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¹H and ¹³C chemical shift–structure effects in anhydrous β-caffeine and four caffeine–diacid cocrystals probed by solid-state NMR experiments and DFT calculations†

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By using density functional theory (DFT) calculations, we refined the H atom positions in the structures of β-caffeine (C), α-oxalic acid (OA; (COOH)₂), α-(COOH)₂·2H₂O, β-malonic acid (MA), β-glutaric acid (GA), and L-maleic acid (ME), along with their corresponding cocrystals of 2:1 (2C–OA, 2C–MA) or 1:1 (C–GA, C–ME) stoichiometry. The corresponding ¹³C/¹H chemical shifts obtained by gauge including projector augmented wave (GIPAW) calculations agreed overall very well with results from magic-angle-spinning (MAS) nuclear magnetic resonance (NMR) spectroscopy experiments. Chemical-shift/structure trends of the precursors and cocrystals were examined, where good linear correlations resulted for all COO¹H sites against the H···O and/or H···N H-bond distance, whereas a general correlation was neither found for the aliphatic/caffeine-stemming ¹H sites nor any ¹³C chemical shift against either the intermolecular hydrogen- or tetrel-bond distance, except for the ¹³COOH sites of the 2C–OA, 2C–MA, and C–GA cocrystals, which are involved in a strong COOH···N bond with caffeine that is responsible for the main supramolecular stabilization of the cocrystal. We provide the first complete ¹³C NMR spectral assignment of the structurally disordered anhydrous β-caffeine polymorph. The results are discussed in relation to previous literature on the disordered α-caffeine polymorph and the ordered hydrated counterpart, along with recommendations for NMR experimentation that will secure sufficient ¹³C signal-resolution for reliable resonance/site assignments.

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1 Introduction

Efficient administration of active pharmaceutical ingredients (APIs) is often challenging because the powders are either unstable or feature a low solubility in water/body fluids. One option for improving the solubility and/or other physical or chemical properties without recourse to ionic salts is the usage of a *cocrystal*, i.e., a supramolecular compound of an API and another molecular species (or even another API, a so-called drug–drug cocrystal).^{1–3} Consequently, intense efforts have been spent during the past two decades to explore synthetic routes to cocrystal preparation as well as their structures and formation mechanisms.^{2–5}

One widely used API is the stimulant caffeine (1,3,7-trimethylpurine-2,6-dione), whose storage and precise quantitative (physiological) administration is complicated by its

highly hygroscopic nature.^{6–8} At room temperature (RT) in the absence of humidity, anhydrous caffeine exists as the low-temperature “β” modification, which at temperatures >150 °C converts into the metastable high-temperature “α” form, which transforms very slowly into β-caffeine at RT and dry air.⁹ Hence, commercial anhydrous caffeine powders often comprise a mixture of both modifications. Caffeine also exists in a monohydrate form, which unfortunately is very unstable at RT and converts either into phases with <1 water molecule/caffeine moiety or anhydrous β-caffeine (depending on the relative humidity).^{7,9} Hence, large-scale storage, processing and other handling of caffeine powder with a well-defined stoichiometry is not trivial.

Yet, as demonstrated by Trask *et al.*,⁸ cocrystals of caffeine with diacids, such as oxalic acid (OA), malonic acid (MA), glutaric acid (GA), and maleic acid (ME), are stable across wide humidity ranges. Several studies are reported on their formation, stability, and physical properties.^{8,10–14} The present investigation targeted the cocrystal structures obtained from β-caffeine (β-C) with each of OA, MA, GA, and ME, abbreviated as 2C–OA, 2C–MA, C–GA, and C–ME, respectively, according to the corresponding 2:1 and 1:1 caffeine:diacid stoichiometry.

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Although a fixed stoichiometry would be desirable, the C-ME cocrystal is metastable,¹⁰ while 2C-OA, 2C-MA, and C-GA cocrystals were hitherto not reported. Each X-ray diffraction (XRD) derived cocrystal structure is reported,⁸ encompassing the main caffeine-diacid motifs governing their supramolecular organization *via* hydrogen bonds (Section 3). The well-known limitation of XRD to locate precise H-atom positions, however, restricts its capacity of unveiling quantitative details on weak non-covalent interactions, such as H···O/N hydrogen and C···O/N tetrel bonds that account for the intermolecular interactions and structural organization of cocrystals, as well as numerous other supramolecular systems. Then, high-resolution magic-angle-spinning (MAS) ¹H and ¹³C nuclear magnetic resonance (NMR) spectroscopy offer valuable complementary structural insight,^{15–20} in particular for structurally disordered (in)organic phases,^{18,20,21} encompassing β-caffeine.^{14,22–24}

Herein, we present a comprehensive combined MAS NMR and density functional theory (DFT) study, reporting on DFT refinements of previous XRD-derived structures of each 2C-OA, 2C-MA, C-GA, and C-ME cocrystal⁸ together with each β-caffeine,⁹ α-OA,²⁵ α-OA·2H₂O,²⁶ β-MA,²⁷ β-GA,²⁸ and I-ME²⁹ precursor phase. ¹H and ¹³C isotropic chemical shifts calculated by the gauge including projector augmented wave (GIPAW) method^{30–33} were contrasted with those from MAS NMR experiments, aiming at improved insight into the shift-structure trends, notably the *hydrogen bond* (HB)^{34–46}—and for the methyl moieties also *tetrel bond* (TB)^{47–49}—effects on the chemical shifts. While the complete sets of ¹³C chemical shifts/NMR-peak assignments of the 2C-MA and C-GA cocrystals were reported earlier by Vigilante and Mehta,⁵⁰ out of the four cocrystals considered herein, a majority of the ¹H chemical shifts are reported for the first time, along with nearly half of their ¹³C counterparts, such as those for 2C-OA and C-ME. We also present the first complete ¹³C NMR spectral assignment of the ten resonances from the eight unique ¹³C sites of the structurally disordered anhydrous β-caffeine polymorph. The results are discussed in relation to previous ¹³C MAS NMR reports on the ordered hydrated and anhydrous disordered α/β-caffeine and polymorphs, highlighting the role of the magnetic field for unambiguously discriminating between the α- and β-caffeine forms.

2 Materials and Methods

2.1 Cocrystal preparation

Anhydrous β-caffeine (98% purity), oxalic acid (98%), malonic acid (99%), glutaric acid (99%) and chloroform were obtained from Sigma-Aldrich along with maleic acid (99%) from Thermo Fisher. All cocrystals were prepared by solvent drop grinding,^{8,13} using ball milling (Mixer Mill 500 Vario; Retsch) of the two precursor powders that were placed in a 25 mL stainless steel jar and ground with 7 mm stainless steel balls at a rate of 30 Hz for 90 min. All cocrystals were prepared from stoichiometric amounts of the precursors. The 2C-OA (space group *P*₂₁/*c*) and

2C-MA (*Fdd*2) cocrystals⁸ with a 2 : 1 caffeine : diacid stoichiometry resulted by mixing 250 mg of β-caffeine with either 60 mg of α-OA or 70 mg of β-MA, respectively, along with 5 drops of chloroform. Likewise, the 1 : 1 stoichiometric C-GA (form II;⁸ space group *P* $\bar{1}$) cocrystal was prepared by grinding 510 mg of β-caffeine with 350 mg of β-GA and a few drops of chloroform, and likewise for the C-ME (*P*₂₁/*n*) counterpart (250 mg of β-MA and 150 mg of I-ME).⁸

2.2 Powder X-ray diffraction

Powder XRD (PXRD) patterns were collected from all cocrystals and precursors with a Bruker D8 Discover diffractometer equipped with a LYNXEYE position-sensitive detector, using Cu Kα_{1,2} radiation ($\lambda_1 = 154.06$ pm; $\lambda_2 = 154.44$ pm) and a focusing Göbel mirror. Each powder was filled in a 0.7 mm borosilicate glass capillary. All diffractograms were collected at RT over a 2θ range of 5°–50°, employing a step size of 0.02° with either 0.25 s per step (for β-caffeine, OA, β-MA, β-GA, I-ME, 2C-MA, and C-ME) or 0.75 s per step (for 2C-OA and C-GA), giving total measurement times of around 10 and 30 min per sample, respectively. The XRD patterns of the precursors and cocrystals are shown in Fig. S1 and S2 (ESI†), respectively. Rietveld refinements and Pawley fits, performed with the TOPAS software,⁵¹ verified a near-100% purity of all specimens, except for the oxalic acid powder, which comprised a mixture of α-OA (96.1 wt%), α-OA·2H₂O (3.5%), and β-OA (0.39 wt%). The anhydrous β-caffeine purity was ≥99.6%, along with a (statistically unascertained) minute amount of α-caffeine (0.37 wt%). All refined unit cell parameters are listed in Table S1 (ESI†).

2.3 Solid-state NMR

All solid-state NMR experiments were performed at ambient temperature with a Bruker Avance-III spectrometer and a magnetic field (*B*₀) of 14.1 T, which provided the respective ¹H and ¹³C Larmor frequencies of −600.1 MHz and −150.9 MHz. ¹H and ¹³C chemical shifts are quoted relative to neat tetramethylsilane (TMS).

¹H → ¹³C cross polarization (CP) MAS NMR (CPMAS) experimentation was performed with filled 4.0 mm zirconia rotors spinning at the MAS rate (ν_r) of 12.00 kHz, using contact periods (τ_{CP}) of 1.25 ms (except for β-MA; 833 μs and $\nu_r = 9.00$ kHz), and the modified Hartmann-Hahn condition $\nu_H = \nu_C + \nu_r$, where the ¹³C nutation frequency (ν_C) was ramped linearly⁵² by ±4.7 kHz around $\nu_C = 47$ kHz. Throughout, the 90° ¹H pulse prior to CP operated at $\nu_H \approx 100$ kHz and SPINAL-64⁵³ proton decoupling at $\nu_H \approx 78$ kHz (6.4 μs pulses) was applied during signal detection. The relaxation delays (τ_{relax}) varied between 15 s and 30 s. The signal averaging involved 2048–3328 co-added signal transients, except for MA and GA (1024). Single-pulse (“Bloch decay”) ¹H MAS NMR spectra were recorded with 1.3 mm zirconia rotors undergoing fast MAS at 60.00 kHz with 90° rf pulses operating at the ¹H nutation frequency $\nu_H \approx 80$ kHz. 256–512 accumulated NMR-signal transients were recorded with $\tau_{relax} = 3$ s. These relaxation delays were not sufficiently long to ensure NMR intensities



that quantitatively reflect the relative H/C site populations in the structures but do not affect any conclusions from our analysis, which only concerned the chemical shifts. NMR spectral deconvolutions were performed with in-house developed software.⁵⁴

2.4 DFT/GIPAW calculations

DFT energy optimizations were performed on the following published PXRD-derived structures: β -caffeine;⁹ α -OA;²⁵ α -OA·2H₂O;²⁶ β -MA;²⁷ β -GA;²⁸ I-ME,²⁹ whereas all cocrystal structures were from ref. 8; see Table S2 (ESI†). The first-principle DFT energy minimizations were performed with the CASTEP software⁵⁵ (version 22.11) along current standard and well-developed protocols,^{19,32,33} encompassing usage of the local density approximation (LDA) functional⁵⁶ with on-the-fly-generated ultrasoft pseudopotentials⁵⁷ and a plane-wave basis set.⁵⁸ The Tkatchenko and Scheffler method was employed for dispersion corrections.⁵⁹ All structures were initially optimized by solely adjusting the proton positions with fixed positions of all heavier atoms and unit-cell parameters. However, the agreement between the experimental and GIPAW-derived chemical shifts improved slightly for the β -caffeine structure by optimizing *all* cell parameters and atom positions (Table S2 and Section 4.3, ESI†). The ¹H and ¹³C magnetic shielding values were calculated with the GIPAW method^{19,30–33} for all DFT-optimized structures. For both the DFT energy optimizations and the GIPAW shielding-parameter calculations, a Monkhorst–Pack *k*-point grid⁶⁰ was used with a maximum spacing of 0.05 Å^{−1} in the reciprocal space, and a 1000 eV plane-wave energy cutoff to ensure convergence.

The DFT/GIPAW-derived principal values, $\{\sigma_{xx}^{Hj}, \sigma_{yy}^{Hj}, \sigma_{zz}^{Hj}\}$ and $\{\sigma_{xx}^{Cj}, \sigma_{yy}^{Cj}, \sigma_{zz}^{Cj}\}$, of the respective second-rank magnetic shielding tensor of each unique ¹H_j and ¹³C_j site in the structure were converted into the corresponding chemical shift values, $\{\delta_{xx}^{Sj}, \delta_{yy}^{Sj}, \delta_{zz}^{Sj}\}$, by the expression^{19,32,33}

$$\delta_{\alpha\alpha}^{Sj} = \sigma_{\text{ref}}^S - \sigma_{\alpha\alpha}^{Sj}, \quad \text{with } \alpha\alpha = \{xx, yy, zz\} \text{ and } S = \{^1\text{H}, ^{13}\text{C}\}. \quad (1)$$

We employ a chemical shift scale throughout, where low (high) chemical shifts correspond to shielded (deshielded) nuclei, and the isotropic chemical shift is given by^{61–63}

$$\delta_{\text{S}}^j \equiv \delta_{\text{S}}^{\text{iso},j} = \frac{1}{3} \left(\delta_{xx}^{Sj} + \delta_{yy}^{Sj} + \delta_{zz}^{Sj} \right) \quad \text{for } S = \{^1\text{H}, ^{13}\text{C}\}. \quad (2)$$

The shielding-to-shift conversion terms of eqn (1) were $\sigma_{\text{ref}}^{\text{H}} = 29.034$ ppm for ¹H and $\sigma_{\text{ref}}^{\text{C}} = 169.837$ ppm for ¹³C. The values were obtained by linear regressions that minimized the difference between the sets of calculated and experimental isotropic ¹H and ¹³C chemical shifts based on roughly half of each entire set of $\{\delta_{\text{H}}^j\}$ and $\{\delta_{\text{C}}^j\}$ values, and giving correlation coefficients (*R*²) of *R*² = 0.980 and *R*² = 0.999 when evaluated across each entire $\{\delta_{\text{H}}^j\}$ and $\{\delta_{\text{C}}^j\}$ ensemble, respectively.

3 Overview of cocrystals and their precursor structures

Fig. 1 shows the structures of caffeine and the four diacid molecules, where each number represents the ¹³C site index, giving the label “C_j” with 1 ≤ *j* ≤ 22, while all its directly bonded H, O, and N atoms carry the same index *j*. Note that although the C14/C18 sites are crystallographically equivalent in the β -GA structure, they are *inequivalent* in the C-GA cocrystal. For simplicity, we employ a strict C1–C22 labelling because all C/H sites are crystallographically distinct in either/both the diacid/cocrystal structure, except for C9/C10 (H9/H10), which remain equivalent in each α -OA, α -OA·2H₂O, and 2C–OA structure.

Fragments from the by DFT energy-minimized crystal structures (Section 2.4) are shown in Fig. 2 for the diacids (a, b, d, f, h) and their respective cocrystals with caffeine (c, e, g, i). All

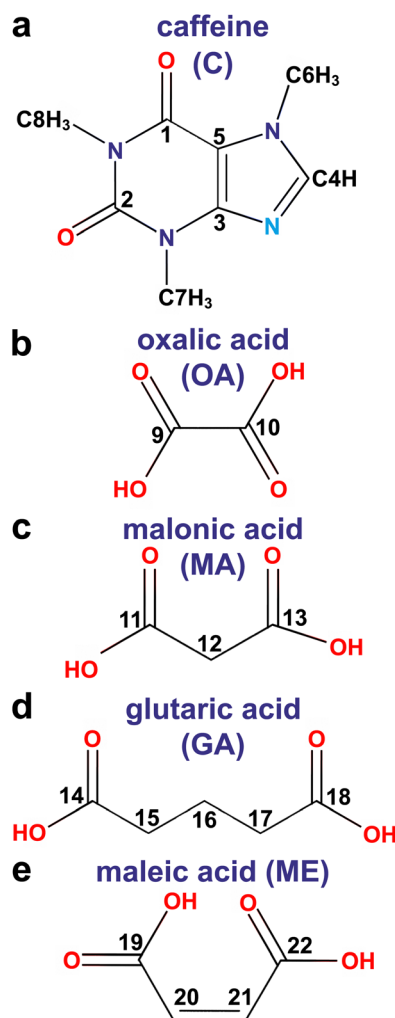


Fig. 1 Molecular structure with atom numbering of (a) caffeine, and the (b) oxalic, (c) malonic, (d) glutaric, and (e) maleic diacids. Each label/index *j* refers to carbon atom “C_j”, whose directly bonded O, N, and H atoms feature the same label (O_j, N_j, and H_j, respectively), except for the nitrogen atom of the imidazolium ring (cyan color) in (a), which is referred to simply as “N”.



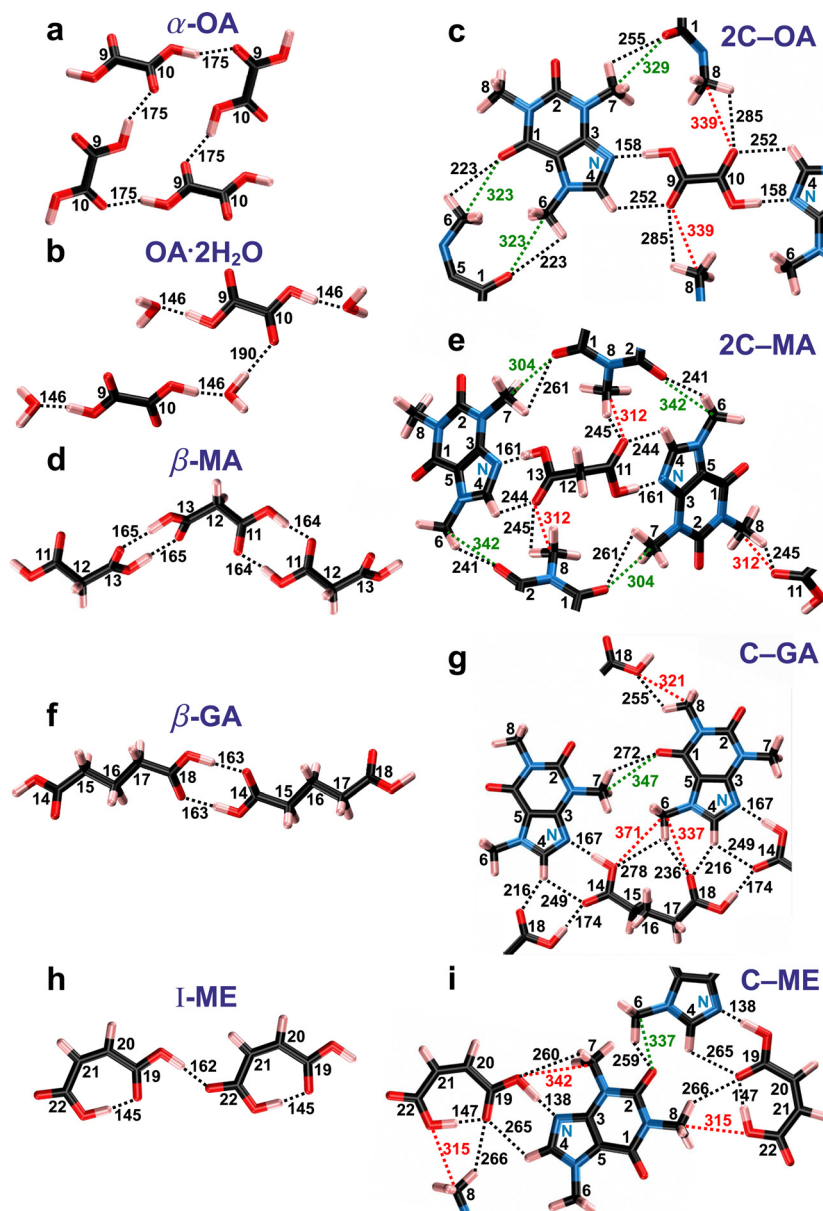


Fig. 2 Fragments from the DFT-refined crystal structures of (a) α -OA, (b) α -OA·2H₂O, (c) 2C-OA, (d) β -MA, (e) 2C-MA, (f) β -GA, (g) C-GA, (h) I-ME, and (i) C-ME. The dotted lines mark intermolecular H and tetrel bonds, with the accompanying number specifying the distance (in pm). The red and green lines/numbers distinguish tetrel bonds between caffeine-diacid and caffeine-caffeine molecules, respectively. Note that each methyl group undergoes a rapid rotation around the C–N bond and that only one H bond is indicated for each of the three equivalent protons.

pristine diacid structures involve strong intermolecular H bonds between the carboxy groups of neighboring molecules, except for I-ME, which besides one expected C19OOH···O22 contact also features a short *intramolecular* C22OOH···O19 HB (Fig. 2h), which remains intact also in the C-ME cocrystal shown in Fig. 2i, within a minute (2 pm) lengthening.

As discussed by Trask *et al.*,⁸ both 2C-OA and 2C-MA cocrystals feature a heteromeric synthon where one diacid molecule is sandwiched between two caffeine units by a strong COOH···N hydrogen bond and a weaker C4H···OOC counterpart (Fig. 2c and e). The alternating caffeine-diacid-caffeine moieties of the supramolecular structure are stabilized

primarily by those H bonds.⁸ The 1:1 caffeine:diacid stoichiometries of C-GA and C-ME, however, yield different intermolecular interactions. Owing to the intact *intramolecular* C22H···O19 bond in the C-ME cocrystal (Fig. 2i), its structure involves strictly alternating caffeine-ME interconnections, with a similar C4H···O19OC and C19OOH···N bond constellation as in the 2C-OA/MA cocrystals, yet with only *one* diacid-COOH group H-bonded to caffeine. Moreover, while the C-GA structure is also stabilized by two N···H14OOC and C4H···O14OC bonds between caffeine and one GA molecule (Fig. 2g), the C4H atom *also* involves another HB with O18 of a second diacid molecule, where moreover C18H is bonded to O14 of the first



diacid unit. Hence, each caffeine molecule in C-GA features three different H bonds to two distinct GA molecules, both of which are additionally interlinked by a HB between their carboxy moieties, as in the parent GA structure.

Although the N $\cdots$ H bond between caffeine and a COOH group of a diacid unit constitutes the strongest intermolecular contact of the cocrystal,⁸ a further minor structure stabilization is arranged by H bonds between the CH₃ groups of caffeine with *either* diacid carboxy moieties *or* the O1/O2 atoms of neighboring caffeine molecules (Fig. 2c, e, g and i). The CH₃ moieties may furthermore form weak CH₃ $\cdots$ O tetrel bonds^{47–49} with the O1/O2 atoms of caffeine, or with the CO counterparts of the diacids, as marked in green and red color, respectively, in Fig. 2c, e, g and i (see Section 4.7). The methyl groups also form intermolecular tetrel and H bonds in the structurally disordered β -caffeine structure (Table S3, ESI[†]). We refer to ref. 9 and 22 for details about the anhydrous β -caffeine structure, whose unit cell comprises five inequivalent molecules, and thereby five NMR signal-contributions to each detected ¹³Cj and ¹Hj resonance.

4 Results and discussion

4.1. ¹³C MAS NMR spectra from cocrystal precursors

Because the chemical shift of a given ¹³C or ¹H nuclear site reflects its electronic environment in the structure, it is often a sensitive probe of the precise location of neighboring atoms in close vicinity. Fig. 3 displays the ¹³C CPMAS NMR spectra recorded from the diacids and β -caffeine cocrystal precursors. They involve resonances ranging from the most deshielded ¹³C nuclei (high chemical shifts) of carboxy groups with $\delta_C \geq 160$ ppm from the diacids (Fig. 3a–d) to the more shielded (lower shifts) aliphatic ¹³C sites with chemical shifts $\delta_C \leq 40$ ppm from β -MA, β -GA, and β -caffeine (Fig. 3b, c and e). The NMR spectral region intermediate of these extreme ¹³C shifts encompasses the comparatively deshielded ¹³C20 and ¹³C21 sites of I-ME (Fig. 3d), together with the ¹³C1–¹³C5 sites of β -caffeine resonating between 105–155 ppm (Fig. 3e).

All cocrystal precursor powders (but OA) were phase pure, where the two crystallographically inequivalent ¹³COOH sites of MA and ME reveal chemical-shift differences of 0.4 ppm and 3.7 ppm, respectively, while both ¹³COOH sites of GA resonate at 181.5 ppm. In contrast, the two ¹³C NMR signals at 160.7 ppm and 163.1 ppm from OA derive from anhydrous α -OA and α -OA $\cdot$ 2H₂O, respectively, each featuring one crystallographically unique ¹³COOH site.^{25,26} From the integrated ¹³C NMR intensities of Fig. 3a, and confirmed by integrating the corresponding ¹H MAS NMR spectrum (Section 4.5.1), we estimated relative OA:OA $\cdot$ 2H₂O amounts of approximately 0.7:1.0 in the oxalic acid mixture. (Although that estimate is only approximate, its precise value is immaterial for our subsequent analyses. The reason for the markedly higher OA $\cdot$ 2H₂O content derived by NMR relative to PXRD is unknown but likely stems from a water uptake of the OA powder prior to the NMR experiments). The ¹³C chemical shifts obtained from Fig. 3

agree well with previous reports from α -OA,^{36,64} α -OA $\cdot$ 2H₂O,³⁶ MA,^{14,65} GA,^{36,65,66} and ME.^{65–67}

4.2. ¹³C NMR-peak assignments of anhydrous β -caffeine

Notwithstanding several ¹³C NMR publications on caffeine-based cocrystals,^{23,24,41,46,49,50} encompassing complete $\{\delta_C\}$ reports thereof,^{23,50} ambiguities prevail about the precise set of ¹³C shifts and their assignments of the anhydrous β -caffeine structure *itself*, which produces two hitherto unassigned ¹³C resonances, none of which are present in MAS NMR spectra from either the structurally ordered caffeine monohydrate (henceforth referred to as C $\cdot$ H₂O) or disordered anhydrous α -caffeine modifications,²² each of which manifests one ¹³C resonance per C1–C8 site. Enright *et al.*²² presented ¹³C MAS NMR spectra from all α / β -caffeine and C $\cdot$ H₂O forms, but precise chemical shifts were only reported for the latter.²² Likewise, δ_C values were not provided along with the ¹³C MAS NMR spectra from the anhydrous caffeine modifications studied in ref. 12, 24, 50 and 68. To the best of our knowledge, Table 1 presents the precise ¹³C chemical shifts *and* complete peak assignments for the anhydrous β -caffeine polymorph for the first time, where all further references herein to “ α / β -caffeine” imply the anhydrous polymorphs. The ¹³C MAS NMR spectrum from β -caffeine of Fig. 3e appears to match very well that reported in ref. 22 while moreover our PXRD analysis confirmed a phase-pure specimen (Section 2.2). Notwithstanding that all eight $\{\delta_C^{21}\}$ values of both α / β -caffeine modifications are (very) similar,²² the presence of two additional ¹³C NMR peaks at 149 ppm and 30 ppm in β -caffeine (Fig. 3e) distinguishes it from its “ α ” and C $\cdot$ H₂O counterparts, which Enright *et al.*²² attributed to ¹³C sites of crystallographically distinct molecules in the β -caffeine unit cell. That very plausible suggestion is confirmed below.

Previous solid-state ¹³C NMR-peak assignments of all anhydrous and monohydrate caffeine polymorphs derive either from (i) employing DFT/GIPAW calculations of a caffeine-based cocrystal to assign its ¹³C sites of the caffeine moiety^{23,24,50} or from (ii) the solution-NMR work of Sitkowski *et al.*,⁶⁹ which was exploited by Enright *et al.*²² for the ¹³C-site/NMR-peak identifications of the ordered C $\cdot$ H₂O modification in the solid state (which is greatly facilitated by its narrow ¹³C resonances). Those peak assignments were subsequently assumed in ref. 14 and 70. Although such straightforward ¹³C site/shift-mappings are also sufficiently reliable for application to α -caffeine, which involves one resonance per ¹³C site, the peak-assignment strategies (i) and (ii) are precluded for the two additional resonances at around 149/30 ppm from β -caffeine and labelled as C3'/C78 in Fig. 3e. To the best of our knowledge, both remain unidentified in current literature. Nonetheless, for each ¹³C1–¹³C8 resonance assignment of Fig. 3e, the DFT/GIPAW-derived ¹³C chemical shifts of Table 1 corroborate previously employed NMR-peak assignments made for both C $\cdot$ H₂O and α -caffeine modifications.^{14,22–24,70} The modeled chemical shifts, which were averaged over the 5 distinct caffeine molecules in the unit cell, reproduce the ¹³C NMR results from β -caffeine (very) well (Table 1), which for every asymmetric peakshape represents the *center-of-gravity* (CG) shift ($\bar{\delta}_C$) obtained from NMR-peak deconvolutions into two components



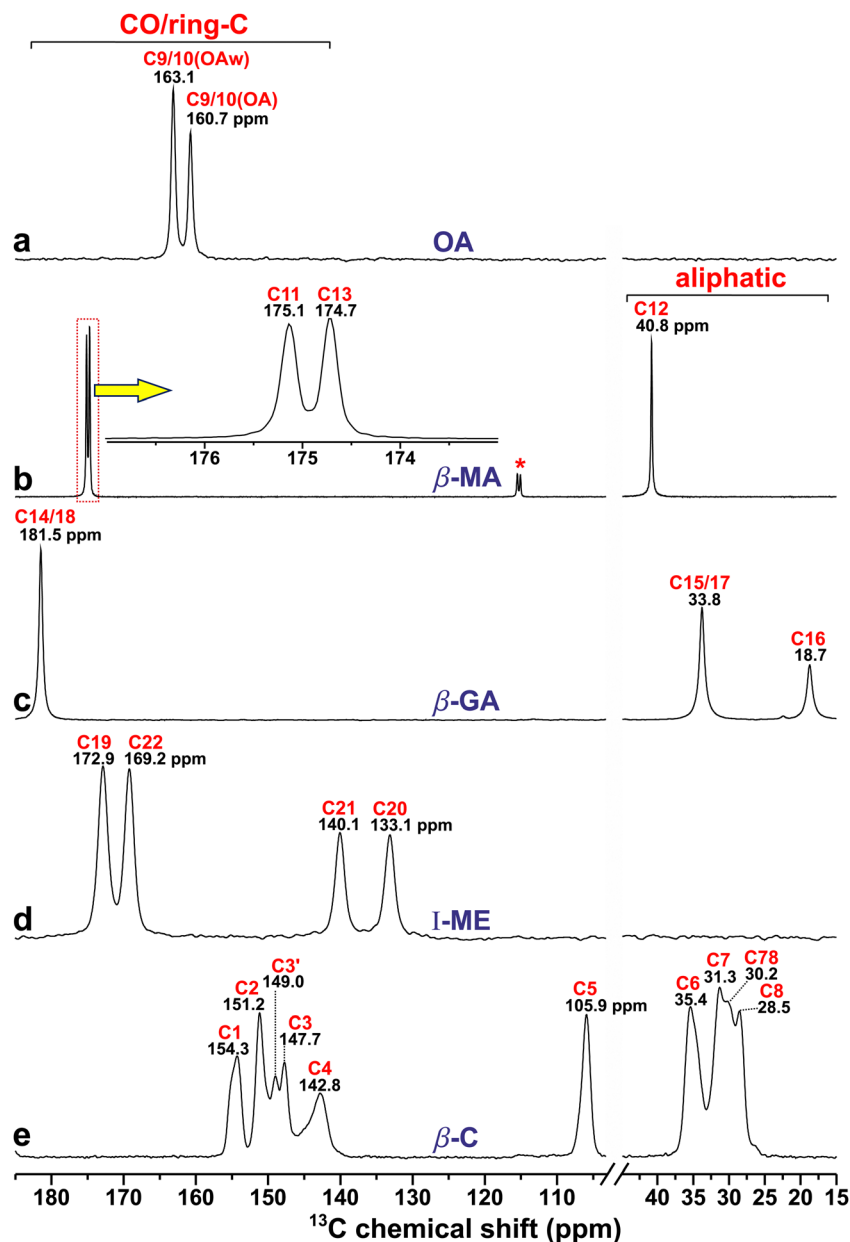


Fig. 3 ^{13}C CPMAS NMR spectra recorded at 14.1 T and a MAS rate of 12.00 kHz from (a) α -OA/ α -OA $\cdot$ 2H $_2$ O mixture ("OA"), (b) malonic acid (β -MA), (c) glutaric acid (β -GA), (d) maleic acid (I-ME), and (e) anhydrous β -caffeine (β -C). Here and in other figures: NMR signals from the α -OA and α -OA $\cdot$ 2H $_2$ O phases are abbreviated by "OA" and "OAw", respectively, whereas the red and black number above each NMR peak marks the C label of Fig. 1 and the chemical shift (in ppm) at the peak maximum, respectively. The inset of (b) is a zoom of the NMR spectral region marked by the dotted rectangle, and the asterisk marks a spinning sideband from the use of a lower MAS rate (9.00 kHz) for this experiment.

(Fig. S3, ESI †) rather than the shift at the peak-maximum amplitude (δ_C^i). Note that (I) $\bar{\delta}_C^i$ and δ_C^i represent the *average* and the *most probable* value over the chemical-shift distribution of $^{13}\text{C}_j$, respectively; (II) invoking CG shifts is only required for the structurally disordered β -caffeine structure, as opposed to those of any cocrystal or diacid precursor considered herein, for which $\bar{\delta}_C^i = \delta_C^i$ throughout.

Despite an overall good agreement between the experimental and modeled ^{13}C chemical shifts of β -caffeine, the DFT/GIPAW calculations alone *cannot uniquely* identify the hitherto unassigned resonances at 149.0 ppm and 30.2 ppm

(Fig. 3e). By spectral deconvolution, however, the integrated ^{13}C NMR-peak intensities at around {154.3, 151.2, 149.0, 147.7} ppm were found to relate as 1.00:1.44:0.27:0.74 (Fig. S3, ESI †). That observation naturally attributes the resonance at 149.0 ppm to C3 (it is therefore referred to as C3' in Fig. 3e), leading to $\bar{\delta}_C^i$ values of {154.5(C1), 150.8(C2), 148.0(C3)} ppm with relative intensities 1.00:1.44:1.01. The deviations from the expected exact 1:1:1 ratios (where the discrepancy is only significant for C2) mainly reflect that integrated ^{13}C NMR-peak intensities obtained by $^1\text{H} \rightarrow ^{13}\text{C}$ CPMAS NMR experiments do generally not exactly reproduce the corresponding ^{13}C site

Table 1 NMR/DFT-derived ^{13}C chemical shifts of β -caffeine^a

Site	$\delta_{\text{C}}^{\text{[NMR]}}$ (ppm)	$\bar{\delta}_{\text{C}}^{\text{[NMR]}}$ (ppm)	$\bar{\delta}_{\text{C}}^{\text{[DFT]}}$ (ppm)	$\delta_{\text{C}}^{\text{[DFT]}}$ (ppm)				
C1	154.3	154.5 $\pm$ 0.2	153.0 $\pm$ 0.3	152.5	152.7	152.8	152.9	154.1
C2	151.2	150.8 $\pm$ 0.2	149.4 $\pm$ 0.3	148.3	148.9	149.5	149.7	150.5
C3 ^b	149.0/147.7	148.0 $\pm$ 0.2	147.5 $\pm$ 0.6	146.5	146.7	147.4	147.7	149.4
C4	142.8	143.5 $\pm$ 0.4	146.0 $\pm$ 1.3	142.9	143.9	145.0	148.9	149.3
C5	105.9	106.1 $\pm$ 0.2	107.8 $\pm$ 0.6	106.2	106.9	107.9	108.1	109.7
C6	35.4	35.0 $\pm$ 0.2	35.3 $\pm$ 0.6	33.7	34.8	35.4	35.7	36.7
C7 ^c	31.3/30.2	30.7 $\pm$ 0.2	29.6 $\pm$ 0.9	28.0	28.3	29.3	30.5	32.2
C8 ^c	30.2/28.5	29.6 $\pm$ 0.2	29.5 $\pm$ 0.6	27.7	28.6	29.2	29.9	32.0

^a ^{13}C chemical shift at the NMR peak maximum, $\delta_{\text{C}}^{\text{[NMR]}}$, along with the center-of-gravity (average) chemical shift obtained by deconvolution of the MAS NMR spectrum (Fig. S3, ESI), $\bar{\delta}_{\text{C}}^{\text{[NMR]}}$, or by DFT calculations, $\bar{\delta}_{\text{C}}^{\text{[DFT]}}$; the latter values are averages over the ^{13}C chemical shift values of five distinct caffeine molecules in the unit cell, each of which is given in the five rightmost columns. ^b The C3 sites produces two resonances at 149.0 ppm and 147.7 ppm, which are labelled by “C3'” and “C3” in Fig. 3e. ^c The “C78” ^{13}C resonance intensity at 30.2 ppm (Fig. 3e) involves equal contributions from the $^{13}\text{C7}$ and $^{13}\text{C8}$ sites.

populations in the structure. Likewise, the $^{13}\text{CH}_3$ -associated resonance at $\bar{\delta}_{\text{C}} = 30.2$ ppm, which is labelled “C78” in Fig. 3e, was deduced to involve equal contributions from $^{13}\text{C7}$ and $^{13}\text{C8}$. Upon distributing half of the NMR-signal intensity at $\bar{\delta}_{\text{C}} = 30.2$ ppm to each of $^{13}\text{C7}$ ($\bar{\delta}_{\text{C}} = 31.3$ ppm) and $^{13}\text{C8}$ ($\bar{\delta}_{\text{C}} = 28.5$ ppm), the as-obtained relative integrated NMR-signal intensities of the C6 : C7 : C8 sites become 1.00 : 1.03 : 0.96, which is in excellent agreement with the expected 1 : 1 : 1 ratios (Fig. S3, ESI†).

Further support for the spectral-deconvolution-derived attribution of the “C3'” and “C78” NMR peaks to the $^{13}\text{C3}$ site and equal contributions to $^{13}\text{C7}/^{13}\text{C8}$, respectively, is provided by the resulting very close $\{\bar{\delta}_{\text{C}}^3, \bar{\delta}_{\text{C}}^7, \bar{\delta}_{\text{C}}^8\} = \{148.0, 30.7, 29.6\}$ ppm values observed relative to the $\{\bar{\delta}_{\text{C}}^3, \bar{\delta}_{\text{C}}^7, \bar{\delta}_{\text{C}}^8\} = \{147.8, 30.6, 29.0\}$ ppm counterparts of the structurally ordered C-H₂O polymorph.²² The excellent agreement between the average ^{13}C chemical shifts (*i.e.*, $\bar{\delta}_{\text{C}}^{\text{[NMR]}}$) of the disordered β -caffeine structure obtained by “down-projecting” its 10 ^{13}C NMR peaks to the unique 8 ^{13}C resonance-mapping of the C-H₂O polymorph²² is very gratifying. Moreover, almost all $\bar{\delta}_{\text{C}}^{\text{[NMR]}}/\bar{\delta}_{\text{C}}^{\text{[DFT]}}$ discrepancies between β -caffeine and C-H₂O are well within the experimental uncertainties. For instance, the $\{\bar{\delta}_{\text{C}}^1, \bar{\delta}_{\text{C}}^5, \bar{\delta}_{\text{C}}^6\}$ values of Table 1 agree within 0.2 ppm with the $\{\bar{\delta}_{\text{C}}^1, \bar{\delta}_{\text{C}}^5, \bar{\delta}_{\text{C}}^6\}$ counterparts of C-H₂O.²² Significant chemical-shift differences between the two polymorphs are *only* observed for the $^{13}\text{C2}$ and $^{13}\text{C4}$ carbonyl sites, both of which are ≈ 1.0 ppm higher for β -caffeine than those of C-H₂O, which is attributed to distinct HB scenarios between the two structures. DFT calculations (not shown) reveal that the crystallographically unique C4H site of C-H₂O involves one C4H...OH₂ HB (201 pm), whereas the five distinct H4 sites of β -caffeine feature a range of HB distances to the O1 (198; 235; 250 pm), O2 (212 pm), and N (236 pm) sites of neighboring molecules. They contribute strongly to the “local” disorder of the $^{13}\text{C4}$ environment, and thereby to its sizable fwhm value relative to any other ^{13}Cj site of β -caffeine, while also explaining the accompanying larger $\bar{\delta}_{\text{C}}^4 - \bar{\delta}_{\text{C}}^{\text{[DFT]}}$ difference (Table 1).

4.3 NMR/GIPAW-derived ^{13}C chemical shifts of cocrystals and precursors

Fig. 4 contrasts the ^{13}C CPMAS NMR spectra recorded from β -caffeine and its four cocrystals. The successful completion of each cocrystal reaction is evidenced both from our PXRD

analyses (Section 2.2) and by the unique ^{13}Cj site/chemical-shift mapping resulting for all cocrystals, for which $\bar{\delta}_{\text{C}}^{\text{[NMR]}}$ of each $^{13}\text{C1} - ^{13}\text{C22}$ site differs only slightly relative to that of its precursor phases (Table 2). In particular, each of the 8 caffeine-moiety-related ^{13}C resonances is readily identified for each cocrystal, meaning that no cocrystal manifests NMR peaks traceable to either “C3'” or “C78” of Fig. 4e, thereby eliminating all NMR-peak assignment ambiguities of β -caffeine. Hence, all ^{13}C resonance identifications for each cocrystal (Fig. 4) readily follow from those established for its parent phases (Fig. 3), as corroborated further by contrasting the experimental and DFT/GIPAW-generated $\{\bar{\delta}_{\text{C}}^{\text{[NMR]}}\}$ values listed in Table 2.

While hitherto no report on any *even partially complete* set of $\{\bar{\delta}_{\text{C}}^{\text{[NMR]}}\}$ data or ^{13}C NMR-peak assignments appears to exist for the 2C-OA and C-ME cocrystals, very notable is the excellent agreement between the ^{13}C chemical shifts obtained herein for 2C-MA and C-GA with those reported earlier by Vigilante and Mehta:⁵⁰ the truly marginal discrepancies of ≤ 0.2 ppm are well within the experimental uncertainties throughout, where systematic ^{13}C MAS NMR shift-referencing errors often yield $\bar{\delta}_{\text{C}}^{\text{[NMR]}}$ discrepancies exceeding 0.5 ppm between studies. For instance, the two ^{13}C chemical shifts reported by Bryce and coworkers for the $^{13}\text{C4}$ (142.9 ppm)⁴⁶ and $^{13}\text{C8}$ (28.3 ppm)⁴⁹ sites of C-ME accord with those of Table 2 within 0.3 ppm and 0.7 ppm, respectively.

Table 2 reveals a typical agreement of 2–3 ppm between the experimental and modeled $\{\bar{\delta}_{\text{C}}^{\text{[NMR]}}\}$ values across all diacid precursors and their corresponding cocrystals with caffeine, where deviations ≤ 1 ppm and ≤ 3 ppm result for 24 and 56 sites out of the entire ensemble of 68 sites/shifts, respectively. Out of the 12 ^{13}Cj sites manifesting a > 3.0 ppm $\bar{\delta}_{\text{C}}^{\text{[NMR]}}$ discrepancy between the model and experiment, however, 8 sites concern the GA and ME diacid precursors and their cocrystals, where the GIPAW calculations of the GA-associated phases consistently overestimate the $^{13}\text{COOH}$ shifts, whereas the $^{13}\text{CH}_2$ counterparts are in excellent agreement with experiments. That scenario is reversed for the ME/C-ME structures, for which the NMR/GIPAW-derived $^{13}\text{COOH}$ chemical shifts accord well, whereas substantial deviations are observed for the ^{13}CH sites (Table 2). Similar systematic errors—but of overall much smaller magnitudes—are observed for the 2C-OA and 2C-MA cocrystals and



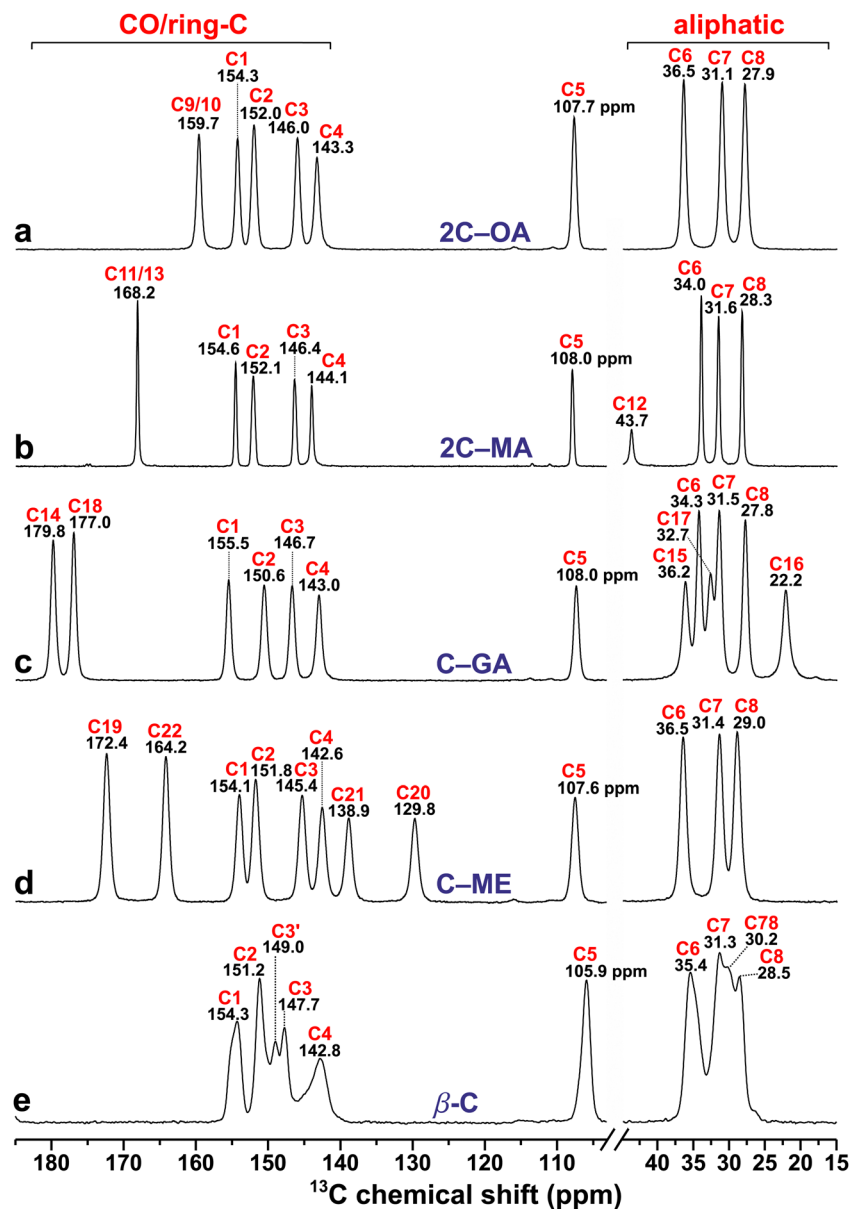


Fig. 4 ^{13}C CPMAS NMR spectra acquired at 14.1 T and 12.00 kHz MAS from cocrystals of β -caffeine and (a) oxalic acid (2C-OA), (b) malonic acid (2C-MA), (c) glutaric acid (C-GA), and (d) maleic acid (C-ME), along with the spectrum from (e) β -caffeine (β -C) for reference.

their OA, OA·2H₂O, and MA precursors, where the modeled $\{\delta_{\text{C}}^{\text{DFT}}\}$ values are consistently higher than their experimental counterparts. While Table 2 also reveals minor systematic discrepancies for the C1–C8 sites of the caffeine moiety in the cocrystals relative to β -caffeine, the GIPAW-derived $\{\delta_{\text{C}}^{\text{DFT}}\}$ values are typically lower than those from NMR, except for C4/C5 that conform to the more typical trend of overestimated $\delta_{\text{C}}^{\text{DFT}}$ values by the calculations.

Because most deviations between the NMR/GIPAW-derived ^{13}C chemical shifts are systematic, they have little/no bearings on our analyses below that target the shift-difference between the cocrystals and precursors (Section 4.6). Nonetheless, the very significant discrepancies between a few experimental/modelled $\delta_{\text{C}}^{\text{DFT}}$ values for the GA and ME moieties in both

precursor and cocrystal structures are reasons for concern. We remind that our $\delta_{\text{C}}^{\text{DFT}}$ values of Table 2 originated from DFT energy minimizations where only the H positions were adjusted, except for β -caffeine, for which all unit-cell parameters and atom coordinates were optimized (Table S2, ESI†). That significantly improved the chemical-shift predictions, as reflected in a root-mean-square deviation (rmsd) of 1.4 ppm relative to the experimental shifts, which may be contrasted with the rmsd(DFT/NMR) = 2.0 ppm outcome that resulted by only adjusting the H atom positions of β -caffeine, or with the corresponding rmsd(DFT/NMR) = 3.0 ppm obtained across the entire $\{\delta_{\text{C}}^{\text{DFT}}\}$ ensemble from all other phases (60 data points). Yet, there were no improvements by employing full-atom/cell optimizations of the {GA, C-GA} and {ME, C-ME} structures (which

Table 2 NMR/DFT-derived ^{13}C chemical shifts of cocrystals and precursors^a

Coformer	Site	Coformer δ_{C} (ppm)	2C-OA δ_{C} (ppm)	2C-MA δ_{C} (ppm)	C-GA δ_{C} (ppm)	C-ME δ_{C} (ppm)
β -Caffeine ^b	C1	154.5(153.0)	154.3(153.1)	154.6(154.8)	155.5(154.5)	154.1(153.1)
	C2	150.8(149.4)	152.0(150.4)	152.1(150.1)	150.6(148.9)	151.8(150.5)
	C3	148.0(147.5)	146.0(147.0)	146.4(146.6)	146.7(148.5)	145.4(145.1)
	C4	143.5(146.0)	143.3(145.2)	144.1(146.6)	143.0(145.0)	142.6(143.2)
	C5	106.1(107.8)	107.7(110.9)	108.0(110.8)	107.4(110.6)	107.6(110.2)
	C6	35.0(35.3)	36.5(36.2)	34.0(33.0)	34.3(33.5)	36.5(36.6)
	C7	30.7(29.6)	31.1(28.5)	31.6(30.0)	31.5(29.1)	31.4(29.7)
	C8	29.6(29.5)	27.9(25.2)	28.3(25.8)	27.8(24.2)	29.0(27.4)
α -OA	C9/10	160.7(162.9)	159.7(162.2)			
α -OA·2H ₂ O	C9/10	163.1(164.5)	159.7(162.2)			
β -MA	C11	175.1(179.0)		168.2(170.1)		
	C12	40.8(39.7)		43.7(44.3)		
	C13	174.7(175.4)		168.2(170.1)		
β -GA	C14	181.5(186.6)			179.8(183.4)	
	C15	33.8(33.8)			36.2(36.0)	
	C16	18.7(16.5)			22.2(22.0)	
	C17	33.8(33.8)			32.7(32.2)	
	C18	181.5(186.6)			177.0(182.8)	
I-ME	C19	172.9(173.4)				172.4(174.2)
	C20	133.1(141.4)				129.8(137.9)
	C21	140.1(148.1)				138.9(146.4)
	C22	169.2(168.5)				164.2(167.2)

^a ^{13}C chemical shifts (δ_{C}) for the $^{13}\text{C}_j$ site labels of Fig. 1, obtained either experimentally from the shift at the NMR-peak maximum, or by DFT/GIPAW calculations (values within parentheses). The δ_{C} uncertainties are around ± 0.30 ppm (DFT/GIPAW) and ± 0.15 ppm (NMR), except for β -caffeine (Table 1). ^b For β -caffeine (only), center-of-gravity ^{13}C chemical shifts ($\bar{\delta}_{\text{C}}$) are reported rather than δ_{C} . The experimental $\{\bar{\delta}_{\text{C}}\}$ data were obtained by deconvolution of the MAS NMR spectrum, while each DFT/GIPAW-derived $\bar{\delta}_{\text{C}}$ value is the average chemical shift over five crystallographically distinct sites/molecules (see Table 1).

manifest the globally largest δ_{C} discrepancies), whose $\{\delta_{\text{C}}\}$ sets revealed an *even larger* rmsd(DFT/NMR) = 6.5 ppm relative to its experimental counterpart, which exceeded that of rmsd(DFT/NMR) = 4.5 ppm resulting by solely optimizing the H atom positions (Table 2). We have no satisfactory explanation for the large discrepancies (≥ 5 ppm) between the models and experiments for a few ^{13}C sites of the GA/ME-related structures, which must originate from unaccounted structural effects.

4.4. Factors governing the ^{13}C resonance widths

Here we discuss three factors expected to primarily govern the full width at half maximum (fwhm) height of the ^{13}C signals observed from the cocrystals and their precursors (Table S4, ESI†), leading to some recommendations about the choice of external magnetic field (B_0) for arranging (sub)optimal ^{13}C MAS NMR spectral resolution. The two primary spectral-resolution-limiting factors involve so-called “inhomogeneous broadening”,⁶¹ *i.e.*, ^{13}C chemical-shift dispersions from either the anisotropic bulk magnetic susceptibility (ABMS)^{24,71–74} or static structural disorder.^{18,20,21} Both scale linearly with B_0 on a frequency scale (in Hz), which complicates their discrimination if only having a ^{13}C (CP)MAS NMR spectrum available.^{24,73,74} Yet, the ABMS-stemming broadening *in ppm* is shared among all nuclear sites in the structure, regardless of their nuclide type or structural origin, meaning that the ABMS remains constant for all $\{^{13}\text{C}_j\}$ and $\{^1\text{H}_j\}$ sites in the sample,^{24,71–74} in contrast to the local structural disorder that varies among

crystallographically distinct $^{13}\text{C}_j$ sites, and thereby giving variable ^{13}C fwhm values (in ppm). Hence, contrasting the (lack of) variations within the $\{^{13}\text{C}_j\}$ fwhm set listed in Table S4 (ESI†) within each precursor/cocrystal helps gauging the potential presence of structural disorder.

Fig. 3 reveals that the ^{13}C resonance widths vary markedly among the various well-ordered cocrystal precursors, which most likely reflects variable ABMS effects. The polycrystalline MA powder produces fwhm values of <30 Hz (≤ 0.2 ppm at 14.1 T; Table S4, ESI†), which are typical for organic molecules in well-ordered crystals with negligible ABMS broadening. In contrast, the OA, OA·2H₂O, and GA powders reveal 3–4 times broader ^{13}C resonances than MA, whereas the ^{13}C sites of the ME and β -caffeine specimens manifest substantially larger fwhm values of ≈ 240 Hz (≈ 1.6 ppm) and 200–435 Hz (1.3–2.9 ppm), respectively. Here, the near-constant peakwidths observed for ME suggests that ABMS broadening mainly limits its ^{13}C NMR spectral resolution, whereas structural disorder is mainly degrading that for β -caffeine, notably so in the CH₃ region (Fig. 3e). Indeed, the markedly larger $^{13}\text{C}_j$ fwhm values of β -caffeine relative to all other specimens (Fig. 3 and 4) primarily stems from a significant (static) structural disorder,^{22,24} where the lowest fwhm value of 1.3 ppm observed for the $^{13}\text{C}_5$ resonance sets an upper limit of the ABMS contribution to all peak widths.

Despite the larger supramolecular aggregate of the cocrystal units, their ^{13}C peak widths remain consistently narrower than



those of β -caffeine (Table S4, ESI[†]), which reflects a higher degree of local structural order of each cocrystal.^{12,14,24} The relative ^{13}C fwhm values increase along the series $2\text{C-MA} \ll 2\text{C-OA} < \text{C-GA} \lesssim \text{C-ME}$, roughly following those of each respective pristine diacids (Table S4, ESI[†]) but with 1.2–6 times wider ^{13}C resonances. Only the peakwidths from C-ME break that trend, where narrower peaks are observed relative to both the ME and β -caffeine crystallites for which ABMS and structural disorder primarily govern the respective ^{13}C fwhm values, both apparently being reduced in the C-ME crystallites.

A third factor known to broaden ^{13}C resonances from molecules featuring direct C–N bonds is the spin-1 ^{14}N nuclide (99.6% natural abundance) that may induce additional NMR-peak splittings/broadenings of nearby ^{13}C sites by ^{14}N – ^{13}C dipolar interactions, whose effects are incompletely suppressed by MAS and scale as B_0^{-1} (in Hz) and B_0^{-2} (in ppm).^{75,76} Indeed, while demonstrated to vastly dominate the ^{13}C MAS NMR-peak widths and the spectral resolution from C–H₂O at a low magnetic field of 4.7 T but being negligible at $B_0 = 21$ T,²² we expect marginal signal broadenings for our experimentation at $B_0 = 14.1$ T, as was also concluded in ref. 50. The data of Table S4 (ESI[†]) confirm those expectations, suggesting a <30 Hz ^{13}C resonance broadening from the ^{14}N – ^{13}C dipolar interactions, as is also supported by the ^{13}C NMR peak-width analysis of ref. 24 performed at $B_0 = 11.7$ T, at which one expects slightly larger effects from ^{14}N – ^{13}C interactions. From the marginal dipolar broadening to the net ^{13}C fwhm values, we conclude that inhomogeneous ABMS and structural-disorder stemming ^{13}C chemical-shift distributions dominate the NMR peak-widths throughout all specimens considered herein, as also suggested by our ^1H NMR results (Section 4.5).

At least for anhydrous β -caffeine, however, ^{13}C NMR experimentation at $B_0 \geq 14.1$ T is advantageous for achieving sufficient resolution. Recalling the two additional C3' and C78 NMR peaks (Section 4.2) of anhydrous β -caffeine relative to its α counterpart,²² a puzzling feature is their apparent absence in several ^{13}C MAS NMR spectra reported from β -caffeine in the literature,^{14,23,24} incidentally rendering those spectra closer to that observed from α -caffeine,²² encompassing some results from “anhydrous caffeine” of unspecified phase identity.^{68,70} Yet all those NMR spectra were recorded at lower magnetic fields of either 9.4 T (ref. 14 and 23) or 11.7 T (ref. 24) relative to those acquired at 14.1 T herein (Fig. 4e) and in ref. 12 along with the high-field result by Enright *et al.*²² at $B_0 = 21$ T. Evidently, only MAS NMR spectra recorded at $B_0 \geq 14.1$ T give discernible ^{13}C resonances at ≈ 30 ppm (C78) and ≈ 149 ppm (C3'), whereas those signals apparently coalesce with those from $^{13}\text{C}7/^{13}\text{C}8$ and $^{13}\text{C}3$, respectively, in NMR spectra obtained at $B_0 < 14.1$ T.^{14,23,24} This effect must stem from the resolution-degradation associated with lower- B_0 experimentation alone and/or its accompanying emphasized ^{13}C resonance broadening from ^{14}N – ^{13}C interactions. Likewise, the ^{13}C MAS NMR spectrum obtained from “anhydrous caffeine” at 9.4 T by Nonappa and Kolehmainen⁶⁸ appears very similar to those reported from β -caffeine in ref. 14, 23 and 24.

We conclude that “high-field” NMR experimentation is beneficial for enabling complete ^{13}C site/NMR-peak assignments of anhydrous β -caffeine, where $B_0 = 14.1$ T appears to be the minimum magnetic field admitting resonance-discrimination between all crystallographically inequivalent ^{13}C sites, potentially enabling ^{13}C NMR quantifications of the α/β -caffeine contributions in a powder mixture without resorting to PXRD.

4.5. ^1H NMR and DFT/GIPAW results

4.5.1. ^1H MAS NMR spectra. Fig. 5 and 6 present the ^1H NMR spectra from the precursors and cocrystals, respectively, all recorded at $B_0 = 14.1$ T and fast MAS (60.0 kHz) to suppress the normally spectral-resolution-limiting homonuclear ^1H – ^1H dipolar interactions.^{15–18} Indeed, the relative

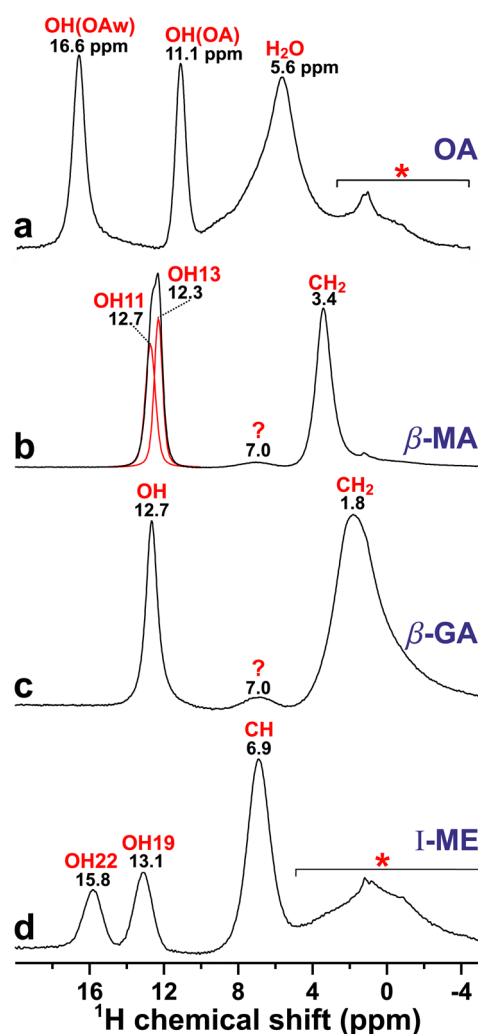


Fig. 5 ^1H MAS NMR spectra recorded at 14.1 T and 60.0 kHz MAS from the (a) α -OA (“OA”) and α -OA-2H₂O (“OAw”) mixture together with powders from the other (b) β -MA, (c) β -GA, and (d) I-ME diacids. The red traces in (b) are deconvolution results of the two overlapping OH11/OH13 resonances. The low-ppm resonance region marked by an asterisk in (a) and (d) reveal background signals from the NMR probehead, and the peak marked by “?” in (b) and (c) stems from an unknown impurity phase of the MA and GA precursors.



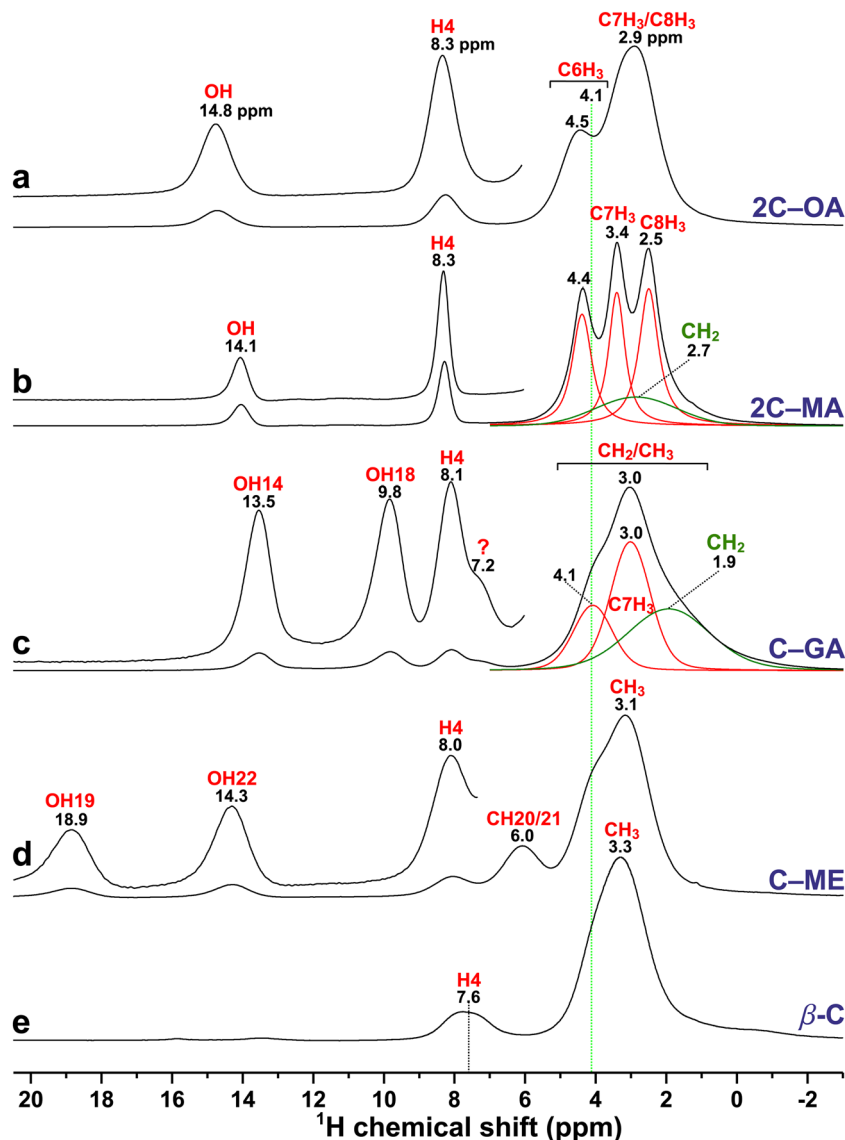


Fig. 6 ^1H MAS NMR spectra recorded at 14.1 T and 60.0 kHz MAS from the (a) 2C–OA, (b) 2C–MA, (c) C–GA, (d) C–ME cocrystals, and (e) anhydrous β -caffeine. The insets show vertically expanded zooms across the high-ppm spectral region. The red/green traces in (b) and (c) are deconvolution results of the overlapping C^1H_3 (red) and C^1H_2 (green) NMR signals. The vertical dotted line at $\delta_{\text{H}} = 4.1$ ppm marks the approximate chemical shift of the C^6H_3 resonance of the caffeine molecule for the C–GA, C–ME, and β -caffeine specimens in (c)–(e), which differ slightly from those of 2C–OA and 2C–MA in (a) and (b). The NMR peak marked by “?” in (c) stems from an unknown impurity of the GA precursor (see Fig. 5c).

^1H NMR peak widths of the cocrystals match well those of the respective ^{13}C NMR spectra (Fig. 4), further underscoring that the $^1\text{H}/^{13}\text{C}$ resonance widths are governed primarily by chemical-shift dispersions stemming from variable ABMS^{24,71–74} and structural disorder^{18,20,21} effects among the specimens (Section 4.4). In the least ordered structure—i.e., that of β -caffeine—all C^1H_3 -stemming ^1H resonances coalesce into a near-Gaussian peak, whereas the C^6H_3 peak is discernible in all NMR spectra from the cocrystals (Fig. 6), despite a further overlap from C^1H_2 signals in those from 2C–MA and C–GA. All three $^1\text{H}6$ – $^1\text{H}8$ resonances are resolved from the most ordered cocrystal structure, i.e., 2C–MA (Fig. 6b). Notably, the present ^1H MAS NMR spectrum from β -caffeine is very similar to those of ref. 23 and 24 where the value of $\delta_{\text{H}}^4 = 7.6$ ppm

reported by Bordignon *et al.*²³ is identical to ours (Table 3), despite that all C^1H_3 resonances coalesced fully^{23,24} (although, for unclear reasons, ref. 23 tabulated distinct $^1\text{H}6$ and $^1\text{H}7/^1\text{H}8$ shifts).

Nonetheless, the reduced ^1H NMR spectral resolution in the aliphatic region, for which broadening from residual ^1H – ^1H interactions is most pronounced, compromises accurate site-specific assessments of the ^1H chemical-shift alterations upon cocrystal formation. Table 3 collects the NMR and DFT/GIPAW derived ^1H chemical shifts, which constitute δ_{H}^i data for all well-resolved signals but CG shifts ($\bar{\delta}_{\text{H}}^i$) for all heavily overlapping resonances. Hence, in what follows, we focus on the $^1\text{H}/j$ resonances that are well resolved in the NMR spectra from both the precursors (Fig. 5) and their corresponding cocrystals



Table 3 NMR/DFT-derived ^1H chemical shifts of cocrystals and precursors^a

Coformer	Site	Coformer $\delta_{\text{H}}/\text{ppm}$	2C-OA $\delta_{\text{H}}/\text{ppm}$	2C-MA $\delta_{\text{H}}/\text{ppm}$	C-GA $\delta_{\text{H}}/\text{ppm}$	C-ME $\delta_{\text{H}}/\text{ppm}$
β -Caffeine ^b	H4	7.6(8.4)	8.3(7.2)	8.3(7.6)	8.1(7.3)	8.0(6.7)
	H6/7/8 ^c	3.3(3.8)	3.3(2.6)	3.4(2.8)	3.3(2.7)	3.4(2.6)
α -OA α -OA·2H ₂ O	HO9/10	11.1(11.2)	14.8(15.0)			
	HO9/10	16.6(17.8)	14.8(15.0)			
	H ₂ O	5.6(5.6)				
β -MA	HO11	12.7(13.7)		14.1(14.8)		
	H12a/b	3.4(2.6)		2.7(1.7)		
	HO13	12.3(13.4)		14.1(14.8)		
β -GA	HO14	12.7(13.7)			13.5(14.0)	
	H15/16/17	1.8(1.2)			1.9(1.0)	
	HO18	12.7(13.7)			9.8(10.6)	
I-ME	HO19	13.1(13.6)				18.9(20.2)
	H20/21	6.9(6.2)				6.0(4.9)
	HO22	15.8(16.2)				14.3(14.6)

^a ^1H chemical shifts (δ_{H}^i) for the Cj¹H site labels of Fig. 1, obtained either experimentally from the shift at the NMR-peak maximum, or by DFT/GIPAW calculations (values within parentheses). The δ_{H} uncertainties are around ± 0.30 ppm (GIPAW) and around ± 0.15 ppm for the experimental values for all well-resolved NMR peaks (± 0.5 ppm for those obtained by spectral deconvolution; Fig. 6). ^b All GIPAW-derived shifts for β -caffeine are averages of the five $\{\delta_{\text{H}}^i\}$ values of the inequivalent molecules in the unit cell. ^c Owing to the complete overlap among the C6H₃, C7H₃, and C8H₃ resonances in β -caffeine, only the average chemical-shift values are reported. However, the NMR spectra from the cocrystals admitted further site/signal-resolution by spectral deconvolution, yielding the following NMR-derived δ_{H}^i (or average CG δ_{H}^i chemical shifts), with the corresponding GIPAW-derived data given within parentheses: for 2C-OA, $\delta_{\text{H}}^6 = 4.5$ (3.7) ppm; $\delta_{\text{H}}^{7/8} = 2.9$ (2.1) ppm; for 2C-MA, $\delta_{\text{H}}^6 = 4.4$ (3.8) ppm; $\delta_{\text{H}}^7 = 3.4$ (2.8) ppm; $\delta_{\text{H}}^8 = 2.5$ (1.8) ppm; for C-GA, $\delta_{\text{H}}^6 = 4.1$ (3.5) ppm; $\delta_{\text{H}}^{7/8} = 3.0$ (2.3) ppm; for C-ME, $\delta_{\text{H}}^6 = 4.1$ (3.2) ppm; $\delta_{\text{H}}^{7/8} = 3.1$ (2.3) ppm.

(Fig. 6), namely COO¹H of the diacid molecules, C4¹H of caffeine, and the equivalent C¹H20/21 sites of ME.

4.5.2. ^1H chemical shifts of carboxy groups. Here, we discuss the set $\{\delta_{\text{H}}^i\}$ of COO¹H chemical shifts, assessed directly from the respective NMR-peak maximum shift values of Fig. 5 and 6 and assigned such that the highest/lowest chemical shift represents the H_j site with shortest/longest H...O distance in the diacid/cocrystal structure. These $^1\text{H}_j/\delta_{\text{H}}^i$ assignments are further corroborated by the DFT/GIPAW calculations (Table 3). The herein observed COO¹H chemical shifts from the α -OA·2H₂O, β -MA, and ME powders agree within 0.3 ppm with the δ_{H}^i values reported by Harris *et al.*³⁴ by using CRAMPS (combined rotation and multiple-pulse spectroscopy) to suppress NMR-peak broadenings from ^1H - ^1H interactions under slow-MAS conditions. This excellent agreement is well within the experimental uncertainties, while moreover δ_{H}^9 of α -OA·2H₂O (16.6 ppm) matches perfectly that reported earlier by Berglund and Vaughan⁷⁷ (16.5 ppm). Notably, the chemical shifts observed by us from MA and ME agree much better with those of Harris *et al.*³⁴ than with ref. 77 but the former study did not include α -OA. Although the experimental shift $\delta_{\text{H}}^{9/10} = 11.1$ ppm (Table 3) is reproduced within 0.1 ppm by our GIPAW calculations, there is a significant deviation with that of 12.6 ppm reported in ref. 77 which is well outside of our experimental uncertainties. We believe that the herein reported $\delta_{\text{H}}^{9/10} = 11.1$ ppm value of α -OA is the hitherto most accurate result.

The COO¹H chemical shifts of the pristine diacids presented in Fig. 5 correlate qualitatively well with the H...O distances of their crystal structures (Fig. 2): the two α -OA·2H₂O and α -OA structures reveal the highest (16.6 ppm) and lowest (11.1 ppm) chemical shifts, respectively, as expected from their

corresponding shortest (146 pm) and longest (175 pm) HB lengths among all diacids (see caveat below). All COO¹H sites of MA and GA along with H19 of ME feature essentially identical H-bond distances of 162–165 ppm (Fig. 2), as is mirrored in very similar δ_{H}^i values between 12.3–13.1 ppm (Fig. 5). Notably, the ^1H NMR signals from the two crystallographically inequivalent OH11 and OH13 sites of MA are discernible in Fig. 5b, despite their very small chemical-shift separation of 0.4 ppm stemming from a minute H...O bond-length difference of 1 pm (Fig. 2d), as also reproduced near-perfectly by the GIPAW calculations (0.3 ppm shift-difference).

Within a consistent but minor overestimation of the DFT/GIPAW-derived ^1H chemical shift within typically $\lesssim 1$ ppm (Table 3), the calculated $\{\delta_{\text{H}}^i\}$ values match the experimental counterparts very well for all COO¹H sites. Note that while all diacid structures feature H...O bonds, all cocrystals involve H...N contacts with the “N” atom of caffeine, except for the OH18...O14 bond in C-GA and OH22...O19 in C-ME (Fig. 2). Each NMR/GIPAW-derived $\{\delta_{\text{H}}^i\}$ dataset gave a good linear correlation with the shortest H...O or H...N distance, denoted $r(\text{H}\cdots\text{O}/\text{N})$, throughout all COO¹H sites. Fig. 7 plots the experimental and calculated ^1H chemical shifts along with the corresponding best-fit results given by

$$\delta_{\text{H}}^i[\text{NMR}]/\text{ppm} = 47.0 - 0.208r(\text{H}\cdots\text{O}/\text{N})/\text{pm} \quad (R^2 = 0.912), \quad (3)$$

$$\delta_{\text{H}}^i[\text{DFT}]/\text{ppm} = 50.0 - 0.222r(\text{H}\cdots\text{O}/\text{N})/\text{pm} \quad (R^2 = 0.919), \quad (4)$$

both of which verify the expected trend^{34,35,39,40,42} of an *increasing* ^1H chemical shift for *decreasing* HB length. Indeed, both the



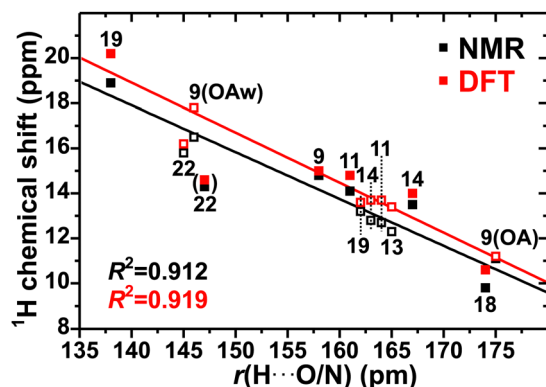


Fig. 7 Experimental (black symbols) and DFT/GIPAW-generated (red) ^1H chemical shifts of the carboxy moieties of the pristine diacids (solid symbols) and the cocrystals (open) plotted against the $\text{H}\cdots\text{O}/\text{H}\cdots\text{N}$ distance, $r(\text{H}\cdots\text{O}/\text{N})$, given in Fig. 2. The number around each data-point represents the proton label of Fig. 1. The lines are best-fit results, eqn (3) and (4), obtained by omitting one outlier data point (δ_{H}^{22}) and yielding the as-indicated R^2 correlations coefficients. All chemical-shift uncertainties are within the symbol sizes.

experimental and modeled $\{\delta_{\text{H}}^j\}$ sets are captured well by their very similar best-fit expressions and nearly equal correlation coefficients. Only the H22 site of ME breaks the trend, manifesting lower experimental and calculated shifts than those predicted from the very short intramolecular $\text{H}\cdots\text{O}$ distance of 145 pm within the diacid unit (*vide supra*).

4.5.3 COO^1H chemical-shift changes upon cocrystal formation. To monitor the ^1H chemical-shift alterations upon cocrystal formation, we define the *difference* between the chemical shift of site $^1\text{H}_j$ in the cocrystal, $\delta_{\text{H}}^j[\text{C-X}]$, relative to that in each pristine coformer, $\delta_{\text{H}}^j[\text{X}]$, where X represents β -caffeine or one of the $\{\text{OA}, \text{OA}\cdot\text{H}_2\text{O}, \text{MA}, \text{GA}, \text{ME}\}$ diacids,

$$\Delta_{\text{H}}^j = \delta_{\text{H}}^j[\text{C-X}] - \delta_{\text{H}}^j[\text{X}], \quad \text{with } 1 \leq j \leq 22, \quad (5)$$

and the index j runs over all proton-bearing C_j sites of Fig. 1. The linear $\delta_{\text{H}}^j/r(\text{H}\cdots\text{O}/\text{N})$ trends of Fig. 7 readily rationalize the observed COO^1H chemical-shift alterations, whose values are presented in Table 4 along with those for the other C^1H_n sites (the latter are discussed in Section S1). Fig. S4 (ESI†) plots the NMR/DFT-derived Δ_{H}^j values against the *difference* between the shortest $\text{H}\cdots\text{O}/\text{N}$ distance of each cocrystal and precursor, which is denoted by $\Delta r(\text{H}\cdots\text{O}/\text{N})$.

For the COO^1H chemical-shift changes (Δ_{H}^j), Table 4 reveals an overall trend of *higher* δ_{H}^j values of the cocrystals (*i.e.*, $\Delta_{\text{H}}^j > 0$), as is witnessed by Fig. 7 and rationalized from the typically *shorter* $\text{H}\cdots\text{O}$ distance encountered in the cocrystal relative to that of the pristine diacid (Fig. 2). Such HB-length effects also account for the higher δ_{H}^9 value of 2C-OA relative to its α -OA counterpart, as well as for the increased δ_{H}^{11} and δ_{H}^{13} values upon 2C-MA cocrystal formation. Likewise, the lower δ_{H}^9 value of 2C-OA relative to α -OA $\cdot$ 2H $_2$ O—along with the lower chemical shift of $^1\text{H}_{18}$ of C-GA compared to the identical values $\delta_{\text{H}}^{14} = \delta_{\text{H}}^{18}$ of the crystallographically equivalent COO^1H sites of β -GA—are readily understood from the longer $\text{H}\cdots\text{O}$ distances in the cocrystals.

4.6. ^{13}C chemical-shift changes upon cocrystal formation

In direct analogy with the ^1H chemical-shift difference between the cocrystal and its precursors [eqn (5)], we define

$$\Delta_{\text{C}}^j = \delta_{\text{C}}^j[\text{C-X}] - \delta_{\text{C}}^j[\text{X}], \quad \text{with } 1 \leq j \leq 22, \quad \text{and} \\ \text{X} = \{\beta\text{-C}, \text{OA}, \text{OA}\cdot 2\text{H}_2\text{O}, \text{MA}, \text{GA}, \text{ME}\}, \quad (6)$$

as the corresponding δ_{C}^j alteration when each $\{2\text{C-OA}, 2\text{C-MA}, \text{C-GA}, \text{C-ME}\}$ cocrystal is formed. Table 5 compiles the $\{\Delta_{\text{C}}^j\}$ data obtained either by NMR experiments or GIPAW calculations. However, as for the ^1H chemical-shift/structure correlations (except those for COO^1H), the δ_{C}^j and Δ_{C}^j data only admits qualitative discussions.

Table 4 ^1H Chemical-shift differences between cocrystals and precursors^a

Coformer	Site	2C-OA $\Delta_{\text{H}}^j/\text{ppm}$	2C-MA $\Delta_{\text{H}}^j/\text{ppm}$	C-GA $\Delta_{\text{H}}^j/\text{ppm}$	C-ME $\Delta_{\text{H}}^j/\text{ppm}$
β -Caffeine	H4	0.6(−1.2)	0.7(−0.8)	0.5(−1.1)	0.4(−1.7)
	H6/7/8 ^b	0.0(−1.2)	0.1(−1.0)	0.0(−1.1)	0.1(−1.2)
α -OA	HO9/10	3.8(3.8)			
α -OA $\cdot$ 2H $_2$ O	HO9/10	−1.8(−2.8)			
β -MA	HO11		1.3(1.1)		
	H12a/b		−0.7(−0.9)		
	HO13		1.8(1.4)		
β -GA	HO14			0.8(0.3)	
	H15/16/17			0.1(−0.2)	
	HO18			−2.9(−3.1)	
I-ME	HO19				5.8(6.6)
	H20/21				−0.9(−1.3)
	HO22				−1.5(−1.6)

^a Chemical-shift differences (Δ_{H}^j) defined by eqn (5) and calculated from δ_{H}^j data with two decimals. ^b Average chemical-shift values over all C6H $_3$, C7H $_3$, and C8H $_3$ sites of the caffeine moiety; see Table 3.



Table 5 ^{13}C Chemical-shift differences between cocrystals and precursors^a

Coformer	Site	2C-OA $\Delta\delta_{\text{C}}/\text{ppm}$	2C-MA $\Delta\delta_{\text{C}}/\text{ppm}$	C-GA $\Delta\delta_{\text{C}}/\text{ppm}$	C-ME $\Delta\delta_{\text{C}}/\text{ppm}$
β -Caffeine	C1	-0.2(0.1)	0.0(1.8)	1.0(1.5)	-0.5(0.1)
	C2	1.2(1.0)	1.3(0.7)	-0.2(-0.5)	1.0(1.1)
	C3	-2.0(-0.6)	-1.6(-0.9)	-1.3(1.0)	-2.7(-2.4)
	C4	-0.1(-0.8)	0.6(0.6)	-0.4(-1.0)	-0.9(-2.8)
	C5	1.7(3.2)	1.9(3.1)	1.4(2.8)	1.5(2.5)
	C6	1.4(0.9)	-1.0(-2.3)	-0.7(-1.7)	1.5(1.4)
	C7	0.4(-1.1)	0.9(0.3)	0.8(-0.6)	0.7(0.0)
	C8	-1.7(-4.3)	-1.3(-3.7)	-1.8(-5.3)	-0.7(-2.0)
α -OA	C9/10	-1.1(-0.7)			
α -OA·2H ₂ O	C9/10	-3.5(-2.3)			
β -MA	C11		-6.9(-8.8)		
	C12		2.9(4.6)		
	C13		-6.5(-5.3)		
β -GA	C14			-1.6(-3.3)	
	C15			2.4(2.2)	
	C16			3.4(5.5)	
	C17			-1.1(-1.6)	
	C18			-4.5(-3.9)	
I-ME	C19				-0.4(0.7)
	C20				-3.3(-3.5)
	C21				-1.1(-1.7)
	C22				-5.0(-1.3)

^a Chemical-shift differences ($\Delta\delta_{\text{C}}$) defined by eqn (6) and calculated from δ_{C} data with two decimals.

In contrast with the $^{13}\text{CH}_2$ shifts of MA and GA, which deshield slightly by a few ppm, significantly decreased δ_{C} values are observed for the $^{13}\text{COOH}$ diacid sites, whose adjacent O atom form an HB with protons of neighboring caffeine/diacid molecules, which along with $\text{COO}^- \rightarrow \text{COOH}$ conversions are well known to decrease the ^{13}C shift in various molecular systems.^{37,38,78–81} However, that trend has remained qualitative without any firm quantitative relationship having thus far been established between the *isotropic* ^{13}C chemical shift and the $^{13}\text{COH} \cdots \text{O}$ or $^{13}\text{CO} \cdots \text{HO}$ distance (the principal value δ_{yy} of the chemical-shift tensor, however, correlates linearly with the HB distance/angle parameters^{36–38}). Indeed, the plot of δ_{C} against $r(\text{H} \cdots \text{O}/\text{N})$ in Fig. 8 reveals a significant scatter for the diacid/cocrystal structures. Although a weak trend of $^{13}\text{COOH}$ shift-reduction with a shortened $\text{COO} \cdots \text{H}$ distance is discernible, our data do not reveal any reasonable $\delta_{\text{C}}/r(\text{H} \cdots \text{O}/\text{N})$ or $\Delta\delta_{\text{C}}/\Delta r(\text{H} \cdots \text{O}/\text{N})$ correlation for either the experimental or modeled data. The only exceptions are the $^{13}\text{COOH}$ sites of the 2C-OA (δ_{C}^9), 2C-MA (δ_{C}^{11}), and C-GA (δ_{C}^{14}) cocrystals, all of which share the feature of a strong $\text{H} \cdots \text{N}$ bond to the N atom of the caffeine molecule (which is expected to constitute the primary stabilization of the supramolecular structure⁸); they establish an excellent linear $\delta_{\text{C}}/r(\text{H} \cdots \text{N})$ relationship (Fig. 8) for both the NMR and GIPAW-derived shifts but with the caveat that only three data points underlie the correlation.

Turning to the δ_{C} changes of the caffeine moiety in the cocrystals relative to β -caffeine, Table 5 reveals the overall largest observed $|\Delta\delta_{\text{C}}|$ values for the {C3, C5, C8} sites upon cocrystal formation (and to a lesser extent, C2 and C6), thereby

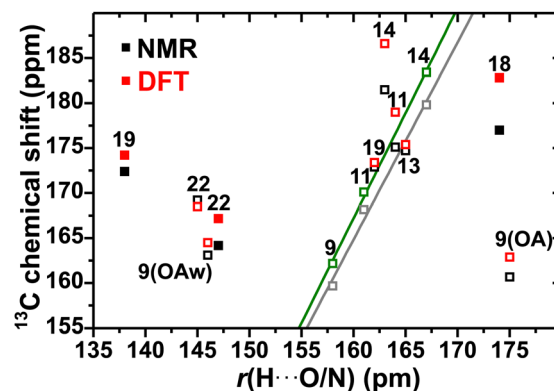


Fig. 8 Experimental (black symbols) and DFT/GIPAW-generated (red) ^{13}C chemical shifts of the carboxy moieties of the pristine diacids (solid symbols) and the cocrystals (open) plotted against the $\text{H} \cdots \text{O}$ or $\text{H} \cdots \text{N}$ distance, $r(\text{H} \cdots \text{O}/\text{N})$, within each structure (Fig. 2). Each number represents the corresponding Cj label of Fig. 1. The symbols set in green and gray color for the NMR and DFT/GIPAW data, respectively, correspond to the results obtained from the 2C-OA (δ_{C}^9), 2C-MA (δ_{C}^{11}), and C-GA (δ_{C}^{14}) cocrystals, all of which are H-bonded to the "N" atom of caffeine. The corresponding lines are best-fit results to the expressions $\delta_{\text{C}}[\text{NMR}]/\text{ppm} = -185.6 + 2.1905r(\text{H} \cdots \text{N})/\text{pm}$ ($R^2 = 0.990$) and $\delta_{\text{C}}[\text{DFT}]/\text{ppm} = -206.5 + 2.336r(\text{H} \cdots \text{N})/\text{pm}$ ($R^2 = 0.998$), respectively. All chemical-shift uncertainties are within the symbol sizes.

suggesting that those local ^{13}C electronic environments alter the most by the intermolecular caffeine-coformer interactions in the cocrystals structure (Fig. 2) relative to the caffeine-caffeine contacts in β -caffeine. The C3 and C5 atoms constitute



a C=C fragment between the pyriminedione and imidazolium rings (Fig. 1). We attribute the (globally largest) chemical-shift alteration of the C3 site of the caffeine molecule to its close proximity to the N atom, whose N...HOOC HB constitutes the primary intermolecular interaction in all cocrystal structures;⁸ see Section 3 and Fig. 2. However, while the ¹³C3 site *shields* upon cocrystal formation ($\Delta_C^3 < 0$), ¹³C5 *deshields* ($\Delta_C^5 > 0$) to a comparable extent, for unknown reasons when considering its remoteness to any atom of a neighboring molecule in either the β -caffeine or cocrystal structures. The ¹³CH₃ shift alterations are discussed in Section 4.7.

As commented in Section 4.5, the C4¹H chemical shift remains near-constant throughout all cocrystals, which is attributed to Δ_C^4 cancellation-effects from variable and complex H-bonding scenarios. Likewise, only minor chemical-shift alterations are observed for ¹³C4H, which besides being directly linked to the H-bonded “N” site of caffeine, *additionally* experiences weak H bonds to the CO atoms of the diacid carboxy groups upon cocrystal formation (Fig. 2). We speculate that the chemical-shift effects from those two competing H bonds might partially cancel each other. We remark that even slightly more negative Δ_C^4 values of {−1.3, −0.5, −1.6, −2.0} ppm apply if the respective {2C-OA, 2C-MA, C-GA, C-ME} cocrystals would be prepared from the ordered C-H₂O phase that features $\delta_C^4 = 144.6$ ppm²² and one C4H...OH₂ bond.

4.7 Tetrel-bonding effects on methyl ¹³C/¹H chemical shifts

Besides the possibility of a weak HB, the electrophilic nature of methyl groups offer the option of a H₃C...O TB to a proximate electronegative and nucleophilic atom.^{47–49} From our as-observed Δ_C^i/Δ_H^i data for the three ¹³CH₃ groups of the caffeine moiety upon cocrystal formation, we discuss the complex HB/TB interplay on their ¹³C/¹H chemical shifts in relation to current literature, which is both sparse and inconclusive. Likewise, our data may only be discussed qualitatively.

Both DFT calculations^{48,49} and NMR experiments⁴⁹ suggest consistently larger δ_C than δ_H alterations when a TB is formed. Scheiner *et al.*⁴⁸ investigated the ¹³CH₃ and C¹H₃ chemical-shift displacements upon CH...O HB or HC...O TB formation by DFT calculations, inferring that both ¹H and ¹³C chemical shifts *increase* for either scenario, with the precise deshielding depending on the TB-length and N-C-O bond angle (θ_{TB}). The largest chemical-shift alterations occurred for a linear TB constellation, for which $\Delta_C^i \leq 6$ ppm was predicted, whereas the ¹H chemical shift only increased marginally (≤ 1 ppm).⁴⁸ Formation of a weak CH...O HB, however, is reported to produce a comparably larger ¹H deshielding than for ¹³C.^{43,46,48} Currently no firm/general experimental correlation is reported of either the ¹H or ¹³C chemical shifts against the CH₃...O or H₃C...O distance. A recent compilation by Southern *et al.*⁴⁹ of experimental ¹³C/¹H chemical shifts from the literature along with GIPAW calculations, suggested very scattered ¹³C/¹H chemical-shift-structure correlations of CH₃ groups involved in tetrel bonds:⁴⁹ although experiments confirm that δ_C^i is often increased for decreasing HC...O distance, no significant dependence was observed *versus* θ_{TB} , nor of δ_H^i

against either the TB distance or θ_{TB} . Moreover, when HB and TB bonds coexist for a CH₃ group, their effects on δ_C^i may either reinforce or counteract each other.⁴⁹

The ¹³C and ¹H chemical-shift data from the current cocrystals accord qualitatively with previous findings of a smaller δ_H^i than δ_C^i change when the TB scenario alters between β -caffeine and its cocrystals. The NMR-derived value $\Delta_H^{6/7/8} \approx 0$ apply throughout all cocrystals. Although the δ_H^i /TB distance-accuracy is compromised by the lack of NMR-signal discrimination among the ¹H6–¹H8 methyl resonances from β -caffeine, the near-constant ¹H chemical shift is also confirmed by the DFT calculations (within a constant shift-displacement of $\Delta_H^{6/7/8} \approx -1$ ppm), for which each δ_H^i and Δ_H^i value is readily determined (Table S5, ESI†). Consistently larger ¹³C chemical-shift displacements occur on cocrystal formation, where Table 2 reveals the largest changes for ¹³C8H₃, all of which are negative and conforming to the experimental range of $-1.8 \leq \Delta_C^8/\text{ppm} \leq -1.3$, whereas more negative values are predicted by the GIPAW calculations: $-5.3 \leq |\Delta_C^8|/\text{ppm} \leq -2.0$. Somewhat lower shift-differences (0.7–1.5 ppm) are observed for the ¹³C6H₃ sites, where 2C-OA and C-ME yield *positive* Δ_C^6 values, while those for 2C-MA and C-GA are *negative* (Table 5). Besides slightly larger *magnitudes* predicted by the GIPAW calculations, all positive/negative Δ_C^6 trends are reproduced, as well as the experimental finding that ¹³C7H₃ consistently manifests the smallest shift-change for each cocrystal.

Unfortunately, the Δ_C^i trends identified above for the three ¹³CH₃ sites of the caffeine moiety are for several reasons very difficult to rationalize: (i) δ_C^i is expected to depend on both H...O/N and C...O/N distance-*changes* between β -caffeine and each cocrystal, whose effects on the chemical shift may augment or cancel partially. (ii) Even if *only* each *shortest* HB/TB distance per ¹³CH₃ site of the cocrystal is considered (which is expected to be most influential on δ_C^i), the presence of five inequivalent molecules in the β -caffeine unit cell implies a complex reference scenario with a *range* of HB/TB lengths relative to the structurally ordered cocrystals. (iii) The current absence of any firm $\delta_C^i/r(\text{CH}_n\cdots\text{O})$ or $\delta_C^i/r(\text{CH}_n\cdots\text{O})$ correlation in conjunction with (i) and (ii), complicate the establishment of even semiquantitative Δ_C^i correlations with the HB/TB-distance alterations accompanying the {2C-OA, 2C-MA, C-GA, C-ME} formation.

The shortest HB and TB distances of the methyl groups in β -caffeine and the cocrystals are compiled in Table S3 (ESI†). We first consider the most pronounced ¹³C chemical-shift change upon cocrystal formation—*i.e.*, Δ_C^8 , assuming that the HB effects are minor because the range of C8H₃...O1/O2 distances (236–294 pm) in β -caffeine span those of the cocrystals (and is thereby expected to give, on the average, a very similar δ_C^8 contribution as for the cocrystals). Hence, the corresponding TB-length alterations are expected to primarily govern Δ_C^8 . Notably, the TB distances are overall shorter in the cocrystals than in β -caffeine (except for 2C-OA). Then given that a TB contraction is expected to *increase* δ_C^8 ,^{48,49} positive values of Δ_C^8 are anticipated throughout all cocrystals, in stark contrast to the *de facto* observed experimental and modeled results. The



lack of an even qualitative $\Delta_C/\Delta r(\text{CH}_3 \cdots \text{O}/\text{N})$ correlation also applies to the $^{13}\text{C}_6\text{H}_3$ site: again, the wide HB-distance range in β -caffeine (207–276 pm; Table S3, ESI†) relative to those of the cocrystals (223–249 pm) suggest minor effects on Δ_C^6 . Notably, however, the $\text{C}_6\text{H}_3 \cdots \text{O}$ bond-lengths in the β -caffeine structure is *consistently shorter* than those of any cocrystal, from which *negative* Δ_C^6 values are predicted throughout,^{48,49} in clear contradiction to our NMR and DFT results. Moreover, no correlation was observed for the set of $^{13}\text{CH}_n$ chemical shifts against the TB angle (data not shown), despite wide θ_{TB} ranges within 71° – 90° ($4 \delta_C^6$ values) and 147° – 178° ($8 \delta_C^6$ values).

Notwithstanding the uncertainties accompanying (i)–(iii) above, we believe that our ^{13}C chemical-shift data suggest that a TB-length contraction may *not necessarily* produce an increased ^{13}C chemical shift, along previous difficulties to establish a general $\delta_C^6/r(\text{CH}_3 \cdots \text{O}/\text{N})$ correlation. Such a relationship is likely obscured for similar reasons as some inconclusive experimental $\delta_C^6/r(\text{CH}_n \cdots \text{O})$ correlations, for which ^{13}C shift-alterations upon $\text{CH} \cdots \text{O}$ bond formation depend both on the precise molecular fragments involved and (longer-range) crystal-structure effects,^{44,45} whose complex interplay may result in ^{13}C deshielding,^{44,46} shielding,⁴³ or either/both.^{42,44}

5 Conclusion

The present comprehensive NMR/DFT study encompassed altogether 68 distinct ^{13}C sites and 33 unique ^1H resonances from altogether four caffeine-based cocrystals and their six precursor structures, with an overall very good agreement between the experimental and modeled chemical shifts. We believe that the results herein may serve as accurate benchmark values for further MAS NMR investigations, given the very good accordance with previous literature for the $^1\text{H}/^{13}\text{C}$ chemical shifts—such as the δ_C data of 2C-MA and C-GA of ref. 50 along with both ^1H and ^{13}C chemical shifts of the diacid precursors—and because a significant fraction of all $\delta_{\text{H}}/\delta_C$ data are presented herein for the first time.

Despite our large ^{13}C and ^1H chemical-shift ensembles and an encouraging NMR/DFT agreement, the herein established linear chemical-shift/distance correlations were confined to the $\text{COO}^1\text{H} \cdots \text{O}/\text{N}$ diacid–diacid and diacid–caffeine hydrogen bonds, along with the $\text{COO}^1\text{H} \cdots \text{N}$ counterparts, which nonetheless are those most instrumental for the supramolecular cocrystal organization. We also examined possible $\text{CH}_3 \cdots \text{O}/\text{N}$ tetrel-bond effects on the $^1\text{H}/^{13}\text{C}$ chemical shifts, which were overall uncertain and inconclusive but suggesting that previously reported (weak) chemical-shift correlations against tetrel-bond parameters might not hold in general, thereby underscoring the need for more research in this very sparsely investigated field. Further efforts towards better confining the H-atom positions of the caffeine–diacid cocrystals by direct ^1H – ^1H and ^1H – ^{13}C internuclear-distance measurements are underway by employing a recent NMR crystallography method.^{81,82}

Except for the methylene groups of MA/GA and their cocrystals, the compromised ^1H NMR spectral resolution (and for β -

caffeine also the ^{13}C NMR counterpart) originates mainly from chemical-shift dispersions stemming from anisotropic bulk magnetic susceptibility and structural disorder rather than the often dominant peak-broadening from ^1H – ^1H interactions. Hence, modest spectral resolution enhancements are expected from faster-MAS (> 60 kHz) ^1H NMR experiments, and let alone by utilizing higher fields ($B_0 > 14.1$ T) because the shift-dispersion scales linearly with B_0 . That sharply contrasts with the ^{13}C NMR scenario, however, where $B_0 \geq 14.1$ T experimentation appears mandatory for resolving all 10 resonances from the 8 distinct ^{13}C sites of anhydrous β -caffeine, where the previously unassigned ^{13}C NMR peaks at 30.2 ppm (C7/C8) and 149.0 ppm (C3) are spectral markers for discriminating the disordered anhydrous β -caffeine polymorph from both its disordered α -caffeine and ordered hydrated $\text{C}\cdot\text{H}_2\text{O}$ counterparts. This insight facilitates further α/β -phase quantifications of mixtures of both polymorphs by ^{13}C MAS NMR alone, which has hitherto only proven possible by PXRD analyses.

Conflicts of interest

There are no conflicts to declare.

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