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Structure of Phase III and Polymorphism of (CH$_3$)$_2$CB

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ABSTRACT: The polymorphism of 2-bromo-2-methyl-propane ((CH$_3$)$_2$CB) has been investigated by both thermal and X-ray powder diffraction experiments. The long time unknown structure of the low-temperature phase III has been determined by X-ray powder diffraction as orthorhombic Pmmn2$_1$, with lattice parameters $a = 19.1350(4)$ Å, $b = 5.7378(1)$ Å, and $c = 10.9626(2)$ Å at 180 K, $Z = 8$ and with an asymmetric unit containing one molecule in a general position and two additional molecules placed in a mirror ($Z' = 1 + 1/2 + 1/2$). The phase transitions between the different phases have been characterized at normal pressure as well as at high pressures (up to 300 MPa). From the $p$ $T$ slopes of the two-phase coexistence lines and density measurements as a function of pressure, the volume variations at the transition points have been obtained and compared with those obtained by means of X-ray powder diffraction at normal pressure. The packing as well as the anisotropy of the intermolecular interactions have been analyzed by means of the thermal-expansion tensor and compared with tert-butyl ((CH$_3$)$_3$CX, X = Cl, CN) related compounds.

1. Introduction

For molecular crystals, the transition from a complete orientational and translational disorder (isotropic liquid phase) toward complete order may be achieved in several stages in such a way that each phase transition involves a stepwise reduction of the disorder. Depending on whether orientational or translational orders appear when cooling the isotropic liquid, one is dealing with liquid crystals or plastic crystals, respectively. While the former are generally found in isotropic liquid, one is dealing with liquid crystals or plastic crystals, respectively.$^{1,2}$ While the former are generally found in molecular materials formed by elongated molecules, plastic crystals are formed by substances whose molecular shapes are symmetrical about one or more axes and, particularly, those where the molecular shape is somewhat globular. Well-known examples of such molecular materials representative of plastic crystals (i.e., displaying orientationally disordered, OD, phases) are the tert-butyl compounds, (CH$_3$)$_3$CX, X = Cl, Br, NO$_2$, CN, SH, COOH ..., because they have molecular shapes sufficiently close to spherical (due to the comparable size of the X groups to the methyl group) to enable orientational disorder in the solid state.$^{1,6}$ Such compounds display a rich polymorphism that has stimulated a large number of experimental investigations as well as theoretical studies in order to rationalize the thermodynamic properties and molecular dynamics of the various solid phases.$^{3-23}$ Polymorphism may be defined as the ability of a compound to exist in different crystalline forms in which the molecules have different arrangements (packing) in the crystal lattice.$^{24}$ Although the space group of the crystal lattice should be equal to or higher than that of the molecular point-group symmetry, nature overcomes such a restriction by forcing molecular reorientations along different axes, and thus making an inseparable link between crystal structure and dynamics which increases the complexity of finding the crystal assemblies. But even for fully ordered systems at the present state of the art, it is not possible to predict crystals structures reliably.$^{25-27}$ Examples of such a situation concern tert-butyl compounds (CH$_3$)$_3$CCl (hereinafter referred to as tBCl, IUPAC name: 2-chloro-2-methyl-propane) and (CH$_3$)$_3$CBr (hereinafter referred to as tBBr, IUPAC name: 2-bromo-2-methyl-propane) for which the general van der Waals close packing should dominate the observed packing of the motifs. Although it might be assumed that owing to the large molecular symmetry similarities (C$_3$ symmetry) of these two tert-butyl compounds they should form isostructural phases (and thus, a close polymorphism sequence), it was early shown that this circumstance is highly unlikely.$^{3,8,16,17}$ Under their equilibrium vapor pressures, both compounds display three solid phases (I, II, and III in order of decreasing temperature)$^{3,5,11,13,17,19,22,23}$ On cooling, the liquid phase of both compounds freeze into an orientationally disordered fcc (face-centered cubic) structure (phase I) with $Z = 4$ (see Table 1 for temperatures). Upon further cooling, this phase transforms to a tetragonal ($P4_{1}mmm$) phase for tBCl$^{25}$ and to a rhombohedral phase for tBBr.$^{28}$ Uniaxial molecular reorientations (3-fold rotations) about the C-X axis ($C_{3}$-axis or dipole axis of the molecule) and fluctuations of the dipole axis are present in both cases.$^{3,8,11,13,17}$ Nevertheless, the fluctuations of the dipole axis are just a few degrees for tBCl while those for the rhombohedral phase of tBBr are really large (ca. 60°), giving to this phase II also the stamp of OD phase.$^{3,5}$ Such characteristic comes up clearly by comparing the enthalpy change of the II I phase transitions, rather low for the case of tBBr (see Table 1). It is worth noting that the puzzle of experimental literature data concerning the symmetry of phase II of tBBr was recently solved by Rudman$^{5,29}$ and lately confirmed.$^{28}$ This phase was found to be isostructural to the stable OD phase of CCl$_4$ and related halogenomethane compounds (as (CH$_3$)$_2$CCl$_2$, (CH$_3$)CCl$_3$, CCl$_3$Br, etc.)
Table 1. Temperature Transitions $T_c$ and Entropy Changes $\Delta S$ Derived from Calorimetric Measurements at Normal Pressure, Slope of the Temperature Pressure Two Phase Equilibria $dT_c/dP$ Derived from Thermal Analysis at High Pressure and Volume Changes Determined from the Clausius–Clapeyron Equation ($\Delta H^\circ$), from X-ray Powder Diffraction Measurements ($\Delta S^\circ$) and from pvT Measurements ($\Delta V^\circ$) for the Different Transitions of (CH$_3$)$_2$CBr,

| $T_c$, K | $\Delta S$, J·K$^{-1}$·mol$^{-1}$ | (dT$_c$/dP), K·MPa$^{-1}$ | $\Delta V^\circ$, cm$^3$·mol$^{-1}$ | $\Delta S^\circ$, cm$^3$·mol$^{-1}$ | $\Delta H^\circ$, cm$^3$·mol$^{-1}$ | ref |
|---------|-----------------|-----------------|-----------------|-----------------|-----------------|--------|
| III−II  | 209.0±0.1       | 28.23±0.80      | 0.206±0.020     | 5.81±0.73       | 5.58±0.21       | 5.50±0.71 |
| II−I    | 209.3±0.5       | 27.95±0.48      | 0.174           | 4.86            | 5.43            | 4.84   |
| IV−I    | 231.8±0.3       | 4.14±0.04       | 0.368           | 1.54            | 1.80            | 4.84   |
| I L     | 254.9±1.1       | 6.77±0.38       | 0.512±0.020     | 3.47±0.33       | 4.02±0.19       | 4.00±0.32 |

*Values in parentheses correspond to tBCl.

and CCl$_3$Br$_2$. The evidence of the structural similarity and even the existence of an isomorphism relationship (the isostructural relationship being then included) between the OD phase II of tBBr and stable phase I of CCl$_4$ was established throughout the existence of continuous mixed crystals for the whole concentration range. Even though the work of Rudman et al. was based on single crystals grown from the liquid state into phase I and lately transformed to phase II without further purification, the disordered character of this phase II and the complexity of the powder pattern and the large number of possible conformations and with a high molecular symmetry (i.e., consists of 4096 channels and enables a simultaneous collection of 1024 data points per second) were not successful, low-temperature high-resolution X-ray powder diffraction was used. As we will prove, the claimed complexity of the phase III structure does not concern the “shattering” number of molecules in the unit cell, but the special motifs forming the asymmetric unit. It has been found that $Z' = (1 + 1/2 + 1/2)$, with one molecule in a general position and two additional molecules placed in a mirror (then contributing as one-half each one). Thus, phase III of tBBr belongs to a reduced “selected group” (ca. 8% according to the Cambridge Structural Database) for which $Z' > 1$. But even more interesting is the fact that the molecular entity within the asymmetric unit is really simple, without different possible conformations and with a high molecular symmetry or, at the very least, no less than those of the tert-butyl compounds, (CH$_3$)$_2$CX.

The structural results as well as the thermodynamics of tBBr and tBCl and tBCN will be compared with each other.

In addition, the concomitant dynamical aspects (generally disregarded) of the polymorphism of these substances, largely explored, will be rationalized as a function of the structures and the pressure-volume-temperature phase diagrams.

2. Experimental Section

tert Butyl bromide was supplied by Aldrich Chemical Co. (purity better than 98%) and always handled under an Ar atmosphere without further purification.

2.1. High Resolution X-ray Powder Diffraction Measurements. High resolution X-ray powder patterns were isothermically recorded by means of a horizontally mounted INEL cylindrical position sensitive detector (CPS120) equipped with a liquid nitrogen cooled 600 series Cryostream Cooler from Oxford Cryosystems with a temperature accuracy of 0.1 K and similar for fluctuations. The detector, used in Debye–Scherrer geometry (transmission mode), consists of 4096 channels and enables a simultaneous recording of the diffraction profile over a 20 range between 2 and 115° (angular step of 0.029° in 20). Monochromatic CuKα ($\lambda = 1.54059$) radiation was selected by means of an asymmetric focusing incident beam curved quartz monochromator. The generator power was set to 1.0 kW (40 kV and 25 mA). Peakoc
application from Diffractinel software was used for the calibration and for the conversion channel angle after an external calibration using the cubic phase Na$_2$Ca$_3$Al$_2$F$_{14}$ mixed with silver behenate. The conversion was done by means of cubic spline fittings in order to correct the deviation from angular linearity in position sensitive detector (PSD) according to recommendations.

The standard measurements were performed at least every 10 K step in the temperature range of phase III, every 2–3 K for phase II, and every 5 K for the orientationally disordered phase I. Acquisition times were no less than 60 min for patterns. The slew rate was 1 K·min$^{-1}$ with a stabilization time of 5 min at each temperature before data collection.

A longer time acquisition pattern of 2 h for phase III (at 180 K) was measured for structure determination and refinement purposes.

The modules Powder Indexing (cell search), Powder Refinement (full profile determination after pseudo Voigt fittings), and Powder Solve of the Material Studio application have been used to resolve the crystal structure.

Liquid samples were sealed in Lindemann capillaries (0.3 mm diameter), which rotated perpendicularly to the X-ray beam during the experiment to improve averaging of the crystallites thus minimizing preferential orientations.

2.2. Thermal Analysis. Thermal analysis measurements at normal pressure were conducted by means of a Q100 thermal analyzer from TA Instruments equipped with a RCS low temperature device. Heating and cooling rates of 2 K·min$^{-1}$ were used. Sample masses around 25 mg were encapsulated into high pressure stainless steel pans with Au covers (from Perkin Elmer) to prevent possible reaction with the container.

The transition temperatures as a function of pressure (up to 300 MPa) were measured in a low temperature high pressure apparatus similar to that described elsewhere. The DTA traces were obtained on heating with scanning rates of about 1 K·min$^{-1}$. The measurements were performed in closed indium capsules containing about 40 mg of the sample. The usual experimental error is ±0.5 K for the temperature and ±0.05 MPa for the pressure values.

2.3. Density Measurements. Liquid density measurements at normal pressure were carried out from 278 to 353 K using an Anton Paar D5000 densimeter with temperature stability of ±0.02 K and uncertainties in density of ca. 5 × 10$^{-5}$ g·cm$^{-3}$. The densimeter was calibrated with bidistilled water and n-hexane (99.9%). The density values were taken from refs 50 and 51.

Density measurements as a function of pressure and temperature were performed by means of a homemade pvT device similar to that described in refs 9 and 52. The uncertainty of the density measurements is around 10$^{-4}$ g·cm$^{-3}$, while for temperature and pressure errors are similar to those of the thermal analyzer as a function of pressure.

3. Results and Data Analysis

3.1. Structure Determination of Phase III of TBr. In order to build up a rigid body molecule, the software Forcite from Materials Studio package together with the Compass force-field was used. The minimization energy process converged to an almost perfect tetrahedral molecular symmetry (C$\text{Br} = 1.948$ Å; C$\text{H} = 1.102$ Å; C$\text{C} = 1.533$ Å; C$\text{H} = 1.075$ Å; C$\text{C} = 1.114$ Å).

Potential solutions of the cell parameters and space groups were obtained using X-Cell software available in the module Powder Indexing of Material Studio.

The cell parameters and space group compatible with the experimental X-ray diffraction pattern were initially determined using a Pawley profile-fitting procedure. The Pawley refinement, which minimizes the weighted $R$-factor, $R_{wp}$, describing the agreement between the experimental pattern and the simulated one, helps to confirm the indexing result and to analyze the systematic absences, and thus to determine the possible space groups. Two space groups were found to be compatible with the Pawley refinement, $Pmmn$ and $Pn2_11$, both involving $Z = 8$ molecules per unit cell and the same systematic absences. Nevertheless, the former one implies one independent molecule and the latter two independent molecules in general positions for reaching $Z = 8$ by density requirements. In both cases, a Powder Solve run did not succeed. Another attempt was performed in space group $Pm$ by placing eight independent molecules in the lattice and applying 90° constraints on the lattice angle values. By this procedure, two clear elements of symmetry emerged: on one side a clear binary axis ($2\overline{1}$) and, on the other hand, the noncentrosymmetric character of the unit cell discarding the $Pmnn$ space group. A new attempt in $P2_1$ space group revealed the solution. Finally in $Pmm2_1$ ($b$ and $c$ axes being permuted), the independent molecules, one in the general position and the two others on the mirror ($Z = 4 + 2 = 8$) were fully positioned and the structure was coherently solved by combining Rietveld refinement and energy lattice minimization (using Forcite with refinement of the molecular geometry). Preferred-orientation corrections of the powder sample were initially searched within the Powder Solve module by means of the March-Dollase function and by the Rietveld-Toraya function for the final Rietveld refinement.

The experimental collected and the final calculated profiles are shown in Figure 1 together with the difference plot between them. The parameters refined in the procedure are collected in Table 2 and the coordinates of the atoms of the asymmetric unit are given in Table 3.

3.2. Lattice Parameter As a Function of Temperature. X-ray powder diffraction experiments were performed on heating from 90 K up to the melting temperature, thus covering all the solid phases in order to determine the volume changes at the phase transitions and the thermal expansion tensor. The lattice parameters were refined at each temperature (Figure 2) as previously described for phase III and using a Pawley profile-fitting procedure.

Figure 1. Experimental (red circles) and calculated (black line) diffraction patterns along with the difference profile (blue line) of orthorhombic phase III of (CH$_2$)$_3$CBr (at 180 K). Inset corresponds to the scale for the data with $2\theta > 38^o$. and $Pm2_11$, both involving $Z = 8$ molecules per unit cell and the same systematic absences.
Table 2. Results from the Rietveld Refinement of Phase III of (CH₃)₃CBr at 180 K

| parameter        | value       |
|------------------|-------------|
| 2θ angular range | 8°–90       |
| 2θ shift (zero correction) | 0.0007 ± 0.0005 |
| profile parameters |            |
| Na               | 0.220 ± 0.021 |
| Nb               | 0.0029 ± 0.0008 |
| lattice parameters |            |
| a/Å              | 19.1350 ± 0.0004 |
| b/Å              | 5.7378 ± 0.0001 |
| c/Å              | 10.9626 ± 0.0002 |
| reliability parameters |       |
| Rwp              | 7.73%       |
| Rp               | 5.56%       |
| peak width parameters |        |
| u                | 0.0492 ± 0.0052 |
| v                | −0.0294 ± 0.0033 |
| w                | 0.0135 ± 0.0005 |
| overall isotropic temperature factor, U/Å² | 0.0446 ± 0.0006 |
| preferred orientation (Rietveld–Toraya function) | |
| a*               | 0.000       |
| b*               | 0.902 ± 0.006 |
| c*               | 0.430 ± 0.013 |
| G2               | −0.882      |
| G1               | 0.328       |

Table 3. Fractional Atomic Coordinates for Phase III of (CH₃)₃CBr at 180 K

| atom   | X      | Y      | Z      |
|--------|--------|--------|--------|
| Br35   | 0.28933(15) | 0.7666(7) | 0.5449(11) |
| C31    | 0.2603(4)   | 0.6181(10) | 0.3933(11)  |
| C32    | 0.2366(4)   | 0.8096(15) | 0.3052(12)  |
| C321   | 0.2791(3)   | 0.9335(12) | 0.2853(11)  |
| H322   | 0.2193(6)   | 0.7346(20) | 0.2177(12)  |
| H323   | 0.1924(2)   | 0.9097(17) | 0.3428(11)  |
| C33    | 0.1999(4)   | 0.4515(13) | 0.4217(12)  |
| H331   | 0.1813(6)   | 0.3645(18) | 0.3382(11)  |
| H332   | 0.2157(4)   | 0.3145(10) | 0.4864(11)  |
| H333   | 0.1551(3)   | 0.5449(15) | 0.4617(12)  |
| C34    | 0.3230(3)   | 0.4843(18) | 0.3413(12)  |
| H341   | 0.3091(8)   | 0.3980(12) | 0.2548(12)  |
| H342   | 0.3674(5)   | 0.6012(7)  | 0.3228(13)  |
| H343   | 0.3411(5)   | 0.3481(6)  | 0.4048(13)  |
| Br5    | 0.00000    | 1.0650(10) | 0.7869(7)   |
| C1     | 0.00000    | 0.9562(10) | 0.6187(7)   |
| C2     | 0.0659(4)  | 1.0488(10) | 0.5558(7)   |
| H21    | 0.0681(5)  | 0.9910(10) | 0.4600(7)   |
| H22    | 0.0674(4)  | 1.2407(10) | 0.5558(7)   |
| H23    | 0.1136(5)  | 0.9860(10) | 0.6013(7)   |
| C3     | 0.00000    | 0.6893(10) | 0.6209(7)   |
| H32    | 0.00000    | 0.6176(10) | 0.5277(7)   |
| H33    | 0.0465(3)  | 0.6206(10) | 0.6661(7)   |
| Br15   | 0.00000    | 0.4661(10) | 0.2886(7)   |
| C11    | 0.00000    | 0.6516(10) | 0.1400(7)   |
| C12    | 0.0660(6)  | 0.8029(10) | 0.1392(7)   |
| H121   | 0.0630(5)  | 0.9202(10) | 0.2187(7)   |
| H122   | 0.0681(4)  | 0.9123(10) | 0.0566(7)   |
| H123   | 0.1138(5)  | 0.6954(10) | 0.1406(7)   |
| C13    | 0.00000    | 0.4840(10) | 0.0312(7)   |
| H132   | 0.00000    | 0.5808(10) | 0.0554(7)   |
| H133   | 0.0465(5)  | 0.3708(10) | 0.0316(7)   |

The most striking features of the structure of phase III of tBBr are the number of molecules forming the asymmetric unit $Z' = (1 + 1/2 + 1/2)$ and the absence of a center of symmetry which gives rise to noticeable differences in the molecular arrangement of the motifs in the lattice structure compared to those of tBCl and tBCN (Figure 7).

For the low-temperature phases of tBCl and tBCN, with space groups $P2_1/m$, the mirror plane of the molecule has to be coincident with the mirror plane of the space group, and thus the molecule can only rotate around $b$ and the $Y_c$ coordinate of the center of the asymmetric unit (formed by one molecule) compared to those of tBCl and tBCN (Figure 7).

The most striking features of the structure of phase III of tBBr are the number of molecules forming the asymmetric unit $Z' = (1 + 1/2 + 1/2)$ and the absence of a center of symmetry which gives rise to noticeable differences in the molecular arrangement of the motifs in the lattice structure compared to those of tBCl and tBCN (Figure 7).
is set as $1/4$. Consequently, the molecules are arranged in a head-to-tail configuration with the 3-fold $C_3$-axis (dipole axis of the molecule) almost parallel to the $c$ direction in the $ac$ plane (tilts are about $2.2^\circ/C_176$ and $4^\circ/C_176$ for tBCl and tBCN, respectively) (see Figure 7b,c). In this way, the adjacent rows of molecules along the $c$ direction (with parallel $C_3$-axis) are inverted with respect to each other, and in spite of the rotation about the 3-fold molecular axis, the inverted molecules get closer with a high packing coefficient.

As far as the structure of phase III of tBBr is concerned, a noticeable difference is inferred. Because of the noncentrosymmetric structure ($Pmn\bar{2}$ space group), all the molecules of the unit lattice point out closely in the same direction (Figure 7a). The four molecules corresponding to the symmetry operations of the molecule in a general position of the asymmetric unit stack with the $C_3$-axis tilted around $31.46^\circ$ with respect to the $c$ axis of the lattice (see Figure 7a). For the two molecules of the asymmetric unit in a special position, the angles of their $C_3$-axis with respect to the $c$-axis are $18.80^\circ$ and $32.75^\circ$.

Such a packing arrangement gives rise to rows of molecules along the crystallographic $c$-axis forming a zigzag and without inversion of molecules and thus, to some extent, to a less effective packing than that of the tBCl within their low-temperature structure ($P2_1/m$). As far as the packing of tBCN is concerned, the elongated CN group gives rise to a noticeably increased distance between the molecules along the $c$-axis (around 6.0 Å for tBCl and 6.7 Å for tBCN, see Figures 7b,c).
Such a difference is strictly coming from the higher van der Waals volume of the CN group in comparison with that of the Cl atom as can be seen by comparing the similar distance between the N or Cl atoms to the central carbon of the closest molecule (4.1 and 4.2 Å, respectively). This fact induces then a considerable reduction of the packing coefficient of tBCN with respect to tBCl.

Figure 8a shows the variation of the packing coefficient \( \eta = V_m/(V/Z) \), in which \( V_m \) is the van der Waals volume of the molecule (tBCl = 93.05 Å\(^3\); tBBBr = 96.97 Å\(^3\); tBCN = 95.80 Å\(^3\)) and \( V/Z \) is the volume occupied by a molecule in the lattice) for the different phases as a function of the scaled temperature with respect to the III to II phase transition (II I for tBCN compound). The molecular van der Waals volumes were computed according to the Kitaigorodsky's method\(^6^1\) with interatomic chemical bonds from the reported structures and atomic van der Waals radii obtained from Material Studio database (C: 1.70 Å; H: 1.20 Å; Cl: 1.75 Å; Br: 1.85 Å; N: 1.54 Å).

As can be seen, the packing coefficients of the low-temperature phases, where the highest intermolecular interactions
are known to be present, are higher than those of the tetragonal (for tBCl and tBCN) and rhombohedral (tBBr) phases, for which interactions have been “softened” due to the appearance of some kind of disorder. This fact is especially noticeable for tBBr, for which the packing variation at the III II transition is the highest, a consequence of the OD character of phase II as clearly indicated by the entropy change at the III to II transition (the highest for tBBr compound, see Table 1) as well as by the large fluctuations of the dipole axis reported from NMR measurements.5,11,12,17 It should be borne in mind that for phase II of tBCl and phase I of tBCN (both of tetragonal symmetry) the disorder of the molecules consists only of small fluctuations (librational motions) of the dipole axes.16

It is worth noting that the packing values of OD fcc phases for tBBr and tBCl (not reported in Figure 8a) are relatively low (ca. 0.57) when compared to other plastic phases for which the existence of dynamical hydrogen bonds has been clearly demonstrated.62,63

As for the low-temperature phase for all these compounds, the a lattice parameter changes faster than b and c with temperature (Figure 2) because the nearest neighbors molecules in this direction have a close packing of the tertiary butyl groups. Nevertheless, this packing should be compatible with the increasing of the reorientational frequency of the fluctuations of the dipole axis, around which (C$_3$-axis) uniaxial molecular reorientations between the three equivalent positions, and thus large thermal expansion occurs in such a direction. This behavior can be straightforwardly seen by analyzing the thermal expansion tensor. The thermal expansion tensor expresses the cell deformation $dU$ due to a temperature change $dT$ by means of a second-rank tensor at a given temperature, $dU = \alpha_i \partial_i T$, $\alpha_i$ being the principal coefficients (eigenvalues) of the thermal-expansion tensor. The principal axes (eigenvectors) of the tensor describe the highest and the weakest deformations related to the directions of the soft and hard directions, respectively, of the intermolecular interactions.64,65 Details of the procedure and method for the calculations of the tensor have been published elsewhere.65

For a monoclinic lattice, the tensor is defined by three principal coefficients and an angle between the direction of one of the principal directions ($\alpha_1$) and one lattice axis, $\alpha_3$ being coincident with the 2-fold axis $b$ of the lattice. For an orthorhombic lattice, the principal directions should be coincident with the lattice axes of the crystal according to the Neumann’s principle (for which the thermal-expansion tensor has to display the point group symmetries of the crystal).

Left panels of Figure 7 display the tridimensional thermal-expansion tensor for the low-temperature phases of tBBr, tBCI, and tBCN. As can be inferred from that figure, the highest deformation (“soft” direction) corresponds closely to the [100] crystallographic direction in low-temperature phases of tBCI and tBCN, and exactly to such a direction for tBBr. On the contrary, the lowest deformation corresponds to the (100) plane (“hard” plane) for all the compounds. In the case of tBCN, the $\alpha_3$ direction (close to [001] crystallographic direction) is even negative (contraction) in the displayed temperature range of the monoclinic phase. For tBCI and especially for tBCN the large thermal expansion along the [100] direction is coupled with the strong interaction along the c axis (parallel to the molecular dipoles) in such a way that the $C_3$-rotation of the tert-butyl group is less impeded with increasing temperature as indicated by the increase of the frequency of reorientations around that axis which cause the molecules to move further apart in the a crystallographic direction as the temperature increases. This is not the case for tBBr, for which the anisotropy of the thermal expansion tensor is lower as can be inferred from the relatively close values of the principal coefficients of the tensor (Figure 8b).

The anisotropy of the intermolecular interactions can be accounted for by means of the aspherism index, defined as

$$A = \left(\frac{2}{3}\right)[1 - (3\beta/\alpha_3^2)]^{1/2}$$

where $\beta = \alpha_1 \alpha_2 + \alpha_2 \alpha_3 + \alpha_3 \alpha_1$.66 This coefficient reaches the zero value when the thermal expansion tensor is isotropic. For the low-temperature phase of the compounds reported in Figure 8c, the aspherism index increases with temperature as a consequence of the increase in the anisotropy of the intermolecular interactions. The highest values are those corresponding to tBCN thus evidencing the strong dipole dipole interaction within the head-to-tail arrangement of the molecules.

The ordered or partially disordered character of the involved phases has consequences in the entropy as well as in the volume changes at the phase transitions. When comparing such volume changes (Figure 6), it comes clear the similarity between the R to FCC transitions for both compounds (even though phase R only exists at high-pressure for tBCl) and the strong difference between T to R (tBCl) and O to R (tBBr), which show that the T phase of tBBr has a partial disorder (librations around the dipole molecular axes) while O phase of tBBr is fully ordered, as the $P2_1/m$ low-temperature phase of tBCl. In this context, tetragonal phase I of tBCN should not be considered as an orientationally disordered phase but a partially ordered phase similar to the tetragonal phase of tBCl.

5. Conclusions

The polymorphism of (CH$_3$)$_3$CBR has been studied from 90 K to the liquid state as well as a function of pressure (until 300 MPa).

The long time unknown structure of the low-temperature phase of (CH$_3$)$_3$CBR has been determined to be orthorhombic with space group Pmn2$_1$. It involves eight molecules per unit cell ($Z = 8$) with an asymmetric unit ($Z'$) formed by one molecule in a general position and two additional molecules placed in a mirror (then contributing as one-half each one), that is, $Z' = 1 + 1/2 + 1/2$. The similarities and differences with related tert-butyl compounds as (CH$_3$)$_3$CCI and (CH$_3$)$_3$CCCN for which the structures of the low-temperatures phases have been previously reported have been highlighted. In particular, through the thermal-expansion tensor, the differences in the role played by the dipole dipole interactions seem to be the main cause of the structural differences.

The pressure temperature phase diagram has been determined within the range of 0.1–300 MPa. The polymorphism sequence at normal pressure is kept whatever the pressure within the studied range. From density measurements as a function of pressure, volume changes have been determined giving a coherent physical image of the determined structure of the orthorhombic phase as well as for the previous lattice determinations of the orientationally disordered phases II (rhombohedral) and I (FCC).
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