Lattice dynamics and methyl rotational excitations of 2-butyne

O. Kirstein and M. Prager
Institut für Festkörperforschung, Forschungszentrum Jülich, D-52425 Jülich, Germany

M. R. Johnson
Institut Laue-Langevin, Avenue des Martyrs, BP 156, 38042 Grenoble Cedex 9, France

S. F. Parker
ISIS facility, Rutherford-Appleton-Laboratory Chilton, Didcot OX11 0QX, United Kingdom

(Received 8 February 2002; accepted 23 April 2002)

On the basis of the recently determined low temperature crystal structure the lattice dynamics of 2-butyne and a single particle methyl rotational potential are calculated using pair potential parameters given by Williams in 1974 within the model of semirigid molecules. In the regime of lattice modes the existence of four methyl librational bands with significant dispersion can explain the measured density of states. The single particle librational energy obtained for the rotational potential from the Schrödinger equation of the methyl rotor coincides well with the mean librational band energy. The calculated single particle tunneling frequency is only 17% lower than observed in experiment. Similarly well the activation energy is reproduced. Although the pair potential parameters developed for aromatic compounds need some scaling they are useful for materials with triple bond carbons. © 2002 American Institute of Physics. [DOI: 10.1063/1.1485729]

I. INTRODUCTION

The theory developed originally to describe rotational tunneling of molecular crystals is a mean field or single particle model (SPM).\(^1\) Despite the more recent detection of systems with coupling and multidimensional tunneling\(^2,3\) the prominent number of molecular crystals can be well described within the original SPM.\(^4\) A data analysis going beyond a phenomenological use of this model has to explain the rotational potential on the basis of the crystal structure and fundamental intermolecular interactions.\(^5,6\) Often intermolecular interactions are modeled as transferable intermolecular pair interactions. There is an increasing number of systems analyzed this way. Their number is mainly limited by the knowledge of a precise low temperature crystal structures of systems whose tunnel splitting is known from high resolution neutron spectroscopy. Another difficulty is due to the reliability of the atom–atom potential ansatz for the intermolecular interaction. The more complex a compound is the more difficult is it to get reliable pair potential parameters from literature. For this second reason we concentrated at the beginning our work on simple systems, mainly pure hydrocarbons as toluene\(^5\) and \(~\)~\)-xylene\(^7\) which belong to a class of materials most intensively studied by the concept of pair potentials.\(^8,9\) In the present paper we want to extend the successful earlier description to an only slightly different material. 2-butyne—also known as dimethylacetylene (DMA)—is again a pure hydrocarbon but with a triple bond between the two central carbon atoms. It is the simplest compound containing at the same time methyl groups but in contrast to other hydrocarbons there are no pair potentials (PP) available from literature for this material. The question arises whether pair potentials of DMA differ from those of aliphatic or aromatic hydrocarbons.

2-butyne is one of the first systems where rotational tunneling was observed by high resolution neutron spectroscopy.\(^10,11\) It was difficult to relate this tunnel splitting to the librational modes measured in a forthcoming experiment.\(^12\) In this paper the missing knowledge of the crystal structure did not allow a mode analysis and the description had finally to be based on an ad hoc model allowing for dispersion of the librational modes.

Thus it became interesting to study this material in a more comprehensive way after the low temperature crystal structure became known,\(^13\) see Fig. 1. The technique of choice is to combine the structural information with transferable intermolecular pair potentials. The basic questions are threefold:

1. Is the assumption of a semirigid molecule justified?
2. How does the phenomenological explanation of the density of states of Ref. 12 compare with a mode analysis based on a lattice dynamics calculation?
3. Do the pair potential parameters need to be modified compared to those used for aromatic hydrocarbons\(^14\) due to triple bond between the center carbons?

II. EXPERIMENTAL RESULTS

A. Excitations

Inelastic neutron scattering was used to study excitations in the regime of phonons and internal excitations. Due to its wide accessible energy range up to 500 meV and its good energy resolution \(\delta E/E \sim 0.02\) in the whole spectral range the inverse time-of-flight spectrometer TFXA of the ISIS spallation source at the Rutherford-Appleton Laboratories, UK, was used.\(^15\) At this crystal analyzer instrument energy transfer and momentum transfer are correlated. The INS spectrum is available from the INS database\(^16\) at www.isis.rl.ac.uk/insdatabase.
A flat protonated sample of volume $30 \times 50 \times 1 \text{ mm}^3$ was cooled down to $T \leq 15 \text{ K}$ and measured for 24 h. This corresponds to an integrated proton dose of 2000 $\mu$Ah. All detectors are summed up in a single spectrum shown in Fig. 2. The spectrum decomposes into two regimes. Above about 20 meV internal molecular vibrations, higher harmonics and combination bands are visible. Below 20 meV we have the regime of phonons. This part of the spectrum was measured already earlier at the direct time-of-flight spectrometer IN4 of the ILL, France. Small differences in peak positions are likely due to the different ways the used instruments integrate over the dispersion curves.

**B. Quasielastic scattering and activation energy**

The quasielastic spectra of DMA had been obtained using the FOCUS spectrometer at the Swiss neutron source SINQ. The wavelength of the incoming neutrons were chosen to be $\lambda = 5.8 \text{ Å}$ with a corresponding elastic energy resolution $\delta E = 50 \mu\text{eV}$. The energy transfer range of interest was between $-2$ and 1 meV. The experimental data were transformed into $S(Q, \omega)$ using standard procedures. Figure 3 shows the broadening of the quasielastic line for two different temperatures. Since all methyl groups are crystallographical equivalent the resulting QNS spectra are described by a single Lorentzian.

The model used to analyze quasielastic spectra is a simple hopping model. Since the potential forces the molecule into the equilibrium position the only possibility for a hydrogen to move is to jump over the barrier into an equivalent position. The scattering function

$$S(Q, \omega) = A_0(Q) \delta(\omega) + A_1(Q) \frac{1}{\pi} \frac{\frac{1}{2} \nu}{(\frac{1}{2} \nu)^2 + \omega}$$

with the prefactors $A_0(Q) = \frac{1}{2}(1+2f_0(Qd))$ and $A_1(Q) = \frac{1}{2}(1-f_0(Qd))$, where $Q$ is the value of the scattering vector and $d$ the jump length, i.e., the proton–proton distance of 1.78 Å inside the CH$_3$ group. The width of the Lorentzian in Eq. (1) is $\Gamma = (3/2) \nu$. Since jumps are thermally activated processes the jump frequency and thus the linewidth of the Lorentzian depends on the temperature and follows an Arrhenius behavior,

$$\Gamma(T) = \Gamma_0 \cdot \exp \left( -\frac{E_a}{kT} \right) = \Gamma_0 \cdot \exp \left( -0.0116 \cdot E_a(\text{meV}) \cdot \Theta[\text{K}^{-1}] \right)$$

with $\Theta = 10^3/T$.

The hopping model allows to derive the activation energy $E_a$, which yields the energy difference between the ground state and the maximum of the methyl rotational potential.

In order to get the activation energy eight quasielastic spectra were collected in the temperature range of $70 \text{ K} \leq T \leq 150 \text{ K}$ (corresponding to $6.7 \text{ K}^{-1} \leq \Theta \leq 14.3 \text{ K}^{-1}$) and analyzed with respect to the temperature dependent FWHM of the Lorentzians. A least squares fit to the data yields $E_a$.
converged to a geometry that was almost D formed, such that one of the C–H bonds is shorter than the point of inversion, however, the molecule is slightly de- antiphase methyl torsion was close to zero. The crystallographic geometry as the initial input were exam- ined. All gave stable structures, although in each case, the semirigidity can be valid. The soft internal modes are now included into the external DOF. Ab initio calculations give a quantitative measure of the internal DOF which have to be included by the lattice dynamical calculation.

A. Ab initio calculations for the isolated molecule

In the gas phase, there is free rotation of the methyl groups, thus the D_{3h} (methyl groups eclipsed) and D_{3d} (methyl groups staggered) structures are equivalent. In the low temperature C2/m phase the center of the molecule lies on a point of inversion, however, the molecule is slightly deformed, such that one of the C–H bonds is shorter than the other two and the symmetry is reduced to C_{2h}.

The vibrational modes of molecules containing up to a few tens of first row atoms can be calculated on a PC using ab initio methods. In the present case, density functional theory (DFT) as implemented in the GAUSSIAN 98 (Ref. 17) package was used with the B3LYP functional and 6-31G, 6-311+G, and 6-311+G(d,p) basis sets. All the methods yielded similar results, with the largest basis set providing the best results.

The D_{3h}, D_{3d} structures and a C_{2h} structure that used the crystallographic geometry as the initial input were examined. All gave stable structures, although in each case, the antiphase methyl torsion was close to zero. The C_{2h} structure converged to a geometry that was almost D_{3d}, this is also shown by the very small (<0.1 meV) splitting of the degenerate modes. This result strongly suggests that the solid state structure is crystallographically imposed.

The observed intensity of an INS band is given by

\[ S(Q,n\omega_i) \propto \frac{(Q \cdot U_i)^{2n}}{n!} \exp\left(-\frac{Q^2 \cdot U_{tot}^2}{\sigma^2}\right) \]

where \( \omega_i \) is the ith mode at frequency \( \omega \), \( n = 1 \) for a fundamental, \( 2 \) for a first overtone or binary combination, \( 3 \) for a second overtone or ternary combination etc., \( Q \) is the momentum transfer defined earlier, \( U_i \) is the root mean square displacement of the atoms in the mode, \( U_{tot} \) is the total root mean square displacement of all the atoms in the modes, both internal and external, \( \sigma \) is the inelastic scattering cross section of the atom. The total scattering cross sections of hydrogen and carbon are 82.0 and 5.6 barns (1 barn = 1 \( \times 10^{-28} \) m\(^2\)), respectively and since the intensity is dependent on the amplitude of vibration, which is larger for light atoms, the spectra will be dominated by motions that involve displacement of hydrogen. The exponential term in Eq. (2) is a Debye–Waller factor. Equation (2) is based on an isolated molecule approximation, thus dispersion is not included. The narrow linewidths of the modes above 25 meV demonstrate that this is a reasonable assumption in the present case.

The output of GAUSSIAN 98 includes the atomic displacements for each mode, which can be used to visualize the mode and also to generate the INS spectrum. The program a-CLIMAX (Ref. 20) was used for this purpose. After minor adjustment of the frequencies to match the observed values and inclusion of events involving up to three quanta the result shown in Fig. 5 is obtained. The agreement between observed and calculated is excellent confirming the assignments given in Table I.

The width of the skeletal bending mode at 39.7 meV is significantly larger than the resolution function which results in an apparent mismatch for this band, although the integrated intensity is approximately correct. This is probably due to factor group splitting since there are two molecules in the unit cell. This frequency is significantly above the phonon energies so justifies the treatment of the molecule as a rigid body in the lattice dynamical calculations described in the following section.
TABLE I. Vibrational modes of DMA as calculated ab initio using the GAUSSIAN 98 program are compared to the excitation energies measured by neutron spectroscopy. Modes are assigned to each excitation energy.

<table>
<thead>
<tr>
<th>Gaussian energies (MeV)</th>
<th>Multiplicity</th>
<th>Mode character</th>
<th>Observed (TFXA energies (meV))</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1</td>
<td>methyl torsion</td>
<td>6.9, 9.2, 12.6, 17.3</td>
</tr>
<tr>
<td>26.4</td>
<td>2</td>
<td>backbone bend</td>
<td>28.5</td>
</tr>
<tr>
<td>47.1</td>
<td>2</td>
<td>backbone twist</td>
<td>47.4</td>
</tr>
<tr>
<td>87.5</td>
<td>1</td>
<td>backbone sym. stretch</td>
<td>87</td>
</tr>
<tr>
<td>132</td>
<td>4</td>
<td>methyl rock</td>
<td>129</td>
</tr>
<tr>
<td>141</td>
<td>1</td>
<td>backbone asym. stretch</td>
<td>172</td>
</tr>
<tr>
<td>176</td>
<td>2</td>
<td>methyl sym. breath</td>
<td>181</td>
</tr>
<tr>
<td>186</td>
<td>4</td>
<td>methyl deformation</td>
<td>290</td>
</tr>
<tr>
<td>378–386</td>
<td>6</td>
<td>proton stretch</td>
<td></td>
</tr>
</tbody>
</table>

The strong mode at 129 meV is the methyl rocking mode and both the INS spectrum and the ab initio results show that the four (two doubly degenerate modes) are coincident. The low temperature Raman spectrum\(^{21}\) shows that the asymmetric methyl bending modes give rise to a broad, ~5 meV, band at 180 meV. The ab initio results suggests that these bands are very close in energy which is why the feature is apparently underestimated. Inspection shows that there is missing intensity in the calculated spectrum at 175 meV, suggesting that one of the modes is calculated too high in energy. Shifting one of the modes to lower energy would also reduce the overestimate of the phonon wings that are responsible for the intensity at ~185 meV.

The higher energy modes are poorly resolved from the phonon wings. This is due to the large Debye–Waller factor that results from a relatively light molecule and the large displacements in the coupled torsional/librational modes. Since the C\(=\)C stretch has little proton motion, it is intrinsically weak in the INS spectrum and is not observed. At large energy transfers, the spectral detail is obliterated by molecular recoil. There is nearly no internal torsional restoring force. The next lowest eigenenergies around 26 meV belongs to a bend of the molecule backbone in two different but equivalent planes. Thus the lowest internal vibrational mode energy is (see below) above the largest phonon energies. Thus we describe the molecule in the following lattice dynamical calculations as completely rigid beside of the internal torsion.

### B. Lattice dynamics of the crystal

Phonon dispersion curves are calculated by the program DYNCAL which was successfully used for similar applications earlier.\(^{5,7,22}\) With DYNCAL phonon energies and eigenvectors are obtained by diagonalizing the dynamical matrix. DMA contains two molecules in the unit cell with one soft internal degree of freedom each, the out-of-phase methyl librations. For otherwise rigid molecules this yields 14 phonon branches. The force constants are derived in harmonic approximation as second derivatives of atom–atom pair potentials. The intermolecular interactions were modeled in the following by a Born–Meyer potential:

\begin{equation}
V_{ij}(r) = k_{ij} \left( \frac{A_{ij}}{(f \cdot r_{ij})^n} + B_{ij} \exp(\alpha_{ij} \cdot (f \cdot r_{ij})) \right),
\end{equation}

where \(ij\) stands for the respective atom-pair (H–H, H–C, and C–C), \(AB\), and \(\alpha\) are pair potential parameters given by Williams\(^{14}\) (referred as Williams’74) and \(k\) and \(f\) are scaling parameters. These parameters allow one to adjust the potential which may be necessary since Williams derived the potential parameters for the class of aromatic molecules not containing carbon–carbon triple bonds. It is obvious that \(V_{ij}(r)\) represents the classical Born–Meyer potential for \(k_{ij} = f = 1\). The values used for the lattice dynamical calculations are given in Table II. They follow the condition of systematic transferability \(k_{ij} = \sqrt{k_{ii} \cdot k_{jj}}\). The electrostatic interaction had been taken into account by an additional potential term

\begin{equation}
V_{ij}^{el}(r) = \frac{n_i \cdot n_j \cdot q_0^2}{r_{ij}}.
\end{equation}

Here \(n_i\) and \(n_j\) are the charges in units of the elementary charge \(q_0\). Values for the partial charges are obtained from the ab initio calculations for the isolated molecule, see above and Fig. 6.

For technical reasons the lattice dynamical program requires the pair potentials as a power series.\(^{23}\) The use of four terms of the series allowed a very good description for interatomic distances above 2.2 (2.0) Å for C–C (H–H, H–H) interactions.

Beside the pair potentials the lattice dynamical program requires the crystal structure to calculate the interatomic distances. The recently measured low temperature crystal structure\(^{13}\) is used.

The molecule is modelled as a linear semirigid one used with negligible internal torsional restoring force. While the ab initio calculation supports this description the observed deformation of the molecule—the carbon-backbone is bent in the crystal by ±1.5°—means that there is a coupling between external and internal modes. An analysis shows, however, that the overall behavior in the regime of lattice modes is unchanged if the molecular bend is frozen or not.

The program allows one to apply an internal restoring force to the methyl libration. This manipulation influences almost only the two librational modes with antiphase rotation of the two methyl groups of the same molecule. The in-phase modes are not shifted in energy under this action. A weak internal methyl–methyl coupling of 0.001 mdyn/Å was chosen. Under the given assumptions the librational modes show energies of 4.2, 11.4, 20.0, 22.8 meV at the Γ-point.

### TABLE II. Williams’74 and scaling parameters used to calculate the lattice dynamics of DMA.

<table>
<thead>
<tr>
<th>Pair</th>
<th>(A_{ij}) (kcal/mol)</th>
<th>(B_{ij}) (10(^{-3}) kcal/mol)</th>
<th>(\alpha_{ij})</th>
<th>(k_{ij})</th>
<th>(f)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H–H</td>
<td>40.1</td>
<td>2.876</td>
<td>3.74</td>
<td>1.012</td>
<td>for all</td>
</tr>
<tr>
<td>H–C</td>
<td>134.2</td>
<td>14.310</td>
<td>3.67</td>
<td>0.700</td>
<td>types</td>
</tr>
<tr>
<td>C–C</td>
<td>449.0</td>
<td>71.430</td>
<td>3.60</td>
<td>0.484</td>
<td>1.016</td>
</tr>
</tbody>
</table>

\(^{6}\)Reference 6.
There is huge dispersion along the [0,1,0] direction, little along [1,0,0]. In both these directions the dispersion curves looks as if the unit cell would be a factor 2, too large. In [0,0,1] direction the phonon branches flatten at the zone border yielding additional van Hove anomalies. Figure 7 shows the dispersion curve along the [001]-direction. The four methyl groups in the unit cell require four modes with a librational character corresponding to an in-phase rotation of methyl groups inside a molecule and an in-phase rotation of the two molecules in the unit cell (in—in) and correspondingly in—out, out—in, and out—out. Figure 8 shows a view of the eigenvectors of the out—out-mode. Numerical results are summarized in Table III. They yield already a qualitative explanation of the multiline phonon spectrum of Fig. 2.

The lattice dynamical program contains all ingredients to calculate the neutron scattering function $S(Q,\omega)$ in the meV regime—or equivalently the DOS—as a weighted average of all excitations,

$$S(Q,\omega) = Q^2 \sum_i \sum_{j} \sigma_i \sum_{l} \langle u_{ij}^2 \rangle \cdot DW(u) \cdot \delta(\omega - \omega_j(q_l)).$$

The summation runs again over all displacements $u$ of 2·10 atoms $i$, the 14 modes $j$ and a homogeneous grid of momentum transfers $q_l$ of the Brillouin zone and includes the Debye–Waller factor $DW(u)$. The only difference to the spectrum of the internal modes, Eq. (2), consists of an additional summation $l$ over the Brillouin zone required by phonon dispersion. Methyl librations are strongly anharmonic and show the largest mean square displacements. Further they invoke the proton with its dominant scattering cross section $\sigma$, and causing the strongest peaks on the DOS. The rotational potential has exactly threefold symmetry since the methyl group relaxes into identical equilibrium orientations after each $2\pi/3$ rotation. If one would rotate the CH$_3$ group as determined crystallographically with all the positional error bars as a rigid unit the proton sites would coincide only after $2\pi$ rotation. Thus when calculating $V(\phi)$ we force the methyl protons to move on the common circle of

$$V(\phi) = V_0 + \sum_{n=1}^{2} \frac{V_n}{2} (1 - \cos(3n \phi + \varphi_n)).$$

The rotational potential has exactly threefold symmetry since the methyl group relaxes into identical equilibrium orientations after each $2\pi/3$ rotation. If one would rotate the CH$_3$ group as determined crystallographically with all the positional error bars as a rigid unit the proton sites would coincide only after $2\pi$ rotation. Thus when calculating $V(\phi)$ we force the methyl protons to move on the common circle of
circumference between equilibrium positions of the two neighboring methyl protons. This procedure makes the protons coincide exactly after 2\(\pi/3\). The resulting potential is shown in Fig. 10 for 0° ≤ \(\varphi\) ≤ 120°. The main contribution to \(V(\varphi)\) stems from atoms within a radius of about 6 Å. Two features are visible in Fig. 10. At first, the potential minimum coincides well with the crystallographic orientation of the methyl group as required for a stable system. Secondly, the potential is exactly symmetric to the maximum. This is due to the presence of the \(a-c\) mirror plane intersecting the methyl group in the \(C_{2v}\) space group. The potential has a minimum flatter than harmonic, a somewhat narrower large maximum indicating a contribution with sixfold symmetry and coinciding maxima of the two Fourier terms.

The potential determines the rotational excitations of the methyl group—tunneling, librations etc.—through the single particle Schrödinger equation of the one-dimensional three-fold rotor,

\[
H = -B \frac{\partial^2}{\partial \varphi^2} + V(\varphi).
\]

\(B = 0.655\) meV is the rotational constant of the rotating methyl group about the rotation axis. The broadened minimum of the potential leads to lowering of the librational modes compared to a pure \(\cos(3\varphi)\) potential. The calculated librational energy of \(E_{01} = 14.16\) meV has no direct counterpart in the observed DOS but the calculated value of \(E_{01}\) can be interpreted as an average of the dispersive librational bands. The value is slightly lower than the experimental value of 15 meV derived by Kollmar and Alefeld\(^{11}\) but still in good agreement. The tunnel frequency of \(\hbar \omega_t = 1.45\) \(\mu\)eV coincides within 17% with the observed tunnel splitting of 1.74 \(\mu\)eV.\(^{10}\) The derived rotational potential is further confirmed by the quasielastic scattering experiment. The measured activation energy \(E_a\) differs from the calculated one by only 6%. Table IV shows the librational and tunnel energies together with the extracted potential parameters. The parameters of the fitted rotational potential are shown in the lowest line of the table.

An artificial increase of the size of the unit cell and a restriction to one molecule per unit cell allows one to simulate the case of the isolated molecule with the lattice dynamical program. In agreement with the GAUSSIAN 98 result the rotational potential is completely removed under these conditions.

**TABLE III.** Calculated libron energies according to Eq. (3) and Table II. The mode characters are described by the phase relation within one molecule and between the two molecules in the unit cell. For example “in–in” means, that all methyl groups rotate in phase.

<table>
<thead>
<tr>
<th>(q)-point</th>
<th>Energy (meV)</th>
<th>Character</th>
</tr>
</thead>
<tbody>
<tr>
<td>(0,0,0)</td>
<td>4.2</td>
<td>in–in</td>
</tr>
<tr>
<td></td>
<td>11.4</td>
<td>out–out</td>
</tr>
<tr>
<td></td>
<td>20.0</td>
<td>out–out</td>
</tr>
<tr>
<td></td>
<td>22.8</td>
<td>in–in</td>
</tr>
<tr>
<td></td>
<td>4.9</td>
<td>in–in</td>
</tr>
<tr>
<td></td>
<td>8.7</td>
<td>out–in</td>
</tr>
<tr>
<td></td>
<td>18.0</td>
<td>in–in</td>
</tr>
<tr>
<td>(0,0,0.5)</td>
<td>18.7</td>
<td>out–out</td>
</tr>
</tbody>
</table>

**TABLE IV.** Comparison between observed energies and calculated ones (tunneling, libration, activation, and potential). Experimental values are boldfaced, calculated ones are italic.

<table>
<thead>
<tr>
<th>(E_{01}) ((\mu)eV)</th>
<th>(V_3) (meV)</th>
<th>(V_6) (meV)</th>
<th>(E_{a}) (meV)</th>
<th>Origin</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.74</td>
<td>45.0</td>
<td></td>
<td></td>
<td>Alefeld and Kollmar(^a)</td>
</tr>
<tr>
<td>1.74</td>
<td>15.2</td>
<td>43.3</td>
<td>3.5</td>
<td>Kollmar and Alefeld(^b)</td>
</tr>
<tr>
<td>6–17.3</td>
<td>14.16</td>
<td>50.5</td>
<td>-3.9</td>
<td>Tilli and Alefeld(^c)</td>
</tr>
<tr>
<td>1.45</td>
<td>33.6</td>
<td>35.2</td>
<td></td>
<td>This work, calculation</td>
</tr>
</tbody>
</table>

\(^a\)Reference 10.  
\(^b\)Reference 11.  
\(^c\)Reference 12.
conditions and the methyl group of the isolated molecule in the lattice dynamical model represents a perfect free rotor.

The case of an isolated dimer can be studied in a similar way. Due to reduced symmetry there are two types of methyl groups in this case. The interesting one is the one with the short distance to the triple bond carbon atoms (the other is again close to a free rotor). Here the potential has reached already 40% of its final strength in the crystal but its minimum is out of phase by ~60° from the crystallographic equilibrium orientation. Only the phase-shifted superposition of interactions with all neighbors leads to consistency. The relatively weak rotational potential results from such compensation effects. An analysis of interatomic distances or the view along the molecular symmetry axis shows that the triple bond carbon belongs to the atoms nearest to the CH₃ group. Thus they give an important contribution to the rotational potential and tunneling probes especially among all others the hydrogen–carbon(3) atom–atom potential.

IV. CONCLUSION

For a long time it was impossible to merge rotational tunneling and librational peaks of the density of states within the single particle model. Based on the measured crystallographical structure the idea that rotor–rotor-coupling causes this discrepancy suggested by Tilli could be confirmed and elaborated. This shows that it may be misleading to identify a peak in the density of states with a single particle methyl librational excitation as it is done often. The analysis of the quasielastic spectra based on classical jump rotation yields the correct barrier height (activation energy $E_a$). Thus it may be better to base an analysis on tunneling and quasielastic scattering instead of tunneling and misinterpretable raw data of the density of states.

The suspicion that DMA requires different pair potentials due to its triple bound carbon atoms was confirmed. Only a modified set of the Williams’74 pair potential parameters used reproduce successfully not only the density of states but also the rotational tunneling, mean librational energy and activation energy of the respective methyl group. Since the triple bound carbons belong to the nearest neighbors of the methyl protons this interaction is really tested with DMA. From this point the Williams parameters seem to be a good starting point to describe hydrocarbon systems which are not single or $\pi$ bonded.

ACKNOWLEDGMENTS

The authors thank J. Tomkinson, RAL, for a first analysis of the TFXA data and M. Monkenbusch, FZJ, and W. Griessl, University of Bayreuth, for the supply of and help with the lattice dynamics program.

15 http://www.isis.rl.ac.uk/molecular/pectroscopy/tfxa/index.htm