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The structure and low-barrier methyl torsion of 3-fluorotoluene

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Abstract

The microwave rotational spectra of 3-fluorotoluene and its seven ¹³C isotopic species have been recorded at natural abundance in the frequency range from 4 to 26 GHz using a pulsed molecular jet Fourier transform microwave spectrometer. The molecular structure comprising bond lengths and angles as well as parameters describing the methyl torsion were determined with high accuracy. Due to the very low torsional barrier of 17 cm⁻¹, the lowest torsional states of the vibrational ground state exhibited large splittings in the spectrum, which were modeled satisfactorily with a modified version of the program *XIAM* and the program *aixPAM*, two programs developed to treat the methyl internal rotation effects. They were also applied to refit the microwave data of 3,4-difluorotoluene to standard deviations close to measurement accuracy.

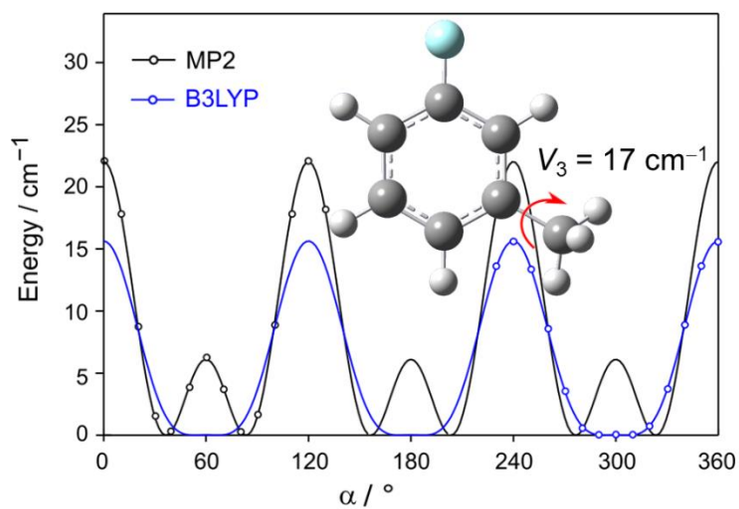
Keywords: large amplitude motion, internal rotation, microwave spectroscopy, structure, dynamics, rotational spectroscopy, 3-fluorotoluene, 3,4-difluorotoluene

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Graphical Abstract



1. Introduction

Toluene and its derivatives have attracted much attention in chemical research including theoretical, spectroscopic, and application fields. Since the first gas phase study of toluene was reported in 1967 by Rudolph et al. [1], the rotational spectrum of toluene has never become an abandoned research target. Further studies have not only reported about an even more precisely determined structure of toluene but also about its interesting internal dynamics arising from the internal rotation of the methyl group with a V_6 potential of 4.837836 cm^{-1} [2-5]. Many halogenated derivatives have subsequently been studied to understand the substitution effects on the structure and the methyl torsion of toluene. Among them, some studies on fluoro-derivatives were reported such as a series of six isomers of difluorotoluene [6-10], as given in Figure 1. The studies have shown that the potential barrier hindering the torsion of the methyl group varies strongly in both, shape and height, depending on the substituted positions of the two fluorine atoms. This is also confirmed in the work on two isomers of trifluorotoluene where the three fluorine atoms attached to the ring significantly affect the methyl internal rotation (see also Figure 1) [11]. Investigations on the three mono-substituted fluorotoluene are also available in the literature [12-14].

While the microwave spectra of all the above mentioned isomers of di- and trifluorotoluene [6-11] as well as 2- [12] and 4-fluorotoluene [14] have been well-understood, there remains a black sheep in the family, 3-fluorotoluene (3FT) which resisted successful characterization. The only microwave study on 3FT by Rudolph and Trinkhaus goes back to 1968, and was a joint-venture in experiment, group theory, and Hamiltonian modeling [13]. This pioneer work using a conventional Stark modulation spectrometer with a frequency accuracy of 100 kHz suffered from problems in the spectral analysis due to the large ratio between V_3 and V_6 terms in the energy potential of the methyl torsion. Though torsional states $m = 0, 1, 2, 3$ of the methyl top could be measured, the very limited number of $m > 0$ transitions led to an ambiguous determination of V_3 and V_6 terms in both value and sign. The combination of the barrier values $15.8(1)$ and $8.0(2) \text{ cm}^{-1}$ for V_3 and V_6 , respectively, (set I in Ref. [13]) was not unique as the transitions could also be reproduced at similar fit quality using respective values of $16.9(1)$ and $-5.3(2) \text{ cm}^{-1}$. Furthermore, while most of the 27 lines from the lowest torsional state $m = 0$ could be fitted at deviations matching the measurement accuracy, the fit quality of the $m > 0$ transitions was unsatisfactory.

Deviations of up to 4.1 MHz were found for the ten $m = 1$ lines, and up to 6.2 MHz for the two $m = 2$ and the five $m = 3$ transitions.

There are two major issues which let 3FT remain a problematic molecule over the last 50 years. First, the energy potential exhibits an extremely low V_3 term accompanied by a V_6 contribution in the same low order of magnitude. In contrast to the potential of 2-fluorotoluene [12], 2,3 [6], 2,4 [7], 2,5-difluorotoluene [8], 2,3,4- [11] and 2,4,5-trifluorotoluene [11], where almost pure V_3 potentials with an intermediate barrier heights close to 200 cm^{-1} dominate, a Hamiltonian model similar to that given in equation (2) of Ref. [13] is no longer sufficient to model simultaneously the $m = 0$ and $m > 0$ states of 3FT. In 4-fluorotoluene [14], 2,6- [10], and 3,5-difluorotoluene [10], the potential exhibits a leading V_6 term by symmetry. The spectrum can be reproduced accurately by including higher order terms available in the program *RAM36* introduced earlier to deal with the rotational spectrum of toluene [5]. The comparable magnitude at very low height of V_3 and V_6 complicates the fitting procedure of 3FT.

Consequently, higher order terms beyond those given in equation (2) of Ref. [13] are required in the Hamiltonian to satisfactorily reproduce the microwave spectrum of 3FT, similar to the case of 2,6- and 3,5-difluorotoluene [10]. Recently, we have modified the program *XIAM*, one of the codes most frequently used to treat the microwave spectra of molecules with methyl internal rotors [15]. *XIAM* uses a combined axis method (CAM), is user-friendly and fast, but often shows difficulties when applied to low torsional barriers. Though some matrix elements are neglected in *XIAM*, a microwave study on *m*-methylanisole has indicated that their effect on the spectral signatures is negligible [16]. Instead, the limited number of high order parameters in *XIAM* seems to be a significant origin for poor fit quality. We have modified the *XIAM* code to deal with the broadband microwave spectrum of 4-methylacetophenone [17]. Two effective higher order parameters D_{3cK} and D_{3c-} , multiplying the potential terms $\cos(3\alpha) P_a^2$ and $\cos(3\alpha) (P_b^2 - P_c^2)$, respectively, have been implemented in addition to the already available D_{3cJ} parameter which is connected with the $\cos(3\alpha) P^2$ term. By using only one of those parameters, D_{3cK} , the root-mean-square (rms) deviation of the 4-methylacetophenone *XIAM* fit was reduced from 99 kHz to 29 kHz, close to the measurement accuracy of about 25 kHz [17]. Applying the modified code to the microwave spectra of the *cis* and *trans* conformer of *m*-methylanisole also decreased the respective deviations from 27.0 and 32.1 kHz to 6.0 and 4.3 kHz, close to the measurement accuracy of 4

kHz, by adding all three D_{3cJ} , D_{3cK} , and D_{3c-} parameters. This again confirms the efficiency of those parameters for low torsional barrier cases [16,18]. The performance of *XIAM* has been cross-checked against the *aixPAM* code which works exclusively in the principal axis system (PAM) and offers several high order effective parameters [16]. We have applied the modified *XIAM* code to test its performance in treating the microwave spectrum of 3FT and check the results against the *aixPAM* code. Furthermore, we also revisited the microwave spectrum of 3,4-difluorotoluene (34DFT), where the performance of the original version of *XIAM* was acceptable, but did not achieve the measurement accuracy [9].

The second remaining issue is the restriction to the main isotopologue in the study of Rudolph and Trinkaus, probably due to the insufficient natural abundance of the minor isotopologues which inhibited the detection of their spectra using a Stark modulated spectrometer [13]. Information on the molecular geometry is limited to the three rotational constants $A = 3715.3(1)$ MHz, $B = 1766.51(1)$ MHz, and $C = 1197.58(1)$ MHz and the angle between the methyl rotor axis and the principal axes of inertia. Alternation of the structure of toluene by inserting substitutions in the ring is known to be important, since geometry parameters such as bond lengths and bond angles change at the point of substitution. The use of pulsed molecular jet Fourier Transform microwave (MJ-FTMW) spectrometers has led to significant improvements in sensitivity and accuracy compared to absorption cells, enabling us to measure and study the isotopic species with even less than 1% natural abundance [21-23]. Beyond remeasuring the microwave spectrum of 3-fluorotoluene, we have also observed the spectra of its seven singly-substituted ^{13}C isotopologues in their natural abundances using an MJ-FTMW spectrometer at Hannover with sub-kHz precision. With such accuracy, we were able to obtain precise molecular parameters for the main isotopologue. Along with those of the ^{13}C isotopologues, the molecular structure of 3FT, including information on bond lengths and bond angles, was derived and compared with those of toluene and other fluoro-substituted toluene derivatives.

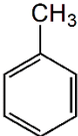
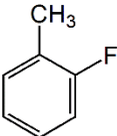
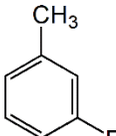
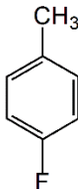
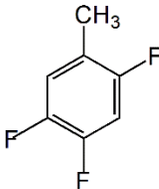
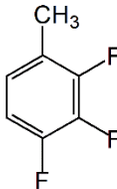
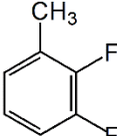
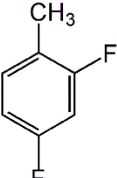
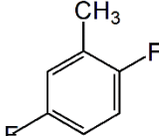
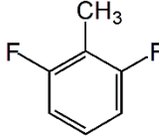
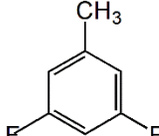
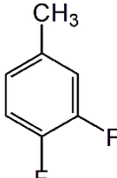
						
(1)	(2) 2FT	(3) 3FT	(4) 4FT	(5) 2,4,5-TFT	(6) 2,3,4-TFT	
V_3 / cm^{-1}	/	227.28(2)	16.8978(3)	/	190.67(20)	216.32(67)
V_6 / cm^{-1}	4.837836(1)		4.8298(54)			
						
(7) 2,3-DFT	(8) 2,4-DFT	(9) 2,5-DFT	(10) 2,6-DFT	(11) 3,5-DFT	(12) 3,4-DFT	
V_3 / cm^{-1}	210.547(13)	234.182(69)	215.7(10)	/	/	33.9624(31)
V_6 / cm^{-1}			12.432(20)	7.156(84)		

Figure 1. A selection of fluoro-derivatives of toluene: (1) toluene [5], (2) 2-fluorotoluene [12], (3) 3-fluorotoluene ([13] and this work), (4) 4-fluorotoluene [14], (5) 2,4,5-trifluorotoluene [11], (6) 2,3,4-trifluorotoluene [11], (7) 2,3-difluorotoluene [6], (8) 2,4-difluorotoluene [7], (9) 2,5-difluorotoluene [8], (10) 2,6-difluorotoluene [10], (11) 3,5-difluorotoluene [10], and (12) 3,4-difluorotoluene ([9] and this work).

2. Theoretical predictions

2.1. Geometry optimizations

To support the experimental work, we used two quantum chemistry methods available in the *Gaussian 16* program package [24]. The first optimization was performed using the Kohn-Sham density functional theory [25] employing Becke's three parameter hybrid exchange functional [26] and the Lee-Yang-Parr correlation functional [27] (B3LYP), which turned out to be adequate for predictive guidance in many previous studies on molecules containing a phenyl ring [28-31]. For comparison, calculations were repeated using the second order Møller-Plesset perturbation method (MP2) [32]. The split-valence triple-zeta basis set 6-311++G(2d,2p) was chosen in combination

with both methods. The equilibrium geometry of 3FT optimized at the MP2/6-311++G(2d,2p) level is illustrated in Figure 2. The Cartesian coordinates of the atoms are given in Table S1 of the Supplementary Material. Vibrational frequency calculations in the harmonic approximation confirmed that the optimized geometry is indeed a minimum and not a saddle point. Anharmonic frequency calculations were carried out to obtain vibrational ground state rotational constants and quartic centrifugal distortion constants.

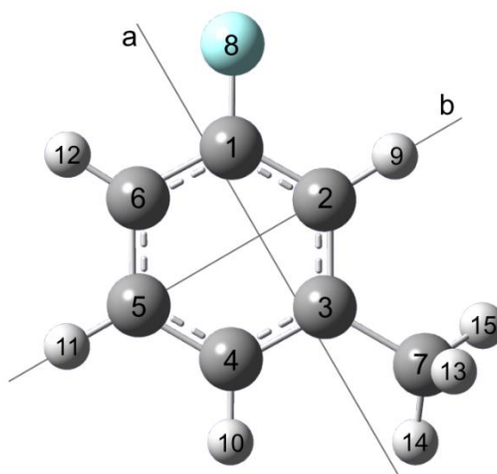


Figure 2. Projection onto the ab -plane of 3FT in its principal axis system. The molecular geometry was optimized at the MP2/6-311++G(2d,2p) level of theory.

2.2. Methyl internal rotation

We started calculating the energy potential function at the MP2/6-311++G(2d,2p) level of theory by varying the dihedral angle $\alpha = \angle(\text{C}_2, \text{C}_3, \text{C}_7, \text{H}_{13})$ on a grid of 10° while all other geometry parameters were optimized. Because of the symmetry of the methyl group, a rotation span of 120° was sufficient. The obtained potential energy points were parameterized with a one-dimensional Fourier expansion and then drawn as a contour plot presented in Figure 3. The potential curve indicates three-fold double minima in the regions $0-120^\circ$, $120-240^\circ$, as well as $240-360^\circ$, i.e. local maxima at $\alpha_{\text{max}} = \alpha_i = 60^\circ, 180^\circ, \text{ and } 300^\circ$ separating the double-minima at $\alpha_{\text{min}} \approx \alpha_i \pm 24^\circ$. The V_3 and V_6 coefficients are 16.30 and 12.60 cm^{-1} , respectively, and a small V_9 term of -0.38 cm^{-1} was also used to fit the potential energy points. In contrast, the potential curve calculated

with the B3LYP method results in a three-fold potential with a dominant $V_3 = 15.61 \text{ cm}^{-1}$ term and only a small V_6 contribution of 4.18 cm^{-1} , leading to single energy minima at α_i . However, we noticed that the minima appear extremely broad and repeated the calculations in the region of 60° around α_i with a finer step size of 2° . The results depicted in the inset of Figure 3 reveals that the B3LYP method indeed predicts somewhat closer lying double minimum which is present at $\alpha_{\text{min}} \approx \alpha_i \pm 10^\circ$, with the dividing barrier being lower than 0.1 cm^{-1} .

We noticed four conformations on the potential curves in Figure 3, each of which appears three times during a full rotation of the methyl group. In the conformation at $\alpha = 0^\circ$, the H_{13} atom is eclipsed with the H_9 atom if we look along the a -axis. The second one at $\alpha \approx 36^\circ$ (MP2 value) has the $\text{C}_7\text{--H}_{14}$ bond roughly perpendicular to the ring plane. In the third conformation with $\alpha = 60^\circ$, the H_{15} atom is eclipsed with the H_{10} atom. The fourth conformation at $\alpha \approx 84^\circ$ has the $\text{C}_7\text{--H}_{13}$ bond roughly perpendicular to the ring plane and is equivalent to the $\alpha \approx 36^\circ$ position, assuming the planarity of the phenyl ring as expected for an aromatic system. Full geometry optimizations following by harmonic frequency calculations confirmed all of them to be stationary points, as summarized in Table 1, with those at $\alpha = 36^\circ$ and $\alpha = 84^\circ$ as the most stable rotamers for the MP2 method as well as $\alpha = 49^\circ$ and $\alpha = 71^\circ$ for the B3LYP method. Both methods predicted the conformers at $\alpha = 0^\circ$ and 60° to correspond to saddle points. The estimation for the V_3 potential including the zero-point energy (ZPE) correction is also given in Table 1. With both methods, the difference of the ZPE corrections for the double-minimum and the barrier dividing it is larger than the value for this barrier, i.e. the zero-point vibrational level lies above the barrier at $\alpha_i = 60^\circ$. Nevertheless, this difference is smaller than the value of the three-fold barrier, indicating that the potential can be effectively described by a V_3 term.

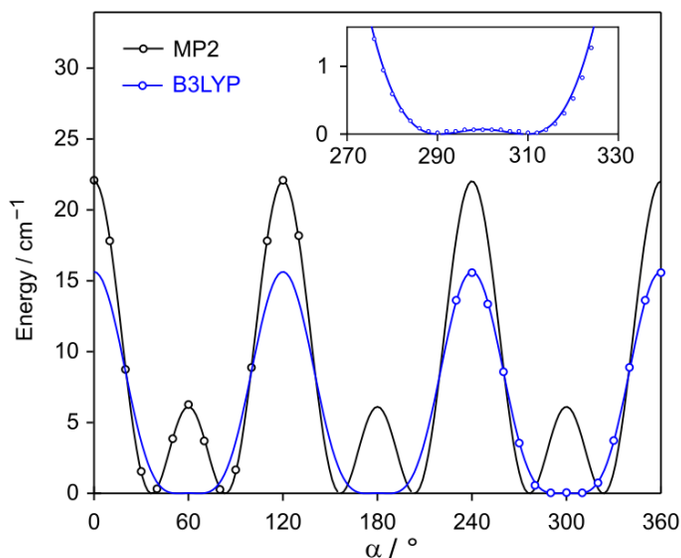


Figure 3. Potential energy curves of the methyl torsion of 3FT obtained by varying the dihedral angle $\alpha = \angle(\text{C}_2, \text{C}_3, \text{C}_7, \text{H}_{13})$ in a grid of 10° , while all other molecular parameters were optimized at the B3LYP/6-311++G(2d,2p) (in blue) and MP2/6-311++G(2d,2p) (in black) levels of theory. The energies are relative to the energetically lowest conformations. Inset: in the range from 270° to 330° , calculations at the B3LYP/6-311++G(2d,2p) level were repeated at a finer step of 2° to capture the double minimum at $\alpha = 290^\circ$ and 310° separated by a very flat local maximum at $\alpha = 300^\circ$.

Table 1. Dihedral angle α , the path α_{path} between the structures, the energy at the stationary points E , the energy difference ΔE between the structures, the energy with vibrational corrections $E+ZPE$, and the energy difference $\Delta(E+ZPE)$ between the structures of 3FT calculated at the B3LYP/6-311++G(2d,2p) and MP2/6-311++G(2d,2p) levels of theory.

Method	$\alpha / ^\circ$	Type	E / Hartree	$E+ZPE$ / Hartree	α_{path}	ΔE / cm^{-1}	$\Delta(E+ZPE)$ / cm^{-1}
B3LYP	0.0	max	-370.9214200	-370.802192	0.0 \rightarrow 10.7	-15.71	-7.24
	10.7	min	-370.9214916	-370.802225	60.0 \rightarrow 10.7	-0.04	6.58
	60.0	max	-370.9214914	-370.802255	60.0 \rightarrow 0.0	15.67	13.83
MP2	0.0	max	-369.9500796	-369.830411	0.0 \rightarrow 23.8	-22.10	10.75
	23.8	min	-369.9501803	-369.830362	60.0 \rightarrow 23.8	-6.28	28.09
	60.0	max	-369.9501517	-369.830490	60.0 \rightarrow 0.0	15.82	17.34

3. Microwave Spectroscopy

3.1. Experimental setup

The microwave spectrum of 3FT was recorded in the frequency range from 4 to 26 GHz using a pulsed supersonic MJ-FTMW spectrometer in Hannover with a coaxially oriented beam-resonator arrangement (COBRA) [33]. The longer transit time of the molecular jet, corresponding to line-shapes that are narrower in frequency, results in an accuracy advantage of the COBRA compared to a perpendicular arrangement. The substance was kept in a small reservoir at the nozzle orifice, and neon or helium as carrier gas was passed over the sample at a stagnation pressure of 1 bar. All frequency measurements were referenced to a GPS-disciplined rubidium frequency standard. The estimated uncertainty is better than 1 kHz for unblended lines while transitions separated less than 5 kHz are not resolved. The adiabatic expansion simultaneously reduces the collisional as well as Doppler line broadening and the molecules experience a strong rovibrational cooling, leaving only the lowest-lying rovibrational levels populated, thus efficiently simplifying the observed spectrum.

The molecular electric dipole moments were determined using the Stark effect. Instead of external electrode plates, we used the spherical reflectors forming the resonator as high-voltage electrodes (Coaxially Aligned Electrodes for Stark effect Applied in Resonators, CAESAR [34]). With the electromagnetic radiation propagating in TEM_{00q} modes, the direction of the static electric Stark field is perpendicular to the polarized microwave field, leading exclusively to the $\Delta M_J = \pm 1$ transitions. Stark effect measurements in the $J = 1 \leftarrow 0$ rotational transitions of the OC³⁶S and ¹⁸OCS isotopologues of OCS were used to calibrate the electric field strength in the cell using a dipole moment value of $\mu = 0.71519(3)$ D [35], as described in Ref. [36].

3.2. Measurements

The rotational constants determined in Ref. [13] were used to predict the rigid rotor spectrum of 3FT in the frequency range of the spectrometer. Since only transitions originating from the ground vibrational state $\nu = 0$ are expected in our jet cooled spectrum, we targeted the associated two lowest torsional state $m = 0$ and $m = 1$, henceforth called the A and the E species. The lines of the A symmetry species follow a rigid rotor pattern and could be fitted well to a standard deviation of 0.7 kHz using an effective Hamiltonian $H = H_r + H_{cd}$ comprising rotational and quartic centrifugal distortion terms of Watson's S reduction in its F' representation, as implemented in the program

SFLAMS written for Separately Fitting of Large Amplitude Motion Species [17,37]. The V_3 potential term and the angle $\angle(i,a)$ between the internal rotor axis and the a -principal axis of inertia reported in Ref. [13] was helpful to identify and assign the E species lines. The assignments of the E species lines were also checked with *SFLAMS* using the additional odd power term

$$H_{op} = (q + q_J P^2 + q_K P_a^2) P_a + (r + r_J P^2 + r_K P_a^2) P_b \quad (1)$$

in the Hamiltonian, where P is the angular momentum operator with its components P_a, P_b, P_c referring to the principal axes of inertia a, b , and c . The parameters q and r are sometimes called D_a and D_b in the literature where the higher order terms q_J, q_K, r_J , and r_K can be also found as D_{aJ}, D_{aK}, D_{bJ} , and D_{bK} . The separate *SFLAMS* fit for the E species achieved a standard deviation of 2.7 kHz. Typical spectra of the A and E symmetry species components of a rotational transition are shown in Figure 4.

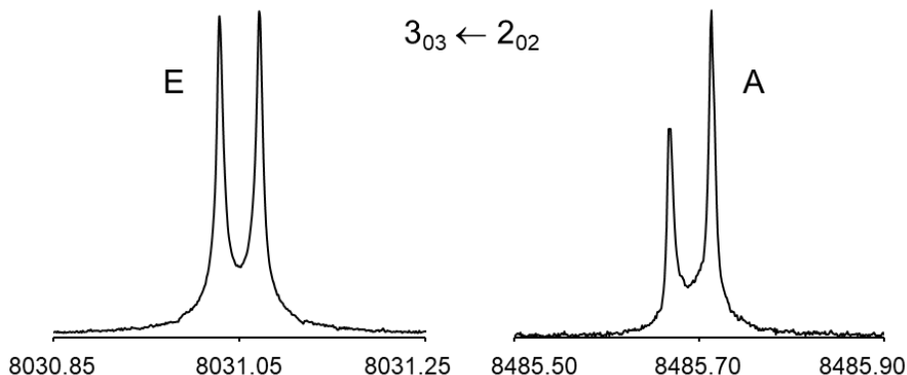


Figure 4. A typical $J''_{K''_a K''_c} \leftarrow J'_{K'_a K'_c}$ rotational transition of 3FT which splits into an A and an E symmetry species. The doublets arise from the Doppler effect; the mean frequency is 8031.0496 MHz for the E species and 8485.6905 MHz for the A species. For these spectra, 771 and 939 free-induction decays, respectively, were co-added.

3.3. Spectral analysis

The rotational transitions of 3FT were fitted using a modified version of the program *XIAM*, henceforward called the *XIAM_{mod}* code [17,18], available at the PROSPE website [38]. The Hamiltonian can be written as

$$H = H_r + H_{cd} + H_{ir} + H_{ird}. \quad (2)$$

The centrifugal distortion term H_{cd} in the Hamiltonian comprises standard fourth order terms for a semi-rigid rotor according to Watson's S-reduction. The standard rigid frame-rigid top Hamiltonian is similar to that given in equation (2) of Ref. [13]:

$$H_r + H_{ir} = AP_a^2 + BP_b^2 + CP_c^2 + F\pi_\alpha^2 + V(\alpha), \quad (3)$$

which contains the structural rotational constants and the reduced rotational constant F of the methyl top. $\pi_\alpha = p_\alpha - \rho_a P_a - \rho_b P_b$ contains the angular momentum p_α of the methyl top and $\rho_g = \lambda_g I_\alpha / I_g$ with $g = a, b$. I_α is the moment of inertia of the methyl top, I_g are the components of principal moments of inertia of the molecule, and λ_g are the direction cosines between the internal rotation axis i and the respective inertial axes. The leading terms of the hindering potential $V(\alpha)$ can be written as in equation (2) of Ref. [13]:

$$V(\alpha) = \frac{V_3}{2}[1 - \cos(3\alpha)] + \frac{V_6}{2}[1 - \cos(6\alpha)]. \quad (4)$$

Generally, higher order potential terms (V_6 , V_9 etc.) cannot be determined from the vibrational ground state alone and thus are normally neglected. In the previous work on 3FT, Rudolph and Trinkaus used a Stark modulated spectrometer and could obtain the spectra in four torsional states $m = 0, 1, 2, 3$ [13]. It was thus possible to derive both the V_3 and the V_6 terms of the potential. Since our study was limited to the ground torsional state because of jet expansion cooling, we could only float either V_3 or V_6 in our fits. In the case of 3FT, only V_3 is floated.

H_{ird} is not included in the Hamiltonian given in Ref. [13]. It gives the empirical internal rotation - overall rotation distortion operator in the principal axis system and includes the two parameters D_{c3K} and D_{c3-} newly implemented in *XIAM*. In the case of 3FT, it can be written as:

$$H_{ird} = 2D_{2\pi J}(p_\alpha - \vec{\rho}^+ \vec{P})^2 P^2 + D_{2\pi K} \left\{ (p_\alpha - \vec{\rho}^+ \vec{P})^2, P_a^2 \right\} + D_{2\pi-} \left\{ (p_\alpha - \vec{\rho}^+ \vec{P})^2, (P_b^2 - P_c^2) \right\} + D_{c3J} \cos(3\alpha) P^2 + D_{c3K} \cos(3\alpha) P_a^2 + D_{c3-} \cos(3\alpha) (P_b^2 - P_c^2). \quad (5)$$

where \vec{P} is the angular momentum vector along the rho axis.

A total of 61 A species and 54 E species lines could be measured and fitted, yielding a quite satisfactory standard deviation of 3.3 kHz. The fitted parameters are collected in the column *XIAM_{mod}* of Table 2. The complete list of all fitted rotational transitions along with their residuals

is given in Table S2 of the Supplementary Material. The original version of *XIAM* without the D_{c3K} and D_{c3-} parameters yielded a deviation of 26.3 kHz. Omitting also the third effective high order parameters D_{c3J} leads to another small increase of the deviation to 27.1 kHz.

We finally applied the *aixPAM* code [16] to cross-check the performance of *XIAM_{mod}*. *aixPAM* works in the principal axis system and can fit the torsion–rotation spectra of molecules with one methyl rotor. Unlike *XIAM*, *aixPAM* does not neglect any matrix elements except for the truncation of the matrix (k_{\max}), and effective high order Hamiltonian terms can be added from the input file. Using a comparable set of parameters, *aixPAM* slightly improved the standard deviation of the *XIAM_{mod}* fit from 3.3 kHz to 1.5 kHz. The fit is shown in the column *aixPAM* of Table 2. The residuals given by *aixPAM* are also presented in Table S2 of the Supplementary Material.

3.4. The ^{13}C isotopologues

Transitions originating from the ^{13}C isotopologues arising from seven inequivalent substitution positions of the carbon skeleton have also been measured in their natural abundance of 1%. The Cartesian coordinates of the optimized structure in Figure 2 were taken, and the mass of the individual carbon atoms was adjusted to calculate the rotational constants of all ^{13}C isotopologues. The predicted ^{13}C -frequencies were corrected with the differences between the frequencies predicted with the B_e constants obtained at the MP2/6-311++G(2d,2p) level and the experimental frequencies of the main species, leading to straightforward assignments of all ^{13}C isotopologues. Since the number of E species lines are small, only the rotational constants, the centrifugal distortion constants, the torsional barrier V_3 , the angle $\angle(i,a)$, and $F_0 = \frac{h}{8\pi^2 I_\alpha}$ were fitted. All higher order terms associating with H_{ird} were kept fixed at the respective values of the main species. The A and E transitions of all ^{13}C isotopologues were also checked by separate fits, unless the smaller number of observed E species lines was insufficient for an independent analysis. The molecular and internal rotation parameters of the ^{13}C species are collected in Table 3. The lists of fitted transitions and their residuals are available in Table S3 of the Supplementary Material.

Table 2. Molecular parameters of 3FT obtained using the modified *XIAM* code and the program *aixPAM*.

Par. ^a	Unit	3-fluorotoluene (3FT)				3,4-difluorotoluene (34DFT)	
		<i>XIAM</i> _{mod} ^b	<i>aixPAM</i> ^c	<i>MP2</i> ^d	<i>B3LYP</i> ^e	<i>XIAM</i> _{mod} ^b	<i>aixPAM</i> ^c
<i>A</i>	MHz	3657.122(32)	3632.76(18)	3654.2	3664.5	3037.5386(14)	3035.870(26)
<i>B</i>	MHz	1762.2985(23)	1762.863(13)	1759.6	1763.8	1290.69640(20)	1293.76(40)
<i>C</i>	MHz	1197.57678(13)	1198.242(14)	1196.5	1199.5	910.87391(12)	908.49(40)
<i>D_J</i>	kHz	0.0676(18)	0.06102(84)	0.0657	0.1377	0.02312(54)	0.02177(47)
<i>D_{JK}</i>	kHz	-0.107(11)	-0.0125(53)	-0.0049	0.0861	0.0704(93)	0.1031(723)
<i>D_K</i>	kHz	-0.76(16)	0.577(88)	0.6762	0.5580	0.588(57)	0.403(44)
<i>d₁</i>	kHz	-0.0231(13)	-0.02399(56)	-0.0244	-0.0256	-0.00792(33)	-0.00772(25)
<i>d₂</i>	kHz	0.00232(74)	-0.00348(56)	-0.0013	0.0335	-0.00136(28)	-0.0015236(26)
<i>D_{π2J}</i>	MHz	0.0495(21)	0.02420(73)				
<i>D_{π2K}</i>	MHz	1.567(28)	0.6671(75)				
<i>D_{π2-}</i>	MHz	0.1457(49)	0.04401(14)			-0.517(38)	-0.203(31)
<i>V₃</i>	cm ⁻¹	17.1785(86)	16.4737(46)	16.30	15.61	32.5042(2)	32.4860(11)
<i>F₀</i>	GHz	168.669(92)	164.302(40)	162.021	161.742	162.116	162.116
<i>V_J/-D_{c3J}</i> ^f	MHz	-0.187(22)	-0.787(16)			-0.2489(17)	-0.4849(27)
<i>V_K/-D_{c3K}</i> ^f	MHz	22.31(30)	28.35(21)			1.225(11)	2.910(37)
<i>V₋/-D_{c3-}</i> ^f	MHz	1.004(50)				-4.49(32)	-3.65(53)
$\angle(i,a)$ ^g	°	29.6131(2)	29.4596(11)	30.0551	30.0151	12.5579(2)	12.5247(37)
<i>N_A/N_E</i> ^h		61/54	61/54			51/33	51/33
σ ⁱ	kHz	3.4	1.5			3.5	2.4

^a All parameters refer to the principal axis system. Watson's S reduction and *I'* representation were used. Standard errors are in the units of the last digits. ^b Fit with the modified version of *XIAM*. ^c Fit with the program *aixPAM*. ^d Calculated at the MP2/6-311++G(2d,2p) level of theory. The rotational constants refer to the equilibrium structure. The vibrational ground state constants are $A_0 = 3625.6$, $B_0 = 1750.3$, and $C_0 = 1189.1$ MHz. Centrifugal distortion constants are obtained from anharmonic frequency calculations. ^e Calculated at the B3LYP/6-311++G(2d,2p) level of theory. The vibrational ground state constants are $A_0 = 3602.9$, $B_0 = 1765.2$, and $C_0 = 1193.3$ MHz. ^f V_J , V_K , V_- in *aixPAM* and D_{3cJ} , D_{3cK} , D_{3c-} in *XIAM*_{mod}. ^g $\angle(i,b) = 90^\circ - \angle(i,a)$. $\angle(i,c)$ was fixed to 90° in all fits due to symmetry. ^h Number of the A and E species lines. ⁱ Standard deviation of the fit.

Table 3. Molecular parameters of the ^{13}C isotopologues of 3FT obtained from the *XIAMmod* fits. Atoms are numbered according to Figure 2. Values of the $D_{\pi^2 J}$, $D_{\pi^2 K}$, $D_{\pi^2 -}$, D_{c3J} , D_{c3K} , and D_{c3-} are fixed to those of the main species.

Par. ^a	Unit	$^{13}\text{C}(1)$	$^{13}\text{C}(2)$	$^{13}\text{C}(3)$	$^{13}\text{C}(4)$	$^{13}\text{C}(5)$	$^{13}\text{C}(6)$	$^{13}\text{C}(7)$
<i>A</i>	MHz	3655.331(20)	3632.090(15)	3655.071(21)	3624.980(26)	3573.091(15)	3624.993(18)	3630.5746(73)
<i>B</i>	MHz	1754.5982(18)	1762.37419(99)	1752.5249(19)	1752.8152(19)	1762.3031(17)	1753.7662(17)	1722.29698(57)
<i>C</i>	MHz	1193.83151(38)	1194.89902(24)	1192.85255(35)	1189.75746(40)	1188.43973(37)	1190.20317(36)	1176.20068(16)
<i>D_J</i>	kHz	0.0157(60)	0.0673(23)	0.0717(58)	0.0640(37)	0.0670(49)	0.0631(51)	0.0658(15)
<i>D_{JK}</i>	kHz	-0.46(15)	-0.107 ^b	-0.107 ^b	-0.107 ^b	-0.107 ^b	-0.107 ^b	-0.107 ^b
<i>D_K</i>	kHz	-0.76 ^b	-0.76 ^b	-2.21(70)	-0.76 ^b	-0.76 ^b	-0.76 ^b	-0.76 ^b
<i>d₁</i>	kHz	-0.0302(45)	-0.0246(17)	-0.0223(43)	-0.0186(28)	-0.0212(35)	-0.0173(35)	-0.0227(11)
<i>d₂</i>	kHz	0.0321(38)	0.00232 ^b	0.0053(21)	0.00232 ^b	0.00232 ^b	0.00232 ^b	0.00232 ^b
<i>V₃</i>	cm ⁻¹	17.1827(44)	17.1792(46)	17.1549(46)	17.2182(87)	17.1393(38)	17.1497(45)	17.1764(21)
<i>F₀</i>	GHz	168.696(50)	169.038(39)	168.394(59)	168.468(70)	168.295(41)	168.417(49)	168.493(20)
$\angle(i,a)$	°	29.7295(2)	29.5937(8)	29.4791(2)	30.1496(12)	29.6344(1)	29.1071(2)	28.6314(3)
<i>N_A/N_E</i> ^c		39/15	38/10	30/12	32/10	26/13	33/13	48/12
σ^d	kHz	6.5	3.3	5.4	5.3	5.1	6.6	3.0

^a All parameters refer to the principal axis system. Watson's S reduction and I' representation were used.

^b Fixed to the value of the main species.

^c Number of the A and E species lines.

^d Standard deviation of the fit .

3.5. Reanalysis of 3,4-difluorotoluene

Though the microwave spectrum of 34DFT has been understood reasonably well, the fit using the original version of *XIAM* has not achieved measurement accuracy [9]. An order of magnitude larger standard deviation of 47 kHz has been reported and attempts to reduce it were not successful.

By separately fitting the A and E species lines with *SFLAMS*, we found that one A symmetry species and three E symmetry species transitions reported in Ref. [9] were misassigned. Eliminating these four transitions did not significantly reduce the standard deviation, which was the reason why these misassignments were not recognized in Ref. [9]. Excluding the misassigned transitions and using the three D_{3c} parameters provided in the *XIAM_{mod}* code reduces the deviation of the fit to 3.5 kHz (given in the column *XIAM_{mod}* of Table 2), while it remains at 40 kHz for the data set reported in Ref. [9].

Applying the *aixPAM* code to the corrected data set using a set of parameters comparable to the *XIAM_{mod}* code, only a slightly better deviation of 3.0 kHz is obtained. The fitted rotational transitions and deviations in both fits are given in Table S4 of the Supplementary Material.

3.6. Electric dipole moment

Rudolph and Trinkaus determined the electric dipole moment components of 3FT to be $\mu_a = 1.71(2)$ D and $\mu_b = 0.62(2)$ D from the Stark satellite of the $3_{03} \leftarrow 2_{02}$ and $3_{22} \leftarrow 2_{21}$ A species transitions [13]. To improve the accuracy of these values, we performed Stark measurement using the CAESAR setup of the spectrometer. The measurements were performed at different electric fields and limited to three low- J transitions given in Table S5. The observed Stark transitions were fitted with the program QSTARK [39], yielding a total dipole moment of $|\mu| = 1.8408(51)$ D with the components $|\mu_a| = 1.7327(23)$ D and $|\mu_b| = 0.6217(86)$ D. The dipole moment component in the c -direction is zero due to symmetry.

4. Microwave structure determinations

The r_s structure of 3FT was derived from multiple isotopic data using the rotational constants of the main species given in Table 2 and those obtained for seven ^{13}C isotopologues collected in Table 3. The position of each substituted carbon atom was first calculated using the substitution method of Kraitchman, which provides the absolute atomic coordinates in the principal axis system

[40,41]. The uncertainties were calculated according to Costain's estimate error [42] and the signs of the atom coordinates were taken to be consistent with the MP2-optimized geometry. The resulting atomic coordinates are given in Table 4. The bond lengths and bond angles are determined with the program EVAL available at the PROSPE website [38] and given in Table 5. For the EVAL input, all values of the c -coordinates were set to zero.

Table 4. Experimental atom positions (substitution r_s and semi-experimental r_0 structures) of 3FT obtained by isotopic substitutions with Kraitchman's equations [40,41] as implemented in the programs KRA and STRFIT [43]. All c -coordinates of the KRA output were set to zero to be consistent with planarity. For comparison, the equilibrium atom positions (r_e structure) obtained from optimizations at the MP2/6-311++G(2d,2p) level of theory are presented.

	r_s		r_0			r_e		
	$a/\text{Å}$	$b/\text{Å}$	$a/\text{Å}$	$b/\text{Å}$	$c/\text{Å}$	$a/\text{Å}$	$b/\text{Å}$	$c/\text{Å}$
C(1)	-1.1243(14)	-0.2615(60)	-1.2777(15)	-0.2813(24)	0.01084(2)	-1.1310	-0.2666	-0.0013
C(2)	0.05916*	-0.9804(16)	-0.0596(25)	-0.9796(19)	0.00831(1)	0.0592	-0.9774	0.0083
C(3)	1.2668(13)	-0.2772(56)	1.1336(20)	-0.2699(24)	-0.00135(2)	1.2743	-0.2838	0.0109
C(4)	1.2434(13)	1.1158(14)	1.1794(22)	1.1198(21)	-0.00746(2)	1.2488	1.1149	0.0073
C(5)	0.03827*	1.80745(86)	-0.0330(26)	1.81279(59)	-0.00214(1)	0.0383	1.8094	-0.0021
C(6)	-1.1785(14)	1.1142(14)	-1.2474(14)	1.1191(21)	0.00727(3)	-1.1741	1.1194	-0.0075
C(7)	2.57923(59)	-1.0306(15)	-2.5815(15)	-1.0364(24)	-0.01085(2)	2.5781	-1.0380	-0.0109
F(8)			2.2918(23)	-0.95995(96)	-0.00073(1)	-2.2980	-0.9561	-0.0007
H(9)			-0.0329(42)	-2.0296(21)	0.01655(1)	0.0297	-2.0582	0.0168
H(10)			-2.1523(24)	1.6531(37)	0.01566(3)	2.1812	1.6630	0.0159
H(11)			-0.0285(43)	2.8630(11)	-0.00192(2)	0.0356	2.8899	-0.0019
H(12)			2.1046(29)	1.6138(37)	-0.0107(3)	-2.1252	1.6295	-0.0108
H(13)			-2.8377(20)	-1.3163(30)	-1.00166(1)	2.8411	-1.3250	-1.0277
H(14)			-2.5198(33)	-1.9214(24)	0.5685(6)	3.3868	-0.4280	0.3828
H(15)			-3.3689(16)	-0.4432(42)	0.3723(4)	2.5153	-1.9461	0.5836

* Imaginary coordinates from the KRA fit. For the determination of bond lengths and angles using EVAL, values from the r_e geometry calculated at the MP2/6-311++G(2d,2p) are taken.

Table 5. Bond lengths (in Å) and bond angles (in degrees) of 3FT deduced from the substitution r_s , semi-experimental r_0 , and equilibrium r_e structures. Standard errors in parentheses are in the unit of the least significant digits.

	r_s	$r_0^{[a]}$	$r_e^{[b]}$
C1–C2	1.3855(35)	1.4041(12)	1.3863
C2–C3	1.3981(32)	1.4041(12)	1.3992
C3–C4	1.3910(58)	1.4041(12)	1.3990
C4–C5	1.3897(14)	1.4041(12)	1.3957
C5–C6	1.4005(14)	1.4041(12)	1.3950
C1–C6	1.3746(62)	1.4041(12)	1.3867
C3–C7	1.5137(32)	1.5068(22)	1.5064
C1–C2–C3	118.42(32)		119.43
C2–C3–C4	119.29(23)		118.68
C3–C4–C5	120.84(11)		120.88
C4–C5–C6	120.45(10)		120.51
C5–C6–C1	117.42(12)		117.86
C6–C1–C2	123.59(25)		122.63
C4–C3–C7	120.86(22)	121.31(22)	120.23

^a All C–H bond lengths and all C–C bond lengths within the phenyl ring are constrained to be the same. The fitted C–H and C–F bond lengths are 1.0503(11) and 1.3482(41) Å, respectively. The fitted F8-C1-C2 angle is 118.47(20) Å.

^b Calculated at the MP2/6-311++G(2d,2p) level of theory.

The r_s structure only represents a partial structure determination since the hydrogen and fluorine related coordinates cannot be determined. Nevertheless, their equilibrium coordinates predicted at the B3LYP/6-311++G(2d,2p) level can be used together with the experimental A_0 , B_0 , and C_0 rotational constants to enable a fit using the program STRFIT [43], yielding the semi-experimental r_0 structure. In the fitting, we allowed the hydrogen and fluorine atom locations to vary, but all C–H bond lengths and all C–C bond lengths within the phenyl ring are constrained to be the same. The atom coordinates, the fitted bond lengths and bond angles of the r_0 structure are also collected in Tables 4 and 5.

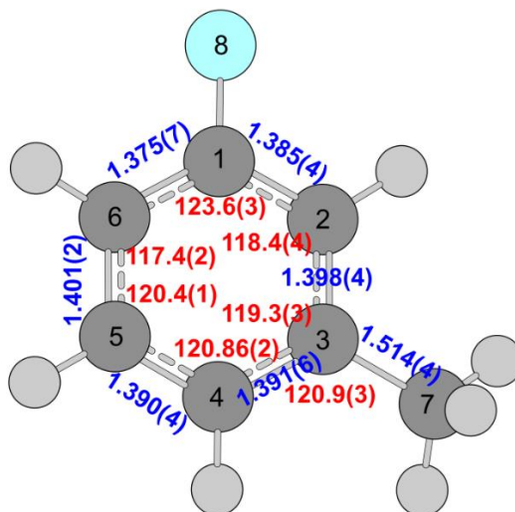


Figure 5. Experimental bond lengths (blue, in Å) and bond angles (red, in degrees) of 3FT obtained from the substitution r_s structure. Standard errors in parentheses are in the unit of the least significant digit.

5. Discussion

5.1. Quality of the fits

Using the modified version of the CAM program *XIAM*, the microwave spectrum of 3FT could be reproduced with a standard deviation of 3.4 kHz, close to measurement accuracy. The deviation obtained with *XIAM_{mod}* was reduced to 1.5 kHz if the program *aixPAM* working in the principle axis system was used.

Involving all three higher order D_{3c} parameters and F_0 is decisive to reduce the deviation in the *XIAM_{mod}* fit, whereas *aixPAM* does not need to float the V_- parameter, but also requires F_0 to be fitted. The correlation between F_0 and V_3 in the *XIAM_{mod}* fit is 1.000, which is frequently observed when the data set only contains transitions from the vibrational ground state. Therefore, F_0 should be fixed to a reasonable value, typically that predicted by quantum chemistry. In several cases, however, fitting F_0 improves significantly the fit, but sometimes it takes values which hardly agree with the moment of inertia of a methyl group [44-48], including the present case of 3FT. The value of 168.669(92) GHz obtained for F_0 is abnormally large compared to that often observed and structurally plausible for a methyl rotor (158-162 GHz) as well as the values indeed predicted by quantum chemistry (see Table 2). Contaminated by absorbing the effects of high order effective

internal rotor parameters that are not explicitly included in the rigid frame-rigid top model, F_0 becomes an effective parameter. With *aixPAM*, the F_0 - V_3 correlation of 0.883 is better. Consequently, the value of 164.302(40) GHz found for F_0 , though still slightly too high, is more reasonable. The F_0 - V_3 correlation also explains different values found for V_3 in the *XIAM_{mod}* and *aixPAM* fits. We note that the errors given in parentheses in the unit of the last significant digits are fitting errors and depend on the model and the set of parameters in use.

The sign of D_K yielded by *XIAM_{mod}* is opposite to the values obtained from *aixPAM* and quantum chemistry. High order effective parameters like D_{c3} severely correlate with geometry parameters, especially the rotational constants but also the centrifugal distortion constants, making the physical meaning of the geometry parameters less clear. For example, we have observed for some molecules with low barriers to methyl internal rotation that the centrifugal distortion constants are contaminated by the internal rotation effects and become themselves effective parameters which no longer solely present the effects of centrifugal distortion [17,49]. The sign of d_2 is also different in the *XIAM_{mod}* and *aixPAM* fits, but for this parameter, calculations at different levels of theory do not agree in sign either.

The internal rotation parameters involved in H_{ird} obtained with *XIAM_{mod}* and *aixPAM* match well. The angle $\angle(i,a)$ is in excellent agreement. The B and C rotational constants deduced from *XIAM_{mod}* and *aixPAM* are similar, but the A rotational constants are not. This has quite often been observed in previous studies where the two approaches (CAM vs PAM) were compared [16,17,50]; all of them suggested the correlation between A and V_K as the main origin.

The standard deviation obtained with *XIAM_{mod}* for the ^{13}C isotopologue fits, except for C(2) and C(7), is slightly higher, but similar to that of the main species fit, probably because the values of high order parameters in H_{ird} are kept fixed at those determined for the main species. Individual adjustments were not possible due to the small number of E species lines available for the ^{13}C isotopologues. Because of the correlation between A and V_K discussed above, we did not perform comparable *aixPAM* fits for the ^{13}C isotopologues, since the rotational constants obtained from those fits would not be reliable for structure determination purposes.

Rudolph and Trinkaus proposed two sets of potentials for 3FT with the V_3/V_6 values of 15.8(1)/8.0(2) cm^{-1} (set I) and 16.9(1)/-5.3(2) cm^{-1} (set II), both of which provided fits with

similar quality [13]. Okuyama et al. later confirmed that set II agrees better with their experimental results obtained in the ground and electronic excited states using fluorescence and dispersed fluorescence spectroscopy [51]. As our data set only contains transitions from the vibrational ground state, V_3 and V_6 cannot reliably be determined simultaneously. However, attempting to fit both terms by $XIAM_{mod}$ simultaneously, values of 17.26(17) and $-1.4(23)$ cm^{-1} (which indeed are in better agreement with set II of Ref. [13]) are obtained at a correlation of -0.999 . Therefore, we decided to fix V_6 to zero, in agreement with the conclusion drawn from the results of quantum chemical calculations (see Section 2.2). Including the $m = 2$ and 3 transitions reported in Ref. [13] to determine V_6 was not successful due to the small number of $m = 2$ (two) and 3 (five) lines and the lack of trust in their assignment.

For 34DFT, both $XIAM_{mod}$ and $aixPAM$ do not require D_{π^2J} , D_{π^2K} , and F_0 to be fitted, but all three D_{c3} (or V_J , V_K , V_-) parameters. We thus fixed F_0 to the value predicted at the MP2/6-311++G(2d,2p) level of theory. The agreement in structural parameters like rotational constants, centrifugal distortion constants, and $\angle(i,a)$ as well as all high order parameters in H_{ird} is much better compared to the 3FT fits. Because F_0 is fixed to the same value, the values of V_3 in the $XIAM_{mod}$ and $aixPAM$ fits match well. The standard deviation of $aixPAM$ is slightly lower compared to that obtained with $XIAM_{mod}$ (2.4 kHz vs 3.5 kHz). The small difference can be traced back to the negligence of some matrix elements in $XIAM$, as discussed in Ref. [16], but also because the parameters V_J , V_K , V_- and D_{c3J} , D_{c3K} , and D_{c3-} refer to different coordinate systems. They are comparable but not equivalent.

5.2. Methyl torsional barrier

The V_3 potential of 17 cm^{-1} found for 3FT is the lowest of all fluoro-substituted toluene derivatives given in Figure 1, followed by 34DFT (32.5 cm^{-1}). All other molecules possess a fluorine atom at a substitution position next to the methyl group, causing steric hindrance of the methyl torsion from which 3FT and 34DFT are free of. As previously discussed [52,53], the more substituents contribute to break the C_{2v} symmetry of toluene, the larger is the V_3 contribution to the potential. In 3FT, the presence of a fluorine atom at the *meta*-position of toluene breaks the C_{2v} symmetry. In the case of 34DFT, two fluorine atoms are attached to the ring and the V_3 value is larger.

Challenging low torsional barrier problems are observed for all *meta*-mono-substituted toluene, such as *m*-chlorotoluene (3.24 cm^{-1}) [54], *m*-methylanisole (55.77 and 36.63 cm^{-1} for the *syn* and the *anti*-conformer, respectively) [16], *m*-toluene nitrile (14.20 cm^{-1}) [55], xylene (4.49 cm^{-1}) [56], *m*-cresole (*syn*: 22.44 cm^{-1} , *anti*: 3.2 cm^{-1}) [57], and *m*-methylbenzaldehyde (*syn*: 35.93 cm^{-1} , *anti*: 4.64 cm^{-1}) [58]. Quite low barriers are also found in 3,4-disubstituted toluenes, such as 34DFT (32.5 cm^{-1}) and 2,4-dimethylanisole (47.65 cm^{-1}) [50].

5.3. Geometry parameters and dipole moments

The B_e rotational constants predicted at both levels of theory match the experimental constants well; also the isotopic dependence is well-reproduced (see Table S6). Though it is not physically meaningful to compare predicted B_e constants with experimental B_0 constants, the MP2 and B3LYP methods in combination with the 6-311++G(2d,2p) basis set offer cost-efficient calculations with sufficient accuracy to be useful in guiding the assignment of vibrational ground state microwave spectra of fluoro-substituted toluenes, as done in many previous investigations [6-11].

The structure of benzene, well-established by X-ray diffraction, possesses a constant C–C–C angle of 120° and a constant C–C distance of 1.39 \AA between six carbon atoms, which is greater than a C–C double bond of 1.35 \AA but shorter than a C–C single bond of about 1.47 \AA . It is known that substitutions on the phenyl ring can affect structural parameters such as bond lengths and bond angles at the substituted positions [59-61]. Comparing the structure of 3FT to the structures of toluene as the prototype [2], 4-fluorotoluene [14], 34DFT [9], and 3,5-difluorotoluene [10] shows that the methyl substitution has little effect on the ring geometry in its surrounding, as long as a fluorine atom is not nearby. The C–C–C angle of the ring at the methyl substituted position remains close to 120° and the C–C bond lengths are approximately 1.39 \AA . On the other hand, the fluorine atom with its $-I$ effect slightly enlarges the C–C–C angle at the fluoro-substituted position. The neighboring C–C–C angle thus becomes smaller. The C–CH₃ bond length varies slightly around 1.5 \AA in all molecules indicated in Figure 1. A more quantitative analysis is not attempted because various fitting approaches as well as sets of fitting parameters were used to treat the effects of methyl internal rotation in different molecules, making the deduced geometry parameters not directly comparable.

The dipole moment components $|\mu_a| = 1.7327(23)$ D and $|\mu_b| = 0.6217(86)$ D determined in the present work agree well with those reported by Rudolph and Trinkaus [13], but are more precise by an order of magnitude. The values of 1.74 and 0.58 D, as well as 1.80 and 0.57 D predicted with the MP2 and B3LYP methods, respectively, agree consistently.

6. Conclusion

The rotational spectra of the main species and all seven ^{13}C isotopologues of 3-fluorotoluene have been studied in a jet expansion, allowing the observation of the lowest-lying torsional states of the vibrational ground state. Each rotational transition splits into two components arising from the internal rotation of the methyl group. The torsional doublings were analyzed with a modified version of the program *XIAM* and the program *aixPAM* to extract the three-fold potential barrier hindering the methyl torsion. The quality of both *XIAM_{mod}* and *aixPAM* fits is very satisfactory. The spectrum of the main species of 3,4-difluorotoluene was refitted, achieving standard deviation close to measurement accuracy. From the spectroscopic constants of the complete set of ^{13}C isotopologues, accurate bond length and bond angles of 3-fluorotoluene could be determined from the r_s and r_0 structure. Stark effect measurements yielded precise dipole moments.

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