Distortion of the Benzene Ring in 1,2,3-Trifluorobenzene; 
A High-Resolution Molecular Beam Fourier Transform Microwave Study

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The results of a molecular beam microwave Fourier transform study of 1,2,3-trifluorobenzene and its monosubstituted $^{13}$C-isotopomers, all observed in natural abundance, are presented. Typically 30 transitions were observed for each isotopomer. Centrifugal perturbation theory up to fourth order was used to analyse the observed rotational spectra. Due to the high sensitivity and resolution of the spectrometer, very accurate vibronic ground state rotational constants can be reported:

\[
\begin{array}{cccccc}
A/\text{MHz} & 2315.26797(10) & 2319.63903(10) & 2298.06711(11) & 2272.17274(13) & 2319.60029(06) \\
B/\text{MHz} & 1757.93383(14) & 1749.22591(17) & 1748.64646(17) & 1757.84791(25) & 1757.83010(08) \\
C/\text{MHz} & 999.16620(09) & 997.15481(08) & 992.96022(09) & 992.96022(09) & 999.93756(10) \\
\end{array}
\]

They are used to derive a partial $r_0$-structure, which shows the distortions of the benzene ring caused by fluorine substitution.

Introduction

We have initiated this study in the context of our continuing interest in the rotational Zeeman effect of substituted aromatic six-membered rings [1, 2, 3, 4]. Accurate structural information is a prerequisite, if one wants to further evaluate the information contained in the measured molecular $g$- and $\zeta$-tensor elements [5].

The rotational spectrum of the most abundant isotopomer of 1,2,3-trifluorobenzene has been carefully reassigned just recently [6]. In the following we report our results for the four monosubstituted $^{13}$C-isotopomers. All spectra were observed in natural abundance by pulsed nozzle molecular beam Fourier transform microwave spectroscopy (MB-FTMW).

Combined with the results for the most abundant isotopomer, a set of effectively ten rotational constants is now available for this presumably planar molecule. If a planar equilibrium geometry with $C_{3v}$-symmetry is assumed, eleven internal coordinates (bond angles and bond distances) are sufficient for a complete description of the molecular structure. In other words, the present experimental data should allow for an almost complete $r_0$-structure [7] determination. Correlations however reduce the number of determinable parameters to seven (see below).

Experimental Details

The sample of 1,2,3-trifluorobenzene (98 % purity) was purchased from Lancaster Synthesis GmbH, Mülheim am Main. It was used after several bulb to bulb distillations without further purification. To observe the spectrum we used our MB-MWFT spectrometer, originally designed for trace gas detection [8]. The spectrometer was improved to include automatic scanning capabilities. The present setup is given in detail in [9, 10]. In Fig. 1 we present a 135 MHz section out of a larger scan.

Trifluorobenzene was deluted (1 %) in argon as carrier gas. The backing pressure was kept below 50 kPa (0.5 atm). Such a comparatively low backing pressure reduces the probability of van der Waals complex formation. The parallel beam arrangement (compare Fig. 2) was used. Due to the large interaction zone in the cavity resonator, this arrangement, which was first introduced by Grabow [11, 12], gives considerably higher sensitivity and resolution than the original Flygare-Balle perpendicular beam arrangement [13].

The spectrometer not only is extremely sensitive but, since high-$J$ transitions can be virtually suppressed,
it is also ideally suited for the analysis and for the assignment of otherwise dense rotational spectra. The reason for the lack of high-J transitions is twofold. First, due to the very effective rotational cooling in the expansion zone immediately downstream of the nozzle, only low-J rotational states are populated at all. Second, since the different M-states within a higher-J rotational transition are polarized at different speeds according to their different dipole matrix elements, M-dephasing during excitation leads to an additional drop in the intensity of high-J free induction decay signals. With increasing J-values the latter effect becomes more and more important, since more M-substates contribute to the corresponding emission signal. To summarize: by the proper choice of the backing pressure (→ cooling) and the duration of the polarizing pulse (→ dephasing) the spectrometer can be set to conditions under which only the lowest rotational transitions are observed. This considerably facilitates the assignment.

Assignment of the \(^{13}\text{C}\)-Spectra

Based on the results of an earlier investigation of 1,2-difluorobenzene, 1,2,4-trifluorobenzene, and 1,2,3,5-tetrafluorobenzene [14, 15], we were able to predict the low-J transition frequencies of the four mono-\(^{13}\text{C}\)-substituted 1,2,3-trifluorobenzenes within better than 1 MHz. All could be observed with sufficient signal to noise ratios. As an example we present a high resolution mode registration of the rotational transition of \(^{13}\text{C}\) in Figure 3. This transition is also included in the broad band scan presented as Figure 1. Spin-rotation coupling splitting caused by the F- and H-nuclei is still below the resolution of the instrument. The observed doublet splitting is caused by the Doppler-effect. (In the parallel beam arrangement the front mirror of the cavity detects a blue-shifted, the rear mirror a red-shifted molecular emission signal and the splitting carries information on the beam velocity. In our case, with argon as carrier gas, the beam velocity was around 560 m/s.)

In Table 1 we list part of the observed transition frequencies. The complete frequency listings are available from the authors. Typically they include about 30 transition frequencies for each isotopomer but for the remaining transitions the rotational quantum numbers are not always the same for all isotopic species.
Fig. 2. General setup of the molecular beam FTMW spectrometer. 1. PC (80386 processor), 2. pulse sequencer (homemade), 3. microwave synthesizer (Systron-Donner 1730B, 2-26.5 GHz), 4. nozzle (General Valve), 5. step motor drive for the movable mirror, 6. Fabry-Perot resonator, 7. power meter (Hewlett Packard 437B), 8. A/D converter for power control (Analog Devices, AD 571), 9. pressure control: baratron (MKS 390), multigas controller (MKS 147C), 10. vacuum system: diffusion pump (Balzers, DIF 250, 3100 l/s), rotary pump (Balzers, DUO 060 A, 68 m³/h), 11. sample preparation and reservoir, 12. transient recorder (TR-AS 102, Dr. Strauss System Elektronik).

Fig. 3. \( J'_{K',K''} - J''_{K',K''} = 2_{20} - 1_{11} \) transition of the \(^{13}\)C₃-isotopomer of 1,2,3-trifluorobenzene. Central frequency of the Doppler doublet: 9113.043 MHz. Doppler splitting: 38 kHz. The given power spectrum was detected using 8 k data points at a sample interval of 10 ns, polarization frequency: 9113.05 MHz.

Derivation of the Molecular Structure

a) Introductory Remarks

The observed effective rotational constants carry information on the molecular geometry. To a high degree of approximation they correspond to vibrational expectation values of the principal moments of inertia [7]:

\[
\begin{align*}
A^{(i)}_{v_1,\ldots,v_{30}} &= \frac{\hbar}{8\pi^2} \cdot \langle v_1,\ldots \rangle \sum m_n \cdot \left( b_{i,n}^2 + c_{i,n}^2 \right) |v_1,\ldots\rangle, \\
B^{(i)}_{v_1,\ldots,v_{30}} &= \frac{\hbar}{8\pi^2} \cdot \langle v_1,\ldots \rangle \sum m_n \cdot \left( c_{i,n}^2 + a_{i,n}^2 \right) |v_1,\ldots\rangle, \\
C^{(i)}_{v_1,\ldots,v_{30}} &= \frac{\hbar}{8\pi^2} \cdot \langle v_1,\ldots \rangle \sum m_n \cdot \left( a_{i,n}^2 + b_{i,n}^2 \right) |v_1,\ldots\rangle.
\end{align*}
\]

\((m_n = \text{atomic mass of the } n-\text{th atom; } a_{i,n}, b_{i,n}, c_{i,n} \text{ its coordinates with respect to the principal inertia})\)
Table 1. Rotational transitions \( J'_{\text{a}} - J''_{\text{a}} \) and experimental frequencies \( \nu_{\text{exp}} / \text{MHz} \) of the \( ^{13}\text{C} \)-isotopomers of 1,2,3-trifluorobenzene. In brackets: \( \nu_{\text{calc}} - \nu_{\text{exp}} \).

<table>
<thead>
<tr>
<th>( J'<em>{\text{a}} - J''</em>{\text{a}} )</th>
<th>( \nu_{\text{exp}} \left( ^{13}\text{C}_2 \right) )</th>
<th>( \nu_{\text{exp}} \left( ^{13}\text{C}_3 \right) )</th>
<th>( \nu_{\text{exp}} \left( ^{13}\text{C}_4 \right) )</th>
<th>( \nu_{\text{exp}} \left( ^{13}\text{C}_5 \right) )</th>
<th>( \nu_{\text{exp}} \left( ^{12}\text{C} \right) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( 2_{10} - 1_{10} )</td>
<td>7944.9666(6)</td>
<td>7956.0679(1)</td>
<td>7887.1582(2)</td>
<td>7807.5417(3)</td>
<td>7958.7344(6)</td>
</tr>
<tr>
<td>( 2_{10} - 1_{10} )</td>
<td>9118.7347(3)</td>
<td>9113.0426(4)</td>
<td>9058.2177(3)</td>
<td>9012.2218(2)</td>
<td>9129.3151(1)</td>
</tr>
</tbody>
</table>

If experimental values for the effective rotational constants were available also for molecules in all singly excited vibrational states, (3) could be solved for the equilibrium values \( A\left( \nu \right), B\left( \nu \right), C\left( \nu \right) \), and these in turn could be used in an iterative non linear least squares fit of the equilibrium internal coordinates. With only vibronic ground state rotational constants available at present, we had to use a different approach. We tried two types of fits of the structural parameters to the observed rotational constants. Both approaches had in common that the rotational constants were treated as if the molecule were rigid and planar (all \( c_{
u,n} \)-coordinates were set to zero), i.e. we treated the rotational constants as if they were

\[ A^{(i)}_{\nu_1 \ldots \nu_{10}} = A^{(i)} - \sum_{s=1}^{10} a^{(i)}_{\nu,s} \left( \nu_s + \frac{1}{2} \right) + \ldots \text{e.t.c.} \quad (3) \]
given by
\[
A^{(i)} = \frac{h}{8\pi^2} \sum m_n \cdot b_{i,n}^2,
\]
\[
B^{(i)} = \frac{h}{8\pi^2} \sum m_n \cdot a_{i,n}^2,
\]
\[
C^{(i)} = \frac{h}{8\pi^2} \sum m_n \cdot (a_{i,n}^2 + b_{i,n}^2).
\]

In the first approach the internal coordinates were directly fitted to the rotational constants. This approach leads to a so-called \( r_0 \)-structure [7]. The second approach, first introduced by the Zürich group [18], involves fits to the differences between observed rotational constants. Specifically, in the second approach, the structural parameters were fitted to the rotational constants of the most abundant isotopomer (the parent molecule) and to the differences between the rotational constants of the \( ^{13}\text{C} \)-isotopomers and the parent molecule, i.e. to
\[
A^{(i)}_{\text{parent}} - A^{(i)}_{\text{isotopomer}},
\]
\[
B^{(i)}_{\text{parent}} - B^{(i)}_{\text{isotopomer}},
\]
\[
C^{(i)}_{\text{parent}} - C^{(i)}_{\text{isotopomer}} \quad (i = 2 \text{ to } 5)
\]

This second approach is assumed to eliminate part of the model errors which are introduced by the neglect of vibrational effects. The reasoning is as follows: Since in the fairly rigid molecule 1,2,3-trifluorobenzene, vibrational averaging is probably very similar in the parent species and in the \( ^{13}\text{C} \)-isotopomers, i.e. since \( \alpha_{\text{parent},i} \approx \alpha_{\text{isotopomer},i} \) (compare (3)), the \( \alpha \)-corrections will partly cancel and the differences in the effective rotational constants will come closer to the differences in the equilibrium values. As a consequence, the second approach may be expected to lead to structural parameters which come closer to the equilibrium values.

### b) Results and Discussion

The fits of the internal coordinates to the observed rotational constants were carried out with Volker Tümpkes program MWSTR1. For the underlying theory compare [18]. If \( C_{2v} \)-symmetry is assumed, the structure of 1,2,3-trifluorobenzene may be described by 11 internal coordinates. Because of correlations, only 7 internal coordinates could be fitted simultaneously, i.e. we had to impose 4 constraints. They were chosen as follows:

1. \( r_{C_4-H_4} = 1.082 \) Å
2. \( r_{C_5-H_5} = 1.082 \) Å
3. \( r_{C_2-F_2} = r_{C_1-F_1} = 0.005 \) Å
4. \( \angle H_{(4/6)}C_{(4/6)}C_{(5/5)} = 121.96^\circ \)

The third constraint, i.e. the 0.005 Å difference between the CF-bonds, was imposed in view of the earlier results obtained for related compounds by Jochims [14] and Stiefvater [19]. It is also in accordance with the results of an ab initio SCF/HF/6-311G equilibrium structure calculation with the GAUSIAN 92 program package [20]. Such a calculation predicts a 0.005 Å difference for the equilibrium values. The fourth constraint was imposed in view of Stiefvaters results obtained for 1,2-difluorobenzene and 1,3-difluorobenzene. We simply assumed that the \( H_{(4/6)}C_{(4/6)}C_{(5/5)} \) bond angle may be approximated as the average of the corresponding values observed in ortho- and meta-difluorobenzene.

In Table 3 and Fig. 4 we present the results of our fits. The differences between the values in the first
Fig. 4. Partial restructure of 1,2,3-trifluorobenzene derived by the fit of 7 structural parameters to the rotational constants of the parent molecule and to the differences between the rotational constants of the parent molecule and the $^{13}$C-isotopomers (* assumed; ** $r_{C(2)-F(2)} = r_{C(1/3)-F(1/3)} - 0.005 \text{ Å}$).

The observed overall pattern of the benzene-ring distortions caused by the fluorine substituents looks familiar. As compared to benzene, the annular bond angles at the $C_{(1)}$ and $C_{(3)}$ positions are enlarged, while the $C_{(1)}C_{(2)}^{-}$ and $C_{(2)}C_{(3)}^{-}$ bonds are slightly shortened. Nygaard et al. [21, 22], in their pioneering work on fluorobenzene, have rationalized both distortions within a simple hybridization model. Within this model, fluorine substitution slightly changes the original local $sp^2$ hybridization at the $C$ atoms in the following way. The $sp$-hybrid pointing towards the electronegative $F$ gains in $p$-character while, as a consequence (orthogonality of the local $sp$-hybrid orbitals), the orbitals pointing towards the neighbouring $C$ atoms gain in $s$-character and become more $sp$-like i.e. linear.

In a subsequent study of the deuterated 1,2,3-trifluorobenzenes we hope to reduce the remaining ambiguities introduced by the constraints which had to be imposed during the fits of the structural parameters. To provide an impression of these ambiguities we give an example. If, for instance, one would assume the CF bond distances to be equal rather than to differ by 0.005 Å, Fit 2 would lead to

$$r_{C_{(1/2)}C_{(2/3)}} = 1.3830 \text{ Å},$$
$$r_{C_{(1/3)}C_{(6/4)}} = 1.3844 \text{ Å},$$
$$r_{C_{(4/5)}C_{(5/6)}} = 1.3954 \text{ Å}$$

for the CC bond distances, and to

$$\angle C_{(1)}C_{(2)}C_{(3)} = 118.850^\circ,$$
$$\angle C_{(2/3)}C_{(3/1)}C_{(4/6)} = 121.500^\circ,$$
$$\angle C_{(3/1)}C_{(4/6)}C_{(5/5)} = 118.692^\circ,$$
$$\angle C_{(4/5)}C_{(5/6)} = 120.767^\circ$$

for the annular bond angles, i.e. the optimized annular bond angles would be changed by 0.1° at most. In the study of the deuterated species we also intend to present the results of high level ab initio calculations which are carried out in parallel to the spectroscopic investigation. While the HF-calculation with the 6-311G - basis produced a planar equilibrium geometry, a higher level partial geometry optimization including electron correlation, in which the geometry of the aromatic ring was fixed to the values given in the second column of Table 3, indicates that the equilibrium configuration of 1,2,3-trifluorobenzene might slightly deviate from planarity [23].

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[23] Ralph Jaquet, University of Siegen, private communication.