61	64	2	62	279.756200	0.0001
63	1	62	64	1	63	279.794717	0.0001
77	11	66	78	11	67	279.896906	0.0001
61	3	58	62	3	59	279.998654	0.0001
62	2	60	63	2	61	280.011813	0.0001
62	1	61	63	1	62	280.060712	0.0001
76	11	65	77	11	66	280.179226	0.0001
60	3	57	61	3	58	280.245841	0.0001
61	2	59	62	2	60	280.266794	0.0001
61	1	60	62	1	61	280.326592	0.0001
59	3	56	60	3	57	280.494620	0.0001
60	2	58	61	2	59	280.521376	0.0001
60	1	59	61	1	60	280.592208	0.0001
74	11	63	75	11	64	280.743499	0.0001
58	3	55	59	3	56	280.745272	0.0001
59	2	57	60	2	58	280.775360	0.0001
59	1	58	60	1	59	280.857797	0.0001
57	3	54	58	3	55	280.997871	0.0001
73	11	62	74	11	63	281.025208	0.0001
58	2	56	59	2	57	281.028820	0.0001
58	1	57	59	1	58	281.123188	0.0001
56	3	53	57	3	54	281.252655	0.0001
57	2	55	58	2	56	281.281772	0.0001
72	11	61	73	11	62	281.307384	0.0001
57	1	56	58	1	57	281.388315	0.0001
55	3	52	56	3	53	281.509816	0.0001
56	2	54	57	2	55	281.534267	0.0001
71	11	60	72	11	61	281.589284	0.0001
56	1	55	57	1	56	281.653146	0.0001

# B. $v_{10}$ band

# B.1. Assigned and fitted transitions from the Jet-AILES spectrum

The full list can be found in the file  $jp0c00623\_si\_003.txt$  where  $v_{10}=1$  is coded at v=6.

11	7	4	12	8	4	991.253455	0.0005
11	7	5	12	8	5	991.253455	0.0005
18	6	12	19	7	12	992.110766	0.0005
18	6	13	19	7	13	992.110766	0.0005
17	6	11	18	7	11	992.339372	0.0005
17	6	12	18	7	12	992.339372	0.0005
15	6	9	16	7	9	992.804081	0.0005
15	6	10	16	7	10	992.804081	0.0005
14	6	8	15	7	8	993.041078	0.0005
14	6	9	15	7	9	993.041078	0.0005
13	6	7	14	7	7	993.279740	0.0005
13	6	8	14	7	8	993.279740	0.0005
12	6	6	13	7	6	993.521450	0.0005
12	6	7	13	7	7	993.521450	0.0005
11	6	5	12	7	5	993.765834	0.0005
11	6	6	12	7	6	993.765834	0.0005
21	5	16	22	6	16	993.882154	0.0005
21	5	17	22	6	17	993.882154	0.0005
10	6	4	11	7	4	994.013307	0.0005
10	6	5	11	7	5	994.013307	0.0005
20	5	15	21	6	15	994.111527	0.0005
20	5	16	21	6	16	994.111527	0.0005
9	6	3	10	7	3	994.263384	0.0005
9	6	4	10	7	4	994.263384	0.0005
19	5	14	20	6	14	994.344398	0.0005
19	5	15	20	6	15	994.344398	0.0005
8	6	2	9	7	2	994.516495	0.0005
8	6	3	9	7	3	994.516495	0.0005
18	5	13	19	6	13	994.579972	0.0005
18	5	14	19	6	14	994.579972	0.0005
7	6	1	8	7	1	994.772359	0.0005
7	6	2	8	7	2	994.772359	0.0005
17	5	12	18	6	12	994.816881	0.0005

17 5 13	18 6 13		0.0005	12	7 6		8 6	991.023893	0.0005
6 6 0 6 6 1	7 7 0 7 7 1		0.0005	10 10	7 3 7 4		8 3 8 4	991.486854 991.486854	0.0005
16 5 11	17 6 11		0.0005	9	7 2		8 2	991.723093	0.0005
16 5 12	17 6 12		.0005	9	7 3		8 3	991.723093	0.0005
15 5 10	16 6 10		.0005	7	7 0		8 0	992.207873	0.0005
15 5 11 14 5 9	16 6 11 15 6 9		0.0005	7 18	7 1 4 14		8 1 5 14	992.207873 997.020643	0.0005 0.0005
14 5 10	15 6 10		0.0005	17	4 13		5 13	997.287492	0.0005
13 5 8	14 6 8		0.0005	17	4 14		5 14	997.287492	0.0005
13 5 9	14 6 9		.0005	16	4 12		5 12	997.543715	0.0005
12 5 7	13 6 7		0.0005	16	4 13		5 13	997.543715	0.0005
12 5 8 11 5 6	13 6 8 12 6 6		0.0005	15 15	4 11 4 12		5 11 5 12	997.797170 997.797170	0.0005
11 5 7	12 6 7		0.0005	14	4 10		5 10	998.049117	0.0005
10 5 5	11 6 5		.0005	14	4 11		5 11	998.049117	0.0005
10 5 6 9 5 4	11 6 6 10 6 4		0.0005	13	4 9		5 9	998.301750	0.0005
9 5 4 9 5 5	10 6 4 10 6 5		0.0005	13 12	4 10 4 8		5 10 5 8	998.301750 998.555949	0.0005
8 5 3	9 6 3		0.0005	12	4 9		5 9	998.555949	0.0005
8 5 4	9 6 4		.0005	12	4 8		5 8	1002.218440	0.0005
7 5 2 7 5 3	8 6 2 8 6 3		0.0005	12 13	4 9 4 9		5 7 5 9	1002.218440	0.0005
6 5 1	7 6 1		0.0005	13	4 10		5 8	1002.246310	0.0005
6 5 2	7 6 2		0.0005	14	4 10		5 10	1002.275860	0.0005
5 5 0	6 6 0		0.0005	14	4 11		5 9	1002.275860	0.0005
5 5 1 11 4 7	6 6 1 12 5 7		0.0005	19 15	2 17 4 11		3 17 5 11	1002.297620 1002.305550	0.0005
11 4 7	12 5 8		0.0005	15	4 12		5 10	1002.305550	0.0005
10 4 6	11 5 6		0.0005	16	4 12		5 12	1002.334540	0.0005
10 4 7	11 5 7		0.0005	16	4 13		5 11	1002.334540	0.0005
9 4 5 9 4 6	10 5 5 10 5 6		0.0005	17 17	4 13 4 14		5 13 5 12	1002.360690	0.0005 0.0005
8 4 4	9 5 4		0.0005	18	2 16		3 16	1002.300090	0.0005
8 4 5	9 5 5		.0005	17	3 15	17	4 13	1004.904400	0.0005
9 5 4	9 6 4		0.0005	18	3 16		4 14	1004.942270	0.0005
9 5 5 18 3 15	9 6 3 19 4 15		0.0005	19 18	3 17 3 15		4 15 4 15	1004.982560 1004.989480	0.0005
10 5 5	10 6 5		0.0005	20	3 18		4 16	1005.024230	0.0005
10 5 6	10 6 4		.0005	19	3 16		4 16	1005.046270	0.0005
11 5 6 11 5 7	11 6 6 11 6 5		0.0005	13 22	1 13 3 20		2 13 4 18	1005.109580	0.0005 0.0005
12 5 7	12 6 7		0.0005	23	3 21		4 19	1005.152320	0.0005
12 5 8	12 6 6		0.0005	17	2 15		3 15	1007.763860	0.0005
13 5 8	13 6 8		0.0005	18	2 16		3 16	1007.879120	0.0005
13 5 9 15 5 10	13 6 7 15 6 10		0.0005	19 20	2 17 2 18		3 17 3 18	1008.004010 1008.136370	0.0005 0.0005
15 5 11	15 6 9		0.0005	19	2 17		1 19	1011.203580	0.0005
16 5 11	16 6 11		.0005	18	2 16		1 18	1011.441510	0.0005
16 5 12 7 4 3	16 6 10 8 5 3		0.0005	17	2 15 0 24		1 17	1011.691360	0.0005
7 4 4	8 5 3 8 5 4		0.0005	24 25	0 24		1 24 1 25	1013.426480 1013.466850	0.0005 0.0005
17 3 14	18 4 14		0.0005	26	0 26		1 26	1013.504110	0.0005
17 5 12	17 6 12		.0005	27	0 27		1 27	1013.539230	0.0005
17 5 13 18 5 13	17 6 11 18 6 13		0.0005	28 29	0 28 0 29		1 28 1 29	1013.572300 1013.603200	0.0005 0.0005
18 5 14	18 6 12		0.0005	24	1 24		0 24	1013.809240	0.0005
19 5 14	19 6 14	999.980921	.0005	23	1 23	23	0 23	1013.823930	0.0005
19 5 15	19 6 13		0.0005	22	1 22		0 22	1013.842510	0.0005
20 5 15 20 5 16	20 6 15 20 6 14		0.0005 0.0005	13 22	3 10 2 21		2 12 1 21	1015.666260 1015.882610	0.0005
16 3 13	17 4 13		0.0005	17	2 15		1 17	1018.998880	0.0005
6 4 2	7 5 2		.0005	25	3 23		2 23	1019.087220	0.0005
6 4 3 15 3 13	7 5 3 16 4 13		0.0005	24 21	3 22 3 19		2 22 2 19	1019.174020 1019.391020	0.0005 0.0005
5 4 1	6 5 1		0.0005	20	3 18		2 18	1019.447710	0.0005
5 4 2	6 5 2		.0005	19	3 17		2 17	1019.496920	0.0005
				18	3 16		2 16	1019.538420	0.0005
				17 8	3 15 4 4		2 15 3 6	1019.572640 1019.577070	0.0005 0.0005
				19	2 17		1 19	1019.577070	0.0005
R 2 Addition	nal linee aeeian	ed in the Jet-AILES spectru	ım (un-	16	3 14	16	2 14	1019.599950	0.0005
fitted)	na mico accign	ica iii tiic det Aleed opcotie	iii (uii	18	3 15		2 17	1020.118440	0.0005
•				19 20	3 16 3 17		2 18 2 19	1020.206640 1020.307140	0.0005 0.0005
		cluded from the fit is reporte	ed here	22	3 19		2 21	1020.546500	0.0005
in the follow	ing format:			23	3 20		2 22	1020.686240	0.0005
	$K_a'' K_c''$ wavenum	ber uncertainty		17 18	2 15 2 16		1 15 1 16	1021.731470	0.0005 0.0005
	er and uncertaint			18	4 9		3 9	1022.046060	0.0005
(wavellullibe	i and uncertaint	., (iii )		13	4 10	13	3 10	1022.220000	0.0005
1/1 7 7	15 0 7	000 574036	0005	13	4 9		3 11	1022.225020	0.0005
14 7 7 14 7 8	15 8 7 15 8 8		0.0005 0.0005	14 14	4 11 4 10		3 11 3 12	1022.243210	0.0005 0.0005
12 7 5	13 8 5		0.0005	15	4 12		3 12	1022.251390	0.0005

17 4 14 17 2 14 1000 200020 0 0005	where $v_{10} = 1$ is coded at $v = 6$ .	
17 4 14 17 3 14 1022.298830 0.0005 19 2 17 18 1 17 1022.370900 0.0005	10	
20 2 18 19 1 18 1022.703510 0.0005	7 0 7 8 1 7	1009.648040 0.0002
17 3 14 16 2 14 1024.462370 0.0005	8 0 8 9 1 8	1009.308150 0.0002
18 3 15 17 2 15 1024.732760 0.0005	9 0 9 10 1 9	1008.959280 0.0002
19     3     16     18     2     16     1024.999570     0.0005       18     3     16     17     2     16     1025.122720     0.0005	11 0 11 12 1 11 12 0 12 13 1 12	1008.229940 0.0002 1007.848580 0.0002
20 3 17 19 2 17 1025.263080 0.0005	13 0 13 14 1 13	1007.454390 0.0002
22 2 21 21 1 21 1025.359170 0.0005	14 0 14 15 1 14	1007.047010 0.0002
19 3 17 18 2 17 1025.472060 0.0005	15 0 15 16 1 15	1006.626690 0.0002
21 3 18 20 2 18 1025.526020 0.0005	16 0 16 17 1 16	1006.191460 0.0002
13 4 9 12 3 9 1025.887130 0.0005	17 0 17 18 1 17	1005.742730 0.0002
13 4 10 12 3 10 1025.887130 0.0005 14 4 10 13 3 10 1026.194060 0.0005	18 0 18 19 1 18 19 0 19 20 1 19	1005.279640 0.0002 1004.803090 0.0002
14 4 11 13 3 11 1026.198880 0.0005	20 0 20 21 1 20	1004.303030 0.0002
24 2 23 23 1 23 1026.306700 0.0005	21 0 21 22 1 21	1003.810180 0.0002
15 4 11 14 3 11 1026.499950 0.0005	24 0 24 25 1 24	1002.237610 0.0002
15 4 12 14 3 12 1026.507500 0.0005	25 0 25 26 1 25	1001.696170 0.0002
22 3 20 21 2 20 1026.562000 0.0005 23 3 21 22 2 21 1026.938090 0.0005	26 0 26 27 1 26 27 0 27 28 1 27	1001.147820 0.0002 1000.593550 0.0002
17 4 13 16 3 13 1027.101830 0.0005	6 0 6 6 1 6	1012.296550 0.0002
18 4 14 17 3 14 1027.388220 0.0005	7 0 7 7 1 7	1012.352500 0.0002
7 7 0 7 6 2 1029.429890 0.0005	8 0 8 8 1 8	1012.413040 0.0002
7 7 1 7 6 1 1029.429890 0.0005	9 0 9 9 1 9	1012.478440 0.0002
8 7 1 8 6 3 1029.466880 0.0005 8 7 2 8 6 2 1029.466880 0.0005	10 0 10 10 1 10 11 0 11 11 1 11	1012.547490 0.0002 1012.619370 0.0002
9 7 2 9 6 4 1029.507230 0.0005	12 0 12 12 1 12	1012.619370 0.0002 1012.692660 0.0002
9 7 3 9 6 3 1029.507230 0.0005	13 0 13 13 1 13	1012.766540 0.0002
10 7 3 10 6 5 1029.552100 0.0005	14 0 14 14 1 14	1012.840110 0.0002
10 7 4 10 6 4 1029.552100 0.0005	15 0 15 15 1 15	1012.912380 0.0002
13 7 6 13 6 8 1029.708350 0.0005 13 7 7 13 6 7 1029.708350 0.0005	16 0 16 16 1 16	1012.982450 0.0002
13 7 7 13 6 7 1029.708350 0.0005 14 7 7 14 6 9 1029.766060 0.0005	17 0 17 17 1 17 18 0 18 18 1 18	1013.050560 0.0002 1013.114750 0.0002
14 7 8 14 6 8 1029.766060 0.0005	19 0 19 19 1 19	1013.175330 0.0002
15 7 8 15 6 10 1029.826540 0.0005	20 0 20 20 1 20	1013.233010 0.0002
15 7 9 15 6 9 1029.826540 0.0005	21 0 21 21 1 21	1013.286840 0.0002
16     7     9     16     6     11     1029.889820     0.0005       16     7     10     16     6     10     1029.889820     0.0005	22	1013.336430 0.0002 1013.382570 0.0002
17 7 10 17 6 12 1029.954640 0.0005	24 0 24 24 1 24	1013.426390 0.0002
17 7 11 17 6 11 1029.954640 0.0005	25 0 25 25 1 25	1013.466570 0.0002
18 7 11 18 6 13 1030.021240 0.0005	26 0 26 26 1 26	1013.503690 0.0002
18 7 12 18 6 12 1030.021240 0.0005	27 0 27 27 1 27	1013.538760 0.0002
19     7     12     19     6     14     1030.091550     0.0005       19     7     13     19     6     13     1030.091550     0.0005	28	1013.571370 0.0002 1013.602480 0.0002
20 7 13 20 6 15 1030.162590 0.0005	30 0 30 30 1 30	1013.631980 0.0002
20 7 14 20 6 14 1030.162590 0.0005	32 0 32 32 1 32	1013.687460 0.0002
22 5 17 21 4 17 1031.283970 0.0005	33 0 33 33 1 33	1013.713990 0.0002
7 7 0 6 6 0 1031.400880 0.0005 7 7 1 6 6 1 1031.400880 0.0005	34 0 34 34 1 34 35 0 35 35 1 35	1013.739170 0.0002 1013.764050 0.0002
8 7 1 7 6 1 1031.719890 0.0005	37 0 37 37 1 37	1013.704030 0.0002
8 7 2 7 6 2 1031.719890 0.0005	38 0 38 38 1 38	1013.837460 0.0002
9 7 2 8 6 2 1032.042970 0.0005	39 0 39 39 1 39	1013.862020 0.0002
9 7 3 8 6 3 1032.042970 0.0005 10 7 3 9 6 3 1032.369460 0.0005	9 1 8 10 2 8 10 1 9 11 2 9	1007.171500 0.0002 1006.958730 0.0002
10 7 4 9 6 4 1032.369460 0.0005	10 1 9 11 2 9 12 1 11 13 2 11	1006.958730 0.0002 1006.533940 0.0002
12 7 5 11 6 5 1033.032860 0.0005	13 1 12 14 2 12	1006.319790 0.0002
12 7 6 11 6 6 1033.032860 0.0005	14 1 13 15 2 13	1006.102940 0.0002
13 7 6 12 6 6 1033.369770 0.0005	15 1 14 16 2 14	1005.881660 0.0002
13 7 7 12 6 7 1033.369770 0.0005 14 7 7 13 6 7 1033.709920 0.0005	17 1 16 18 2 16 18 1 17 19 2 17	1005.419750 0.0002 1005.177550 0.0002
14 7 8 13 6 8 1033.709920 0.0005	19 1 18 20 2 18	1004.925550 0.0002
15 7 8 14 6 8 1034.052200 0.0005	20 1 19 21 2 19	1004.663100 0.0002
15 7 9 14 6 9 1034.052200 0.0005	21 1 20 22 2 20	1004.388500 0.0002
16 7 9 15 6 9 1034.397040 0.0005	22 1 21 23 2 21	1004.101340 0.0002
16 7 10 15 6 10 1034.397040 0.0005 17 7 10 16 6 10 1034.744360 0.0005	23 1 22 24 2 22 24 1 23 25 2 23	1003.800600 0.0002 1003.486330 0.0002
17 7 11 16 6 11 1034.744360 0.0005	25 1 24 26 2 24	1003.155700 0.0002
19 7 12 18 6 12 1035.444720 0.0005	26 1 25 27 2 25	1002.810550 0.0002
19 7 13 18 6 13 1035.444720 0.0005	27 1 26 28 2 26	1002.448500 0.0002
20 7 13 19 6 13 1035.799460 0.0005	28 1 27 29 2 27	1002.069920 0.0002
20 7 14 19 6 14 1035.799460 0.0005 21 7 14 20 6 14 1036.153760 0.0005	29 1 28 30 2 28 30 1 29 31 2 29	1001.673920 0.0002 1001.260740 0.0002
21 7 15 20 6 15 1036.153760 0.0005	31 1 30 32 2 30	1000.830730 0.0002
22 7 15 21 6 15 1036.510630 0.0005	32 1 31 33 2 31	1000.383800 0.0002
22 7 16 21 6 16 1036.510630 0.0005	34 1 33 35 2 33	999.440171 0.0002
	36 1 35 37 2 35	998.436977 0.0002
	37 1 36 38 2 36 38 1 37 39 2 37	997.914997 0.0002 997.381285 0.0002
	39 1 38 40 2 38	996.836785 0.0002
B 2 Assigned transitions in the warm OLO anastrone (confi	40 1 39 41 2 39	996.282122 0.0002
B.3. Assigned transitions in the warm CLS spectrum (unfit-	41 1 40 42 2 40	995.719652 0.0002
ted)	9 1 8 9 2 8 10 1 9 10 2 9	1010.043290 0.0002 1010.136520 0.0002
The full list can be found in the file:	11 1 10 11 2 10	1010.138320 0.0002
jp0c00623_si_004.txt	12 1 11 12 2 11	1010.347330 0.0002
<u> </u>		

13	1 12	13	2	12	1010.464750	0.0002
14	1 13	14	2	13	1010.589600	0.0002
15	1 14	15	2	14	1010.721510	0.0002
16	1 15	16	2	15	1010.859340	0.0002
18	1 17	18	2	17	1011.152460	0.0002
19	1 18	19	2	18	1011.306260	0.0002
20	1 19	20	2	19	1011.463830	0.0002
21	1 20	21	2	20	1011.624100	0.0002
22	1 21	22	2	21	1011.786340	0.0002
23	1 22	23	2	22	1011.949560	0.0002
24	1 23	24	2	23	1012.113400	0.0002
25	1 24	25	2	24	1012.275910	0.0002
26	1 25	26	2	25	1012.436770	0.0002
27	1 26	27	2	26	1012.594730	0.0002
28	1 27	28	2	27	1012.748300	0.0002
29	1 28	29	2	28	1012.897940	0.0002
30	1 29	30	2	29	1013.041970	0.0002

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