Conformational aspects of polymorphs and phases of 2-propyl-1H-benzimidazole

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S1. Crystal preparation by sublimation

Crystal of forms I and II (HT and RT) prepared by sublimation heating at 400 K a sample of purified commercial sample. Experimental device and obtained crystals are shown in Figs S1 and S2.

**Figure S1** Hot stage used to prepared crystals in forms I and II (HT and RT) obtained by sublimation from sample-I type crystals (inside the ring). The sublimated compound condensates on a thin glass plate (not shown) covering the ring.
Needle shape crystals belong to form II<sub>RT</sub> grown on a hot area (T>384 K). The small squared shape crystals of form I grown on a cooler area of the glass plate on covering the heated sample-1.
S2. DSC. Thermal behaviour

Differential scanning calorimetric (DSC) measurements were carried out in a TA Instruments MDSC Q-2000 calorimeter in the temperature range of 140-450 K at heating and cooling rate of 10 deg/min under a helium atmosphere. Three polycrystalline samples of 2-propyl-1H-Benzimidazole of 10 up to 15 mg of mass were encapsulated in aluminum hermetic pans. The calibration of the calorimeter was performed using single-crystal sample of sapphire for calibration of specific heat values and melting point of indium for temperature and heat flow calibration. An uncertainty of 2% in \(\Delta H\) was calculated from calibration with Indium.

In the first cooling run from room temperature down to 120 K, no anomaly was detected DSC. In the heating cycle, heating up to 400 K, an irreversible phase transition is observed at 381 K with enthalpy of 5.1±0.3 kJ/mol (average of 4.8, 5.2 and 5.4 kJ/mol). In the next cycle cooling down 120 K, two reversible anomalies are observed at 361 K and 181 K. The enthalpy of the low temperature transition increases upon cycling and reach a stable value after a few cycles. DSC runs and details of the transition peaks are shown in next figures.

**Figure S3** Flow heat v.s. temperature, positive in cooling cycles and negative in heating cycles. First cycle (red line) starts at room temperature, cool down to 120 K and heat up to 410 K. In the next cycle (blue line), cooling from 410 K down to 120 K. Next cycles are omitted for enhance clarity.
**Figure S4** Flow heat in a first heating-cycle around the $\text{I-\text{II}_{\text{HT}}}$ transitions. The red curve corresponds to a recrystallized sample and the blue one to a commercial sample. The small anomaly hoisted over the blue peak corresponds to the $\text{II}_{\text{RT}}$-$\text{II}_{\text{HT}}$ transition of a small fraction of form $\text{II}_{\text{RT}}$ in the commercial sample.
**Figure S5** DSC curve of a heating cycle between forms $\text{II}_{\text{RT}}$ and $\text{II}_{\text{HT}}$. The blue line is the base line.
**Figure S6** DSC curve of a heating cycle between forms \( \text{II}_{\text{LT}} \) and \( \text{II}_{\text{RT}} \). The blue line is the base line.
**Figure S7** DSC curves showing the evolution of the low temperature transition peak in successive cooling and heating cycles. The flat line corresponds to the first cycle cooling from room temperature. Note the thermal hysteresis of 10 K.

**S3. IR and VCD measurements.**

For the high temperature measurements a Bruker heated sample holder (A599 model), designed for the analysis of solid samples for Raman spectroscopy, was coupled to our VCD spectrometer. Small circular BaF$_2$ windows (13x3 mm) were built in our laboratory starting from other bigger ones (PIKE Technologies, 32x3 mm), in order to fit them in the space available in the heated sample holder above-mentioned. Then, the holder was connected to a power supply and temperatures above 298 K for the crystalline 2PrBzIm samples in fluorolube mineral oil could be achieved. This part of the study could not be carried out in nujol mineral oil because of the lower intensities of the spectral bands in the corresponding useful region (2000-1500 cm$^{-1}$ and 1300-900 cm$^{-1}$). The VCD bands were more intense in the 1550-1420 cm$^{-1}$ range, which can only be measured in fluorolube mull (2000-1300 cm$^{-1}$ useful region). In this mineral oil, the VCD spectral changes according to the temperature could be followed properly.

The IR and VCD spectra were observe at temperatures below and above 378 K. The lack of chirality is evident above 378 K and when cooled again at room temperature.
Supporting information, sup-9

**Figure S8** IR (top) and VCD (bottom) spectra of 2PrBzIm compound in fluorolube mull at different temperatures: a) Form I at room temperature (298 K, black), heating until 343 K (orange) and cooling at room temperature again (blue) and b) Form I at room temperature (298 K, black), form II\(_{\text{HT}}\) at 378 K (red) and form II\(_{\text{RT}}\) at room temperature (cooling after heating, blue).

**S4. X-Ray powder diffraction**

X-Ray powder diffraction (XPD) spectra obtained in Debye geometry with sample in capillary of 0.5 mm. Diffractometer STOE STADIP, with Ge-monochromatized Cu\(\alpha_1\) radiation. Temperature monitored near the capillary with a calibrated thermocouple and controlled with a homemade heater element and Oxford Instrument ITC controller. The variation of the temperature was less that 1\(^\circ\) during the whole exposure time.

First, diffraction patterns were recorded in heating and cooling cycles in the temperature range 333 K-398 K in steps of 5 K and heating rate of 10 K/min. Patterns are show in Fig. S9. The orthorhombic form I is stable up to the high temperature transition. The drastic change observed in the recorded X-ray powder diffraction diagram around 380 K led to conclude that, the first anomaly observed in DSC at 384 K correspond to a reconstructive type transition and lead to a new structure identified as orthorhombic with lattice parameter \(a_{\text{II}} = 8.9957(1), b_{\text{II}} = 21.5226(3)\) and \(c_{\text{II}} = 9.8982(1)\) Å. Note that the transition temperature observed in DSC correspond the maximum of the Cp signal but transitions starts around 370 K. Cooling down to room temperature the recorded spectra does not show major changes so that the transition observed in DSC at 361 K does not involves a big structural modifications. The diffraction pattern at room temperature of polymorph II\(_{\text{RT}}\) was indexed as orthorhombic with lattice parameters \(a_{\text{III}} = 8.975(3), b_{\text{III}} = 21.134(6)\) and \(c_{\text{III}} = 9.823(2)\) Å. XPD patterns of forms I, II\(_{\text{HT}}\) and II\(_{\text{RT}}\) are shown in Figs. S10 and S11. The crystallographic characterization of polymorph II\(_{\text{LT}}\) was done by means of single crystal X-ray diffraction and is described below.
Figure S9  XRD patterns recorded in steps of 5 K in the temperature range 333 K-398 K in (a) heating and (b) cooling cycles respectively.
Figure S10 Superposition of powder diffractograms of form I at room temperature (blue) and form II\text{HT} at 380 K (red).
**Figure S11** Superposition of powder diffractograms of phases II\textsubscript{RT} at room temperature (blue) and II\textsubscript{HT} at 380 K (red).

**S5. Structural refinement**

**S5.1. Rietveld refinement of form II\textsubscript{HT}**

The structural model of 2PrBzIm in the high temperature form II\textsubscript{HT} was refined through the Rietveld method based on the XPD data measured at 380 K. The indexing peaks are compatible with extinctions of the space group $P_\text{cam}$ and $P_\text{ca}c_2_1$ ($Z = 8$, $Z' = 2$). Both space groups were tested using as starting model that obtained for form II\textsubscript{RT} (see below). Refinement with the non-centrosymmetric space groups becomes unstable and was discarded. In the centrosymmetric space group the molecules are disordered between equally populated positions symmetry related by the mirror plane. The refinement was performed including geometrical restrictions for planar benzimidazole and regular six carbon-atoms ring. Hydrogen atoms bonded to the imidazole fragment were assigned to the N atom with the largest intra-molecular C-N-C' angle. The H-atoms were calculated in geometrically restricted in idealized positions using riding model (C-N=0.87, Å C-H = 0.98 Å) and their isotropic displacements parameters set to 1.2Ueq of the parent N or C atom. The structure was refined using isotropic thermal displacements restricted to a unique value. Agreement between observed and calculated patterns is presented in Fig. S12. Crystal data and details of the refinement are given in Table 2.
Supporting information, sup-13

**Figure S12** Rietveld refinement (JANA 2006) of the X-ray powder diffraction pattern of form II$_{HT}$ at 380 K.

**S5.2. Single crystal X-ray refinement of forms II$_{RT}$ and II$_{LT}$**

Suitable single crystals in form II$_{RT}$ were obtained by sublimation as described in S1. The selected crystal used to collect intensities at 200 K (in II$_{RT}$) was also used for measurements at 150 K (II$_{LT}$). Experimental details are given in Table 2. The global diffraction pattern of form II$_{RT}$ was interpreted as orthorhombic with systematic absences compatible with the Pcam and Pca2$_1$ space groups (Z=8, $Z'$=2). The structural solution (SIR2002, Burla et al., 2003) confirms the non-centrosymmetric space group.

The diffraction pattern of polymorph II$_{LT}$ at 150 K shows new weak diffraction spots (superstructure reflections) indicating a doubling the period along the $a$ axis. The global diffraction pattern was interpreted as orthorhombic with systematic absences compatible with the Pna2$_1$ space group (Z = 16, $Z'$ = 4) confirmed with the structural refinement. The starting model in the refinement was generated as a subgroup of the parent form II$_{RT}$. Both structures (II$_{RT}$ and II$_{LT}$) were refined with the same type of intra-molecular restriction. In the final steps of the refinements all the individual atomic coordinates free to vary but restricting the six carbon-atoms rings to a regular shape and the benzimidazole in flat configuration. At the end of the refinement anisotropic thermal displacements were refined for all non-hydrogen atoms. The H-atoms bonded to N atoms were located in difference electron density maps and their positions, together with the rest of the H-atoms, were calculated in idealized positions using riding model (C-N=0.87, Å C-H = 0.98 Å) and their isotropic displacements parameters set to 1.2Ueq of the
parent N or C atom. Crystal data, data collection and refinement details for $\text{II}_{\text{RT}}$ and $\text{II}_{\text{LT}}$ are given in Table 2. Ortep plots showing the molecular configuration in forms $\text{II}_{\text{RT}}$ and $\text{II}_{\text{LT}}$ are shown in Fig. S13. Projections showing the molecular packing are shown in Figs. S14 and S15.

**Figure S13** Ortep plots of molecules in the asymmetric unit in forms $\text{II}_{\text{RT}}$ and $\text{II}_{\text{LT}}$. Configurations of molecules in polymorph $\text{II}_{\text{HT}}$ are similar to that of $\text{II}_{\text{RT}}$ and are not represented. The molecular labels “Mol n” are added to help identification in Table S1.
Figure S14 Schematic representation of the packing in the unit cell of polymorph II\textsubscript{RT} (a) polymorph II\textsubscript{LT} (b). Symmetry independent molecules are colored coded. Origins in structures II\textsubscript{RT} and II\textsubscript{LT} are shifted.
Figure S15 Projections along the [100]-direction showing the N-H..N’ bonds scheme in the (a) polymorph II_{RT} (b) polymorph II_{LT}. H-bonds not involved in N-H…N’ bridges are omitted for enhanced clarity.
S5. Computational studies on 2-propyl-1H-benzimidazole (2PrBzIm)

**Table S1** Summary of conformation torsion angles $\tau_1$ and $\tau_2$ of molecules labeled as “Mol n” in Fig. S11. Torsion angles values of independent molecules are typeset in black and values typeset in grey correspond to symmetry related molecules.

| Torsion angle $\tau_1$ | I (295)$^\S$ | II_HHT (380) | III_RT (200) | II_LT (150) |
|-------------------------|--------------|--------------|--------------|--------------|
| Mol1: N1—C1—C8—C9     | 45(1)        | -76 (2)      | -81.9(2)     | -81.5(2)     |
| Mol2: N3—C11—C18—C19 | -46(1)       | -39 (2)      | -47.4(2)     | -44.6(3)     |
| Mol3: N5—C21—C28—C29 | -50(1)       | 76 (4)       | 81.9(2)      | 84.2(2)      |
| Mol4: N7—C31—C38—C39 | 47(1)        | 39 (2)       | 47.4(2)      | 47.6(3)      |

| Torsion angle $\tau_2$ | I (295)$^\S$ | II_HHT (380) | III_RT (200) | II_LT (150) |
|-------------------------|--------------|--------------|--------------|--------------|
| Mol1: C1—C8—C9—C10    | 168.3(8)     | -85 (2)      | -71.1(2)     | -72.4(3)     |
| Mol2: C11—C18—C19—C20 | 177.4(8)    | 180 (2)      | -177.4(2)    | -67.1(3)     |
| Mol3: C21—C28—C29—C30 | -172.6(8)   | 85 (2)       | 71.1(2)      | 75.0(3)      |
| Mol3: C31—C38—C39—C40 | 168.5(8)    | 180 (2)      | -177.4(2)    | 176.4(2)     |

$^\S$ Structural data from Cabildo et al. (2015)

**Table S2** Intra-molecular coordinates of the conformation molecular energies minima obtained from calculations with B97D hybrid functional with a 6-311G** Pople basis set. Energy values are given with respect to the absolute minimum.

| Conformer | $\tau_1$(°) | $\tau_2$(°) | E (kJ/mol) |
|-----------|-------------|-------------|------------|
| CT-       | 257.0615    | 63.3726     | 0.0        |
| CL        | 282.3384    | 181.387     | +2.15829506491 |
| CT+       | 288.5064    | 297.9062    | +0.0992386565596 |
Figure S16 Perspective view of the conformational energy surface (PES) of the 2PrBzIm molecule calculated with B97D functional.