SPECTROSCOPIC EVIDENCE FOR A CONTINUOUS CHANGE IN MOLECULAR AND CRYSTAL STRUCTURE: DEFORMATION OF BIPHENYL IN THE LOW TEMPERATURE SOLID

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A Raman temperature study of biphenvl-10 and biphenvl-d10 crystals reveals an unusual spectral change over a wide temperature region. Many internal and external infrared-active modes appear gradually as the temperature is lowered from about 75 to 1.5°K. We interpret this as a mild and gradual change in both crystal and molecular structure: a loss of center of inversion, probably involving the partial return of the molecule to its non-planar shape (D2) in the vapor phase.

We report here what appears to be a non-monotonic behavior of the molecular structure of biphenvl in its condensed phases. Biphenyl has been reported to be non-planar in the gas phase [1-3], in solution [2, 3] and in the melt [2]. However, X-ray crystallography [4] (room temperature) shows the molecule to be planar (D2h) and the crystal centrosymmetric (Csh). We present below spectroscopic evidence for a continuous structural change occurring in the crystal at low temperature, characterized by the molecule becoming non-centrosymmetric and implying the existence of a corresponding continuous phase transition.

In the Raman spectrum six phonon bands are expected, as the biphenvl crystal at room temperature has two interchange equivalent molecules (site symmetry C1) per primitive cell [4]. At 55°K (fig. 1) we see five phonon bands of external vibrations (determined from the amalgamation limit criterion [5, 6], at frequencies of 41, 67, 71, 107 and 117 cm⁻¹. The sixth band is suspected to be at around 30 cm⁻¹. At 15°K the high temperature features remain essentially unaltered, while at least three additional weak bands appear reproducibly at 49, 58 and 78 cm⁻¹. These weak bands being in the amalgamation limit [5, 6] must also be phonons (or torsions). The possibility that they are phonon overtones and/or combination bands is ruled out on the following grounds: (i) overtones or combination bands being due to a multi-phonon process are expected to grow in intensity as the temperature is increased, but we observe the reverse; (ii) the 49 cm⁻¹ peak is too low in energy to be accounted for as either an overtone or combination band of the observed phonon bands; (iii) in this molecular crystal, overtones and combinations are expected to have a two particle excitation character [8] and should thus be broad compared to one-phonon peaks. This also is not observed.

In the intramolecular region the major Raman bands are again very similar at both high and low temperature (fig. 1). Even the doublet at 740 cm⁻¹, which we have established to be due to Davydov splitting (from our isotopic mixed crystal criterion [9]), remains nearly the same at both temperatures. Yet at low temperature (15-40°K) at least five additional peaks (not due to overtones and/or combination bands) appear in the Raman spectrum, this time at 136, 188, 389, 87, 5 and 961 cm⁻¹. A careful temperature dependence study shows an extremely wide transition region, with new bands appearing gradually. One example is shown in fig. 2, where we see that the strongest new feature is...
discernible already at 65°K. We stress that all our data are completely reproducible and reversible with temperature cycling.

In summary, we observe that as the crystal is cooled, the high temperature Raman bands continue to retain their identity, however, in the low temperature range (15—40°K), additional new bands gradually emerge both in the phonon and intramolecular regions (biphenyl-\(d_{10}\) crystal shows an analogous behaviour [7] to biphenyl-\(d_{10}\)). This suggests that at low temperature some sort of structural change of very mild character is taking place, for if a large change in the crystal structure were occurring, it would be expected to significantly alter both the appearance of the phonon spectrum [10] and of the Davydov splitting.

Of the eight low temperature (15°K) Raman bands so far described, the three appearing at 78, 136 and 188 cm\(^{-1}\) correlate closely with the far-infrared peaks [7] seen by us (preliminary data) at 76, 133 and 188 cm\(^{-1}\) (47°K). These infrared frequencies are in general agreement with previous work [11, 12]. Also, the (new) Raman bands appearing at about 389, 825 and 961 (15°K) correlate with \(A_u\) infrared bands [2, 11, 13], located around 401, 839 and 968 cm\(^{-1}\) (93°K) [11]. This correspondence between Raman and infrared bands indicates that biphenyl has lost its center of inversion\(^\dagger\).

Of the three most feasible non-centrosymmetric static molecular deformations (long axis twist, out-of-plane ring—ring bending, in-plane ring—ring bending) we expect the one caused by a long axis twist (D\(_3\)) to be the most likely one taking place, because the long axis twist involves the softest force field. Also, if either of the two ring—ring bending molecular deformations (C\(_{2v}\)) were occurring, a large frequency shift would be expected in the corresponding bending vibrations (133 and 188 cm\(^{-1}\) at 47°K). No such effect has been observed.

Our temperature dependence study shows that the structural change occurs gradually (also reversibly and reproducibly) as \(kT\) varies by a factor of three or more\(^*\).

We stress here that the observed continuous change involves at least three external and five internal modes (as opposed to just a single degree of freedom). This seems to us to be unprecedented.

The 2°K fluorescence spectrum of the isotopic mixed

\(^\dagger\) Lecolier et al. [12] have recently reported the far-infrared spectrum of the biphenyl crystal at several temperatures. A peak appears in their 15°K spectrum at 67 cm\(^{-1}\), which is not seen at 80°K. We attribute this low temperature peak to a Raman allowed phonon, being made infrared active through a loss of the molecular center of inversion.

\(^*\) Heat capacity measurements are underway. We observe at 1.5°K essentially the same Raman spectrum as at 15°K.
crystal (1% biphenyl-$d_{10}$ in biphenyl-$d_{10}$) shows a splitting of approximately 11 cm$^{-1}$ on virtually every major vibronic band [7, 14] and cannot be explained by the room temperature crystal structure. Such a splitting would be consistent with our proposed non-centrosymmetric biphenyl structure, in that a pair of such deformed biphenyl molecules can have two possible mutual orientations of slightly different energy.

References