The true structural periodicities and superspace group
descriptions of the prototypical incommensurate composite
materials: Alkane/urea inclusion compounds

Michel Couzi¹(a), François Guillaume¹(b), Kenneth D. M. Harris²(c), Benjamin A. Palmer²,³, Kirsten Christensen⁴,⁵ and Stephen P. Collins⁴

¹ Université de Bordeaux, CNRS, ISM UMR 5255 - 351 cours de la Libération, F-33405 Talence Cedex, France
² School of Chemistry, Cardiff University - Park Place, Cardiff CF10 3AT, Wales, UK
³ Department of Structural Biology, Weizmann Institute of Science - Rehovot 7610001, Israel
⁴ Diamond Light Source, Harwell Science and Innovation Campus - Didcot, Oxfordshire OX11 0DE, England, UK
⁵ Inorganic Chemistry Laboratory - South Parks Road, Oxford OX1 3QR, England, UK

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Abstract – The prototypical family of incommensurate composite materials are the n-alkane/urea inclusion compounds, in which linear chain n-alkane guest molecules are arranged in a periodic manner along one-dimensional tunnels in a urea host structure, with an incommensurate relationship between the periodicities of the host and guest substructures along the tunnel. We develop interpretations of the structural periodicities, superspace group descriptions and symmetry properties of the low-temperature phases of n-alkane/urea inclusion compounds, based in part on a high-resolution synchrotron single-crystal X-ray diffraction study of n-nonadecane/urea. Specifically, we prove that, on passing from phase I to phase II, the C-centering of the orthohexagonal unit cell is lost for both the host and guest substructures, and that the symmetries of all phases I, II and III are described completely by (3+1)-dimensional superspace groups.

Aperiodic crystals are defined as materials that do not have 3-dimensional translational periodicity, yet exhibit sharp Bragg maxima in their X-ray diffraction patterns. Among the various classes of aperiodic materials, incommensurate solids [1] and quasicrystals [2] have received particular attention both from the fundamental perspectives of diffraction physics and structural science and with regard to their potential applications [3,4].

Host/guest composite materials are constructed from two interpenetrated substructures. In many cases, the host and guest subsystems share common crystallographic axes in three-dimensional space, and are commensurate structures. However, in some cases, there is an irrational relationship between the periodicities of the host and guest substructures, at least in one direction, and such materials are classified as incommensurate structures. The prototypical family of incommensurate composite materials is the n-alkane/urea inclusion compounds [5–11], in which linear chain n-alkane guest molecules are arranged in a periodic manner along one-dimensional tunnels (diameter ca. 5.25 Å) that exist within the urea host structure, which is constructed from a helical hydrogen-bonded arrangement of urea molecules (fig. 1(a)).

Urea inclusion compounds exhibit a wealth of interesting physico-chemical properties that are derived from their unique structural characteristics including crystal growth processes [12–16], X-ray dichroism [17] and X-ray birefringence [18]. A consequence of the incommensurateness of

¹E-mail: michel.couzi@u-bordeaux.fr
²E-mail: francois.guillaume@u-bordeaux.fr
³E-mail: HarrisKDM@cardiff.ac.uk
urea inclusion compounds is that the set of guest molecules may, in principle, slide along the host tunnel with essentially zero activation energy; this phenomenon has been harnessed in molecular transport experiments [19–22]. In order to fully understand the properties of these materials, it is imperative to establish an accurate description of their aperiodic structures.

The n-alkane/urea inclusion compounds are characterized by the following features at ambient temperature: i) a hexagonal urea tunnel structure [5,6,8], ii) an incommensurate relationship [6,8,11,23–43] between the periodicities of the host and guest substructures along the tunnel axis, and iii) dynamic disorder of the guest molecules [32,44–49] (reorientation about the tunnel axis and translations along the tunnel axis).

Focusing on the incommensurate properties of n-alkane/urea inclusion compounds, the misfit parameter $\gamma$ is defined as $\gamma = c_{\text{host}}/c_{\text{guest}}$, where $c_{\text{host}}$ and $c_{\text{guest}}$ are the periodicities of the host and guest substructures along the tunnel direction, respectively (fig. 1(b)). Clearly, $c_{\text{host}}$ depends on the pitch of the urea helix and $c_{\text{guest}}$ depends on the length of the n-alkane guest molecule. For all n-alkane/urea inclusion compounds at ambient pressure, $c_{\text{host}} \approx 11.02 \text{Å}$ [5,6].

In this paper, we focus on the urea inclusion compounds containing n-nonadecane [CH$_3$(CH$_2$)$_{17}$CH$_3$] and n-hexadecane [CH$_3$(CH$_2$)$_{14}$CH$_3$] guest molecules.

At ambient temperature, the urea host substructure has space group $P6_122$ (or $P6_522$) [5,6]. The composite crystal achieves 6-fold symmetry as a consequence of the n-alkane guests being distributed statistically in at least six equi-probable orientations. Under these conditions, the n-alkane substructure is described by space group $P622$. This assignment is based on the fact [7,29,50] that the n-alkane guest molecules exhibit 3-dimensional ordering in which guest molecules in adjacent tunnels are located at the same set of positions ($z$-coordinates) along the tunnel direction: i.e., the offset ($\Delta_g$), along the tunnel direction, between the positions of guest molecules in adjacent tunnels (defined in ref. [29]) is $\Delta_g = 0$.

It follows that the symmetry of the incommensurate composite material is described by the (3 + 1)-dimensional superspace group $P6_122(00\gamma)$ [28–30] and four integer indices $(h, k, l, m)$ are required to index all Bragg peaks in the diffraction pattern:

$$Q_{hklm} = h a^* + k b^* + l c^*_{\text{host}} + m c^*_{\text{guest}}. \quad (1)$$

The reflections $(h, k, l, m)$ can be subdivided into three classes according to whether $l$ and $m$ are zero or non-zero: i) $m = 0$: “main reflections” from the host substructure, which primarily contain information on the basic host structure, but also contain information on the incommensurate modulations within the guest substructure; ii) $l = 0$: “main reflections” from the guest substructure, which primarily contain information on the basic guest structure, but also contain information on the incommensurate modulations within the host substructure; iii) $l \neq 0$ and $m \neq 0$: “satellite reflections” that arise due to the inter-modulation of the two substructures. Each $(h, k, 0, 0)$ reflection is the superposition of a main reflection from each substructure, representing the common $a^*b^*$ reciprocal lattice plane for the host and guest substructures. The satellite reflections with $l \neq 0$ and $m \neq 0$ are typically very weak in comparison to the other types of reflection, and it is essential to use the high intensity of a synchrotron source in order to be able to observe and measure these satellite reflections in X-ray diffraction data.

Many techniques have been used to explore structural phase transitions in n-alkane/urea inclusion compounds [8,25,28–30,32,50–57]. Early studies reported that n-alkane/urea inclusion compounds undergo a single phase transition below ambient temperature [56] from the hexagonal phase I at ambient temperature to an orthorhombic phase II with space group $P2_12_12_1$ at low temperature. It was later suggested [58] that this phase transition is governed by a simple pseudospin-phonon model due to an antiferro-ordering of the n-alkane guests affecting both the host shearing and guest orientation. A theoretical analysis [29] described the phase transitions within the framework of a (3 + 1)-dimensional superspace group description of all possible group-subgroup related host and guest sub-lattices. It followed that phase I is described by superspace group $P6_122(00\gamma)$ and phase II is described by superspace group $P2_12_12_1(00\gamma)$. In both phases I and II, the host and guest substructures have the same periodicities in the $ab$-plane, and the misfit parameter $\gamma$ along the $c$-axis is the same. In the $ab$-plane, the unit cell of phase II closely approximates the orthorhombic description of the hexagonal unit cell of phase I (with orthorhombic lattice parameters $a_0$ and $b_0$ related by: $a_0 \approx b_0\sqrt{3}$, but with loss of C-centering).

A recent series of papers [11,36–43] has argued against this simple generalized description of the phase transi-
tions in \(n\)-alkane/urea inclusion compounds. First, for \(n\)-nonadecane/urea [11,36,38,42], in addition to the “classical” \(1\rightarrow 2\) phase transition (discussed above) at temperature \(T_1\), another phase transition at a lower temperature \(T_2\) was reported, corresponding to a weak thermal event in DSC data [59]. The new phase below \(T_2\) is denoted phase III. Furthermore, it was reported that, for phase II, no “main reflections” (i.e., \((h, k, l, m)\) with \(l = 0\) and/or \(m = 0\)) were observed for which \(h + k\) is odd, leading to the conclusion that this phase is based on a C-centred orthorhombic unit cell. Surprisingly, however, several satellite reflections (i.e., \((h, k, l, m)\) with \(l \neq 0\) and \(m \neq 0\)) were observed for values of \(h\) and \(k\) for which \(h + k\) is odd [38,39,42], implying that the C-centering is actually lost for the incommensurate inter-modulations. The positions of these satellite reflections along the \(c^*\)-axis were rationalized by introducing an additional misfit parameter \(\delta = 0.090\) that coexists with the misfit parameter \(\gamma = 0.418\) at ambient pressure [11,36,38,42] (note that the value \(\gamma = 0.428\) has been reported at the higher pressure of 0.5 GPa [38]), necessitating the description of the symmetry properties of phase II in terms of a \((3+2)\)-dimensional superspace. In phase III, reflections for which \(h + k\) is odd, attributed to the loss of C-centering of the orthorhombic unit cell, were observed both for main reflections and for satellite reflections.

These observations raise a seemingly impossible physical conundrum: how does the material undergo a phase transition from phase I to phase II involving a change in the inter-modalisation of the two substructures (i.e., the loss of C-centering) that is not reflected by a corresponding change in the basic host structure or the basic guest structure (for which C-centering is retained), particularly as it is the interactions between the two basic structures that ultimately cause the incommensurate inter-modalisations?

The \((3+2)\)-dimensional superspace groups proposed [38,42] for phase II and phase III were \(C222_1(00\gamma)\) \((108)\) and \(P2_12_12_1(00\gamma)(00\delta)\), respectively, with five independent indices \((h, k, l, m, n)\) required to index each reflection:

\[
Q_{hklmn} = h \ a^* + k \ b^* + l \ c_{\text{host}}^* + m \ c_{\text{guest}}^* + n \ c_i^*.
\]  

(2)

The indices \(m\) and \(n\) relate to \(\gamma\) and \(\delta\), respectively, with \(c_{\text{guest}}^* = \gamma \ c_{\text{host}}^*\) and \(c_i^* = \delta \ c_{\text{host}}^*\).

Similar results were later reported [39] for \(n\)-hexadecane/urea, with phase I assigned to the \((3+1)\)-dimensional superspace group \(P6_122(00\gamma)\). However, in contrast to \(n\)-nonadecane/urea, phase II was reported to exhibit main reflections (i.e., \((h, k, l, m)\) with \(l = 0\) and/or \(m = 0\)) with \(h + k\) odd, but not to exhibit satellite reflections (i.e., \((h, k, l, m)\) with \(l \neq 0\) and \(m \neq 0\)) with \(h + k\) odd, and phase II was described by the \((3+1)\)-dimensional superspace group \(P2_12_12_1(00\gamma)\) with \(\gamma = 0.486\). In phase III, many satellite reflections were observed with \(h + k\) odd and were indexed in the \((3+2)\)-dimensional superspace group \(P2_12_12_1(00\gamma)(00\delta)\), involving the additional misfit parameter \(\delta = 0.058\) coexisting with the misfit parameter \(\gamma\) along the \(c\)-axis [39].

In the present paper, we develop alternative interpretations of the superspace group descriptions and phase transitions in \(n\)-nonadecane/urea and \(n\)-hexadecane/urea, based in part on a high-resolution synchrotron single-crystal X-ray diffraction study of \(n\)-nonadecane/urea. Significantly, we demonstrate that \((3+1)\)-dimensional superspace groups are sufficient to provide a complete description of the symmetry properties of the low-temperature phases, in contrast to the unnecessarily elaborate \((3+2)\)-dimensional superspace descriptions proposed previously [11,36–42].

First, we consider the assertion [36,38,42] that a \((3+2)\)-dimensional superspace group representation is required to describe the symmetry properties of the low-temperature phases of \(n\)-nonadecane/urea (phases II and III) and \(n\)-hexadecane/urea (phase III). For \(n\)-nonadecane/urea, the temperature-independent misfit parameters were determined [36,38,42] to be \(\gamma = 0.418\) and \(\delta = 0.090\) at ambient pressure. The X-ray diffraction data for \(n\)-nonadecane/urea recorded in the present work can also be indexed on the basis of the same \((3+2)\)-dimensional description, with the same values of the misfit parameters \(\gamma\) and \(\delta\). However, on careful consideration of the indexing of reflections according to the \((3+2)\)-dimensional superspace group representation, we have recognized that a single relation actually exists between the values of the two misfit parameters \(\gamma\) and \(\delta\), specifically: \(−2 + 5\gamma = \delta\). It then follows that the reciprocal lattice basis vectors \(c_{\text{host}}^*\), \(c_{\text{guest}}^*\), and \(c_i^*\) (see eq. (2)) are actually related by

\[
c_i^* = −2c_{\text{host}}^* + 5c_{\text{guest}}^*.
\]  

(3)

On this basis, it is sufficient to describe the reciprocal space completely using a \((3+1)\)-dimensional superspace description, with the following relationships between the indices in the \((3+2)\)-dimensional \((h, k, l, m, n)\) and \((3+1)\)-dimensional \((h, k, l', m')\) superspace descriptions:

\[
l' = l − 2n\text{ and } m' = m + 5n.
\]  

(4)

In the case of \(n\)-hexadecane/urea at ambient pressure, the superspace groups assigned previously [39] to phase II and phase III were \(P2_12_12_1(00\gamma)\) and \(P2_12_12_1(00\gamma)(00\delta)\), respectively, and the temperature-independent misfit parameters were reported [39] to be \(\gamma = 0.486\) and \(\delta = 0.058\). Again, within experimental errors, we have identified a simple relation between \(\gamma\) and \(\delta\), in particular: \(2−4\gamma = \delta\). Within the \((3+2)\)-dimensional superspace description for phase III, the reciprocal lattice basis vectors \(c_{\text{host}}^*, c_{\text{guest}}^*, c_i^*\) (see eq. (2)) are actually related by

\[
c_i^* = 2c_{\text{host}}^* − 4c_{\text{guest}}^*.
\]  

(5)

Again, a \((3+1)\)-dimensional superspace description is sufficient, with the following relationships between the indices
Fig. 2: (Colour online) One-dimensional $l$-scans through reciprocal space for $n$-nonadecane/urea at 147 K (phase II) with (a) $h = 3$, $k = -2$, and (b) $h = 3$, $k = 1$. The position ($q$) along the $c^*\text{-axis}$ is given as $q = l_{\text{meas}} c^*_{\text{host}}$. The major observed reflections are labelled by the ($l'$, $m'$) indices in the ($3+1$)-dimensional superspace description. In the ($3+2$)-dimensional ($h$, $k$, $l$, $m$, $n$) and ($3+1$)-dimensional ($h$, $k$, $l'$, $m'$) superspace descriptions is specified in table 1, verifying that all reflections are correctly indexed in the ($3+1$)-dimensional superspace description.

Fig. 3: (Colour online) One-dimensional $l$-scans through reciprocal space for $n$-nonadecane/urea at 100 K (phase III) with (a) $h = -3$, $k = -2$, and (b) $h = 3$, $k = 1$. The position ($q$) along the $c^*$-axis is given as $q = l_{\text{meas}} c^*_{\text{host}}$. The major observed reflections are labelled by the ($l'$, $m'$) indices in the ($3+1$)-dimensional superspace description.

These simple relationships suggest that, for the low-temperature phases of both $n$-hexadecane/urea and $n$-nonadecane/urea, the ($3+2$)-dimensional ($h$, $k$, $l$, $m$, $n$) and ($3+1$)-dimensional ($h$, $k$, $l'$, $m'$) superspace group descriptions are not necessary.

In our single-crystal X-ray diffraction study of $n$-nonadecane/urea, data were recorded for phase II at 147 K (i.e., just below the I$\leftrightarrow$II transition temperature $T_1 = 157$ K [59]) and for phase III at 100 K (i.e., well below the II$\leftrightarrow$III transition temperature $T_2 = 140$ K [59]). Representative one-dimensional $l$-scans through reciprocal space, parallel to the $c^*_{\text{host}}$ and $c^*_{\text{guest}}$ axes, are shown for phase II in fig. 2 and for phase III in fig. 3 (all $l$-scans were determined, using standard data analysis techniques, from experimental measurements of the complete reciprocal space; see the “Experimental details” section). For the $l$-scan shown in fig. 2(a), the indexing of the reflections based on both the ($3+2$)-dimensional ($h$, $k$, $l$, $m$, $n$) and ($3+1$)-dimensional ($h$, $k$, $l'$, $m'$) superspace descriptions is specified in table 1, verifying that all reflections are correctly indexed in the ($3+1$)-dimensional superspace description.

For phase II, surprisingly, the most intense peaks in the scan with $h + k$ odd (fig. 2(a)) correspond to satellite reflections; significantly, however, weak main reflections of both ($h$, $k$, $l'$, 0) and ($h$, $k$, 0, $m'$) types are also observed. In the scan with $h + k$ even (fig. 2(b)), main reflections of both ($h$, $k$, $l'$, 0) and ($h$, $k$, 0, $m'$) types are observed, as well as satellite reflections ($h$, $k$, $l'$, $m'$) with $l' \neq 0$ and $m' \neq 0$.

For phase III, the corresponding scans with $h + k$ odd (fig. 3(a)) or $h + k$ even (fig. 3(b)) reveal a significant increase in the intensities of the main reflections of both ($h$, $k$, $l'$, 0) and ($h$, $k$, 0, $m'$) types, but there are no marked changes in the intensities of the satellite reflections.

The major difference between our data and those reported previously [36,38,42] is that, in phase II, we ob-
serve main reflections of both \((h, k, l', 0)\) and \((h, k, 0, m')\) types with \(h + k\) odd, although, as noted above, we emphasize that the main reflections with \(h + k\) odd generally have lower intensities than the satellite reflections with the same \(h\) and \(k\) values. Clearly, we conclude that, on passing from phase I to phase II, the \(C\)-centering of the orthohexagonal unit cell is lost for both the basic host structure and the basic guest structure.

Referring momentarily to the \((3 + 2)\)-dimensional superspace group description, we note that the reflection condition \([60]\) \(((h, k, l, m, n)\): \(h + k + n = 2N; N = \text{integer})\) for the superspace group \(C222_1(00\gamma)(10\delta)\) proposed previously \([38,42]\) for phase II of \(n\)-nonadecane/urea is not supported by our X-ray diffraction data. For example, the satellite reflections indexed in the \((3 + 2)\)-dimensional system as \((3, -2, 0, 0, 2)\) and \((3, -2, 0, 1, -2)\), which should be absent for superspace group \(C222_1(00\gamma)(10\delta)\), are clearly observed in our X-ray diffraction data. These reflections correspond (using the relations in eq. (4)) to \((3, -2, -4, 10)\) and \((3, -2, -4, -9)\), respectively, in the \((3 + 1)\)-dimensional description used to index the reflections in fig. 2(a).

In the frame of the \((3 + 2)\)-dimensional superspace proposed previously for \(n\)-nonadecane/urea \([38,42]\), the observed reflections in phase II are actually compatible with superspace group \(P2_12_12_1(00\gamma)(006)\) \([60]\), rather than superspace group \(C222_1(00\gamma)(10\delta)\). We note that \(P2_12_12_1(00\gamma)(006)\) is the same superspace group as phase III, and is a subgroup of \(C222_1(00\gamma)(10\delta)\) \([29]\).

However, as we have already shown that a \((3 + 1)\)-dimensional superspace is sufficient to describe the symmetry properties of these incommensurate materials, we propose that, for both \(n\)-nonadecane/urea and \(n\)-hexadecane/urea, phase II and phase III are both fully described by the \((3 + 1)\)-dimensional superspace group \(P2_12_12_1(00\gamma)\). For each material, the primary difference between phase II and phase III is that the intensities of the main reflections \(((h, k, l', 0)\) and \((h, k, 0, m')\)) and the satellite reflections \(((h, k, l', m')\) with \(l' \neq 0\) and \(m' \neq 0\)) have significantly different temperature dependences when \(h + k\) is odd.

We emphasize that our conclusions are fully corroborated by the analysis of several other one-dimensional \(l\)-scans for phases II and III of \(n\)-nonadecane/urea, in addition to those shown in figs. 2 and 3. We also note that most reflections in table I have large values of the indices \((l', m')\) in the \((3 + 1)\)-dimensional superspace description, whereas the values of the indices \((l, m, n)\) in the \((3 + 2)\)-dimensional superspace description are smaller. Examples of incommensurate systems with high indices for satellite reflections arise when the modulation function becomes discontinuous and a non-analytic “soliton regime” can exist, allowing higher harmonics to be observed \([61,62]\).

Importantly, in the present case, the modulation function is discontinuous by construction, as it is interrupted by the specific periodic \(\text{CH}_3 \cdot \text{H}_2\text{C}\) interactions that exist between the ends of adjacent \(n\)-alkane guests.

| \(l_{\text{meas}}\) | \(l_{\text{calc}}\) | \((l', m')\) | \((l, m, n)\) |
|----------------|----------------|----------------|----------------|
| 0.095          | 0.090          | \((-2, 5)\)     | \((0, 0, 1)\)  |
| 0.189          | 0.180          | \((-4, 10)\)    | \((0, 0, 2)\)  |
| 0.232          | 0.238          | \((4, -9)\)     | \((0, 1, -2)\) |
| 0.327          | 0.328          | \((2, -4)\)     | \((0, 1, -1)\) |
| 0.914          | 0.910          | \((3, -5)\)     | \((1, 0, -1)\) |
| 0.999          | 1.000          | \((1, 0)\)      | \((1, 0, 0)\)  |
| 1.093          | 1.090          | \((-1, 5)\)     | \((1, 0, 1)\)  |
| 1.913          | 1.910          | \((4, -5)\)     | \((2, 0, -1)\) |
| 2.922          | 2.926          | \((0, 7)\)      | \((2, 2, 1)\)  |
| 2.996          | 3.000          | \((3, 0)\)      | \((3, 0, 0)\)  |
| 3.090          | 3.090          | \((1, 5)\)      | \((3, 0, 1)\)  |

In contrast to our \((3 + 1)\)-dimensional superspace, we emphasize that the \((3 + 2)\)-dimensional superspace gives a huge number of possible solutions for indexing each experimentally observed reflection. For example, the reflection observed in fig. 2(a) (see table I) at \(l_{\text{meas}} = 0.914\) is indexed unambiguously and uniquely in our \((3 + 1)\)-dimensional superspace as \((3, -2, 3, -5)\), with \(l_{\text{calc}} = 0.910\), but can be indexed in the \((3 + 2)\)-dimensional superspace by any of the reflections \((3, -2, 1, 0, -1)\), \((3, -2, 3, -5, 0)\), \((3, -2, -1, 5, -2)\) or \((3, -2, 5, -10, 1)\), etc., all of which give the same value of \(l_{\text{calc}} = 0.910\). Clearly, such ambiguity and non-uniqueness in the indexing of reflections is another unsatisfactory feature of the \((3 + 2)\)-dimensional superspace description.

In conclusion, we have shown that rationalization of the diffraction patterns of \(n\)-nonadecane/urea and \(n\)-hexadecane/urea inclusion compounds in their low-temperature phases does not require the introduction of a fifth dimension in superspace, as proposed previously, and does not require new concepts, such as “hidden degrees of freedom in superspace” \([36]\), to be invented. As a consequence, the temperature-independent misfit parameter \(\gamma\) is the only aperiodicity in these materials, and, furthermore, the misfit parameter \(\gamma\) is a common feature in all phases. The most probable mechanism responsible for the phase transitions is then related to “classical” degrees of freedom of the rigid \(n\)-alkane guest molecules inside the urea tunnels \((i.e.,\) reorientational motions of the guest molecules about the host tunnel axis
Experimental details. — Crystals of the n-nonadecane/urea inclusion compound were prepared by the following method. An excess of n-nonadecane was added to a saturated solution of urea in methanol in a conical flask at 55˚C. The mixture was added dropwise until the solution was homogeneous. Any nonadecane/urea inclusion compound were prepared along the n-nonadecane/urea and n-hexadecane/urea. The development of this model will be presented in a forthcoming paper.

Single-crystal X-ray diffraction data were recorded for n-nonadecane/urea on beamline I19 (EH1) at Diamond Light Source, with temperature controlled by an Oxford Cryosystems Cryostream Plus open-flow nitrogen gas cryostat. The data were measured using a Rigaku Crystal Logic Kappa goniometer with a Saturn 724+ detector at the zirconium absorption edge (λ = 0.6889 Å). At 180 K, 147 K and 100 K, a full sphere of data to a resolution of 0.8 Å was collected with a scan width of 0.3°. At ambient temperature, only indexing frames were collected. Raw frame data were processed using CrysAlisPro from Agilent Technologies (now Rigaku Oxford Diffraction). For the data recorded at 147 K (phase II) and 100 K (phase III), after peak hunting and indexing, a cell unit with orthorhombic metric symmetry (a ≈ 14.1 Å, b ≈ 8.15 Å, c ≈ 10.99 Å) was used (with some variation in unit cell parameters at the different temperatures). From the collected data, reconstructed precession images were generated for the regions of reciprocal space of interest. To generate one-dimensional l-scans through reciprocal space along the c*-axis (as shown in figs. 2 and 3), an appropriate hkl layer in reciprocal space (with one of the values of h or k fixed) was reconstructed to a resolution of 2 Å or 3 Å. From the reconstructed layer, a line drag was performed (using CrysAlisPro) at fixed h and k, moving from positive to negative values of l, giving a continuous plot of intensity vs. l. Within these reconstructed l-scans, the spacing between adjacent digitized data points corresponds to Δl ≈ 0.01 for fig. 2(a) and Δl ≈ 0.007 for fig. 2(b), 3(a) and 3(b); thus, the experimental error in assessing the positions of peak maxima in these l-scans is of the order of Δl = ±0.005 and 1/Δl = ±0.0035, respectively.

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REFERENCES

[1] Bak P., Rep. Prog. Phys., 45 (1982) 587.
[2] Shechtman D., Blech I., Gratias D. and Cahn J. W., Phys. Rev. Lett., 53 (1984) 1951.
[3] Dubois J. M., Useful Quasicrystals (World Scientific, Singapore) 2005;
[4] Steurer W. and Sutter-Widmer D., J. Phys. Appl. Phys., 40 (2007) R229.
[5] Smith A. E., Acta Crystallogr., 5 (1952) 224.
[6] Harris K. D. M. and Thomas J. M., J. Chem. Soc. Faraday Trans., 86 (1990) 2985.
[7] Harris K. D. M., J. Solid State Chem., 106 (1993) 83.
[8] Hollingsworth M. D. and Harris K. D. M., in Comprehensive Supramolecular Chemistry, edited by Atwood A. L., Davies J. E. D., MacNicol D. D. and Vogtle F., Vol. 6 (Pergamon Press, Oxford) 1996, pp. 177–237.
[9] Guillaume F., J. Chim. Phys. Phys.-Chem. Biol., 96 (1999) 1295.
[10] Harris K. D. M., Supramol. Chem., 19 (2007) 47.
[11] Mariette C., Guérin L., Rabiller P., Ecolivet C., García-Orduna P., Bourges P., Bosak A., de Sanctis D., Hollingsworth M. D., Janssen T. and Toudic B., Phys. Rev. B, 87 (2013) 104101.
[12] Hollingsworth M. D., Brown M. E., Hillier A. C., Santarsiero B. D. and Chaney J. D., Science, 273 (1996) 1355.
[13] Lee S. O. and Harris K. D. M., Chem. Phys. Lett., 307 (1999) 327.
[14] Kelly N. E., Lee S. O. and Harris K. D. M., J. Am. Chem. Soc., 123 (2001) 12682.
[15] Palmer B. A., Harris K. D. M. and Guillaume F., Angew. Chem., Int. Ed., 49 (2010) 5096.
[16] Palmer B. A., Le Comte A., Harris K. D. M. and Guillaume F., J. Am. Chem. Soc., 135 (2013) 14512.
[17] Collins S. P., Laundy D., Harris K. D. M., Kariuki B. M., Bauer C. L., Brown S. D. and Thompson P., J. Phys.: Condens. Matter, 14 (2002) 123.
