Tackling the stacking disorder of melon—structure elucidation in a semicrystalline material[†]

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In this work we tackle the stacking disorder of melon, a layered carbon imide amide polymer with the ideal composition ($C_6N_7(NH)(NH_2)$). Although its existence has been postulated since 1834 the structure of individual melon layers could only recently be solved *via* electron diffraction and high-resolution ¹⁵N solid-state NMR spectroscopy. With only weak van der Waals interactions between neighboring layers its long range stacking order is poorly defined preventing an efficient use of diffraction techniques. We, therefore, rely on a combination of solid-sate NMR experiments and force field calculations. The key information is obtained based on heteronuclear $(^{1}H-^{13}C)$ and homonuclear $(^{1}H-^{1}H)$ second moments M_{2} acquired from $^{1}H-^{13}C$ cross polarization experiments. To allow for an interpretation of the polarization transfer rates the resonances in the ¹³C MAS spectra have to be assigned and the hydrogen atoms have to be located. The assignment was performed using a two-dimensional ¹⁵N-¹³C *iDCP* experiment. For the determination of the position of the hydrogen atoms NH and HH distances were measured via ¹H-¹⁵N Lee-Goldburg CP and ¹H-¹H double-quantum build-up curves, respectively. Furthermore, the homogeneity of the material under examination was investigated exploiting ¹⁵N spin-diffusion. Based on force field methods 256 structure models with varying lateral arrangements between neighboring layers were created. For each model the M_2 were calculated allowing them to be ranked by comparing calculated and measured M_2 as well as via their force field energies. This allows the creation of markedly structured hypersurfaces with two distinctly favored shift vectors for the displacement of neighboring layers.

1. Introduction

During the last two decades scientific research on carbon nitride materials underwent a renaissance sparked by theoretical calculations predicting a 3D form of carbon nitride to be harder than diamond.^{1,2} Since then, a plethora of structure models were developed, based either on 3D networks or graphite-like layered structures.³ While for the latter, both triazine (C_3N_3) and heptazine (C_6N_9) subunits have been put forward as elementary building blocks, evidence for the heptazine model keeps accumulating.4,5 Structure elucidation of carbon nitride materials allows for the design of optimized synthesis protocols and is, arguably, the key to their macroscopic properties. Recently, analogies have been revealed between the first continuous 2D carbon nitride network and crystalline triazine-based covalent organic frameworks, dubbed "CTFs".^{6,7} The framework topology of poly(heptazine imide), (C₆N₇)₂(NH)₃·C₃N₃(NH₂)₃, though featuring "closed" pores occupied by the template melamine, bodes well for the

templated synthesis of novel, highly porous carbon nitride architectures with tailor-made properties.

Despite the recent progress made towards highly condensed CN_x materials, experimental access to carbon nitrides remains a challenge due to their metastability with respect to the formation of graphite and nitrogen.^{3,8} Since most synthetic routes invoke the pyrolysis of molecular precursors,⁹⁻¹² moreover, the incorporation of additional hetero atoms or hydrogen can hardly be prevented. This was demonstrated recently by Thomas et al. who reported on the synthesis of a microcrystalline, largely hydrogen free graphitic carbon nitride-type material by ionothermal treatment of dicyandiamide.¹³ Nevertheless, reports on promising properties and applications of carbon nitride materials abound in the literature. These imply mechanical hardness and elasticity,^{1,3,14,15} fire-retarding and field-emission properties,¹⁶ as well as potential in energy conversion,^{3,14} hydrogen¹⁷ and Li¹⁸ storage, in solar cells,¹⁹ as humidity and gas sensors²⁰ as well as semiconductors.^{18,21} Along these lines, a recent break-through was achieved by the use of polymeric carbon nitride as an efficient and inexpensive photocatalyst for water splitting.²² Moreover, the same material was demonstrated to be active as a metal-free heterogeneous catalyst in Friedel-Crafts reactions, the activation of CO₂ and in the cyclotrimerisation of triple bonds.²³⁻²⁶

In these applications mainly layered materials are employed which are prepared as bulk materials, ^{15,27–35} as films, ^{16,17,19,20,36–39}

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as nano- or mesoporous samples,^{14,23,26,33,40} and as nanotubes.^{18,41} For nearly all cases hydrogen contents well above 1.6 wt-% were observed, which indicates the presence of oligo- or polymeric melon-type materials rather than of binary carbon nitride.^{4,29,30,35,42–44} The acquisition of accurate data beyond stoichiometry and composition remains largely elusive due to an inherent poor crystallinity and a high degree of disorder. X-ray powder patterns usually indicate a pronounced stacking disorder of neighboring layers with an interlayer distance amounting to about 3.2 Å.^{4,9,13,23,42} Besides X-ray diffraction techniques,^{15,27–31,34,40,42–44} infrared (IR) and Raman spectroscopy^{15,20,27,28,32,33,40,45} as well as X-ray photoelectron spectroscopy $(XPS)^{16,19,26,29,30}$ are the most commonly employed characterization techniques. As these mostly probe the local environment rather than intermediate or long range properties, no information about the stacking disorder of melon-type materials have been obtained. However, real world applications of carbon nitrides necessitate an understanding of the structure-property relationship and mechanistic aspects of their catalytic performance, both of which are intrinsically linked to their 3D architecture.

The current work is thus dedicated to unravelling the 3D arrangement of neighboring layers in highly condensed graphitic carbon nitride materials. Due to the inherent disorder we rely on a combination of quantitative multinuclear solid-state NMR spectroscopic experiments and chemical modeling based on force fields. Within the restraints of the diffraction data the force field calculations allow for a fast buildup of structure models and the determination of homo- and heteronuclear second moments provides a straightforward and accurate verification.⁴⁶ As a case study we used Liebig's melon,⁴⁷ which is a 1D carbon amide imide nitride polymer with an ideal composition $C_6N_9H_3$.⁹ The nanocrystalline material was recently investigated by X-ray and electron diffraction, solid-state NMR spectroscopy, and theoretical calculations.⁹

Melon consists of a layer structure well-ordered in two dimensions, but with a pronounced stacking disorder in the third. The structure of the layers was solved by projection from electron diffraction data (plane group p2gg with a = 16.7 Å, b = 12.4 Å, $\gamma = 90^{\circ}$ and $Z = 4)^{9}$ and is shown in Fig. 1. It is composed of infinite 1D chains of NH-bridged heptazine (C₆N₇(NH₂)₃)) monomers with one terminal NH₂ group. The chains exhibit a zigzag-type geometry and form a hydrogen bonded, 2D planar array. In agreement with other carbon nitrides (see above), the interlayer distance amounts to 3.2 Å.

The arrangement between neighboring layers exhibits a high degree of lateral disorder, which may be attributed to the much weaker dispersion forces dominating the interlayer arrangement. However, it is not completely stochastic as the X-ray powder diffraction data reveal asymmetric reflections besides the 00*l* series.⁹ That certain layer arrangements are more favored than others is also indicated by the modulated intensity of the "streaking" observed in the SAED (selective area electron diffraction) patterns viewed perpendicular to c* (see Fig. 1 (bottom)). The missing cone-like smearing in the *hk*0 plane implies disorder originating from translational planar defects rather than rotational ones.⁹ Therefore, energetically similar stacking modes with various, yet not

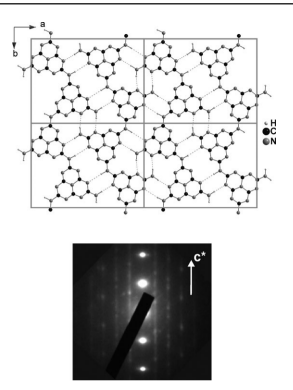


Fig. 1 (Top) Structure model for individual layers of melon.⁹ (bottom) SAED pattern perpendicular to the c axis of the reciprocal lattice.

statistically distributed translational layer offsets are likely to contribute to the overall 3D structure.

2. Experimental

2.1 Sample

For a detailed description of the sample preparation for the work presented here refer to ref. 9. The sample was enriched in ^{15}N to about 25%.

2.2 Solid-state NMR spectroscopy

Solid-state NMR experiments were recorded on a commercial Avance II spectrometer (Bruker) operating at a proton frequency of 300 MHz. Samples were contained in 4 mm ZrO₂ rotors for the ¹⁵N and ¹³C measurements and in 2.5 mm ZrO₂ rotors for the ¹H experiments. Both were mounted in standard triple resonance MAS probes (Bruker). ¹³C and ¹H shifts are reported with respect to TMS, ¹⁵N shifts were referenced to nitromethane. Data collection of the ¹⁵N and ¹³C experiments was performed under broadband proton decoupling using a SPINAL-64 sequence with a nutation frequency of about 70 kHz.⁴⁸ The repetition times were set between 5 and 10 s. In the 2D experiments quadrature detection of the *F*₁ domain was achieved *via* TPPI and a SPINAL-64 decoupling (nutation frequency about 70 kHz) was applied during *t*₁.

The 2D fpRFDR experiment was implemented as described in ref. 49 (see Fig. 2(a)). A spinning frequency of 12.5 kHz was used which corresponds to a 180° pulse length of 24 µs for the fpRFDR mixing pulses. The overall fpRFDR mixing time

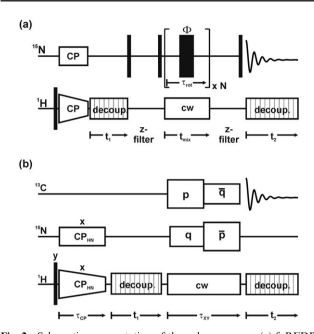


Fig. 2 Schematic representation of the pulse sequences (a) fpRFDR and (b) *i*DCP. *cw* denotes constant-wave decoupling times and *decoup*. composite pulse decoupling times (for further details refer to the text).

 (τ_{mix}) was set to 115 ms and an XY-16 phase cycle (Φ) was applied for the 180° pulses.⁵⁰ As preparation a ramped CP block was employed with a contact time of 3 ms. Two additional delays of 30 ms, before and after the fpRFDR mixing block, act as *z*-filters. t_1 was incremented rotor-synchronized (80 µs) collecting 128 FIDs with 128 transients each. During the fpRFDR mixing time a cw decoupling with a nutation frequency of 85 kHz was applied on the proton channel.

The 2D-*i*DCP experiment was performed as presented in ref. 51 (see Fig. 2(b)). The spinning frequency was set to 5 kHz, the ¹H–¹⁵N CP contact time to 6 ms and the overall ¹⁵N–¹³C contact time to 10 ms. As shown in Fig. 2(b) the *i*DCP pulse sequence of the ¹⁵N–¹³C contact block τ_{XY} is divided into two parts: during the first half the nutation frequency of ¹⁵N accounts to $\nu^1(^{15}N) = q \cdot \nu_{Rot}$ and of ¹³C to $\nu^1(^{13}C) = p \cdot \nu_{Rot}$ where |p - q| = 1. Then nutation frequencies are switched to $\nu^1(^{15}N) = p \cdot \nu_{Rot}$ and $\nu^1(^{13}C) = q \cdot \nu_{Rot}$. Simultaneously the phase of both channels is switched by 180° (indicated by the bar in Fig. 2(b)). For our experiment *p* and *q* were set to 7 and 8, respectively. A cw decoupling with a nutation frequency of 100 kHz was used during the complete ¹⁵N–¹³C contact time. Overall 124 experiments with 512 transients each were performed with a t_1 increment of 100 µs.

For the determination of the hydrogen substructure of melon we employed the strategy described previously in ref. 46. Details for the implementation of the LG-CP, the R14 pulse and the CP sequences can be found therein. For the LG-CP experiments^{52,53} a spinning frequency of 12 kHz was used. The nutation frequency of the proton channel during the spin-lock was set to 60 kHz, to which the LG offset as well as the ¹³C nutation frequency were optimized. The ¹⁵N excitation frequency was set to the middle of the NH and NH₂ signals. A build-up curve of 36 1D experiments with 96 transients each

was recorded. The LG-CP time τ_{CP} was thereby increased in steps of 20 µs up to a maximum of 700 µs.

The symmetry-based ¹H-R14⁶ double-quantum experiments^{54,55} were set-up with an excitation-reconversion- $\pi/2$ —acquisition scheme. For symmetric build-up curves the excitation and the reconversion time were incremented simultaneously ($\tau_{exc} = \tau_{rec}$) while for the constant-time (CT) procedure the reconversion time was decreased starting from τ_{total} with equal timesteps as the excitation time was increased $(\tau_{exc} + \tau_{rec} = \tau_{total})$. For the experiments at a spinning frequency of 30 kHz a simple 180°_{Φ} pulse was used for the R block of the $R14_2^6$ sequence. In contrast, at a spinning frequency of 15 kHz the composite pulse $90^{\circ}_{\Phi} - 270^{\circ}_{-\Phi}$ was applied for a better chemical shift compensation. The phase Φ was optimized to 77.04° for $\nu_{\rm rot} = 30$ kHz and to 76.29° for $\nu_{\rm rot}$ = 15 kHz. A total of 32 1D experiments were measured for the symmetric build-up curves while for the CT curves 17 and 9 1D experiments were performed for $\nu_{rot} = 30$ and 15 kHz, respectively. The increment of the excitation time in subsequent 1D experiments accounted for one RR' element which corresponds to 9.52 μ s at $\nu_{rot} = 30$ kHz and to 19.05 μ s at $\nu_{\rm rot} = 15$ kHz. 16 repetitions for each 1D experiment were performed.

For the cross polarization (CP) experiments a standard CP experiment was used.⁵⁶ The spinning frequency was set to 4 kHz in order to prevent suppression of spin-diffusion. For the optimization of the Hartmann-Hahn match attention was paid to ensure a Hartmann–Hahn condition of $n = \pm 1$. Therefore nutation frequencies were preoptimized using 90° pulses and then fine-tuned in the CP experiment. 22 1D CP-experiments were measured incrementing the CP contact time from 0.1 to 20 ms. 1024 transients were recorded for each FID. In advance of the measurement of the CP build-up curves we checked the dephasing behavior of the magnetization of ¹H and ¹³C during the spin-locks using the pulse sequences presented in ref. 56. These revealed no significant dephasing in the relevant timescale of 20 ms ($T_{1\rho}$ was estimated to some hundred milliseconds for both ¹H and ¹³C). $T_{1\rho}$ could thus be neglected in the analysis of the CP data.

2.3 Solid-state NMR simulations

Simulations for the LG-CP and the R14⁶₂ experiments were performed with the simulation package SIMPSON.⁵⁷ For the fits of the simulated curves to experimental data least-square fits were performed using the fit routine *lsqcurvefit* of MATLAB⁵⁸ in a self-developed script which internally calls the program SIMPSON.

For the simulations of the LG-CP pulse sequence spinsystems consisting of one NH and NH₂ group were built. CSA parameters were found to have only minor influence on the build-up curves and were therefore omitted. For the same reason the H–N–H angle of the NH₂ group could be set fixed to the value derived from force field calculations. However, for a correct simulation of the LG-CP conditions the proton isotropic chemical shift was set to the required LG offset of –42426 Hz corresponding to the spinning frequency of 12 kHz and to the ¹H nutation frequency of 60 kHz. As the build-up curves for the two NH signals as well as for the two

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 NH_2 signals run identically their intensities were summed up in each case. For both intensities (NH and NH_2) the optimal N–H distance was then determined with MATLAB as described above under variation of the N–H distance and an overall scaling factor.

The procedure of the extraction of the H–H distances from the $R14_2^6$ data was described in detail in ref. 46. A spin-system consisting of two protons and their common dipole coupling was used.

For the calculation of the 2nd moments (M_2) (see Results) a MATLAB script was developed. For each structure model both the heteronuclear $M_2^{\rm HC}$ and the homonuclear $M_2^{\rm HH}$ were calculated including atoms within a sphere with a radius of 30 Å around each crystallographically independent atom. For the $M_2^{\rm HH}$ the individual homonuclear moments for all hydrogen atoms of the unit cell were summed up as all hydrogen atoms participate in the polarization transfer during the CP spinlock. The individual $M_2^{\rm HC}$ of the CN₃ and of the CN₂ (NH_x) atoms were added separately to give the heteronuclear $M_2^{\rm HC}$ (CN₃) and $M_2^{\rm HC}$ (CN₂(NH_x)), respectively.

2.4 Computational methods

The force field calculations were carried out with the program package MS modeling 4.0^{59} using the Dreiding force field. As a starting point for the layer displacements the force field optimized unit cell from ref. 9 was taken as well as the Hirschfeld charges determined from a cluster calculation on the DFT level. For the calculation of the energy, structure models with lateral displacements were generated by shifting these layers in steps of 4% of the cell constants with a fixed layer distance of 3.19 Å. The energies were then calculated without further optimization.

For the geometry optimization of the minima of the force field hypersurface the unit cells with shift vectors $\vec{S} = [16\% \ 12\%]$, $\vec{S} = [0\% \ 24\%]$ and $\vec{S} = [36\% \ 36\%]$ were fully relaxed (optimization of all fractional coordinates and all cell constants) in space group *P*1 with the Quasi-Newton algorithm, the Dreiding force field and Hirschfeld charges.

3. Results

3.1 Phase homogeneity

Fig. 3(a) shows a 15 N CP-MAS spectrum of melon. Due to a strong and very well optimized proton decoupling we can resolve two signals each for both the NH and the NH₂ groups. The 2D structure solution for the layers from the electron diffraction data however exhibits only one crystallographically independent nitrogen atom for both the NH and the NH₂ groups. Either these separated signals originate from different phases or the 2D symmetry and thus the equality of the N atoms of the electron diffraction data is not conserved in three dimensions. The latter may be caused by a non-eclipsed arrangement of the layers.

Therefore we examined the phase homogeneity of our sample with a ¹⁵N-fpRFDR spin-diffusion experiment (see Fig. 3(b)). With a mixing time of 115 ms the signals have common cross signals if the involved nuclei are positioned within a range of several nanometres and thus are in the same

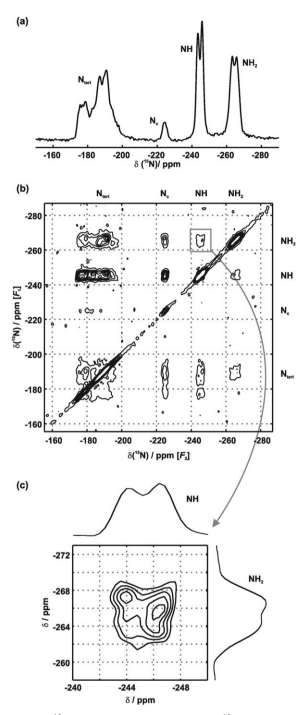


Fig. 3 (a) ¹⁵N CP-MAS spectrum of melon; ¹⁵N-fpRFDR spindiffusion spectrum with a mixing time of 115 ms: complete spectrum (b) and extension of the NH₂ \rightarrow NH correlation peak (c).

phase. As preparation block for this spectrum a short CP mixing time of 3 ms was used which guarantees a high excitation of the NH and the NH₂ signals. For an effective excitation of the central nitrogen atom (N_c) a longer CP mixing time would have been necessary which is why in the F_1 dimension the corresponding signal is very weak ($\delta(N_c) \approx 225$ ppm). Nevertheless, the spectrum shows cross signals for all other signals in the F_1 dimension with the N_c signal in the F_2 dimension. This reflects an intensity transfer

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from all other nuclei to the N_c nucleus and thus proves that all phases of the sample must exhibit a heptazine subunit for which the N_c atom is unique.

The enlarged depiction of the correlation signals of the NH and NH₂ groups in Fig. 3(c) reveals that both NH signals correlate with both NH₂ signals and *vice versa*. Additionally, all tertiary nitrogen signals (N_{tert}) show doublets as correlation signals with the NH_x signals. Thus, the NH, NH₂ and N_{tert} groups must be positioned in the same phase. The split NH and NH₂ signals thus have to represent crystallographically inequivalent NH and NH₂ groups within one phase.

3.2 ¹³C-Signal assignment

In the following section we will employ a strategy suited to derive information about the three-dimensional structure of melon based on ¹H-¹³C CP rates.⁴⁶ The prerequisite for this analysis, however, is an exact knowledge of the hydrogen substructure which will be determined in the next sections. Additionally, it considerably improves the expressiveness of the rate experiments if all ¹³C signals are assigned so that no sum intensities of signals have to be used. The ¹³C CP-MAS spectrum of melon in Fig. 4(a) reveals two signal groups at 157 and 164 ppm which were previously assigned to the CN_3 and the $CN_2(NH_x)$ carbon atoms, respectively.⁹ However the origin for the shoulder in the signal at 164 ppm could not vet be determined. In order to differentiate between the CN₂(NH) and the CN₂(NH₂) carbon signals we performed a 2D ¹⁵N-¹³C correlation experiment. Fig. 4 shows the 2D-iDCP experiment of melon with ¹⁵N-¹³C mixing time of 10 ms which guarantees an excitation of bonded N-C connectivities only.

As expected the spectrum shows correlations with both ¹³C signals for the outer tertiary nitrogen atoms (N_{tert}), as each N_{tert} atom is bonded to a CN₃ and a CN₂(NH_x) carbon atom. The central nitrogen atom (N_c) at δ (¹⁵N) = -225 ppm correlates only with the CN₃ carbon atom in agreement with the chemical structure. This signal is very weak due to the poor CP excitation of the N_c signal (see fpRFDR experiment). Additionally, for the CN₂(NH_x) signal correlates with the main resonance of the CN₂(NH_x) signal (166 ppm) while the NH signal reveals a common signal with both the main signal (166 ppm) and the shoulder (163 ppm).

This distribution of ¹³C chemical shifts however impedes an extraction of individual intensities for the $CN_2(NH_2)$ ¹³C signals, as in the main signal both overlap heavily. In the analysis of the ¹H–¹³C cross-polarization experiment we therefore have to restrict it to the analysis of two build-up curves, namely for the CN_3 and the $CN_2(NH_x)^{13}C$ signals.

3.3 Localization of hydrogen atoms

We recently proved that the exact knowledge of the hydrogen substructure is a prerequisite for a reasonable quantitative analysis of CP rates.⁴⁶ We demonstrated that even the accuracy of theoretical calculations with DFT methods is not sufficient. Therefore we established a strategy for an accurate localization of the hydrogen atoms which includes the measurement of

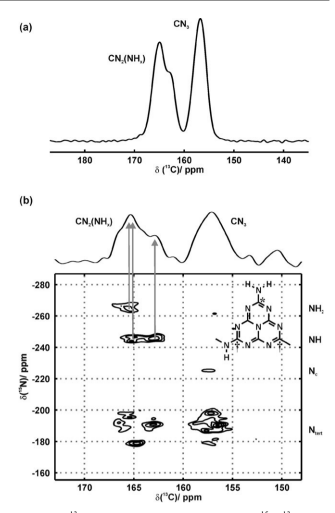


Fig. 4 (a) ¹³CP-MAS spectrum of melon; (b) 2D ¹⁵N–¹³C *i*DCP spectrum⁵¹ of melon with a ¹⁵N–¹³C contact time of 10 ms. The correlation peaks indicate through-bond ¹⁵N–¹³C proximities. The gray arrows mark the correlations of the NH and NH₂ signals with the CN₂(NH_x) signals. The insert shows the subunit of the structure of melon where * denotes CN₂(NH₂) and † CN₂(NH) groups.

N–H and H–H distances *via* NMR. For melon we used a Lee–Goldburg cross-polarization (LG-CP) experiment for measuring the NH distances in the NH and the NH₂ group and a R14⁶₂ pulse sequence for determining the H–H distance in the NH₂ group. The experiments were performed as outlined in ref. 46. These data were subsequently incorporated into the force field structure under retention of the torsion angles for the NH_x groups.

3.3.1 Determination of N–H distances. Fig. 5 shows the ¹⁵N LG-CP build-up curves for the NH and the NH₂ groups of melon. As demonstrated in ref. 46 the evaluation of these curves can be performed without taking into account further remote protons of the structure if only the first oscillation is utilized. This results in an N–H distance of 1.024(1) Å for the NH group and of 1.043(3) Å for the NH₂ group.

3.3.2 Determination of H–H distances. The determination of the H–H distance allows for a calculation of the H–N–H angle of the NH_2 group when combined with the N–H

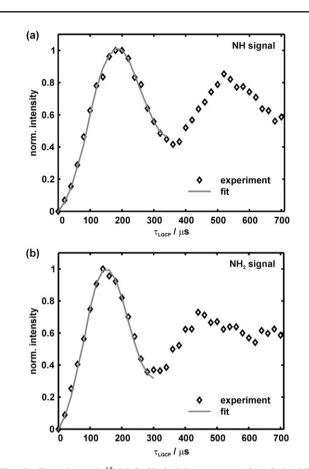


Fig. 5 Experimental ¹⁵N-LG-CP build-up curves (\diamond) of the NH (a) and the NH₂ (b) signal of melon with fit (line) using a 2- and a 3-spin system for the NH and NH₂ curve, respectively.

distance. It was derived with the symmetry-based doublequantum sequence R14⁶₂. Both symmetric and constant-time build-up curves were measured at spinning frequencies (ν_{rot}) of 15 and 30 kHz. Especially the averaged value of these four experiments yields a very accurate H–H distance. Fig. 6(a) shows the ¹H spectrum of melon with its deconvoluted resonances. The ¹H signal of the NH group can be found at 9.5 ppm while the signal of the NH₂ arises at 10 ppm. The two further minor signals are to be assigned to impurities from the probe. This assignment is corroborated since in the doublequantum excitation curve only the short H–H distance of the NH₂ group results in fast oscillations while the NH signal shows a slow build-up without oscillations. This is presented exemplarily in Fig. 6(b) for the symmetric build-up curve at $\nu_{rot} = 30$ kHz.

Our previous studies on melem revealed that the influence of weaker H–H couplings from remote hydrogen atoms have to be considered for long excitation times only.⁴⁶ Restricting the analysis however to the first oscillations simulations with 2-spin systems can be employed for a reasonable analysis of build-up curves. We could show for melem (C₆N₇(NH₂)₃) that this is fulfilled with an excitation time of 80 µs in symmetric and for an overall time $\tau_{total} = \tau_{exc} + \tau_{rec}$ of 160 µs in constant-time build-up curves. Since in melon the hydrogen atoms are more diluted than in melem these values are appropriate for melon as well.

Table 1 ${}^{1}H^{-1}H$ distance of the NH₂ group of melon from the R14⁶₂ experiments

$\nu_{\rm rot}$	30 kHz	15 kHz
Symmetric	1.77(2) Å	1.82(2) Å
Constant-time	1.75(1) Å	1.80(1) Å

In Fig. 6(b)–(e) the experimental symmetric and constanttime build-up curves of the NH₂ signal of melon, each with $\nu_{rot} = 15$ and 30 kHz, are shown. Additionally, the Figure depicts the 2-spin simulations matching best the experimental data. The thus obtained distances are given in Table 1, their average value amounts to 1.785 Å. In combination with an N–H distance of 1.04 Å this corresponds to an H–N–H angle of 118.2° for the NH₂ group.

3.4 ¹H–¹³C cross-polarization experiments

The successful determination of the hydrogen substructure of melon allowed for the extraction of structure information with cross-polarization (CP) build-up curves which will be described in the following.⁴⁶ The initial slope for each signal can be extracted *via* the logarithmic plot of a normalized CP build-up curve. It represents the experimental mean CP rate constant $\langle k \rangle$. On the other hand $\langle k \rangle$ can be calculated from the rates $k_{\rm CP}$ of all crystallographic independent nuclei (*j*) overlapping to the corresponding signal in the ¹³C MAS spectrum:

$$\langle k \rangle = \frac{1}{N} \sum_{j}^{N} k_{\rm CP}(j) \tag{1}$$

The rates k_{CP} reflect the transfer of polarization from ¹H to ¹³C (or any other I = 1/2-nucleus) and are connected with the homo- and heteronuclear second moments M_2^{HH} and M_2^{HC} via

$$k_{\rm CP} = C_{\rm CP} \frac{M_2^{\rm HC}}{\sqrt{M_2^{\rm HH}}} \tag{2}$$

where the constant C_{CP} depends solely on physical constants and the experimental setup but contains no structural information. The second moment M_2^{XY} can be calculated from a structure model by

$$M_2^{XY} = \left(\frac{1}{2\pi}\right)^2 \frac{4}{5} \left(\frac{1}{2}\gamma_I \gamma_S \hbar \frac{\mu_0}{4\pi}\right)^2 \sum_i (r_i^{XY})^{-6}$$
(3)

Thus, for the verification of a structure model $\langle k \rangle$ can be determined for each observed signal while the second moments are computed from the model. The structure model used is then confirmed if the constant $C_{\rm CP}$ is found to be equal for all signals.^{46,60} In the following we will employ this condition for the determination of the lateral arrangement of the layers in melon.

In melon we can differentiate the two resolved ¹³C signals $CN_2(NH_x)$ and CN_3 as shown above. So it is possible to evaluate structure models with different lateral arrangements by the equality of $C_{CP}(CN_2(NH_x))$ and C_{CP} (CN_3) or, as it will be done in the following, by their ratio $C_{CP}(CN_3)/C_{CP}(CN_2(NH_x))$ which must equal 1 for the most likely models.

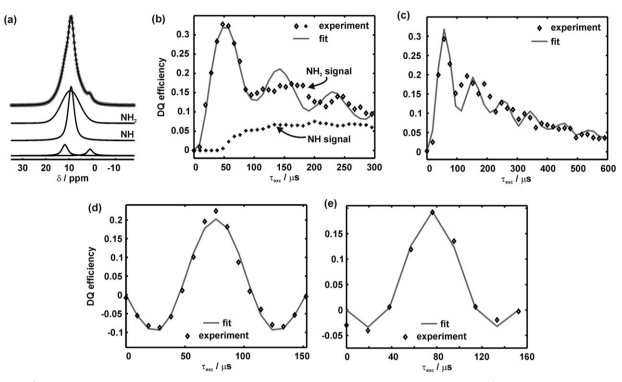


Fig. 6 (a) ¹H spectrum of melon (\bullet) with fit (top) and deconvoluted signals (bottom); (b)–(e) Experimental R14⁶₂ build-up curves (\diamond) of the NH₂ signal with a fit of a 2-spin systems (line): (b) symmetric curve at $\nu_{rot} = 30$ kHz; (c) symmetric curve at $\nu_{rot} = 15$ kHz; (d) constant-time curve at $\nu_{rot} = 30$ kHz; (e) constant-time curve at $\nu_{rot} = 15$ kHz. In (b) the experimental build-up curve of the NH signal is additionally shown (+).

Fig. 7(a) shows the logarithmic plot of the normalized CP build-up curves for the ¹³C signals of melon. From this data the initial slopes were determined (see lines in Fig. 7(a)) which result in the averaged rates $\langle k \rangle$ of 0.86×10^3 Hz for the CN₃ and of 2.59×10^3 Hz for CN₂(NH_x) signal.

For the calculation of the second moments the force field optimized structure from ref. 9 was utilized. The N–H distances and the H–N–H angle for the NH and the NH₂ groups were corrected according to the results of the NMR experiments presented above. Subsequently, structure models with varying lateral shifts in *a* and *b* were generated which in the following will be characterized by the shift vector $\mathbf{S} = [\Delta a \, \Delta b]$ where Δa and Δb are the lateral shifts along the *a* and *b* axis, respectively. Thus, the used models correspond to an AAA stacking of the layers of melon. For all these structures the corresponding second moments and, in combination with the experimental values for $\langle k \rangle$, the constants $C_{\rm CP}$ were computed for each of the two signals.

Fig. 7(b) depicts the dependency of their ratio $C_{CP}(CN_3)/C_{CP}(CN_2(NH_x))$ on the shift vector **S**. This hypersurface is symmetric in Δa and Δb due to the symmetry of the layers (see Fig. 1). Therefore, the following discussion will only refer to shift vectors between 0 and 50% for Δa and Δb covering the "asymmetric unit" of the hypersurface.

In the hypersurface the dark areas represent structure models with a ratio close to 1 which are the more suitable models. A wide range of ratios from 1.19(1) to 1.63(2) is covered. This shows that there are layer arrangements which reproduce the experimental data better than others and

thus, that from the CP data preferred lateral shifts can be identified. The best results with a ratio of 1.19 can be found for the shift vectors $\mathbf{S} = [0\% 50\%]$, $\mathbf{S} = [50\% 50\%]$ and $\mathbf{S} = [25\% 0\%]$. However, none of the models used reaches a ratio of 1. This can be explained by the fact that the structure of melon is disordered and we employ a well-ordered model which only approximately reflects the reality. The largest ratios arise for the shift vectors $\mathbf{S} = [0\% 0\%]$, $\mathbf{S} = [50\% 0\%]$ and $\mathbf{S} = [25\% 50\%]$.

In order to examine the influence of free relaxation of the structures on the ratio the structure models of the minima were optimized with force field methods. These optimizations resulted in minor changes of the geometry of the layers. Correspondingly, the ratio $C_{\rm CP}({\rm CN}_3)/C_{\rm CP}({\rm CN}_2({\rm NH}_x))$ of optimized cells differ only up to 0.02 from the previously used fixed layers which justifies their utilization for the hypersurface from Fig. 7(b).

Finally, we examined if more elaborate structure models result in a ratio closer to 1. Instead of the used AAA model we employed an ABA model with two different shift vectors for $A \rightarrow B$ and $B \rightarrow A$. Thus, layers possess unequal lateral arrangements with their two neighboring layers. However, with these models the minimal ratio once more amounted to 1.19. The structures of these minima represent combinations of the structural motifs of the minima in the AAA model, *e.g.* a shift of S = [0% 50%] for $A \rightarrow B$ in combination with a shift of S = [25% 0%] for $B \rightarrow A$. This demonstrated that primarily the adjacent layers contribute to the second moments while there is only a minor influence from more distant layers.

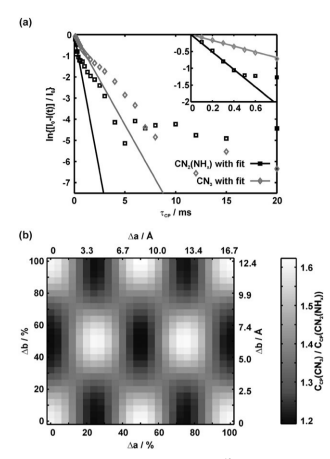


Fig. 7 (a) Logarithmic plot of the normalized ¹³C-CP build-up curves of melon (symbols) with linear fits of the initial gradient (solid line); (b) CP hypersurface: dependency of the ratio $C_{CP}(CN_3)/C_{CP}(CN_2(NH_x))$ on the lateral shift vector **S** = [$\Delta a \Delta b$].

3.5 Force field energies

The calculations of relative force field energies were used as complementary information for the determination of preferred lateral shifts in the layered structure of melon. Since the energy changes expected are caused by van der Waals interactions only, the variation on the energy hypersurface will be small and may even be smaller than the absolute error which arises in force field calculations. Even modern force fields usually do not reach an absolute accuracy better than 1-4 kcal mol^{-1,61,62}. To ensure the significance of the following analysis we, therefore, validated the Dreiding force field for use with melon. Already in a previous study⁹ it was demonstrated that this force field reproduces the lattice parameters ($a = 16.7-16.8 \text{ \AA}/b =$ 12.8 Å) as well as the interlayer distance (c = 3.19 Å) within the experimental error. Especially the observed planarity of the individual layers caused by the tight N-H···N hydrogen bond network is retained in excellent agreement. Shifting mainly rigid layers against each other, however, leaves the error in the absolute energies constant. By turning to relative energies this error is mainly cancelled which allows to discuss the energy hypersurface as well as structure models derived from the force field calculations based on laterally shifted layers.

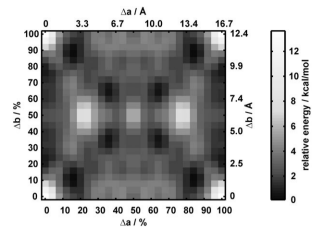


Fig. 8 Energy hypersurface: dependency of the relative force field energy on the lateral shift vector $\mathbf{S} = [\Delta a \ \Delta b]$.

We employed the same models as for the analysis of the CP rates but left the positions of the hydrogen atoms unchanged in order to guarantee consistency within the force field calculations. Fig. 8 shows the resulting energy hypersurface which is symmetric with respect to Δa and Δb as is the CP hypersurface. The color coding was chosen in a way to label those lateral shifts in black to dark gray which are up to 2 kcal mol⁻¹ above the global minimum and are thus energetically accessible at ambient temperature.

As most regions of the hypersurface are colored black to dark grey we can conclude that the lateral displacement of adjacent layers leads to a very flat energy profile. This is in agreement with the experimental results where a distribution of lateral shifts and thus a disorder is found. However, some lateral shifts are definitely unfavored with energies up to 12 kcal mol⁻¹ above the minimum. This explains why the disorder is not completely statistical but exhibits preferred arrangements.

The maximum on the hypersurface occurs at $\mathbf{S} = [0\% \ 0\%]$. This corresponds to the ecliptic position of the layers. Further local maxima are positioned at about $\mathbf{S} = [20\% \ 50\%]$ as well as at $\mathbf{S} = [50\% \ 50\%]$. Here the heptazine units in adjacent layers are arranged partially ecliptic.

The minima are situated at about S = [0% 24%], S = $[16\% \ 12\%]$ and S = $[36\% \ 36\%]$. For an evaluation of the quality of hypersurface structures these minima were optimized freely. During the optimization the cell constants and angles showed only minor changes (see Table 2) and the interlayer distance d with 3.2-3.3 Å reproduces very well the value obtained from the diffraction data. In addition, the geometry of the heptazine strands remained unaltered though they slightly tilt out of the ab plane (see Fig. 9). After the optimization the minima on the hypersurface (Fig. 8) at S = [0% 24%], S = [16% 12%] and S = [36% 36%] shift to S = [0% 22.8%], S = [12.4% 7.2%] and S = [35.0% 37.2%],respectively. The small deviations demonstrate that the energy surface based on single point calculations (Fig. 8) reproduces the energetic situation in the structure of melon quite well. The deepest minimum occurs at $S = [16\% \ 12\%]$ and is roughly one kcal mol^{-1} lower in energy than the other two minima.

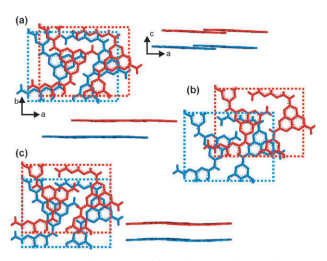


Fig. 9 Optimized structures with minimal energies at shift vectors (a) $\mathbf{S} = [12.4\% \ 7.2\%]$, (b) $\mathbf{S} = [35.0\% \ 37.2\%]$ and (c) $\mathbf{S} = [0.0\% \ 22.8\%]$. They correspond to the three minima on the energy hypersurface at $\mathbf{S} = [16\% \ 12\%]$, $\mathbf{S} = [36\% \ 35\%]$ and $\mathbf{S} = [0\% \ 24\%]$ (compare Fig. 8).

4. Discussion

In the following we will compare the results for the lateral arrangement in melon from the CP experiment and the force field calculations. To simplify this comparison in both cases the dark areas of Fig. 7 and 8 represent shift vectors of favored structures. It becomes clear that these two hypersurfaces show a similar profile. The only disagreement between the two methods can be identified for $\mathbf{S} = [50\% 50\%] = [8.4 \text{ \AA } 6.4 \text{ \AA}]$ which is favored by the NMR data but has an energy which is about 12 kcal mol^{-1} higher than the global minimum and is, thus, definitely disfavored from energetic considerations. Therefore, it was discarded as a reasonable lateral arrangement. A possible explanation for the contradictory preference of the NMR data for this structure can be found in the use of sum intensities of ¹³C signals which had to be employed for the CP rate analysis. As we showed recently for homonuclear spindiffusion experiments, such overlapping signals can lead to a good agreement of experimental and calculated build-up curves even for incorrect structure models.63 The structure models discussed in the following however agree for the two employed methods (CP rates and energies) and thus represent authentic maxima and minima.

Both the NMR and the energy hypersurfaces favor shifting along the *b*-axis while the shift parameter for the *a*-axis remains zero (Fig. 7(b) and 8). The minimum for the force field energy is located at $\mathbf{S} = [0\% 20\%] = [0.0 \text{ Å} 2.6 \text{ Å}]$ while for the CP NMR data it is found at $\mathbf{S} = [0\% 50\%] = [0.0 \text{ Å} 6.4 \text{ Å}]$. However, at about $\mathbf{S} = [0\% 40\%] = [0.0 \text{ Å} 5.1 \text{ Å}]$ in both hypersurfaces the values lie only slightly above the minima. Thus, for further discussion this shift vector is taken as a mean minimum (compare Fig. 7(b) and 8). The second region where both the NMR and energy hypersurface share minima is situated at $\mathbf{S} = [22\% 5\%] = [3.7 \text{ Å } 0.6 \text{ Å}]$. In this case the position of the minima is nearly identical for both hypersurfaces. Clearly disfavored in both hypersurfaces is the ecliptic arrangement with $\mathbf{S} = [0\% 0\%]$. Further light-colored and thus unfavored areas are found around $\mathbf{S} = [50\% 0\%] = [8.4 \text{ Å } 0.0 \text{ Å}]$ and $\mathbf{S} = [25\% 50\%] = [4.2 \text{ Å } 6.4 \text{ Å}]$.

Fig. 10(a) and (b) show the structures of the two mean minima at S = [0% 40%] = [0.0 Å 5.1 Å] and S = [22% 5%] =[3.7 Å 0.6 Å]. Both reveal geometries where the heptazine units of adjacent layers are arranged in a staggered way. Fig. 10(c) and (d) depict the layer arrangement for two of the three mean maxima on the hypersurfaces at $S = [50\% \ 0\%]$ = [8.4 Å 0.0 Å] and S = [25% 50%] = [4.2 Å 6.4 Å]. The ecliptic arrangement at $S = [0\% \ 0\%]$ was omitted in Fig. 10 due to the perfect congruence of both layers. The structures displayed in Fig. 10(c) and (d) as well as the ecliptic arrangement have in common that at least two out of four heptazine units per unit cell are arranged in an ecliptic mode with adjacent layers. Thus, from both the minima and the maxima structure models it may be concluded that the preferred relative positions of the layers in melon are dominated by the purpose of obtaining as little overlap for the heptazine rings as possible in order to maximize the π - π interactions.

5. Conclusion

In the presented publication we utilized a combination of solid-state NMR experiments and force field calculations to derive information about the stacking disorder of a layered material.

Based on a ¹⁵N spin-diffusion experiment the phase homogeneity of the sample could be established on the nanometre scale. Thus the two ¹⁵N-signals for the NH and the NH₂ group must arise from a reduction of the 2D layer symmetry by the 3D arrangement. For a further assignment of the overlapping resonances of the $CN_2(NH_x)$ ¹³C-signal we performed an *i*DCP experiment. We could differentiate between the $CN_2(NH_2)$ and the two $CN_2(NH)$ signals. However, as one of the $CN_2(NH)$ signals and the $CN_2(NH_2)$ signal overlap tightly a determination of their isolated intensities was not possible. This implied for the following CP experiment that the sum intensity of the $CN_2(NH_x)$ signals had to be employed wherefore only two CP build-up curves (CN_3 and $CN_2(NH_x)$ could be evaluated.

Table 2 Relevant parameters for the three minima of the energy hypersurface (Fig. 8): lateral shift vector ($\mathbf{S}_0 = [\Delta a \ \Delta b]$) before optimization and cell parameters (*a*, *b*, *c*, α , β , γ), lateral shift vector ($\mathbf{S} = [\Delta a \ \Delta b]$), layer distance (*d*) and force field energy (*E*) for the optimized structures

\mathbf{S}_{0} [%]	$a/ m \AA$	$b/{ m \AA}$	$c/{ m \AA}$	α (°)	β (°)	γ (°)	S [%]		$d/{ m \AA}$	$E^a/\text{kcal mol}^{-1}$
0 24	16.75	12.83	4.38	48.16	90.0	90.0	0.0	22.8	3.26	-2.09
16 12	16.73	12.83	3.95	76.48	58.27	90.1	12.4	7.2	3.23	-3.38
36 36	16.77	12.83	8.24	54.6	44.6	89.9	35.0	37.2	3.27	-2.06
^a Energies	are given rela	tive to the gl	obal minimu	um of the ener	rov hypersurf	ace in Fig 8				

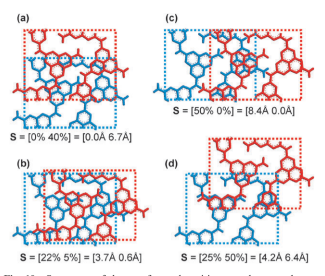


Fig. 10 Structures of the two favored positions on the mean hypersurface: (a) $\mathbf{S} = [0\% \ 40\%]$ and (b) $\mathbf{S} = [22\% \ 5\%]$. Layer arrangements for disfavored stacking scenarios: (c) at $\mathbf{S} = [50\% \ 0\%]$ and (d) $\mathbf{S} = [25\% \ 50\%]$.

For the CP experiment additionally an accurate knowledge of the position of the protons in melon was necessary. We determined the NH distances with the LG-CP experiment to 1.02 Å for the NH group and to 1.04 Å for the NH₂ group. The H–N–H angle of the NH₂ group was measured via the H–H distance with the $R14_2^6$ pulse sequence. It was found to be 118.2°. These values were incorporated into the force field structures under retention of the torsion angles in order to generate structure models for the evaluation of the CP build-up curves. In addition, we computed the force field energies for structures with various lateral shifts along the a and b axis of the 2D structure. Thus, both methods gave information about preferred lateral arrangements of neighboring layers in melon. The comparison of the corresponding hypersurfaces revealed a very good agreement of these completely independent approaches. Structures with (partially) eclipsed arrangements gave maxima on the hypersurfaces and are thus disfavored. These are located at the shift vectors $\mathbf{S} = [0\% \ 0\%]$, S = [25% 50%] = [4.2 Å 6.4 Å] and around S = [50% 0%] =[8.4 Å 0.0 Å]. The lateral shifts preferred in both methods are situated at S = [20%-10%] = [3.3 Å 0-1.3 Å] and around $\mathbf{S} = [0\% 40\%] = [0.0 \text{ Å } 6.7 \text{ Å}]$. Both correspond to structures with a displacement for the heptazine rings in neighboring lavers.

To conclude, we were able to obtain information about the 3D structure of melon although the material is disordered and no information can be gained from diffraction techniques. Due to the convincing results from solid-state NMR spectroscopy we are confident that this method will increasingly be utilized for the derivation of structure information in C/N materials which are mostly semicrystalline or even amorphous.

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