

Formation of one-dimensional quantum crystals of molecular deuterium inside carbon nanotubes

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Abstract

Crystallization under stringent cylindrical confinement leads to novel quasi-one-dimensional materials. Substances with strong cohesive interactions can eventually preserve the symmetries of their bulk phase compatible with the restricted geometry, while those with weak cohesive interactions develop qualitatively different structures. Frozen molecular deuterium (D_2), a solid with a strong quantum character, is structurally held by weak dispersive forces. Here, the formation of one-dimensional D_2 crystals under carbon nanotube confinement is reported. In contradiction with its weak cohesive interactions, their structures, scrutinized using

neutron scattering, correspond to definite cylindrical sections of the hexagonal close-packed bulk crystal. The results are rationalized on the grounds of numerical calculations, which point towards nuclear quantum delocalization as the physical mechanism responsible for the stabilization of such outstanding structures.

Keywords: Quantum Solids, 1D crystals, Confinement, Molecular Hydrogen, Neutron Scattering

1. Introduction

Carbon nanotubes are used to study crystallization under cylindrical constraints as well as to obtain a plethora of one-dimensional (1D) materials [1, 2, 3]. These new materials emerge, owing to the geometrical restriction, as crystalline phases with structures unlike their bulk counterparts (see chapter 5 in [3]). Some substances with strong cohesive interactions can retain their bulk crystallinity down to the strict one-dimensional limit, in which all the crystal constituents are simultaneously bulk and surface [4]. However, in general, for a certain level of geometrical restriction, crystalline arrangements with symmetries alien to the bulk phases develop [5]. As expected, this is the rule if the cohesive interactions are weak, for instance, hydrogen bonding in the archetypal case of water [6, 7, 8].

For purely dispersive interactions, molecular hydrogen (H_2) is an outstanding case owing to the strong quantum character of its condensed phases [9, 10, 11]. In particular, crystalline H_2 and its deuterium isotopologue (D_2) are prominent examples of quantum solids. Such crystals are characterized by a quantum nuclear delocalization sizable compared to the intermolecular distance, eventually rendering the harmonic crystal approximation useless [12]. The effect is maximized in the case of H_2 , where quantum fluctuations are so dominant that make the crystal insensitive to thermal variations [10, 11]. Furthermore, in both crystals the molecules behave as quantum free rotators, which implies that the nuclear orientational degrees of freedom are fully delocalized at low temperatures. Both crystalline states are van der Waals solids with an interaction potential well of just -2.77 meV, while that of the binding potential to a graphite surface amounts to $\simeq -50$ meV [13]. Confinement within narrow carbon nanotubes can roughly double this depth.

Calculations without including the quantum nuclear delocalization predict new ordered arrangements not present in the bulk [14, 15, 16], some of them reminiscent of spiral configurations observed in the iodine 1D case [17, 18, 19]. The search for superfluid behavior has spurred intense theoretical efforts to deal with the quantum nature of the H_2 molecular movements [20, 21, 22, 23, 24]. The results, not free of controversy, seem to favor a quantum solid resembling neither the spiral arrangements nor the hexagonal close-packed (HCP) structure of the bulk phase [23]. In the case of molecular deuterium, nuclear wave function calculations hint at the possibility of HCP structures within narrow carbon nanotubes [25].

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On the experimental side, as far as solidification under strong cylindrical confinement is concerned, the results are scarce. In carbon nanotubes, there is only evidence of transitions from one- to two-dimensional arrangements of D_2 on the external grooves of carbon nanotube bundles [26, 27], but no results about crystal structures within the nanotubes. There are hints of HCP crystallization of D_2 in the cylindrical mesopores of silica-based MCM-41 materials [28], but with a pore diameter as wide as 2.5 nm, quite far from the 1D limit. In these studies D_2 is preferred because the strong incoherent neutron scattering of H_2 makes diffraction measurements impractical.

In this work, by resorting to neutron diffraction (ND), crystallization of molecular deuterium within multi-walled carbon nanotubes (MWCNT) is studied. We have succeeded in resolving the crystalline structure down to the strict 1D limit. In spite of the weakness of the cohesive interactions mentioned above, the 1D crystals preserve the structure of its bulk counterpart. The strong quantum character of the bulk crystalline D_2 is also preserved in the 1D confined phases: the large delocalization of the nuclear positions as well as the full quantum nature of the rotational degrees of freedom are manifest in the measured neutron diffraction. The structures of the discovered 1D crystals correspond to maximally symmetric cylindrical cuts of the bulk HCP crystal. Such structures were addressed after the failure in reproducing the measurements by fully atomistic molecular dynamics (MD) simulations as well as by dispersive density functional theory (DFT) calculations corresponding to our experimental conditions. Instead, DFT and MD approaches yield spiral-like arrangements in full agreement with energy considerations as well as consistent with previously reported calculations. Since such calculations do not deal with quantum nuclear effects, given the strong quantum nature of the observed structures we conclude that quantum delocalization in the nuclear degrees of freedom is essential to stabilize the discovered 1D crystals.

The paper is organized as follows: in section 2 the experimental procedures and setups are described. The results, in particular the finally resolved crystalline structures, are presented and discussed in section 3. In section 4 our findings are summarized and the main conclusions exposed. To lighten the exposition, the details of materials and methods as well as of the theoretical developments used in the data analysis are given as appendixes. Likewise, the large series of diffraction patterns explored along with a detailed description of the nonlinear fitting analysis used to resolve the structures are presented in a Supplementary Material file. For the sake of reproducibility, details of custom-made experimental setups and specifics of the numerical runs made are also given there. Finally, figures corresponding to some complementary measurements and tables of numerical parameters complete the Supplementary Material.

2. Experimental

Our main goal is to study the structural evolution of molecular deuterium during solidification under extreme cylindrical confinement. In short, we measured neutron diffraction (ND) in runs of 350 seconds

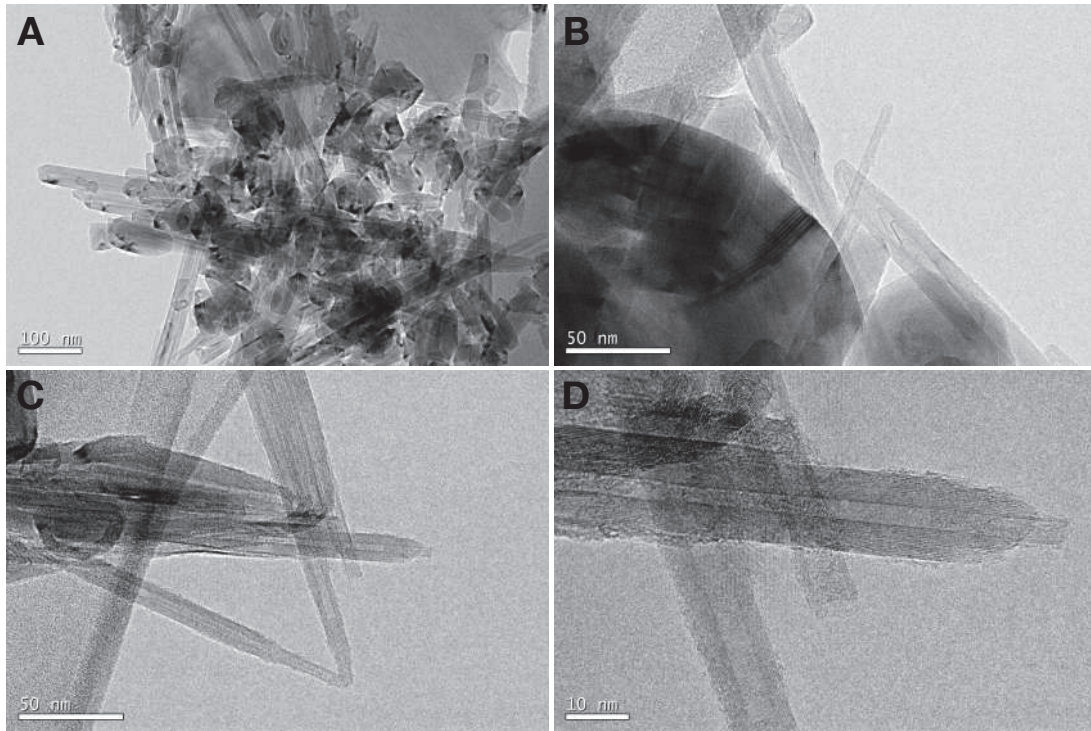


Figure 1: Some TEM pictures of the final processed samples. A corresponds to X 30000, B and C to X 100000 and D to X 300000

60 while submitting a sample of open MWCNTs to thermal cycles (between 24.5 K and 2 K) at four increasing loads of adsorbed D_2 at 20 K, corresponding to the equilibrium pressures 62 hPa, 175 hPa, 225 hPa, and 284 hPa. The details of the experiment design, the experimental setup as well as of the materials and methods used are given in Appendix B.

We synthesized MWCNTs by the arc discharge method. Like in the single-walled case, the tubes in the raw material are closed. In order to open them, we submitted the samples to an oxidation treatment (for details, see Appendix B.2). Transmission electron microscopy (TEM) of the final material (see Fig. 1) shows the expected structure for typical arc discharge MWCNT samples: It is quite heterogeneous, showing a wide variety of MWCNTs among a plethora of irregular polyhedral closed nanostructures. The surfaces do not present eroded scars or holes other than the openings of some nanotubes.

70 The adsorption of the sample was characterized through an isothermal D_2 adsorption/desorption measurement at 20 K using a custom-made setup. The isotherm in Fig. 2 shows during desorption a hysteresis beginning at the bulk vapor pressure. This is expected in this kind of samples. It comes from macro/mesopores, i.e., the interstitials among the various structures observed in the TEM. Consistently with the opening of a portion of the MWCNTs, the isotherm also presents a small, but noticeable, almost
 75 vertical increase at zero pressure, a feature revealing the presence of micropores of molecular dimensions

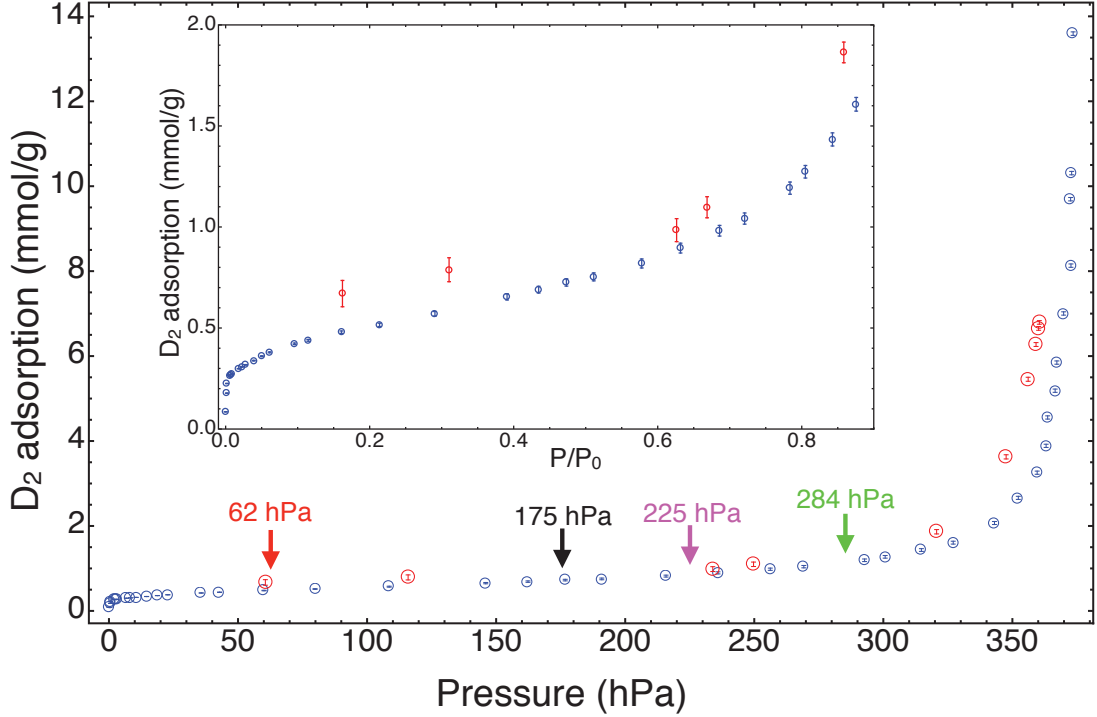


Figure 2: D₂ adsorption (blue)/desorption (red) isotherm at 20.23 ± 0.03 K for the sample of open MWCNTs. The amount of adsorbed gas is given in millimole per gram of sample. The arrows label the initial working points of the thermal cycles during the neutron diffraction measurements as described in the text. The inset is a zoom of the same data, focusing in the lower adsorption range in units of the bulk vapor pressure at 20.23 K (364.9 hPa [31]).

[30]. At sufficiently low loads of D₂, which correspond to low pressures in the isotherm, only the MWCNTs with the narrowest opened inner tubes are filled.

The chosen instrument was D20 at the Laue-Langevin Institute (ILL) in Grenoble, France, since it is a diffractometer of very high flux which allows to address very small quantities of scatterers in reasonable acquisition times (see details in Appendix B.6).

With respect to the neutron scattering, given the lightness of D₂ molecule, energy transfers between the incident neutron and the molecule affect the diffraction signal in a non-perturbative way. Usually this inelastic component is obtained heuristically since a proper correction requires a knowledge of the complete scattering law. Based on the neutron scattering theory for diatomic molecules developed by Sears [32], we have worked out an ansatz for the energy-integrated signal accounting for the observed $I_D(Q)$ that incorporates inelasticity without relying in any heuristic correction. From such an ansatz, the inelastic correction is given by a background, $bg(Q^2)$, expressible as an expansion in Q^2 (up to second order in practical terms). Physically, $I_D(Q)$ decomposes in a coherent and an incoherent part. The incoherent part stems from the random distribution of nuclear spin states along the sample. Formally is like the

self-scattering contribution coming out from the scatterers. The coherent part of $I_D(Q)$ depends on the orientationally-averaged structure factor of the molecular center-of-mass (COM) specific arrangement of D_2 molecules. It can be calculated efficiently from the pair distribution function, i.e., the table of COM pair distances $d_{ij} = |\langle \mathbf{R}_i \rangle - \langle \mathbf{R}_j \rangle|$ and its associated multiplicities m_{ij} , where \mathbf{R}_i is the position vector of the i molecule. An additional key ingredient to reproduce the ND coherent signal is a form factor corresponding to a spherical shell of diameter equal to the distance between the molecular nuclei, even in the solid phases at low temperature. This is a consequence of the nuclear orientational full delocalization. Likewise, the translational degrees of freedom also present a very significant quantum delocalization, which induces a large root mean square (rms) COM displacement. As explained in the next section, the observed diffraction signal reveals also the presence of oriented small crystals (or crystal) through barely discernible Bragg peaks. Although such precursor peaks can be ignored, we must consider the corresponding diffuse scattering, i.e., the diffraction signal between Bragg peaks in a crystal, which arises from such oriented crystallites. This is well approximated by a term of the form $1 - \exp(-Q^2 u_{3D}^2/3)$ plus an incoherent part, where u_{3D} is the rms molecular displacement of the molecules belonging to such crystallite. There is also an instrumental background whose major contribution to $I_s(Q)$ is a flat noise accounted for in $bg(Q^2)$. With these basic constituents we have constructed a theoretical model of the expected ND as a function of the pair distribution functions of the confined structures with the corresponding rms COM displacement, the proportion of external crystallite scatterers, and its associated rms COM displacement. The detailed derivation is provided in Appendix A.1. The final expression for a concentration (mole fraction) C_{1D} of 1D confined molecules with a specific structure and a concentration C_{3D} of molecules in the 3D crystallite reads,

$$I_D(Q) = C_{1D} (4a_d^2 j_0(Qd_{D_2}/2)^2 (S_{1D}^M(Q, u_{1D}) - 1) + F_{inc}(Q, u_{1D})) \\ + C_{3D} \left(F_{inc}(Q, u_{3D}) - 4a_d^2 j_0(Qd_{D_2}/2)^2 \exp \left[-\frac{(Q u_{3D})^2}{3} \right] \right) + bg(Q^2), \quad (1)$$

where a_d is the neutron coherent scattering length of the deuteron, j_0 is the spherical Bessel function of zeroth order, d_{D_2} is the distance between the two deuterons, u_{1D} and u_{3D} are the corresponding rms molecular displacements, and the background reduces to $bg(Q^2) = bg_0 + bg_1 Q^2 + bg_2 Q^4$. $S_{1D}^M(Q, u_{1D})$ is the structure factor of the molecular COMs of the 1D crystal. The factor $j_0(Qd_{D_2}/2)^2$ is the spherical form factor mentioned above. The $F_{inc}(Q, u)$ terms are the incoherent part of the corresponding crystals. As explained in Appendix A.1, they depend not only in the temperature but in the concentration of ortho molecules. In practical terms our scattering intensity is not normalized to absolute units, so that C_{1D} and C_{3D} are scale factors proportional to the real concentrations.

With respect to the interference part $I_{DC}(Q)$, by its very nature, its most prominent contribution is around 1.8 \AA^{-1} where the main MWCNTs Bragg peak is located, being negligible for larger values of Q

since they correspond to spatial scales shorter than the D₂-C distances. This is confirmed a posteriori by the success of $I_D(Q)$ in reproducing the observed ND above 1.8 \AA^{-1} . An atomistic description of $I_{DC}(Q)$ is exceedingly expensive in computational terms; instead, we have assumed the nanotubes as continuous and infinite to reduce the calculation to sums of analytical expressions as detailed in Appendix A.2. The approximated $I_{DC}(Q)$ depends on the inner radius of the cylinder where the confined phase forms, r_i , on the dispersion of the distribution of external MWCNTs diameters, σ_n , on the average interlayer distance within the MWCNTs, h , and on an effective rms displacement, u_{eff} , to take into the account the disorder in the involved pair distances. We remark that its intensity with respect to $I_D(Q)$ is not free but determined by the calculation. Around 1.8 \AA^{-1} , the $I_{DC}(Q)$ profile and its derivative change wildly in a few points, so that iterative minimization algorithms fail to converge. Instead, $I_{DC}(Q)$ is added as a correction to the fitted $I_D(Q)$, the corresponding free parameters chosen by visual inspection comparing series of plots of the modeled $I_s(Q)$ against the measurements. More specifically, u_{eff} is chosen to the minimum value that does not perturb the independently fitted $I_D(Q)$. The value of h essentially controls the position of the “dispersive” profile around 1.8 \AA^{-1} . A value of 3.41 \AA works well for all the explored cases. The inner radius, r_i , is the most relevant parameter with respect to the profile shape. As shown below our $I_{DC}(Q)$ reproduces qualitatively the “dispersive” profile around 1.8 \AA^{-1} .

3. Results and Discussion

Fig. 3 displays the ND signal $I(Q)$ as a function of the scattering vector modulus Q of the empty MWCNT sample (black) superimposed to that of the 284 hPa load (green) at low temperature. From now on and for the sake of simplicity, we label the loads by their equilibrium pressures at 20 K. The red rectangle encloses the four main observed Bragg peaks. The normalized area of $I(Q)$ within the rectangle during a thermal cycle for the two lower loads is shown in the inset. A reversible change of the slope appears at around 13.5 K in both cases, revealing a liquid/solid transition at that temperature. Given that the bulk triple point is at 18.73 K, and that we are under SVP conditions, such a temperature reduction indicates that the transition is happening under confinement [33]. Furthermore, since the transition temperature depends of the pore size, its reversibility implies confinement under a rather narrow distribution of pore sizes. On the contrary, the curves of the two larger loads show hysteresis, as expected from the heterogeneous distribution in size and shape of the interstitials observed in the TEM pictures at meso/macroscales (see Fig. S3 in the Supplementary Material for the hysteretic cases).

Fig. 4 displays the ND of the solid phases at the four loads after subtraction of the empty MWCNT signal, denoted as $I_s(Q)$. Here, solid phases refer to those below 11.5 K, a value chosen well under the 13.5 K where the liquid/solid transition manifests. Consistently, the ND of all the independent runs below 11.5 K do not show any discernible changes, so that all the runs with $T < 11.5 \text{ K}$ have been accumulated. Details

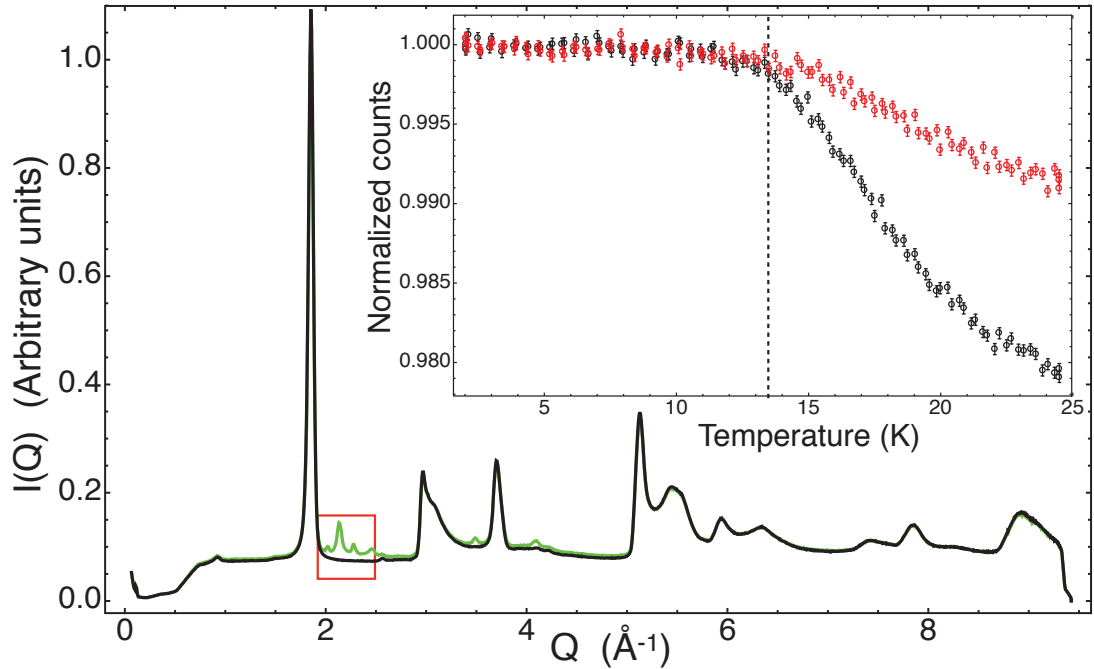


Figure 3: Neutron diffraction pattern, $I(Q)$, from the pristine MWCNT sample (black) superimposed to that corresponding to the 284 hPa D_2 load (green) at low temperature. The inset shows the evolution during a thermal cycle of the area (normalized to one at 2 K) under $I(Q)$ within the red rectangle for the 62 hPa load (red circles) and for the 175 hPa case (black circles).

of the neutron scattering data reduction are given in Appendix B.7. The 225 hPa and the 284 hPa loads
 125 present clear crystalline peaks identified as corresponding to both hexagonal close-packed (HCP) and face-
 centered cubic (FCC) phases. As shown in the figure, both kinds of peaks match to an excellent precision
 with strictly closed-packed arrangements with the same nearest-neighbor distance, namely, $d = 3.605 \text{\AA}$,
 a value in agreement with those reported for the bulk solids [34, 35]. The HCP phase is the fundamental
 crystal corresponding to the energy minimum of solid D_2 , but FCC crystallization can also be observed as
 130 soon as any perturbation is introduced in the crystallization environment, as for example, the heterogeneous
 nature of our MWCNTs sample.

D_2 tends to crystallize in rather large crystal domains. Since the ND instrument relies on a horizontal
 narrow strip detector geometry (see Appendix B.6), the non-uniform distribution of crystal orientations
 leads to distorted peak heights and the extinction of peaks with decreasing D_2 load. At first glance the
 135 175 hPa and 62 hPa patterns resemble those of amorphous substances with no Bragg peaks. A closer look,
 presented in the inset of Fig. 4, reveals the presence of a couple of precursors of bulk crystalline peaks at
 175 hPa and one at 62 hPa. This can be appreciated better in Fig. 5. Since these Bragg peak precursors
 appear in both thermal cycles of 62 hPa and 175 hPa, they reveal oriented crystal growth somewhere in the
 sample-can system, probably owing to an inhomogeneity-related nucleation site. The “pseudo-amorphous”

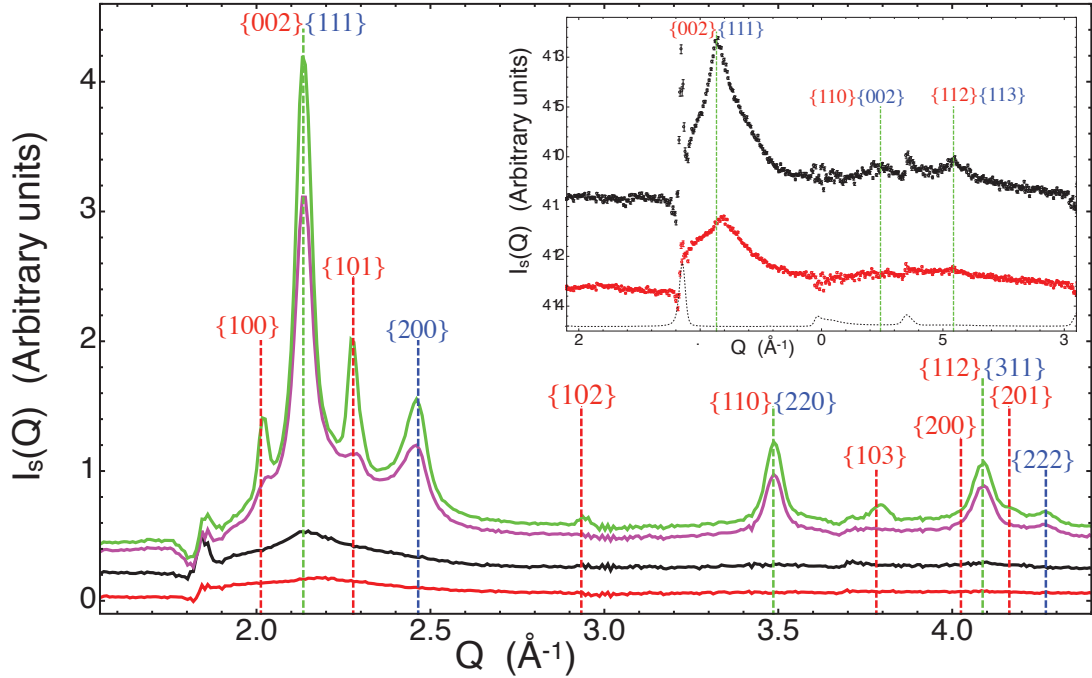


Figure 4: $I(Q)$ after subtraction of the MWCNT sample contribution, $I_s(Q)$, for the 62 hPa (red), 175 hPa (black), 225 hPa (magenta), and 284 hPa (green) D_2 loads. The vertical dashed lines mark the position of the bulk solid D_2 crystalline peaks. Dashed blue lines are for peaks present only in the HCP crystal, red dashed lines for those only present in the FCC crystal and green dashed line for peaks belonging to both. On top of the dashed lines the corresponding Miller indexes are specified, in red for the HCP structure and in blue for the FCC case. The inset is a zoom of the two lowest loads, i.e., 62 hPa (red) and 175 hPa (black) along with a scaled-down version of the MWCNT sample $I(Q)$ (dotted line).

140 part necessarily corresponds to confined phases, in view of the behavior of the $I(Q)$ area of each load shown in the inset of Fig. 3. Within this respect, they depart significantly from the observed broader patterns in solid D_2 (10 K) adsorbed in SWCNT bundles [26]. There, the fits ascribe the ND to the presence of chains and stripes of D_2 molecules expected to be within the interstitials (chains) and external grooves (chains and stripes) of the bundles. These two loads are, therefore, the object of our main interest. The scattering from
 145 D_2 pairs, $I_D(Q)$, contributes dominantly to $I_s(Q)$. There is also a contribution from D_2 and carbon pairs, $I_{DC}(Q)$, relevant mainly below around 2 \AA^{-1} . We focus first in modeling the dominant deuterium part, $I_D(Q)$.

Using the $I_D(Q)$ theoretical model described in section 2, a nonlinear fitting analysis detailed in the Supplementary Material was performed along a family of constrained nanostructures. We naturally started
 150 from structures obtained by MD simulations including MWCNTs up to five walls and effective potentials derived from state-of-the-art ab-initio calculations. Not surprisingly, they match the expected arrangements from energy minimization arguments: tubular shells adsorbed to the inner wall of the nanotubes that can be seen as rolled portions of the D_2 HCP crystal basal plane. As a matter of fact, they are explicit realizations

of the tubular compact packing of spheres studied in the context of biological structures [36]. However, they
155 are not compatible with our measurements as illustrated in Fig. S4, Supplementary Material.

Such disagreement drove us to consider other 1D crystalline structures corresponding to sections of
maximal cylindrical symmetry of both the HCP and FCC bulk crystals with increasing diameter. For these
crystals we fixed d to the value obtained from the 225 hPa and 284 hPa loads, i.e., 3.605 Å. Figure 5
presents the final results of our nonlinear model fitting analysis. Let us consider first the 62 hPa load (Fig.
160 5A). The agreement is excellent just using a log-normal distribution in length (see inset) of the remarkable
strict 1D crystal displayed in the right side of the figure. Differences of $I_D(Q)$ with increasing length become
negligible (see Supplementary Material Figures S10 and S11). Consequently, there is not enough information
in the diffraction signal to discern the 1D crystal length distribution; in fact, a non-parametric fit of the
crystal lengths only needs to populate a few isolated lengths to achieve excellent results. Lacking further
165 information, we have relied in the log-normal distribution because it is usually observed in growth processes
with preferential direction [37, 38]. With respect to the ortho concentration, the fits need a proportion
corresponding to room temperature, i.e., there was no para-ortho conversion during the experiment. The
result is not only of a sheer beauty in its simplicity — the simplest nontrivial strict 1D compact crystal
made out of spheres — but incompatible with theoretical predictions, as discussed above. It corresponds to
170 a cylindrical cut of the minimal diameter which allows more than one molecule in the horizontal plane with
the axis perpendicular to the hexagonal basal plane of the HCP fundamental bulk crystal, and centered so
that it passes through the molecular centers at alternate layers. The six-fold symmetry of the parent HCP
bulk crystal is reduced to three-fold (Fig. 5A, top right panel), so we call it triangular-close packed (TCP)
1D crystal.

175 Consistently, in the 175 hPa load (Fig. 5B) the nanocrystalline structure is just a radially enlarged version
of the TCP 1D crystal. Again, there is a three-fold symmetry, as it can be seen in the transversal cut of
the crystal, upper right panel of Fig. 5B. Since the crystal consists of alternating triangular and hexagonal
layers, we denote it as THCP 1D crystal. Both 1D crystals display large rms COM displacements, around
0.33 Å in the TCP case, and 0.45 Å in the THCP counterpart, to be compared with the 0.46 Å obtained
180 from neutron scattering of the D₂ bulk crystal [39] (see Tables SI and SII in Supplementary Material).
Interestingly, there is no hint in the 175 hPa ND of the pattern corresponding to the 62 hPa load. Indeed,
the fittings yield a zero population of the TCP phase, i.e., any finite population of the TCP phase worsens
the fits. Should the TCP 1D crystals form also during the 175 hPa load in quantities comparable to the
62 hPa case, they would also manifest in the 175 hPa ND. Therefore, either the THCP crystals grew out
185 of previously formed TCP seeds, or the MWCNTs of the smallest diameters are not filled during the 175
hPa load. In the first hypothesis, the TCP crystals grown at 62 hPa would be formed within MWCNTs of
large enough inner diameter, whereas in the latter the two crystals would grow within MWCNTs of different
inner diameters. Fortunately, the interference part, $I_{DC}(Q)$, carries information about the inner MWCNTs

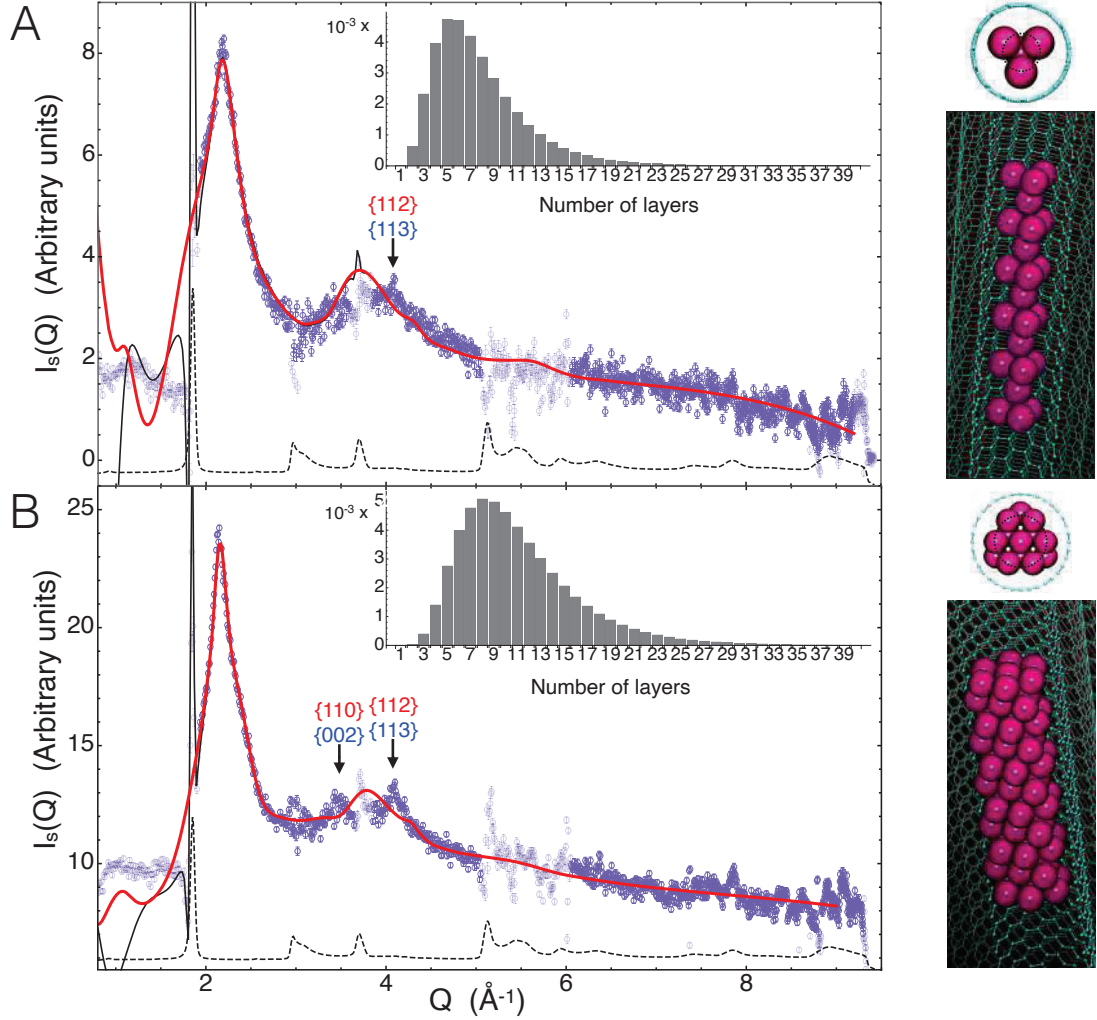


Figure 5: D₂ 1D crystals fitting the ND data. (A) D₂ load corresponding to the 62 hPa case. (B) D₂ load corresponding to the 175 hPa case. ND data points displayed in light blue are not included in the fittings (points below 2 \AA^{-1} , areas too affected by MWCNT peak subtraction, and some outliers). Solid red curves are the fitted $I_D(Q)$ component. Thin black curves show the final $I_s(Q)$ including the $I_{DC}(Q)$ part. The dashed lines are a down-scaled representation of the pristine MWCNT sample ND. Arrows mark the positions of some relevant Bragg peaks of the bulk crystal. At the right side, a representative structure giving rise to the corresponding $I_s(Q)$ is shown in perspective. The small spheres have a diameter equal to the deuteron-deuteron distance of the D₂ molecule (0.74 \AA). The diameter of the semitransparent larger spheres equals $d = 3.605 \text{ \AA}$. In the top views only the inner nanotubes are shown, an (11,4) single-walled CNT (A) and an (18,0) (B). Dotted circumferences mark the position of the minimum of the adsorption potential. Insets display the fitted log-normal distributions of 1D crystal lengths in terms of the number of molecular layers perpendicular to the MWCNT axis.

diameters so we can use it to settle this question. Fig. 5 shows in solid black the complete NDs including
 190 the $I_{DC}(Q)$ contribution. By construction, it coincides with $I_D(Q)$ above $\sim 1.8 \text{ \AA}^{-1}$ so that it falls below
 the red line representing $I_D(Q)$. As shown in the top right panels of Fig. 5, the chosen inner diameters
 $2r_i$ (10.53 \AA for the TCP and 14.09 \AA for the THCP) accommodate well the D_2 molecules in terms of
 the estimated locus [40] of the adsorption potential minima. However, inner diameters around 14 \AA in the
 TCP case render $I_{DC}(Q)$ profiles incompatible with the measured ND at 62 hPa. We are therefore left to
 195 conclude that the MWCNTs of the narrowest inner diameters are not significantly filled during the 175 hPa
 load. Since we are referring to pore sizes at the molecular limit (below two molecular diameters), steric
 molecular hindrance as well as anomalous capillary and condensation effects [41, 42, 43], dependent on the
 D_2 load, seem plausible. Within this respect, the irregular shapes and the functional groups at the entrance
 of the opened nanotubes could play a relevant role. For instance, funnel-like attractive potentials at the
 200 entrance would probably enhance the possibility of molecular jamming at 175 hPa owing to the different
 kinetics and density with respect to the 62 hPa case. Given the large de Broglie wavelength of D_2 , quantum
 wave-like behavior contributing to these effects, as observed with He [44, 45, 46], are also conceivable. At any
 rate, here we can only speculate about the microscopic mechanisms involved in this interesting anomalous
 behavior since their elucidation requires a study by its own.

205 Taking into the account that MD simulations do not describe accurately dispersive interactions, which are
 crucial in this kind of systems, we have performed advanced dispersive DFT calculations validated against
 experiments, in order to shed light on the discrepancy between the observed structures and theoretical
 predictions (details of the calculations are given in Appendix B.9 and Supplementary Material). Due to the
 large computational demand of this approach, we restrict ourselves to SWCNTs of a diameter compatible
 210 with the TCP 1D crystal. More specifically, the nanotube of chiral vector (10,5) has an adequate unit cell
 length while its diameter of 10.36 \AA is just 2% below that of the (11,4) tube shown in Fig. 5A. The number
 of D_2 molecules is chosen to be eleven, corresponding to five layers of the 1D TCP crystal, the most probable
 length (inset of Fig. 5A). Fig. 6 summarizes the DFT results. Spiral geometries similar to those obtained by
 MD are once more the most stable configurations, at variance with the experimental results (see Fig. S4A of
 215 Supplementary Material). Remarkably, the observed 1D TCP structure, presented in Fig. 6B, is metastable
 within this approach, but the next most plausible 1D crystal derived from the FCC bulk structure is not
 stable under the DFT approximation (depicted in Fig. S5B of Supplementary Material). The calculated
 $I_D(Q)$ s are also shown in Fig. 6 along with those corresponding to the equivalent TCP 1D structure obtained
 from the fits. Additionally, notice that the DFT calculations yield an effective nearest-neighbor distance
 220 slightly shorter than the experimental one. This is a consequence of the quantum delocalization of the
 rotational degrees of freedom of the nuclei, not taken into account by the DFT approach, which renders
 the molecular interactions essentially isotropic. On the contrary, within the DFT approximation the D_2
 molecules retain its dumbbell-like anisotropic character and tend to order in a parallel, more compact,

configuration.

225 The present results tell of a van der Waals molecular compound that crystallizes retaining the bulk crystal
structure down to the strict 1D limit. This is in stark contrast with results pertaining to other materials
with weak cohesive energies, which do not show the marked quantum behavior in their translational and/or
rotational degrees of freedom as our 1D crystals do, conspicuously revealed by their neutron scattering
response. Our DFT calculations, with reliable dispersive functionals, make clear that the missing quantum
230 treatment of the nuclear motions are one key ingredient for the stabilization of these 1D crystals. On the
other hand, the fact that the TCP 1D crystal is weakly metastable under the DFT calculations suggest
that a precise account of the quantum behavior of the electronic cloud in the description of the dispersive
interactions could also be relevant. In fact, the development of dispersive DFT techniques during the last two
decades is characteristic of the nanoscience revolution: adsorption phenomena within nanostructures depend
235 too subtly on the dispersive interactions to be accommodated within the traditional effective potential
approach. The challenge now is to incorporate the nuclear quantum effects accurately, a subject attracting
an increasing interest [47]. At present, inclusion of translational quantum nuclear delocalization effects
in condensed matter can be tackled with computationally very demanding quantum Monte Carlo or path
integral methods [23, 24] but we remark that, both, the nuclear rotational delocalization and the dispersive
240 forces are treated in the same effective manner as in our MD simulations. Within this context, the structures
here discovered stand out for testing future theoretical developments since they are dispersive molecular
solids displaying a non-trivial quantum nuclear behavior at the same time being structurally simple.

4. Conclusion

Using neutron diffraction we have discovered 1D quantum crystals of molecular deuterium within MWC-
245 NTs. Unexpectedly for a molecular solid with very weak cohesive interactions, the 1D crystals preserve the
structure of the HCP bulk D_2 crystal. Neither atomistic molecular dynamics simulations nor advanced dis-
persive DFT calculations predicts such a possibility, signaling the strong quantum nuclear effects as essential
to the emergence of the discovered 1D crystals.

These 1D crystals are the simplest nontrivial condensed matter systems in which the quantum nature,
250 not only of the electronic cloud, but also of the nuclear translational and rotational degrees of freedom,
plays an essential role. Our results are deemed to be of special relevance, since they provide a testbed to
confront theoretical approaches aimed to understand and incorporate accurately the full quantum nature of
real interactions, including the nuclear degrees of freedom. Such advances will have a bearing on the design
and simulation of novel (possibly quantum) materials.

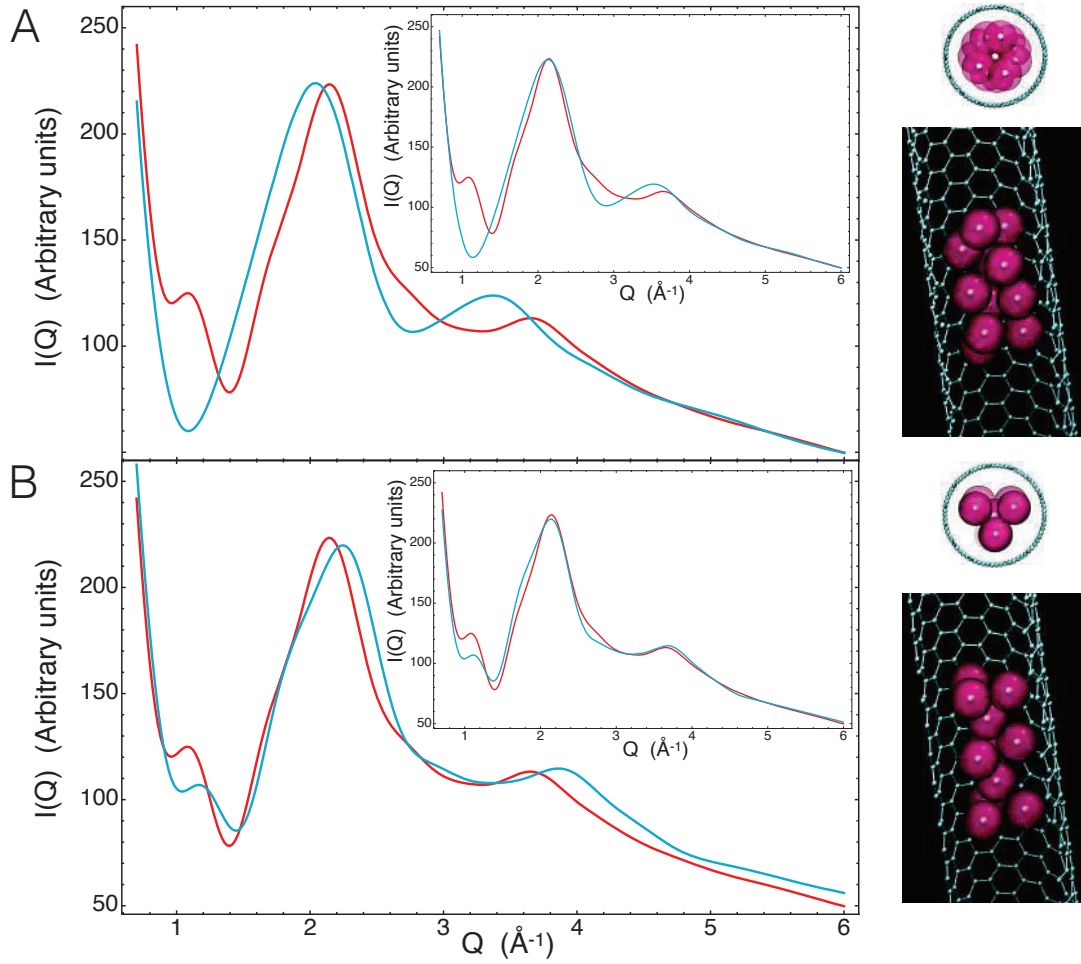


Figure 6: 1D structures obtained by dispersive DFT calculations. (A) Fundamental structure corresponding to the lowest energy minimum. (B) Structure of the 1D TCP kind corresponding to a metastable energy minimum. Their associated $I_D(Q)$ s are also shown (cyan curves) along with that corresponding to the experimentally-determined equivalent 1D TCP crystal (red curves). Since the DFT calculations do not reproduce the nearest-neighbor distance accurately, properly Q -rescaled versions of the $I_D(Q)$ s are displayed in the insets.

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Appendix A. Theoretical developments

270 *Appendix A.1. Theory of the measured neutron scattering*

In a neutron diffraction experiment at a continuous source installation such as the ILL, the intensity of scattered neutrons by the sample out of a mono-energetic incident neutron beam is measured as a function of the scattering angle. The relevant differential scattering cross section is then given by

$$\frac{d\sigma}{d\Omega} = \int_{-\infty}^{E_i} \frac{d^2\sigma}{d\Omega dE_f} dE_f, \quad (\text{A.1})$$

where the Ω denotes solid angle, E_i the energy of the incident neutrons and E_f the final detected neutron energy. Following the full quantum calculations of Sears [32], the double differential cross section in the case of a homonuclear diatomic molecule under the assumption of free rotation, can be expressed as

$$\frac{d^2\sigma}{d\Omega dE_f} = N \frac{k_f}{k_i} \left(4 a_d^2 j_0(Q d_{D_2}/2)^2 S_{int}(\mathbf{Q}, E) + \sum_{J, J'} a^2(Q; J, J') S_{self}(Q, E - E_{J, J'}) \right). \quad (\text{A.2})$$

Here N is the number of nuclei in the sample, $k_{i,f}$ are the initial and final neutron angular wavenumbers, a_d is the coherent neutron scattering length of the deuteron, j_0 the spherical Bessel function of zero order, d_{D_2} the distance between the nuclei in the molecule, \mathbf{Q} is the scattering angular wavevector (Q its norm), the J s denote the nuclear orbital angular momentum quantum levels, $E_{J, J'}$ the corresponding energy jumps among J levels and $E = E_i - E_f$ is the energy transfer. The $a^2(Q; J, J')$ factors are quite complex expressions involving linear combinations of Bessel spherical functions dependent on the bosonic or fermionic character

of the nuclei [32]. They depend on all the rotational levels excited by the neutrons at the given incident energy and, as such, incorporate the corresponding orientational form factors. Transitions between the ortho (even J angular momentum) and para (J odd) levels are forbidden so that thermalization of the populations proceeds independently. The $J = 1$ state is highly metastable so that equilibration at low temperatures downwards the fundamental $J = 0$ level is extremely slow in the bulk [12] but it could be accelerated by the interaction with the substrate. Otherwise, thermal equilibration proceeds unhindered so that the model depends on the temperature and the concentration of ortho molecules [32].

$S_{int}(\mathbf{Q}, E)$ is the interference part of the dynamical structure factor of the molecular COMs, i.e., the part not involving single particle self-scattering. Such self-scattering part (of the COMs) corresponds to $S_{self}(Q, E)$ (Sears uses the subscript *inc* from incoherent). It is a consequence of the free rotation assumption that both $S_{int}(\mathbf{Q}, E)$ and $S_{inc}(Q, E)$ in Eq. (A.2), refer only to the dynamics of the COMs of the molecules. In order to calculate Eq. (A.1), we need a model of such dynamical structure factors. Since their contribution to Eq. (A.1) is through integration in E , we will take advantage of general sum rules, i.e., moments in E , that must be satisfied independently of the system dynamics.

Focusing on the interference part, the relevant sum rule is $\int_{-\infty}^{\infty} S_{int}(\mathbf{Q}, E) dE = S(\mathbf{Q}) - 1$, where the $S(\mathbf{Q})$ is the structure factor. This is the quantity of interest, since it yields a structural characterization of the system and it is amenable to theoretical modeling. It is given by

$$S(\mathbf{Q}) = 1 + \frac{1}{N} \sum_{i \neq j} \langle \exp [i(\mathbf{R}_i - \mathbf{R}_j) \cdot \mathbf{Q}] \rangle, \quad (\text{A.3})$$

where N is the number of scatterers, \mathbf{R}_j the position vector of the j molecular COM and $\langle \rangle$ denotes quantum-mechanical thermal average. In a solid, thermal fluctuations are well approximated by a Gaussian distribution of displacements around the mean positions $\langle \mathbf{R}_j \rangle$. In such a case, after orientational and thermal average, Eq. (A.3) reduces to

$$S(Q) = 1 + \frac{1}{N} \sum_{i \neq j} j_0(|\langle \mathbf{R}_i \rangle - \langle \mathbf{R}_j \rangle|Q) e^{-\frac{1}{3}u^2 Q^2} = 1 + \frac{1}{N} \sum_{d_{ij}} m_{ij} j_0(d_{ij}Q) e^{-\frac{1}{3}u^2 Q^2}, \quad (\text{A.4})$$

where the final sum is over all the different pair distances $d_{ij} = |\langle \mathbf{R}_i \rangle - \langle \mathbf{R}_j \rangle|$, m_{ij} are the corresponding multiplicities and u is the rms displacement around the $\langle \mathbf{R}_j \rangle$, assumed here, isotropic and independent of the position. In a solid, the vast majority of the interference scattering is elastic, i.e., it accumulates around $E = 0$, so that our model for the interference part is $S_{int}(Q, E) = (S(Q) - 1) \delta(E)$, where $\delta(E)$ denotes the Dirac delta.

With respect to the self part the elastic term amounts to

$$S_{self}^e(Q, E) = e^{-\frac{1}{3}u^2 Q^2} \delta(E). \quad (\text{A.5})$$

However, it is not so dominant as in the interference case, so we need to proceed further. The relevant

moments to develop a model of the inelastic part are [48]

$$\int_{-\infty}^{\infty} S_{self}(Q, E) dE = 1, \quad (\text{A.6})$$

$$\int_{-\infty}^{\infty} E S_{self}(Q, E) dE = \frac{\hbar^2 Q^2}{2M} = E_r, \quad (\text{A.7})$$

$$\int_{-\infty}^{\infty} E^2 S_{self}(Q, E) dE = \frac{4}{3} E_r \bar{K} + E_r^2, \quad (\text{A.8})$$

where M is the mass of the molecule, E_r is the so-called recoil energy and \bar{K} is the mean kinetic energy. The zeroth order moment is always satisfied, the first order moment is satisfied provided that the interaction does not depend on the molecular velocities, and the second is true for isotropic systems. The elastic part does not contribute to the first and second moment; decomposing the incoherent part as $S_{self}(Q, E) = S_{self}^e(Q, E) + S_{self}^{ine}(Q, E)$ and using Eq. (A.5), the zeroth moment implies

$$\int_{-\infty}^{\infty} S_{self}^{ine}(Q, E) dE = 1 - e^{-\frac{1}{3}u^2 Q^2}. \quad (\text{A.9})$$

On the other hand, from the first and the second moments, the energy dispersion is obtained:

$$\sigma_E^2 = \langle E^2 \rangle - \langle E \rangle^2 = \frac{4}{3} E_r \bar{K}. \quad (\text{A.10})$$

The mean kinetic energy at low temperature is expressed in terms of the rms displacement as [49, 50],

$$\bar{K} = \frac{9}{16} \Theta = \frac{81}{64} \frac{\hbar^2}{Mu^2}, \quad (\text{A.11})$$

where Θ is the Debye temperature, the parameter usually quoted in the experimental reports. We are now in disposition to write up a model for the incoherent part preserving all the moments up to the second:

$$S_{self}^M(Q, E) = e^{-\frac{1}{3}u^2 Q^2} \delta(E) + (1 - e^{-\frac{1}{3}u^2 Q^2}) \frac{1}{\sqrt{2\pi\sigma_E^2}} \exp\left[-\frac{(E - E_r)^2}{2\sigma_E^2}\right]. \quad (\text{A.12})$$

where the M superscript serves to recall it refers only to molecular COM contribution. The chosen Gaussian shape for the inelastic scattering is a natural choice to warrant the integral properties up to second order.

Such a profile must be understood as a kind of smoothing of the real signal since it does not incorporate the details of the density of vibrational modes. Only in the limit of very large momentum transfer the same kind of profile would be exact (impulse approximation) but for the nuclei themselves, not for the molecular COMs. Interestingly, in the case of H_2 , given the lightness of the molecule, this profile for the COM dynamics has been observed at least from Q 's as low as 5 \AA^{-1} [51].

In order to calculate Eq. (A.1) the factor $k_f/k_i = k_i \sqrt{1 - E/E_i}$ must be also included in the inelastic part. Conservation of energy and momentum establishes a relation among Q , E , and, ϕ , the angle between \mathbf{k}_i and \mathbf{k}_f , namely,

$$Q^2 = k_i^2 + k_i^2 \left(1 - \frac{E}{E_i}\right) - 2k_i^2 \sqrt{1 - \frac{E}{E_i}} \cos \phi. \quad (\text{A.13})$$

305 Substituting Q in terms of E and ϕ , the integration of Eq. (A.2) in E yields the measured cross section Eq. (A.1). Notice that direct integration of Eq. (A.2) does not yield the single cross section, since it is defined (and measured) at constant ϕ , not at constant Q . However, usually the ND is expressed in terms of Q , but only at its elastic value using the relation $Q_e = 2k_i \sin \phi/2$. As a rule, the elastic condition is not explicitly stated but the same symbol Q is used. We adhere here to such a practice. At any rate, in many experiments 310 E_i is sufficiently high for the static approximation to be valid [48], where all the scattering can be considered elastic.

As far as the fitting is concerned, we first generate a two-dimensional numerical table in u and Q of the expression

$$F_{inc}(Q, u) = \int_0^\infty \frac{k_f}{k_i} \sum_{J, J'} a^2(Q; J, J') S_{self}(Q, E - E_{J, J'}) dE, \quad (\text{A.14})$$

where the suffix *inc* remarks that this expression includes not only the coherent self-contribution but the incoherent scattering. In the actual fittings a two-dimensional interpolation of the table is used to avoid the computational load of a numerical integration at each fitting step.

With respect to the diffuse coherent scattering of the external crystallite, performing only the thermal average in Eq. (A.3), we have

$$S(\mathbf{Q}) = 1 + \frac{e^{-\frac{1}{3}u^2 Q^2}}{N} \sum_{i \neq j} e^{i(\langle \mathbf{R}_i \rangle - \langle \mathbf{R}_j \rangle) \cdot \mathbf{Q}} = 1 + e^{-\frac{1}{3}u^2 Q^2} \left(\frac{1}{N} \sum_i \left| e^{i\langle \mathbf{R}_i \rangle \cdot \mathbf{Q}} \right|^2 - 1 \right) \quad (\text{A.15})$$

$$= 1 - e^{-\frac{1}{3}u^2 Q^2} + e^{-\frac{1}{3}u^2 Q^2} \frac{1}{N} \sum_i \left| e^{i\langle \mathbf{R}_i \rangle \cdot \mathbf{Q}} \right|^2 = 1 - e^{-\frac{1}{3}u^2 Q^2} + S_B(\mathbf{Q}). \quad (\text{A.16})$$

In a crystal, and in the limit of $N \rightarrow \infty$, $S_B(\mathbf{Q})$ gives rise to the Dirac delta peaks characteristic of Bragg scattering. The rest is obviously the scattering among the Bragg peaks, i.e., the diffuse scattering. This remains true also at finite N , where the peaks broaden. Therefore, the diffuse scattering from external crystallite contributes with an interference part given by

$$S_{int}(\mathbf{Q}, E) = -e^{-\frac{1}{3}u^2 Q^2} \delta(E), \quad (\text{A.17})$$

315 while the incoherent part is again given by $F_{inc}(Q, u)$, but this time with u corresponding to that of the external crystallite.

Altogether, for a number concentration C_{1D} of 1D confined molecules and a C_{3D} number concentration of molecules in the 3D external small crystal, gives,

$$I_D(Q) = C_{1D} \left(4a_{coh}^2 j_0(Qd_{D_2}/2)^2 (S_{1D}^M(Q, u_{1D}) - 1) + F_{inc}(Q, u_{1D}) \right) + C_{3D} \left(F_{inc}(Q, u_{3D}) - 4a_{coh}^2 j_0(Qd_{D_2}/2)^2 \exp \left[-\frac{(Q u_{3D})^2}{3} \right] \right), \quad (\text{A.18})$$

where $S_{1D}^M(Q, u_{1D})$ is the structure factor of the molecular COMs as given by Eq. (A.4) and u_{1D} and u_{3D} are the corresponding rms displacements.

Besides Eq. (A.18) there must be an instrumental flat background, bg_0 . With such an addition, Eq. (A.18) produces convergent fits to the measured $I(Q)$. That is, it captures adequately most of the ND signal. A final refinement is now expressible as series in Q^2 . A background second order in Q^2 , $bg_0 + bg_1Q^2 + bg_2Q^4$, is enough to obtain excellent fits.

Appendix A.2. Modeling of the C-D₂ diffraction interference term

In terms of the positions of the D₂ COMs, \mathbf{R}_i and of the carbon atoms, r_j , the ND signal corresponding to the interference between carbon atoms and D₂ molecules reads,

$$\begin{aligned} I_{DC}(Q) &= C_{1D} a_c 2 a_d j_0(Qd_{D_2}) \frac{1}{N} \sum_i \sum_j \langle e^{-i\mathbf{Q}\cdot(\mathbf{R}_i - \mathbf{r}_j)} \rangle + c.c. \\ &= C_{1D} a_c 2 a_d j_0(Qd_{D_2}) e^{-\frac{1}{6}Q^2(u_{1D}^2 + u_c^2)} \frac{1}{N} \sum_i \sum_j \langle e^{-i\mathbf{Q}\cdot(\langle \mathbf{R}_i \rangle - \langle \mathbf{r}_j \rangle)} \rangle + c.c., \quad (\text{A.19}) \end{aligned}$$

where *c.c.* denotes complex conjugate, N is the number of molecules, a_c is the coherent neutron scattering length of carbon nuclei, u_c their rms mean displacement and in the second line we have performed thermal average, again assumed Gaussian and isotropic. Consequently, $\langle \rangle$ reduces to only an orientational average. Here, $2 a_d j_0(Qd_{D_2})$ plays the role of the coherent scattering length of the D₂ molecule, dependent on Q owing to the non point-like molecular structure.

An atomistic description of the MWCNTs is computationally too expensive. However, assuming a continuous distribution of the carbon atoms along infinite tubes, we can take advantage of some analytical calculations. Given a microscopic density of carbon atoms, $\rho_c(\mathbf{r})$, a general term of the sums above is written as

$$\left\langle e^{-i\mathbf{Q}\cdot\langle \mathbf{R}_i \rangle} \int_V e^{-i\mathbf{Q}\cdot\mathbf{r}} \rho_c(\mathbf{r}) d\mathbf{r}^3 \right\rangle = \frac{1}{4\pi} \int d\Omega_Q e^{-i\mathbf{Q}\cdot\langle \mathbf{R}_i \rangle} \int e^{i\mathbf{Q}\cdot\mathbf{r}} \rho_c(\mathbf{r}) d\mathbf{r}^3, \quad (\text{A.20})$$

where $d\Omega_Q$ is the solid angle corresponding to a direction of the scattering vector and the integrals extend to all the directions and the whole space respectively. For an infinite tube of diameter R_{CNT} , the density is expressed in the cylindrical coordinates r, φ, z , with z axis that of the cylinder as $\rho_c(\mathbf{r}) = \rho_A \delta(r - R_{CNT})$, being ρ_A the areal density of carbon in the nanotube. Analytical integration in the cylindrical coordinates yields

$$\int e^{i\mathbf{Q}\cdot\mathbf{r}} \rho_c(\mathbf{r}) d\mathbf{r}^3 = 4\pi^2 \rho_A R_{CNT} J_0(QR_{CNT}) \delta(Q_z), \quad (\text{A.21})$$

where J_0 is the Bessel function of order zero and Q_z is the z component of the scattering vector. Notice that for an infinite tube the symmetry leads to cancellation of the dependence along the z axis. Substituting Eq. (A.21) in Eq. (A.20), again analytical integration can be done, this time using spherical coordinates, so that

$$\left\langle e^{-i\mathbf{Q}\cdot\langle \mathbf{R}_i \rangle} \int_V e^{-i\mathbf{Q}\cdot\mathbf{r}} \rho_c(\mathbf{r}) d\mathbf{r}^3 \right\rangle = 2\pi^2 \rho_A R_{CNT} J_0(QR_{CNT}) J_0(QR_i^\perp) / Q, \quad (\text{A.22})$$

where R_i^\perp is the modulus of the projection of $\langle R_i \rangle$ onto the plane perpendicular to the tube axis. Now the interference term for a SWCNT amounts to

$$I_{DC}(Q) = 8\pi^2 C_{1D} a_d \rho_A j_0(Qd_{D_2}) e^{-\frac{1}{6}Q^2(u_{1D}^2 + u_c^2)} R_{CNT} J_0(QR_{CNT}) \frac{1}{N_c Q} \sum_i J_0(QR_i^\perp), \quad (\text{A.23})$$

where i runs from 1 to N_c , being N_c the number of D_2 molecules of a unit cell of the corresponding 1D crystal.

For a distribution of MWCNTs of well-defined internal radius r_i and m layers separated by a distance h , the term $R_{CNT} J_0(QR_{CNT})$ transforms into

$$\sum_{n=1}^{m-1} (r_i + h(n-1)) J_0(Q(r_i + h(n-1))). \quad (\text{A.24})$$

In our case the m distribution is not narrow, but those of r_i and h must be quite. Assuming r_i fixed, the average over the ensemble of MWCNTs modulates the terms in the sum over layers so that they are scaled down as n increases, since the statistics of larger n decreases because the instances of large m also decreases. Furthermore, for n small there are almost as many cases as for $n = 1$, that is, the histogram of n must start horizontally. A half-Gaussian decay of the n terms matches such specification and emerges as a natural choice. Altogether it yields

$$I_{DC}(Q) = 8\pi^2 C_{1D} a_d \rho_A j_0(Qd_{D_2}) e^{-\frac{1}{3}Q^2 u_{eff}^2} \frac{1}{N_c Q} \sum_i J_0(QR_i^\perp) \sum_{n=1}^{\infty} \exp\left[-\frac{h^2(n-1)^2}{2\sigma_n^2}\right] (r_i + h(n-1)) J_0(Q(r_i + h(n-1))), \quad (\text{A.25})$$

where we have defined an effective rms displacement, u_{eff} , and σ_n is the dispersion of the half-Gaussian decay. In practical terms a value of 50 layers worked well for σ_n , for which a termination of the n sum at 250 were enough to reach convergence in a short cpu time. In general, there are far more sources of disorder than just the thermal fluctuations around the COMs. For instance, in the real MWCNTs h presents also dispersion with some systematic evolution with the number of layers [52]. All of these effects are here boldly represented through u_{eff} , chosen as the minimum needed to maintain the line shape of the main ND peak (1.2 Å for the 62 hPa load and 1.5 Å for the 175 hPa case). The parameter h controls the location in Q of the dispersive profile. A value of 3.41 Å was consistent with both the ND measurements and the values reported in the literature [52]. Finally, the areal density of any nanotube is given by $\rho_A = 4/(3\sqrt{3}d_{cc}^2)$, where d_{cc} is the distance of the C–C bond in graphene taken here as 1.42 Å.

Appendix B. Materials and Methods

Appendix B.1. Experimental design

As for the confining system, we rely on samples of multi-walled carbon nanotubes (MWCNTs). For our
345 interests, they present several advantages compared to single-walled nanotubes. To begin with, they are
far stiffer. Given the weakness of the dispersive interactions among the molecules, tube bending should be
avoided as much as possible to facilitate unperturbed crystallization. Besides, since the external diameters
show a large dispersion in a typical MWCNT sample, they do not form so tightly and well ordered ropes, as
their single-walled counterparts tend to do [29]. On the other hand, the interstitial sizes are typically much
350 larger than those of the smaller inner diameters because the distribution of external diameters is centered
at much higher values than those of the inner diameters (much larger also than the typical mean diameter of
good quality SWCNTs samples). In such a case, it can be expected that the adsorption inside the smaller of
the multi-walled tubes will not compete appreciably with 1D-like adsorption in the interstitials or external
grooves of the ropes.

355 To select the smaller inner diameters as our confining systems we rely in the physics of adsorption. The
smaller open inner diameter tubes produce the stronger attractive potential wells for the molecules so that
they are filled first. For sufficiently small loads these micropores dominate the distribution of adsorbed
molecules.

As for the probe to study the system, neutrons are uncharged, so that they can penetrate deeply into
360 the target. Thus, they can explore the structure and dynamics of atomic and molecular arrangements
buried into nanostructured materials, while at the same time making possible the use of complex sample
environments such as cryogenic and gas handling equipment. Contrary to X-rays, they strongly interact with
hydrogen. Since we are interested in structural information, we resort to the deuterium isotope in order to
avoid the very large incoherent scattering of molecular hydrogen that does not contribute to the diffraction
365 signal. Incoherent scattering stems from the random distribution of nuclear spin states along the sample,
and implies the interchange of the spin state between the incident neutron and the nuclei. It can also be
avoided in H₂ if prepared in its fundamental para-state, that only scatters coherently. However, the neutron
energy must be well below 14.7 meV in order not to excite the molecular rotational levels responsible for
the very strong incoherent scattering. This would limit the range of the scattering vector norm Q below 4
370 \AA^{-1} , too short to properly explore the spatial scales involved.

Regarding the measurement protocol, we started by measuring the diffraction of the nanotubes sample
(2 g) during a slow cooling/heating cycle (0.1 K decrement/increment each 70 s) between ≈ 24.5 K and
2 K while neutron diffraction signal was acquired in runs of 350 s. Then we proceeded by injecting very
small quantities of D₂ into the sample can at 20 K until we observed a barely discernible liquid state
375 diffraction signal. The equilibrium pressure at that point was 62 hPa. After heating at ≈ 24.5 K, the same

cooling/heating cycle as before was done. The same thermal protocol was repeated increasing the D_2 load in each cycle up to a total of four loads, namely, 62 hPa, 175 hPa, 225 hPa, 284 hPa. Before the 175 hPa thermal cycle, in order to improve the statistics of the measurements at 62 hPa, a second cooling ramp was performed down to 2 K where neutron diffraction was measured in runs of 600 s up to a total of 31 runs. 380 The cooling/heating rate during the thermal cycles was chosen as slow as it was feasible in order to remain as near as possible to thermodynamic equilibrium, corresponding in this case to those of saturated vapor pressure (SVP). From the adsorption isotherm in Fig. 2 it is clear that the four loads are far from the bulk condition at 20 K.

Appendix B.2. Multi-walled carbon nanotubes synthesis and treatment

385 The raw MWCNTs were prepared by the evaporation of pure graphite anodes in an electric arc discharge under a helium atmosphere of 66 kPa applying a current of 60 A and a voltage of 25 V [53]. The raw material was oxidized by heating to 973 K in open air during 30 minutes. After an annealing at 1223 K in Ar atmosphere during 10 hours, around 2 g of the open MWCNT sample was obtained.

Appendix B.3. Deuterium

390 The deuterium gas was taken directly from a 95 liter bottle of Air Liquide N30 Deuterium (purity $\geq 99.9\%$; ~ 10 ppm of D_2O and O_2 and ~ 25 ppm of N_2).

Appendix B.4. Electron microscopy

The structure of the obtained MWCNT material was explored by TEM at the National Center for Electron Microscopy (CNME) in Madrid, Spain. Drops taken from a dispersion of the samples in ethanol 395 were deposited over copper grids with holey carbon support films. Transmission electron micrographs were taken with a JEOL JEM2100 HT microscope operating at 200 kV.

Appendix B.5. Adsorption isotherm

The D_2 adsorption of the MWCNT sample was characterized by a measurement of the adsorbed moles as a function of the pressure at constant temperature (adsorption isotherm) using a standard volumetric 400 procedure [30]. The chosen temperature was 20 K (D_2 triple point at 18.73 K). In essence, given a known volume, the adsorbed moles are obtained from the decrease in pressure they cause with respect to the same volume with no adsorbant. A home-made gas handling manifold was built to accurately manage the gas dosage. A layout of the system is shown in Figure S1. All pipes and connections in the manifold are made of steel and Swagelok valves were used to connect the different volumes in the system. The pressure was 405 determined with a Baratron[®] MKS 690A absolute manometer with a full range scale of 1000 Torr and an accuracy of 0.12% of the reading. The reference volume V_R is provided by a calibrated steel vessel of

volume $1045.7 \pm 2.3 \text{ cm}^3$. The MWCNT sample was located in an aluminium cylindrical can of 15 mm of inner diameter and 56 mm of height. The can was connected vertically to a specially built insertion stick (used to introduce the sample can in the cryostat) with an inner capillary for the D_2 injection. The stick was designed to match the standard ILL “Orange” cryostat [54, 55]. This is the same kind of vertical cryostat used in the neutron scattering experiment. The inner capillary of the stick was connected through a valve to the end of the manifold labeled as “sample” in Figure S1. Using the known value of V_R all the needed volumes are determined by expansion of He loads. In particular, the volume comprising from “sample” valve (closed) down to the sample can (not shown in the Figure) was $20.64 \pm 0.15 \text{ cm}^3$. Once the system volumes were determined, 0.521 g of the MWCNT sample were loaded in the sample can, the sample stick was introduced in the cryostat and the temperature set to 20 K. From repeated D_2 loads in V_{inj} and subsequent expansions, the adsorbed moles were determined [30]. The sample temperature during the adsorption/desorption isotherm was $T = \langle T_S^i \rangle = 20.23 \pm 0.03 \text{ K}$.

Appendix B.6. Neutron scattering experimental setup

The neutron scattering instrument of choice was D20 at the ILL in Grenoble, a reactor-based neutron facility (continuous neutron beam). The most characteristic feature of D20 is its very high neutron flux, which allows for real-time studies of many processes. The setup was chosen to maximize the neutron flux corresponding to a neutron wavelength of 1.30 \AA . D20 is a powder diffractometer with a circular detection zone 1.47 m of radius, 4 m long and 15 cm high with the sample holder at the center of the circle (see [56] for details about D20). For the setup chosen the resolution remains below 0.02 \AA^{-1} until $Q = 5.5 \text{ \AA}^{-1}$, reaching 0.1 \AA^{-1} at $Q = 9 \text{ \AA}^{-1}$. Such resolution values are high enough to avoid resolution integrals in the data treatment. The gas handling manifold, described in Fig. S2, was located outside the neutron beam safety zone connected to the cryostat insertion stick through a 5-m-long 1-mm-diameter steel capillary. The injection volume, V_{inj} , used during the 20 K loads remained open during the neutron scattering measurements as a safety expansion volume. The sample can was a standard sample holder for the D20 instrument, namely, a 0.1 mm thick vanadium cylinder of 60 mm high and 8 mm of inner diameter. The neutron beam size was 8 mm width and 4 cm high. It was centered at the axis of the vanadium cylinder and the lower vertically edge of the beam located at the bottom part of the vanadium can. All the neutron irradiated volume was filled with MWCNTs sample (2 g). The pressure was monitored with a MKS 640A manometer (1000 Torr range) and a MKS 627B manometer (15000 Torr range). Since V_{inj} was open, the moles of D_2 irradiated by the neutron beam increased by a factor of around 1.4 from the equilibrium after load at 20 K to zero pressure solid at 2 K.

Appendix B.7. Neutron scattering data reduction

The raw data were corrected from detector efficiency and transformed to scattering vector reciprocal units using the LAMP software package freely available at the ILL facility [57, 58]. Self-shielding and neutron

absorption correction were estimated using the DAVE software package [59, 60]. As expected, given the small D_2 quantities and the cylindrical geometry of the sample can, self-shielding and neutron absorption amounts to a small and essentially Q -independent correction. Likewise, multiple scattering corrections are again Q -independent [61]. Both can be ignored for our purposes, which do not depend on absolute scattering values. An important point to take into the account is the variation in the number of scatterers irradiated by the beam. During a thermal cycle, due to the varying thermodynamic conditions (thermal expansion/contraction, D_2 condensation and diffusion, etc), some grains of the MWCNT powder move in and out the irradiated volume. In absolute terms the effect is small but not negligible with respect to the smaller D_2 loads. To compensate the effect the $I(Q)$ s of each run are divided by their corresponding total neutron count, which is proportional to the number of scatterers. In order to improve the statistics, all the runs corresponding to the pristine MWCNTs were added together, while for the low temperature $I(Q)$ s of the loaded MWCNTs, only those below 11.5 K (well below solidification temperature) were considered.

With respect to the subtraction of the MWCNT signal, since self-shielding and neutron absorption can be considered Q -independent, $I_C(Q)$ is just given by an attenuation transmission factor applied to the $I(Q)$ corresponding to the empty MWCNTs. The porous nature of the D_2 distribution within the MWCNT sample does not allow for a theoretical calculation of the effective transmissions. Instead, we have just heuristically adjusted the corresponding values, namely, 0.994 at 62 hPa, 0.9675 at 175 hPa, 0.940 at 225 hPa and 0.930 at 284 hPa.

Appendix B.8. Numerical methods. Molecular Dynamics

Molecular Dynamics (MD) simulations used in our nonlinear fitting analysis have been carried out using Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) code [62]. Taking into account the quantum delocalization of the nuclear rotational degrees of freedom we have considered for the D_2 – D_2 interaction an isotropic effective potential [12] adapted to the LAMMPS code. For the C–C interaction we have relied in the Adaptive Intermolecular Reactive Empirical Bond Order (AIREBO) potential described in [63] as it is included in the LAMMPS package. In the case of the D_2 –C interaction, we have relied in the anisotropic pair potential developed in [25] out of accurate *ab initio* calculations using Density Functional Theory (DFT) based on Symmetry Adapted Perturbation Theory (SAPT(DFT)), again adapted to LAMMPS. For details, including the parameterization used, see Supplementary Material in [25]. Thermalization has been achieved through the Berendsen thermostat [64], which presents a large thermal stability suitable for small samples. The chosen time step is one fs. The specifics of the procedure in the present calculations are given in the Supplementary Information.

Appendix B.9. Numerical methods. DFT

For the DFT calculations we have relied on the Perdew, Burke, and Ernzerhof (PBE) approach [65] as implemented in the CASTEP code [66]. The pseudopotentials are those corresponding to the 00PBE_OP.recpot

475 family as given by the Materials Studio package. The dispersive interactions are treated using the semi-
empirical dispersion correction scheme (DFT-SEDC) [67] with Tkatchenko and Scheffler (TS) correction [68]
(hydrogen atoms corrected to deuterium mass). The energy cutoff of the plane wave basis set was 900 eV,
only one k point was used and the maximum force tolerance per atom was set to 0.006 eV/Å. The CNT
containing the D₂ molecules is a (10,5) SWCNT with a length of 22.6211727 Å and 280 carbon atoms. The
480 specifics of the procedure are given in the Supplementary Material and in Table SIII therein.

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Formation of one-dimensional quantum crystals of molecular deuterium inside carbon nanotubes

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Abstract

Crystallization under stringent cylindrical confinement leads to novel quasi-one-dimensional materials. Substances with strong cohesive interactions can eventually preserve the symmetries of their bulk phase compatible with the restricted geometry, while those with weak cohesive interactions develop qualitatively different structures. Frozen molecular deuterium (D_2), a solid with a strong quantum character, is structurally held by weak dispersive forces. Here, the formation of one-dimensional D_2 crystals under carbon nanotube confinement is reported. In contradiction with its weak cohesive interactions, their structures, scrutinized using

neutron scattering, correspond to definite cylindrical sections of the hexagonal close-packed bulk crystal. The results are rationalized on the grounds of numerical calculations, which point towards nuclear quantum delocalization as the physical mechanism responsible for the stabilization of such outstanding structures.

Key words: Quantum Solids, 1D crystals, Confinement, Molecular Hydrogen, Neutron Scattering

1. Introduction

Carbon nanotubes are used to study crystallization under cylindrical constraints as well as to obtain a plethora of one-dimensional (1D) materials [1–3]. These new materials emerge, owing to the geometrical restriction, as crystalline phases with structures unlike their bulk counterparts (see chapter 5 in [3]). Some substances with strong cohesive interactions can retain their bulk crystallinity down to the strict one-dimensional limit, in which all the crystal constituents are simultaneously bulk and surface [4]. However, in general, for a certain level of geometrical restriction, crystalline arrangements with symmetries alien to the bulk phases develop [5]. As expected, this is the rule if the cohesive interactions are weak, for instance, hydrogen bonding in the archetypal case of water [6–8].

For purely dispersive interactions, molecular hydrogen (H_2) is an outstanding case owing to the strong quantum character of its condensed phases [9–11]. In particular, crystalline H_2 and its deuterium isotopologue (D_2) are prominent examples of quantum solids. Such crystals are characterized by a quantum nuclear delocalization sizable compared to the intermolecular distance, eventually rendering the harmonic crystal approximation useless [12]. The effect is maximized in the case of H_2 , where quantum fluctuations are so dominant that make the crystal insensitive to thermal variations [10, 11]. Furthermore, in both crystals the molecules behave as quantum free rotators, which implies that the nuclear orientational degrees of freedom are fully delocalized at low temperatures. Both crystalline states are van der Waals solids with an interaction potential well of just -2.77 meV, while that of the binding potential to a graphite surface amounts to $\simeq -50$ meV [13]. Confinement within narrow carbon nanotubes can roughly double this depth.

Calculations without including the quantum nuclear delocalization predict new ordered arrangements not present in the bulk [14–16], some of them reminiscent of spiral configurations observed in the iodine 1D case [17–19]. The search for superfluid behavior has spurred intense theoretical efforts to deal with the quantum nature of the H_2 molecular movements [20–24]. The results, not free of controversy, seem to favor a quantum solid resembling neither the spiral arrangements nor the hexagonal close-packed (HCP) structure of the bulk phase [23]. In the case of molecular deuterium, nuclear wave function calculations hint at the possibility of HCP structures within narrow carbon nanotubes [25].

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On the experimental side, as far as solidification under strong cylindrical confinement is concerned, the results are scarce. In carbon nanotubes, there is only evidence of transitions from one- to two-dimensional arrangements of D_2 on the external grooves of carbon nanotube bundles [26, 27], but no results about crystal structures within the nanotubes. There are hints of HCP crystallization of D_2 in the cylindrical mesopores of silica-based MCM-41 materials [28], but with a pore diameter as wide as 2.5 nm, quite far from the 1D limit. In these studies D_2 is preferred because the strong incoherent neutron scattering of H_2 makes diffraction measurements impractical.

In this work, by resorting to neutron diffraction (ND), crystallization of molecular deuterium within multi-walled carbon nanotubes (MWCNT) is studied. We have succeeded in resolving the crystalline structure down to the strict 1D limit. In spite of the weakness of the cohesive interactions mentioned above, the 1D crystals preserve the structure of its bulk counterpart. The strong quantum character of the bulk crystalline D_2 is also preserved in the 1D confined phases: the large delocalization of the nuclear positions as well as the full quantum nature of the rotational degrees of freedom are manifest in the measured neutron diffraction. The structures of the discovered 1D crystals correspond to maximally symmetric cylindrical cuts of the bulk HCP crystal. Such structures were addressed after the failure in reproducing the measurements by fully atomistic molecular dynamics (MD) simulations as well as by dispersive density functional theory (DFT) calculations corresponding to our experimental conditions. Instead, DFT and MD approaches yield spiral-like arrangements in full agreement with energy considerations as well as consistent with previously reported calculations. Since such calculations do not deal with quantum nuclear effects, given the strong quantum nature of the observed structures we conclude that quantum delocalization in the nuclear degrees of freedom is essential to stabilize the discovered 1D crystals.

The paper is organized as follows: in section 2 the experimental procedures and setups are described. The results, in particular the finally resolved crystalline structures, are presented and discussed in section 3. In section 4 our findings are summarized and the main conclusions exposed. To lighten the exposition, the details of materials and methods as well as of the theoretical developments used in the data analysis are given as appendixes. Likewise, the large series of diffraction patterns explored along with a detailed description of the nonlinear fitting analysis used to resolve the structures are presented in a Supplementary Material file. For the sake of reproducibility, details of custom-made experimental setups and specifics of the numerical runs made are also given there. Finally, figures corresponding to some complementary measurements and tables of numerical parameters complete the Supplementary Material.

2. Experimental

Our main goal is to study the structural evolution of molecular deuterium during solidification under extreme cylindrical confinement. In short, we measured neutron diffraction (ND) in runs of 350 seconds

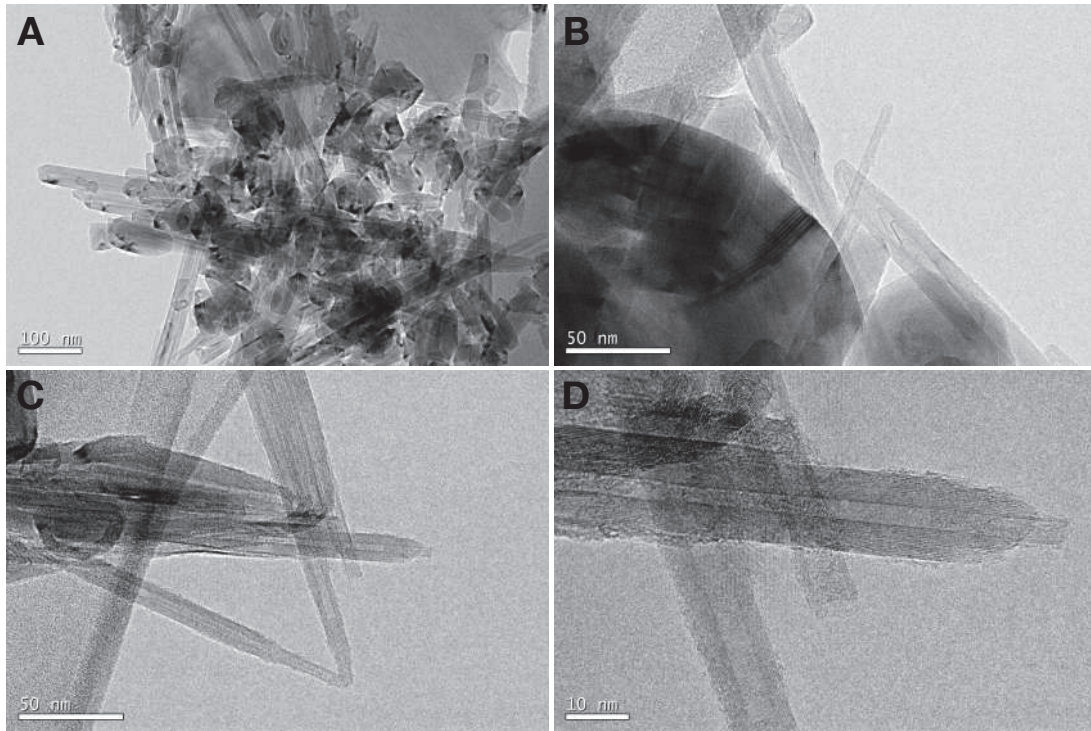


Figure 1: Some TEM pictures of the final processed samples. A corresponds to X 30000, B and C to X 100000 and D to X 300000

60 while submitting a sample of open MWCNTs to thermal cycles (between 24.5 K and 2 K) at four increasing loads of adsorbed D_2 at 20 K, corresponding to the equilibrium pressures 62 hPa, 175 hPa, 225 hPa, and 284 hPa. The details of the experiment design, the experimental setup as well as of the materials and methods used are given in Appendix B.

We synthesized MWCNTs by the arc discharge method. Like in the single-walled case, the tubes in the raw material are closed. In order to open them, we submitted the samples to an oxidation treatment (for details, see Appendix B.2). Transmission electron microscopy (TEM) of the final material (see Fig. 1) shows the expected structure for typical arc discharge MWCNT samples: It is quite heterogeneous, showing a wide variety of MWCNTs among a plethora of irregular polyhedral closed nanostructures. The surfaces do not present eroded scars or holes other than the openings of some nanotubes.

70 The adsorption of the sample was characterized through an isothermal D_2 adsorption/desorption measurement at 20 K using a custom-made setup. The isotherm in Fig. 2 shows during desorption a hysteresis beginning at the bulk vapor pressure. This is expected in this kind of samples. It comes from macro/mesopores, i.e., the interstitials among the various structures observed in the TEM. Consistently with the opening of a portion of the MWCNTs, the isotherm also presents a small, but noticeable, almost vertical increase at zero pressure, a feature revealing the presence of micropores of molecular dimensions

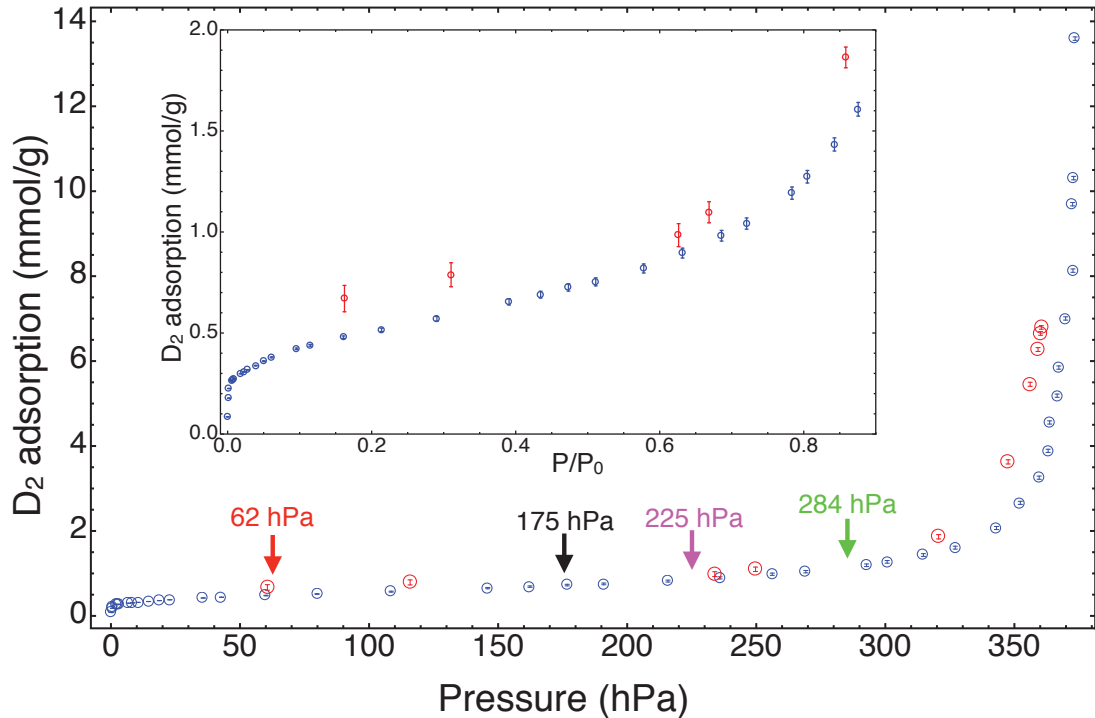


Figure 2: D₂ adsorption (blue)/desorption (red) isotherm at 20.23 ± 0.03 K for the sample of open MWCNTs. The amount of adsorbed gas is given in millimole per gram of sample. The arrows label the initial working points of the thermal cycles during the neutron diffraction measurements as described in the text. The inset is a zoom of the same data, focusing in the lower adsorption range in units of the bulk vapor pressure at 20.23 K (364.9 hPa [30]).

[29]. At sufficiently low loads of D₂, which correspond to low pressures in the isotherm, only the MWCNTs with the narrowest opened inner tubes are filled.

The chosen instrument was D20 at the Laue-Langevin Institute (ILL) in Grenoble, France, since it is a diffractometer of very high flux which allows to address very small quantities of scatterers in reasonable acquisition times (see details in Appendix B.6).

With respect to the neutron scattering, given the lightness of D₂ molecule, energy transfers between the incident neutron and the molecule affect the diffraction signal in a non-perturbative way. Usually this inelastic component is obtained heuristically since a proper correction requires a knowledge of the complete scattering law. Based on the neutron scattering theory for diatomic molecules developed by Sears [31], we have worked out an ansatz for the energy-integrated signal accounting for the observed $I_D(Q)$ that incorporates inelasticity without relying in any heuristic correction. From such an ansatz, the inelastic correction is given by a background, $bg(Q^2)$, expressible as an expansion in Q^2 (up to second order in practical terms). Physically, $I_D(Q)$ decomposes in a coherent and an incoherent part. The incoherent part stems from the random distribution of nuclear spin states along the sample. Formally is like the

self-scattering contribution coming out from the scatterers. The coherent part of $I_D(Q)$ depends on the orientationally-averaged structure factor of the molecular center-of-mass (COM) specific arrangement of D_2 molecules. It can be calculated efficiently from the pair distribution function, i.e., the table of COM pair distances $d_{ij} = |\langle \mathbf{R}_i \rangle - \langle \mathbf{R}_j \rangle|$ and its associated multiplicities m_{ij} , where \mathbf{R}_i is the position vector of the i molecule. An additional key ingredient to reproduce the ND coherent signal is a form factor corresponding to a spherical shell of diameter equal to the distance between the molecular nuclei, even in the solid phases at low temperature. This is a consequence of the nuclear orientational full delocalization. Likewise, the translational degrees of freedom also present a very significant quantum delocalization, which induces a large root mean square (rms) COM displacement. As explained in the next section, the observed diffraction signal reveals also the presence of oriented small crystals (or crystal) through barely discernible Bragg peaks. Although such precursor peaks can be ignored, we must consider the corresponding diffuse scattering, i.e., the diffraction signal between Bragg peaks in a crystal, which arises from such oriented crystallites. This is well approximated by a term of the form $1 - \exp(-Q^2 u_{3D}^2/3)$ plus an incoherent part, where u_{3D} is the rms molecular displacement of the molecules belonging to such crystallite. There is also an instrumental background whose major contribution to $I_s(Q)$ is a flat noise accounted for in $bg(Q^2)$. With these basic constituents we have constructed a theoretical model of the expected ND as a function of the pair distribution functions of the confined structures with the corresponding rms COM displacement, the proportion of external crystallite scatterers, and its associated rms COM displacement. The detailed derivation is provided in Appendix A.1. The final expression for a concentration (mole fraction) C_{1D} of 1D confined molecules with a specific structure and a concentration C_{3D} of molecules in the 3D crystallite reads,

$$I_D(Q) = C_{1D} (4a_d^2 j_0(Qd_{D_2}/2)^2 (S_{1D}^M(Q, u_{1D}) - 1) + F_{inc}(Q, u_{1D})) + C_{3D} \left(F_{inc}(Q, u_{3D}) - 4a_d^2 j_0(Qd_{D_2}/2)^2 \exp \left[-\frac{(Q u_{3D})^2}{3} \right] \right) + bg(Q^2), \quad (1)$$

where a_d is the neutron coherent scattering length of the deuteron, j_0 is the spherical Bessel function of zeroth order, d_{D_2} is the distance between the two deuterons, u_{1D} and u_{3D} are the corresponding rms molecular displacements, and the background reduces to $bg(Q^2) = bg_0 + bg_1 Q^2 + bg_2 Q^4$. $S_{1D}^M(Q, u_{1D})$ is the structure factor of the molecular COMs of the 1D crystal. The factor $j_0(Qd_{D_2}/2)^2$ is the spherical form factor mentioned above. The $F_{inc}(Q, u)$ terms are the incoherent part of the corresponding crystals. As explained in Appendix A.1, they depend not only in the temperature but in the concentration of ortho molecules. In practical terms our scattering intensity is not normalized to absolute units, so that C_{1D} and C_{3D} are scale factors proportional to the real concentrations.

With respect to the interference part $I_{DC}(Q)$, by its very nature, its most prominent contribution is around 1.8 \AA^{-1} where the main MWCNTs Bragg peak is located, being negligible for larger values of Q

since they correspond to spatial scales shorter than the D₂-C distances. This is confirmed a posteriori by the success of $I_D(Q)$ in reproducing the observed ND above 1.8 \AA^{-1} . An atomistic description of $I_{DC}(Q)$ is exceedingly expensive in computational terms; instead, we have assumed the nanotubes as continuous and infinite to reduce the calculation to sums of analytical expressions as detailed in Appendix A.2. The approximated $I_{DC}(Q)$ depends on the inner radius of the cylinder where the confined phase forms, r_i , on the dispersion of the distribution of external MWCNTs diameters, σ_n , on the average interlayer distance within the MWCNTs, h , and on an effective rms displacement, u_{eff} , to take into the account the disorder in the involved pair distances. We remark that its intensity with respect to $I_D(Q)$ is not free but determined by the calculation. Around 1.8 \AA^{-1} , the $I_{DC}(Q)$ profile and its derivative change wildly in a few points, so that iterative minimization algorithms fail to converge. Instead, $I_{DC}(Q)$ is added as a correction to the fitted $I_D(Q)$, the corresponding free parameters chosen by visual inspection comparing series of plots of the modeled $I_s(Q)$ against the measurements. More specifically, u_{eff} is chosen to the minimum value that does not perturb the independently fitted $I_D(Q)$. The value of h essentially controls the position of the “dispersive” profile around 1.8 \AA^{-1} . A value of 3.41 \AA works well for all the explored cases. The inner radius, r_i , is the most relevant parameter with respect to the profile shape. As shown below our $I_{DC}(Q)$ reproduces qualitatively the “dispersive” profile around 1.8 \AA^{-1} .

3. Results and Discussion

Fig. 3 displays the ND signal $I(Q)$ as a function of the scattering vector modulus Q of the empty MWCNT sample (black) superimposed to that of the 284 hPa load (green) at low temperature. From now on and for the sake of simplicity, we label the loads by their equilibrium pressures at 20 K. The red rectangle encloses the four main observed Bragg peaks. The normalized area of $I(Q)$ within the rectangle during a thermal cycle for the two lower loads is shown in the inset. A reversible change of the slope appears at around 13.5 K in both cases, revealing a liquid/solid transition at that temperature. Given that the bulk triple point is at 18.73 K, and that we are under SVP conditions, such a temperature reduction indicates that the transition is happening under confinement [32]. Furthermore, since the transition temperature depends of the pore size, its reversibility implies confinement under a rather narrow distribution of pore sizes. On the contrary, the curves of the two larger loads show hysteresis, as expected from the heterogeneous distribution in size and shape of the interstitials observed in the TEM pictures at meso/macroscales (see Fig. S3 in the Supplementary Material for the hysteretic cases).

Fig. 4 displays the ND of the solid phases at the four loads after subtraction of the empty MWCNT signal, denoted as $I_s(Q)$. Here, solid phases refer to those below 11.5 K, a value chosen well under the 13.5 K where the liquid/solid transition manifests. Consistently, the ND of all the independent runs below 11.5 K do not show any discernible changes, so that all the runs with $T < 11.5 \text{ K}$ have been accumulated. Details

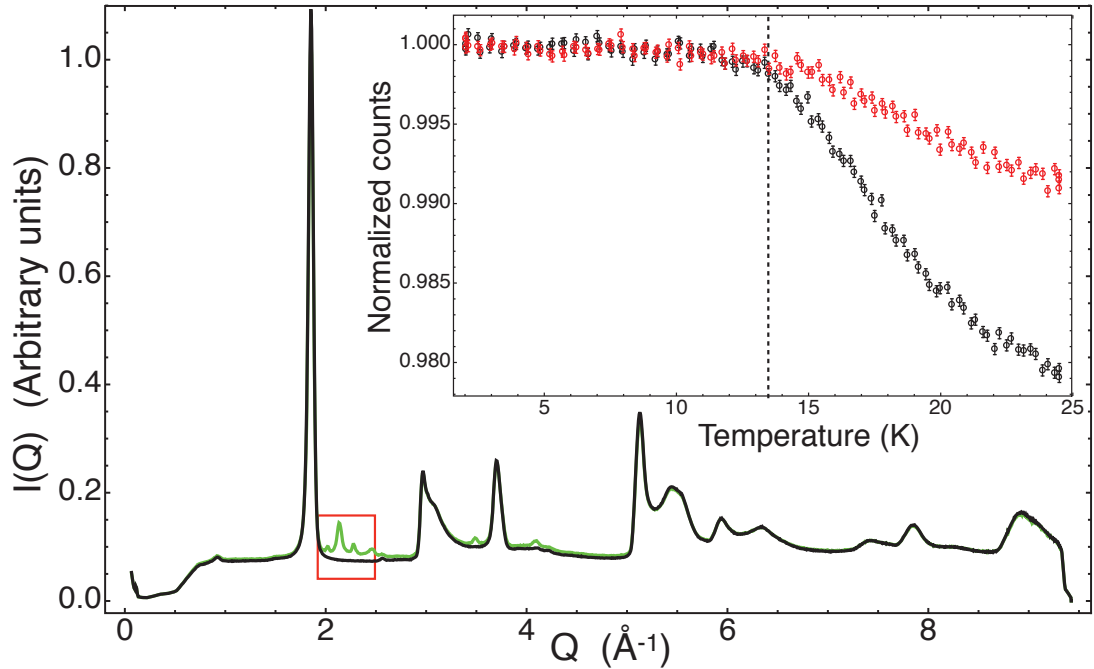


Figure 3: Neutron diffraction pattern, $I(Q)$, from the pristine MWCNT sample (black) superimposed to that corresponding to the 284 hPa D_2 load (green) at low temperature. The inset shows the evolution during a thermal cycle of the area (normalized to one at 2 K) under $I(Q)$ within the red rectangle for the 62 hPa load (red circles) and for the 175 hPa case (black circles).

of the neutron scattering data reduction are given in Appendix B.7. The 225 hPa and the 284 hPa loads
 125 present clear crystalline peaks identified as corresponding to both hexagonal close-packed (HCP) and face-
 centered cubic (FCC) phases. As shown in the figure, both kinds of peaks match to an excellent precision
 with strictly closed-packed arrangements with the same nearest-neighbor distance, namely, $d = 3.605 \text{\AA}$,
 a value in agreement with those reported for the bulk solids [33, 34]. The HCP phase is the fundamental
 crystal corresponding to the energy minimum of solid D_2 , but FCC crystallization can also be observed as
 130 soon as any perturbation is introduced in the crystallization environment, as for example, the heterogeneous
 nature of our MWCNTs sample.

D_2 tends to crystallize in rather large crystal domains. Since the ND instrument relies on a horizontal
 narrow strip detector geometry (see Appendix B.6), the non-uniform distribution of crystal orientations
 leads to distorted peak heights and the extinction of peaks with decreasing D_2 load. At first glance the
 135 175 hPa and 62 hPa patterns resemble those of amorphous substances with no Bragg peaks. A closer look,
 presented in the inset of Fig. 4, reveals the presence of a couple of precursors of bulk crystalline peaks at
 175 hPa and one at 62 hPa. This can be appreciated better in Fig. 5. Since these Bragg peak precursors
 appear in both thermal cycles of 62 hPa and 175 hPa, they reveal oriented crystal growth somewhere in the
 sample-can system, probably owing to an inhomogeneity-related nucleation site. The “pseudo-amorphous”

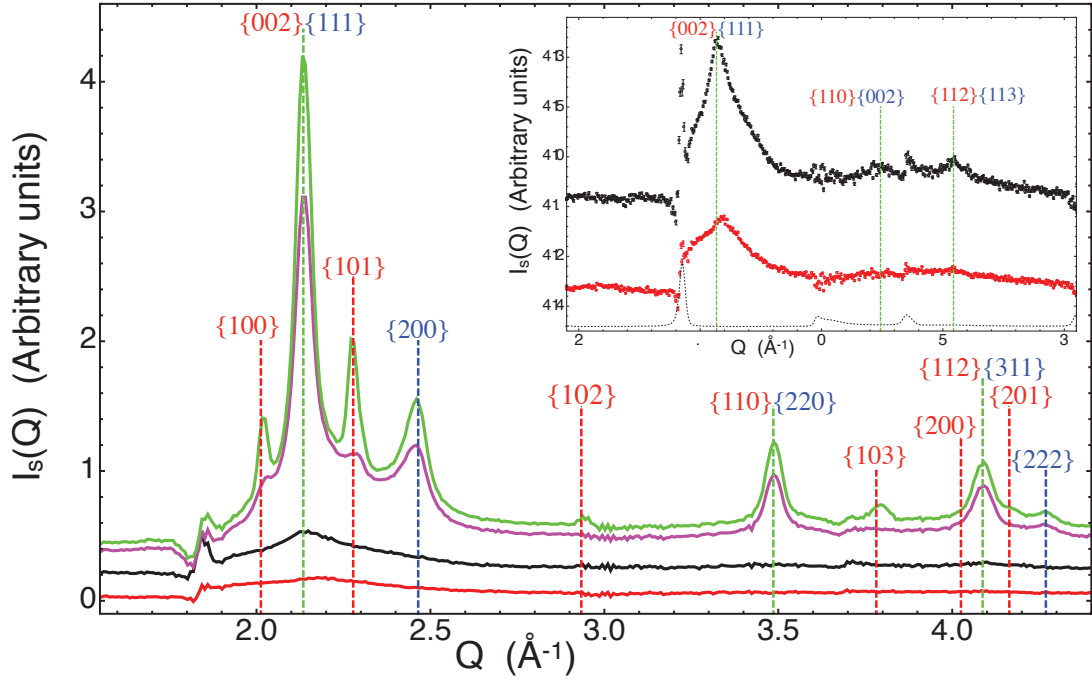


Figure 4: $I(Q)$ after subtraction of the MWCNT sample contribution, $I_s(Q)$, for the 62 hPa (red), 175 hPa (black), 225 hPa (magenta), and 284 hPa (green) D_2 loads. The vertical dashed lines mark the position of the bulk solid D_2 crystalline peaks. Dashed blue lines are for peaks present only in the HCP crystal, red dashed lines for those only present in the FCC crystal and green dashed line for peaks belonging to both. On top of the dashed lines the corresponding Miller indexes are specified, in red for the HCP structure and in blue for the FCC case. The inset is a zoom of the two lowest loads, i.e., 62 hPa (red) and 175 hPa (black) along with a scaled-down version of the MWCNT sample $I(Q)$ (dotted line).

part necessarily corresponds to confined phases, in view of the behavior of the $I(Q)$ area of each load shown in the inset of Fig. 3. Within this respect, they depart significantly from the observed broader patterns in solid D_2 (10 K) adsorbed in SWCNT bundles [26]. There, the fits ascribe the ND to the presence of chains and stripes of D_2 molecules expected to be within the interstitials (chains) and external grooves (chains and stripes) of the bundles. These two loads are, therefore, the object of our main interest. The scattering from D_2 pairs, $I_D(Q)$, contributes dominantly to $I_s(Q)$. There is also a contribution from D_2 and carbon pairs, $I_{DC}(Q)$, relevant mainly below around 2 \AA^{-1} . We focus first in modeling the dominant deuterium part, $I_D(Q)$.

Using the $I_D(Q)$ theoretical model described in section 2, a nonlinear fitting analysis detailed in the Supplementary Material was performed along a family of constrained nanostructures. We naturally started from structures obtained by MD simulations including MWCNTs up to five walls and effective potentials derived from state-of-the-art ab-initio calculations. Not surprisingly, they match the expected arrangements from energy minimization arguments: tubular shells adsorbed to the inner wall of the nanotubes that can be seen as rolled portions of the D_2 HCP crystal basal plane. As a matter of fact, they are explicit realizations

of the tubular compact packing of spheres studied in the context of biological structures [35]. However, they
155 are not compatible with our measurements as illustrated in Fig. S4, Supplementary Material.

Such disagreement drove us to consider other 1D crystalline structures corresponding to sections of
maximal cylindrical symmetry of both the HCP and FCC bulk crystals with increasing diameter. For these
crystals we fixed d to the value obtained from the 225 hPa and 284 hPa loads, i.e., 3.605 Å. Figure 5
presents the final results of our nonlinear model fitting analysis. Let us consider first the 62 hPa load (Fig.
160 5A). The agreement is excellent just using a log-normal distribution in length (see inset) of the remarkable
strict 1D crystal displayed in the right side of the figure. Differences of $I_D(Q)$ with increasing length become
negligible (see Supplementary Material Figures S10 and S11). Consequently, there is not enough information
in the diffraction signal to discern the 1D crystal length distribution; in fact, a non-parametric fit of the
crystal lengths only needs to populate a few isolated lengths to achieve excellent results. Lacking further
165 information, we have relied in the log-normal distribution because it is usually observed in growth processes
with preferential direction [36, 37]. With respect to the ortho concentration, the fits need a proportion
corresponding to room temperature, i.e., there was no para-ortho conversion during the experiment. The
result is not only of a sheer beauty in its simplicity — the simplest nontrivial strict 1D compact crystal
made out of spheres — but incompatible with theoretical predictions, as discussed above. It corresponds to
170 a cylindrical cut of the minimal diameter which allows more than one molecule in the horizontal plane with
the axis perpendicular to the hexagonal basal plane of the HCP fundamental bulk crystal, and centered so
that it passes through the molecular centers at alternate layers. The six-fold symmetry of the parent HCP
bulk crystal is reduced to three-fold (Fig. 5A, top right panel), so we call it triangular-close packed (TCP)
1D crystal.

Consistently, in the 175 hPa load (Fig. 5B) the nanocrystalline structure is just a radially enlarged version
of the TCP 1D crystal. Again, there is a three-fold symmetry, as it can be seen in the transversal cut of
the crystal, upper right panel of Fig. 5B. Since the crystal consists of alternating triangular and hexagonal
layers, we denote it as THCP 1D crystal. Both 1D crystals display large rms COM displacements, around
0.33 Å in the TCP case, and 0.45 Å in the THCP counterpart, to be compared with the 0.46 Å obtained
180 from neutron scattering of the D₂ bulk crystal [38] (see Tables SI and SII in Supplementary Material).
Interestingly, there is no hint in the 175 hPa ND of the pattern corresponding to the 62 hPa load. Indeed,
the fittings yield a zero population of the TCP phase, i.e., any finite population of the TCP phase worsens
the fits. Should the TCP 1D crystals form also during the 175 hPa load in quantities comparable to the
62 hPa case, they would also manifest in the 175 hPa ND. Therefore, either the THCP crystals grew out
185 of previously formed TCP seeds, or the MWCNTs of the smallest diameters are not filled during the 175
hPa load. In the first hypothesis, the TCP crystals grown at 62 hPa would be formed within MWCNTs of
large enough inner diameter, whereas in the latter the two crystals would grow within MWCNTs of different
inner diameters. Fortunately, the interference part, $I_{DC}(Q)$, carries information about the inner MWCNTs

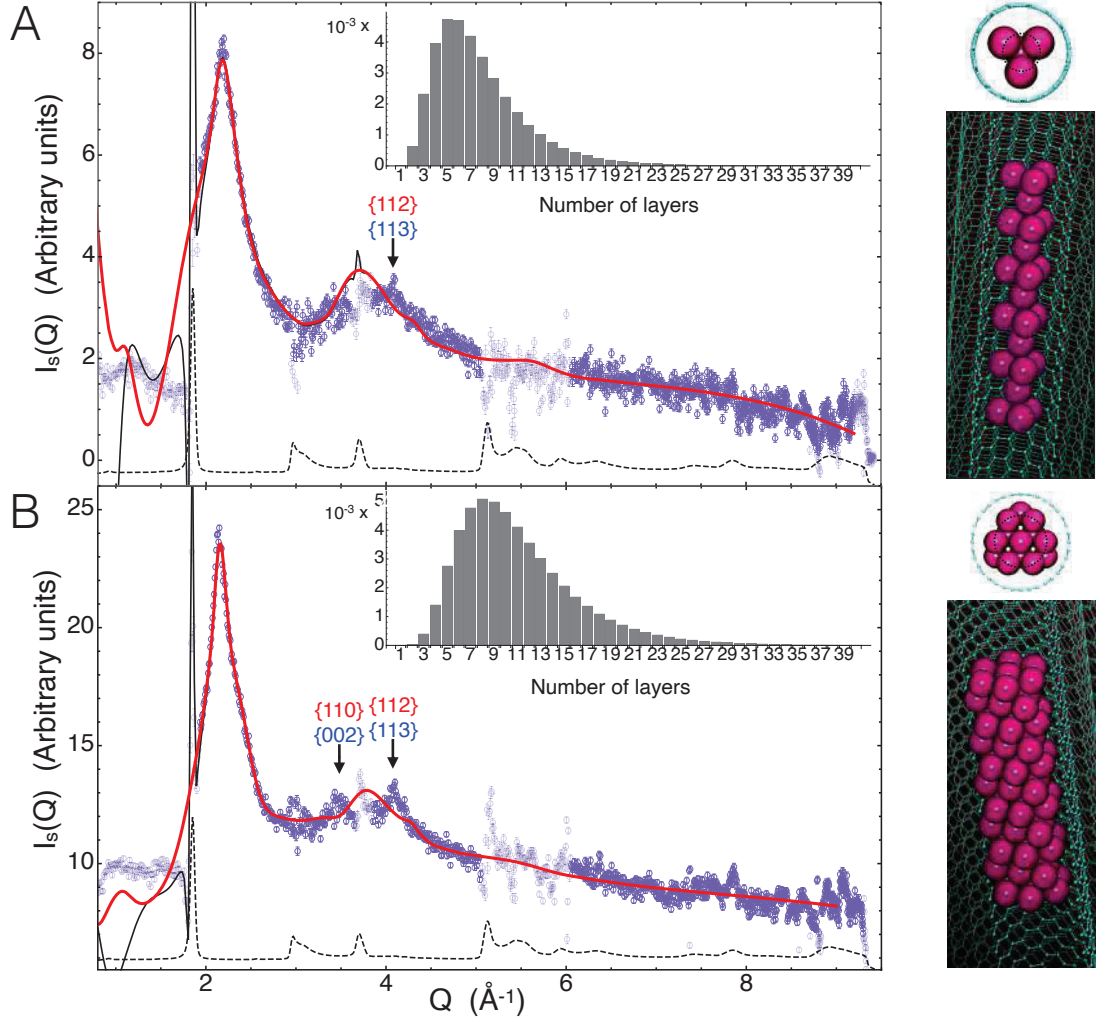


Figure 5: D₂ 1D crystals fitting the ND data. (A) D₂ load corresponding to the 62 hPa case. (B) D₂ load corresponding to the 175 hPa case. ND data points displayed in light blue are not included in the fittings (points below 2 \AA^{-1} , areas too affected by MWCNT peak subtraction, and some outliers). Solid red curves are the fitted $I_D(Q)$ component. Thin black curves show the final $I_s(Q)$ including the $I_{DC}(Q)$ part. The dashed lines are a down-scaled representation of the pristine MWCNT sample ND. Arrows mark the positions of some relevant Bragg peaks of the bulk crystal. At the right side, a representative structure giving rise to the corresponding $I_s(Q)$ is shown in perspective. The small spheres have a diameter equal to the deuteron-deuteron distance of the D₂ molecule (0.74 \AA). The diameter of the semitransparent larger spheres equals $d = 3.605 \text{ \AA}$. In the top views only the inner nanotubes are shown, an (11,4) single-walled CNT (A) and an (18,0) (B). Dotted circumferences mark the position of the minimum of the adsorption potential. Insets display the fitted log-normal distributions of 1D crystal lengths in terms of the number of molecular layers perpendicular to the MWCNT axis.

diameters so we can use it to settle this question. Fig. 5 shows in solid black the complete NDs including
 190 the $I_{DC}(Q)$ contribution. By construction, it coincides with $I_D(Q)$ above $\sim 1.8 \text{ \AA}^{-1}$ so that it falls below
 the red line representing $I_D(Q)$. As shown in the top right panels of Fig. 5, the chosen inner diameters
 $2r_i$ (10.53 \AA for the TCP and 14.09 \AA for the THCP) accommodate well the D_2 molecules in terms of
 the estimated locus [39] of the adsorption potential minima. However, inner diameters around 14 \AA in the
 TCP case render $I_{DC}(Q)$ profiles incompatible with the measured ND at 62 hPa. We are therefore left to
 195 conclude that the MWCNTs of the narrowest inner diameters are not significantly filled during the 175 hPa
 load. Since we are referring to pore sizes at the molecular limit (below two molecular diameters), steric
 molecular hindrance as well as anomalous capillary and condensation effects [40–42], dependent on the D_2
 load, seem plausible. Within this respect, the irregular shapes and the functional groups at the entrance
 of the opened nanotubes could play a relevant role. For instance, funnel-like attractive potentials at the
 200 entrance would probably enhance the possibility of molecular jamming at 175 hPa owing to the different
 kinetics and density with respect to the 62 hPa case. Given the large de Broglie wavelength of D_2 , quantum
 wave-like behavior contributing to these effects, as observed with He [43–45], are also conceivable. At any
 rate, here we can only speculate about the microscopic mechanisms involved in this interesting anomalous
 behavior since their elucidation requires a study by its own.

205 Taking into the account that MD simulations do not describe accurately dispersive interactions, which are
 crucial in this kind of systems, we have performed advanced dispersive DFT calculations validated against
 experiments, in order to shed light on the discrepancy between the observed structures and theoretical
 predictions (details of the calculations are given in Appendix B.9 and Supplementary Material). Due to the
 large computational demand of this approach, we restrict ourselves to SWCNTs of a diameter compatible
 210 with the TCP 1D crystal. More specifically, the nanotube of chiral vector (10,5) has an adequate unit cell
 length while its diameter of 10.36 \AA is just 2% below that of the (11,4) tube shown in Fig. 5A. The number
 of D_2 molecules is chosen to be eleven, corresponding to five layers of the 1D TCP crystal, the most probable
 length (inset of Fig. 5A). Fig. 6 summarizes the DFT results. Spiral geometries similar to those obtained by
 MD are once more the most stable configurations, at variance with the experimental results (see Fig. S4A of
 215 Supplementary Material). Remarkably, the observed 1D TCP structure, presented in Fig. 6B, is metastable
 within this approach, but the next most plausible 1D crystal derived from the FCC bulk structure is not
 stable under the DFT approximation (depicted in Fig. S5B of Supplementary Material). The calculated
 $I_D(Q)$ s are also shown in Fig. 6 along with those corresponding to the equivalent TCP 1D structure obtained
 from the fits. Additionally, notice that the DFT calculations yield an effective nearest-neighbor distance
 220 slightly shorter than the experimental one. This is a consequence of the quantum delocalization of the
 rotational degrees of freedom of the nuclei, not taken into account by the DFT approach, which renders
 the molecular interactions essentially isotropic. On the contrary, within the DFT approximation the D_2
 molecules retain its dumbbell-like anisotropic character and tend to order in a parallel, more compact,

configuration.

225 The present results tell of a van der Waals molecular compound that crystallizes retaining the bulk crystal
structure down to the strict 1D limit. This is in stark contrast with results pertaining to other materials
with weak cohesive energies, which do not show the marked quantum behavior in their translational and/or
rotational degrees of freedom as our 1D crystals do, conspicuously revealed by their neutron scattering
response. Our DFT calculations, with reliable dispersive functionals, make clear that the missing quantum
230 treatment of the nuclear motions are one key ingredient for the stabilization of these 1D crystals. On the
other hand, the fact that the TCP 1D crystal is weakly metastable under the DFT calculations suggest
that a precise account of the quantum behavior of the electronic cloud in the description of the dispersive
interactions could also be relevant. In fact, the development of dispersive DFT techniques during the last two
decades is characteristic of the nanoscience revolution: adsorption phenomena within nanostructures depend
235 too subtly on the dispersive interactions to be accommodated within the traditional effective potential
approach. The challenge now is to incorporate the nuclear quantum effects accurately, a subject attracting
an increasing interest [46]. At present, inclusion of translational quantum nuclear delocalization effects
in condensed matter can be tackled with computationally very demanding quantum Monte Carlo or path
integral methods [23, 24] but we remark that, both, the nuclear rotational delocalization and the dispersive
240 forces are treated in the same effective manner as in our MD simulations. Within this context, the structures
here discovered stand out for testing future theoretical developments since they are dispersive molecular
solids displaying a non-trivial quantum nuclear behavior at the same time being structurally simple.

4. Conclusion

Using neutron diffraction we have discovered 1D quantum crystals of molecular deuterium within MWC-
245 NTs. Unexpectedly for a molecular solid with very weak cohesive interactions, the 1D crystals preserve the
structure of the HCP bulk D_2 crystal. Neither atomistic molecular dynamics simulations nor advanced dis-
persive DFT calculations predicts such a possibility, signaling the strong quantum nuclear effects as essential
to the emergence of the discovered 1D crystals.

These 1D crystals are the simplest nontrivial condensed matter systems in which the quantum nature,
250 not only of the electronic cloud, but also of the nuclear translational and rotational degrees of freedom,
plays an essential role. Our results are deemed to be of special relevance, since they provide a testbed to
confront theoretical approaches aimed to understand and incorporate accurately the full quantum nature of
real interactions, including the nuclear degrees of freedom. Such advances will have a bearing on the design
and simulation of novel (possibly quantum) materials.

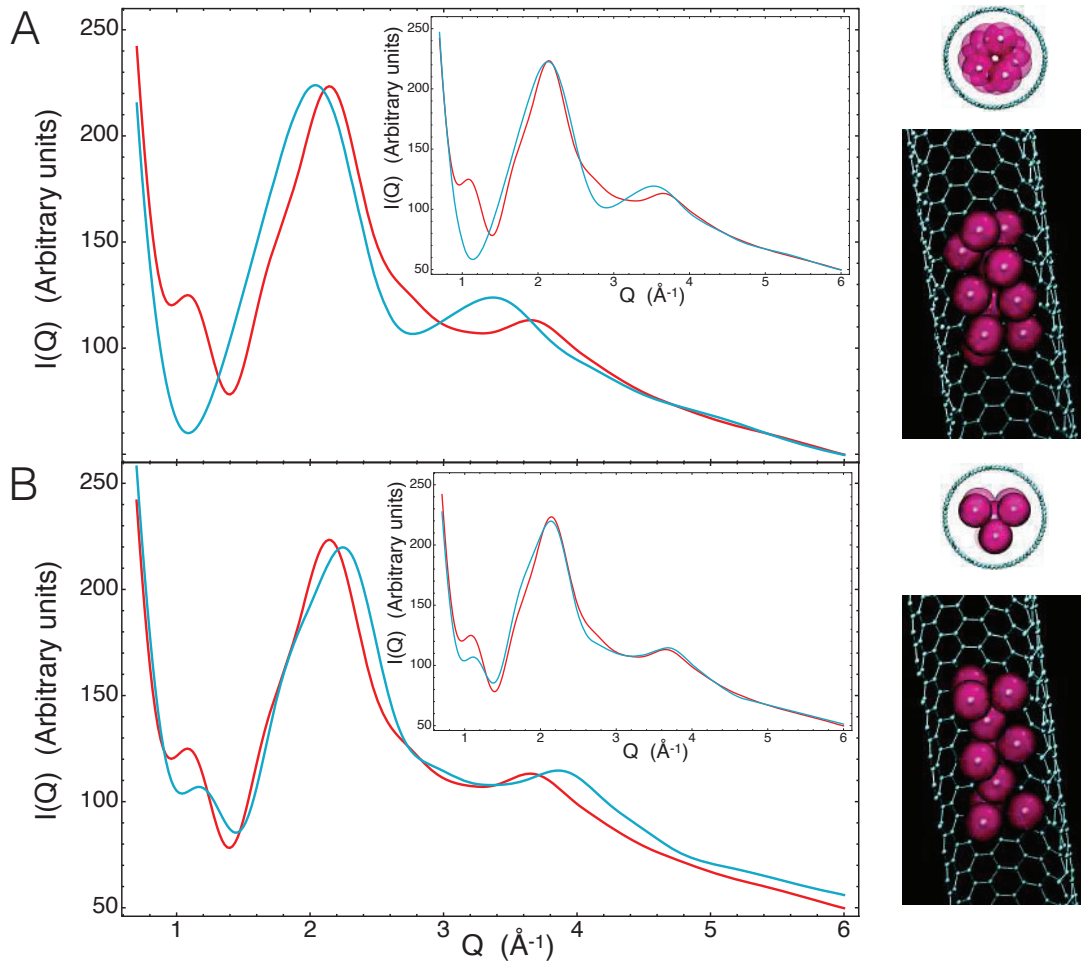


Figure 6: 1D structures obtained by dispersive DFT calculations. (A) Fundamental structure corresponding to the lowest energy minimum. (B) Structure of the 1D TCP kind corresponding to a metastable energy minimum. Their associated $I_D(Q)$ s are also shown (cyan curves) along with that corresponding to the experimentally-determined equivalent 1D TCP crystal (red curves). Since the DFT calculations do not reproduce the nearest-neighbor distance accurately, properly Q -rescaled versions of the $I_D(Q)$ s are displayed in the insets.

255 **Acknowledgements**

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Appendix A. Theoretical developments

270 *Appendix A.1. Theory of the measured neutron scattering*

In a neutron diffraction experiment at a continuous source installation such as the ILL, the intensity of scattered neutrons by the sample out of a mono-energetic incident neutron beam is measured as a function of the scattering angle. The relevant differential scattering cross section is then given by

$$\frac{d\sigma}{d\Omega} = \int_{-\infty}^{E_i} \frac{d^2\sigma}{d\Omega dE_f} dE_f, \quad (\text{A.1})$$

where the Ω denotes solid angle, E_i the energy of the incident neutrons and E_f the final detected neutron energy. Following the full quantum calculations of Sears [31], the double differential cross section in the case of a homonuclear diatomic molecule under the assumption of free rotation, can be expressed as

$$\frac{d^2\sigma}{d\Omega dE_f} = N \frac{k_f}{k_i} \left(4 a_d^2 j_0(Q d_{D_2}/2)^2 S_{int}(\mathbf{Q}, E) + \sum_{J, J'} a^2(Q; J, J') S_{self}(Q, E - E_{J, J'}) \right). \quad (\text{A.2})$$

Here N is the number of nuclei in the sample, $k_{i,f}$ are the initial and final neutron angular wavenumbers, a_d is the coherent neutron scattering length of the deuteron, j_0 the spherical Bessel function of zero order, d_{D_2} the distance between the nuclei in the molecule, \mathbf{Q} is the scattering angular wavevector (Q its norm), the J s denote the nuclear orbital angular momentum quantum levels, $E_{J, J'}$ the corresponding energy jumps among J levels and $E = E_i - E_f$ is the energy transfer. The $a^2(Q; J, J')$ factors are quite complex expressions involving linear combinations of Bessel spherical functions dependent on the bosonic or fermionic character

of the nuclei [31]. They depend on all the rotational levels excited by the neutrons at the given incident energy and, as such, incorporate the corresponding orientational form factors. Transitions between the ortho (even J angular momentum) and para (J odd) levels are forbidden so that thermalization of the populations proceeds independently. The $J = 1$ state is highly metastable so that equilibration at low temperatures downwards the fundamental $J = 0$ level is extremely slow in the bulk [12] but it could be accelerated by the interaction with the substrate. Otherwise, thermal equilibration proceeds unhindered so that the model depends on the temperature and the concentration of ortho molecules [31].

$S_{int}(\mathbf{Q}, E)$ is the interference part of the dynamical structure factor of the molecular COMs, i.e., the part not involving single particle self-scattering. Such self-scattering part (of the COMs) corresponds to $S_{self}(Q, E)$ (Sears uses the subscript *inc* from incoherent). It is a consequence of the free rotation assumption that both $S_{int}(\mathbf{Q}, E)$ and $S_{inc}(Q, E)$ in Eq. (A.2), refer only to the dynamics of the COMs of the molecules. In order to calculate Eq. (A.1), we need a model of such dynamical structure factors. Since their contribution to Eq. (A.1) is through integration in E , we will take advantage of general sum rules, i.e., moments in E , that must be satisfied independently of the system dynamics.

Focusing on the interference part, the relevant sum rule is $\int_{-\infty}^{\infty} S_{int}(\mathbf{Q}, E) dE = S(\mathbf{Q}) - 1$, where the $S(\mathbf{Q})$ is the structure factor. This is the quantity of interest, since it yields a structural characterization of the system and it is amenable to theoretical modeling. It is given by

$$S(\mathbf{Q}) = 1 + \frac{1}{N} \sum_{i \neq j} \langle \exp [i(\mathbf{R}_i - \mathbf{R}_j) \cdot \mathbf{Q}] \rangle, \quad (\text{A.3})$$

where N is the number of scatterers, \mathbf{R}_j the position vector of the j molecular COM and $\langle \rangle$ denotes quantum-mechanical thermal average. In a solid, thermal fluctuations are well approximated by a Gaussian distribution of displacements around the mean positions $\langle \mathbf{R}_j \rangle$. In such a case, after orientational and thermal average, Eq. (A.3) reduces to

$$S(Q) = 1 + \frac{1}{N} \sum_{i \neq j} j_0(|\langle \mathbf{R}_i \rangle - \langle \mathbf{R}_j \rangle|Q) e^{-\frac{1}{3}u^2 Q^2} = 1 + \frac{1}{N} \sum_{d_{ij}} m_{ij} j_0(d_{ij}Q) e^{-\frac{1}{3}u^2 Q^2}, \quad (\text{A.4})$$

where the final sum is over all the different pair distances $d_{ij} = |\langle \mathbf{R}_i \rangle - \langle \mathbf{R}_j \rangle|$, m_{ij} are the corresponding multiplicities and u is the rms displacement around the $\langle \mathbf{R}_j \rangle$, assumed here, isotropic and independent of the position. In a solid, the vast majority of the interference scattering is elastic, i.e., it accumulates around $E = 0$, so that our model for the interference part is $S_{int}(Q, E) = (S(Q) - 1) \delta(E)$, where $\delta(E)$ denotes the Dirac delta.

With respect to the self part the elastic term amounts to

$$S_{self}^e(Q, E) = e^{-\frac{1}{3}u^2 Q^2} \delta(E). \quad (\text{A.5})$$

However, it is not so dominant as in the interference case, so we need to proceed further. The relevant

moments to develop a model of the inelastic part are [47]

$$\int_{-\infty}^{\infty} S_{self}(Q, E) dE = 1, \quad (\text{A.6})$$

$$\int_{-\infty}^{\infty} E S_{self}(Q, E) dE = \frac{\hbar^2 Q^2}{2M} = E_r, \quad (\text{A.7})$$

$$\int_{-\infty}^{\infty} E^2 S_{self}(Q, E) dE = \frac{4}{3} E_r \bar{K} + E_r^2, \quad (\text{A.8})$$

where M is the mass of the molecule, E_r is the so-called recoil energy and \bar{K} is the mean kinetic energy. The zeroth order moment is always satisfied, the first order moment is satisfied provided that the interaction does not depend on the molecular velocities, and the second is true for isotropic systems. The elastic part does not contribute to the first and second moment; decomposing the incoherent part as $S_{self}(Q, E) = S_{self}^e(Q, E) + S_{self}^{ine}(Q, E)$ and using Eq. (A.5), the zeroth moment implies

$$\int_{-\infty}^{\infty} S_{self}^{ine}(Q, E) dE = 1 - e^{-\frac{1}{3}u^2 Q^2}. \quad (\text{A.9})$$

On the other hand, from the first and the second moments, the energy dispersion is obtained:

$$\sigma_E^2 = \langle E^2 \rangle - \langle E \rangle^2 = \frac{4}{3} E_r \bar{K}. \quad (\text{A.10})$$

The mean kinetic energy at low temperature is expressed in terms of the rms displacement as [48, 49],

$$\bar{K} = \frac{9}{16} \Theta = \frac{81}{64} \frac{\hbar^2}{Mu^2}, \quad (\text{A.11})$$

where Θ is the Debye temperature, the parameter usually quoted in the experimental reports. We are now in disposition to write up a model for the incoherent part preserving all the moments up to the second:

$$S_{self}^M(Q, E) = e^{-\frac{1}{3}u^2 Q^2} \delta(E) + (1 - e^{-\frac{1}{3}u^2 Q^2}) \frac{1}{\sqrt{2\pi\sigma_E^2}} \exp\left[-\frac{(E - E_r)^2}{2\sigma_E^2}\right]. \quad (\text{A.12})$$

where the M superscript serves to recall it refers only to molecular COM contribution. The chosen Gaussian shape for the inelastic scattering is a natural choice to warrant the integral properties up to second order.

300 Such a profile must be understood as a kind of smoothing of the real signal since it does not incorporate the details of the density of vibrational modes. Only in the limit of very large momentum transfer the same kind of profile would be exact (impulse approximation) but for the nuclei themselves, not for the molecular COMs. Interestingly, in the case of H_2 , given the lightness of the molecule, this profile for the COM dynamics has been observed at least from Q 's as low as 5 \AA^{-1} [50].

In order to calculate Eq. (A.1) the factor $k_f/k_i = k_i \sqrt{1 - E/E_i}$ must be also included in the inelastic part. Conservation of energy and momentum establishes a relation among Q , E , and, ϕ , the angle between \mathbf{k}_i and \mathbf{k}_f , namely,

$$Q^2 = k_i^2 + k_i^2 \left(1 - \frac{E}{E_i}\right) - 2k_i^2 \sqrt{1 - \frac{E}{E_i}} \cos \phi. \quad (\text{A.13})$$

305 Substituting Q in terms of E and ϕ , the integration of Eq. (A.2) in E yields the measured cross section Eq. (A.1). Notice that direct integration of Eq. (A.2) does not yield the single cross section, since it is defined (and measured) at constant ϕ , not at constant Q . However, usually the ND is expressed in terms of Q , but only at its elastic value using the relation $Q_e = 2k_i \sin \phi/2$. As a rule, the elastic condition is not explicitly stated but the same symbol Q is used. We adhere here to such a practice. At any rate, in many experiments 310 E_i is sufficiently high for the static approximation to be valid [47], where all the scattering can be considered elastic.

As far as the fitting is concerned, we first generate a two-dimensional numerical table in u and Q of the expression

$$F_{inc}(Q, u) = \int_0^\infty \frac{k_f}{k_i} \sum_{J, J'} a^2(Q; J, J') S_{self}(Q, E - E_{J, J'}) dE, \quad (\text{A.14})$$

where the suffix *inc* remarks that this expression includes not only the coherent self-contribution but the incoherent scattering. In the actual fittings a two-dimensional interpolation of the table is used to avoid the computational load of a numerical integration at each fitting step.

With respect to the diffuse coherent scattering of the external crystallite, performing only the thermal average in Eq. (A.3), we have

$$S(\mathbf{Q}) = 1 + \frac{e^{-\frac{1}{3}u^2 Q^2}}{N} \sum_{i \neq j} e^{i(\langle \mathbf{R}_i \rangle - \langle \mathbf{R}_j \rangle) \cdot \mathbf{Q}} = 1 + e^{-\frac{1}{3}u^2 Q^2} \left(\frac{1}{N} \sum_i \left| e^{i\langle \mathbf{R}_i \rangle \cdot \mathbf{Q}} \right|^2 - 1 \right) \quad (\text{A.15})$$

$$= 1 - e^{-\frac{1}{3}u^2 Q^2} + e^{-\frac{1}{3}u^2 Q^2} \frac{1}{N} \sum_i \left| e^{i\langle \mathbf{R}_i \rangle \cdot \mathbf{Q}} \right|^2 = 1 - e^{-\frac{1}{3}u^2 Q^2} + S_B(\mathbf{Q}). \quad (\text{A.16})$$

In a crystal, and in the limit of $N \rightarrow \infty$, $S_B(\mathbf{Q})$ gives rise to the Dirac delta peaks characteristic of Bragg scattering. The rest is obviously the scattering among the Bragg peaks, i.e., the diffuse scattering. This remains true also at finite N , where the peaks broaden. Therefore, the diffuse scattering from external crystallite contributes with an interference part given by

$$S_{int}(\mathbf{Q}, E) = -e^{-\frac{1}{3}u^2 Q^2} \delta(E), \quad (\text{A.17})$$

315 while the incoherent part is again given by $F_{inc}(Q, u)$, but this time with u corresponding to that of the external crystallite.

Altogether, for a number concentration C_{1D} of 1D confined molecules and a C_{3D} number concentration of molecules in the 3D external small crystal, gives,

$$I_D(Q) = C_{1D} \left(4a_{coh}^2 j_0(Qd_{D_2}/2)^2 (S_{1D}^M(Q, u_{1D}) - 1) + F_{inc}(Q, u_{1D}) \right) + C_{3D} \left(F_{inc}(Q, u_{3D}) - 4a_{coh}^2 j_0(Qd_{D_2}/2)^2 \exp \left[-\frac{(Q u_{3D})^2}{3} \right] \right), \quad (\text{A.18})$$

where $S_{1D}^M(Q, u_{1D})$ is the structure factor of the molecular COMs as given by Eq. (A.4) and u_{1D} and u_{3D} are the corresponding rms displacements.

Besides Eq. (A.18) there must be an instrumental flat background, bg_0 . With such an addition, Eq. (A.18) produces convergent fits to the measured $I(Q)$. That is, it captures adequately most of the ND signal. A final refinement is now expressible as series in Q^2 . A background second order in Q^2 , $bg_0 + bg_1Q^2 + bg_2Q^4$, is enough to obtain excellent fits.

Appendix A.2. Modeling of the C–D₂ diffraction interference term

In terms of the positions of the D₂ COMs, \mathbf{R}_i and of the carbon atoms, r_j , the ND signal corresponding to the interference between carbon atoms and D₂ molecules reads,

$$\begin{aligned} I_{DC}(Q) &= C_{1D} a_c 2 a_d j_0(Qd_{D_2}) \frac{1}{N} \sum_i \sum_j \left\langle e^{-i\mathbf{Q}\cdot(\mathbf{R}_i - \mathbf{r}_j)} \right\rangle + c.c. \\ &= C_{1D} a_c 2 a_d j_0(Qd_{D_2}) e^{-\frac{1}{6}Q^2(u_{1D}^2 + u_c^2)} \frac{1}{N} \sum_i \sum_j \left\langle e^{-i\mathbf{Q}\cdot(\langle \mathbf{R}_i \rangle - \langle \mathbf{r}_j \rangle)} \right\rangle + c.c., \quad (\text{A.19}) \end{aligned}$$

where *c.c.* denotes complex conjugate, N is the number of molecules, a_c is the coherent neutron scattering length of carbon nuclei, u_c their rms mean displacement and in the second line we have performed thermal average, again assumed Gaussian and isotropic. Consequently, $\langle \rangle$ reduces to only an orientational average. Here, $2 a_d j_0(Qd_{D_2})$ plays the role of the coherent scattering length of the D₂ molecule, dependent on Q owing to the non point-like molecular structure.

An atomistic description of the MWCNTs is computationally too expensive. However, assuming a continuous distribution of the carbon atoms along infinite tubes, we can take advantage of some analytical calculations. Given a microscopic density of carbon atoms, $\rho_c(\mathbf{r})$, a general term of the sums above is written as

$$\left\langle e^{-i\mathbf{Q}\cdot\langle \mathbf{R}_i \rangle} \int_V e^{-i\mathbf{Q}\cdot\mathbf{r}} \rho_c(\mathbf{r}) d\mathbf{r}^3 \right\rangle = \frac{1}{4\pi} \int d\Omega_Q e^{-i\mathbf{Q}\cdot\langle \mathbf{R}_i \rangle} \int e^{i\mathbf{Q}\cdot\mathbf{r}} \rho_c(\mathbf{r}) d\mathbf{r}^3, \quad (\text{A.20})$$

where $d\Omega_Q$ is the solid angle corresponding to a direction of the scattering vector and the integrals extend to all the directions and the whole space respectively. For an infinite tube of diameter R_{CNT} , the density is expressed in the cylindrical coordinates r, φ, z , with z axis that of the cylinder as $\rho_c(\mathbf{r}) = \rho_A \delta(r - R_{CNT})$, being ρ_A the areal density of carbon in the nanotube. Analytical integration in the cylindrical coordinates yields

$$\int e^{i\mathbf{Q}\cdot\mathbf{r}} \rho_c(\mathbf{r}) d\mathbf{r}^3 = 4\pi^2 \rho_A R_{CNT} J_0(QR_{CNT}) \delta(Q_z), \quad (\text{A.21})$$

where J_0 is the Bessel function of order zero and Q_z is the z component of the scattering vector. Notice that for an infinite tube the symmetry leads to cancellation of the dependence along the z axis. Substituting Eq. (A.21) in Eq. (A.20), again analytical integration can be done, this time using spherical coordinates, so that

$$\left\langle e^{-i\mathbf{Q}\cdot\langle \mathbf{R}_i \rangle} \int_V e^{-i\mathbf{Q}\cdot\mathbf{r}} \rho_c(\mathbf{r}) d\mathbf{r}^3 \right\rangle = 2\pi^2 \rho_A R_{CNT} J_0(QR_{CNT}) J_0(QR_i^\perp) / Q, \quad (\text{A.22})$$

where R_i^\perp is the modulus of the projection of $\langle R_i \rangle$ onto the plane perpendicular to the tube axis. Now the interference term for a SWCNT amounts to

$$I_{DC}(Q) = 8\pi^2 C_{1D} a_d \rho_A j_0(Qd_{D_2}) e^{-\frac{1}{6}Q^2(u_{1D}^2 + u_c^2)} R_{CNT} J_0(QR_{CNT}) \frac{1}{N_c Q} \sum_i J_0(QR_i^\perp), \quad (\text{A.23})$$

where i runs from 1 to N_c , being N_c the number of D_2 molecules of a unit cell of the corresponding 1D
330 crystal.

For a distribution of MWCNTs of well-defined internal radius r_i and m layers separated by a distance h , the term $R_{CNT} J_0(QR_{CNT})$ transforms into

$$\sum_{n=1}^{m-1} (r_i + h(n-1)) J_0(Q(r_i + h(n-1))). \quad (\text{A.24})$$

In our case the m distribution is not narrow, but those of r_i and h must be quite. Assuming r_i fixed, the average over the ensemble of MWCNTs modulates the terms in the sum over layers so that they are scaled down as n increases, since the statistics of larger n decreases because the instances of large m also decreases. Furthermore, for n small there are almost as many cases as for $n = 1$, that is, the histogram of n must start horizontally. A half-Gaussian decay of the n terms matches such specification and emerges as a natural choice. Altogether it yields

$$I_{DC}(Q) = 8\pi^2 C_{1D} a_d \rho_A j_0(Qd_{D_2}) e^{-\frac{1}{3}Q^2 u_{eff}^2} \frac{1}{N_c Q} \sum_i J_0(QR_i^\perp) \sum_{n=1}^{\infty} \exp\left[-\frac{h^2(n-1)^2}{2\sigma_n^2}\right] (r_i + h(n-1)) J_0(Q(r_i + h(n-1))), \quad (\text{A.25})$$

where we have defined an effective rms displacement, u_{eff} , and σ_n is the dispersion of the half-Gaussian decay. In practical terms a value of 50 layers worked well for σ_n , for which a termination of the n sum at 250 were enough to reach convergence in a short cpu time. In general, there are far more sources of disorder than just the thermal fluctuations around the COMs. For instance, in the real MWCNTs h presents also
335 dispersion with some systematic evolution with the number of layers [51]. All of these effects are here boldly represented through u_{eff} , chosen as the minimum needed to maintain the line shape of the main ND peak (1.2 Å for the 62 hPa load and 1.5 Å for the 175 hPa case). The parameter h controls the location in Q of the dispersive profile. A value of 3.41 Å was consistent with both the ND measurements and the values reported in the literature [51]. Finally, the areal density of any nanotube is given by $\rho_A = 4/(3\sqrt{3}d_{cc}^2)$,
340 where d_{cc} is the distance of the C–C bond in graphene taken here as 1.42 Å.

Appendix B. Materials and Methods

Appendix B.1. Experimental design

As for the confining system, we rely on samples of multi-walled carbon nanotubes (MWCNTs). For our
345 interests, they present several advantages compared to single-walled nanotubes. To begin with, they are
far stiffer. Given the weakness of the dispersive interactions among the molecules, tube bending should be
avoided as much as possible to facilitate unperturbed crystallization. Besides, since the external diameters
show a large dispersion in a typical MWCNT sample, they do not form so tightly and well ordered ropes, as
their single-walled counterparts tend to do [52]. On the other hand, the interstitial sizes are typically much
350 larger than those of the smaller inner diameters because the distribution of external diameters is centered
at much higher values than those of the inner diameters (much larger also than the typical mean diameter of
good quality SWCNTs samples). In such a case, it can be expected that the adsorption inside the smaller of
the multi-walled tubes will not compete appreciably with 1D-like adsorption in the interstitials or external
grooves of the ropes.

355 To select the smaller inner diameters as our confining systems we rely in the physics of adsorption. The
smaller open inner diameter tubes produce the stronger attractive potential wells for the molecules so that
they are filled first. For sufficiently small loads these micropores dominate the distribution of adsorbed
molecules.

As for the probe to study the system, neutrons are uncharged, so that they can penetrate deeply into
360 the target. Thus, they can explore the structure and dynamics of atomic and molecular arrangements
buried into nanostructured materials, while at the same time making possible the use of complex sample
environments such as cryogenic and gas handling equipment. Contrary to X-rays, they strongly interact with
hydrogen. Since we are interested in structural information, we resort to the deuterium isotope in order to
avoid the very large incoherent scattering of molecular hydrogen that does not contribute to the diffraction
365 signal. Incoherent scattering stems from the random distribution of nuclear spin states along the sample,
and implies the interchange of the spin state between the incident neutron and the nuclei. It can also be
avoided in H₂ if prepared in its fundamental para-state, that only scatters coherently. However, the neutron
energy must be well below 14.7 meV in order not to excite the molecular rotational levels responsible for
the very strong incoherent scattering. This would limit the range of the scattering vector norm Q below 4
370 \AA^{-1} , too short to properly explore the spatial scales involved.

Regarding the measurement protocol, we started by measuring the diffraction of the nanotubes sample
(2 g) during a slow cooling/heating cycle (0.1 K decrement/increment each 70 s) between ≈ 24.5 K and
2 K while neutron diffraction signal was acquired in runs of 350 s. Then we proceeded by injecting very
small quantities of D₂ into the sample can at 20 K until we observed a barely discernible liquid state
375 diffraction signal. The equilibrium pressure at that point was 62 hPa. After heating at ≈ 24.5 K, the same

cooling/heating cycle as before was done. The same thermal protocol was repeated increasing the D_2 load in each cycle up to a total of four loads, namely, 62 hPa, 175 hPa, 225 hPa, 284 hPa. Before the 175 hPa thermal cycle, in order to improve the statistics of the measurements at 62 hPa, a second cooling ramp was performed down to 2 K where neutron diffraction was measured in runs of 600 s up to a total of 31 runs. 380 The cooling/heating rate during the thermal cycles was chosen as slow as it was feasible in order to remain as near as possible to thermodynamic equilibrium, corresponding in this case to those of saturated vapor pressure (SVP). From the adsorption isotherm in Fig. 2 it is clear that the four loads are far from the bulk condition at 20 K.

Appendix B.2. Multi-walled carbon nanotubes synthesis and treatment

385 The raw MWCNTs were prepared by the evaporation of pure graphite anodes in an electric arc discharge under a helium atmosphere of 66 kPa applying a current of 60 A and a voltage of 25 V [53]. The raw material was oxidized by heating to 973 K in open air during 30 minutes. After an annealing at 1223 K in Ar atmosphere during 10 hours, around 2 g of the open MWCNT sample was obtained.

Appendix B.3. Deuterium

390 The deuterium gas was taken directly from a 95 liter bottle of Air Liquide N30 Deuterium (purity $\geq 99.9\%$; ~ 10 ppm of D_2O and O_2 and ~ 25 ppm of N_2).

Appendix B.4. Electron microscopy

The structure of the obtained MWCNT material was explored by TEM at the National Center for Electron Microscopy (CNME) in Madrid, Spain. Drops taken from a dispersion of the samples in ethanol 395 were deposited over copper grids with holey carbon support films. Transmission electron micrographs were taken with a JEOL JEM2100 HT microscope operating at 200 kV.

Appendix B.5. Adsorption isotherm

The D_2 adsorption of the MWCNT sample was characterized by a measurement of the adsorbed moles as a function of the pressure at constant temperature (adsorption isotherm) using a standard volumetric 400 procedure [29]. The chosen temperature was 20 K (D_2 triple point at 18.73 K). In essence, given a known volume, the adsorbed moles are obtained from the decrease in pressure they cause with respect to the same volume with no adsorbant. A home-made gas handling manifold was built to accurately manage the gas dosage. A layout of the system is shown in Figure S1. All pipes and connections in the manifold are made of steel and Swagelok valves were used to connect the different volumes in the system. The pressure was 405 determined with a Baratron[®] MKS 690A absolute manometer with a full range scale of 1000 Torr and an accuracy of 0.12% of the reading. The reference volume V_R is provided by a calibrated steel vessel of

volume $1045.7 \pm 2.3 \text{ cm}^3$. The MWCNT sample was located in an aluminium cylindrical can of 15 mm of inner diameter and 56 mm of height. The can was connected vertically to a specially built insertion stick (used to introduce the sample can in the cryostat) with an inner capillary for the D_2 injection. The stick was designed to match the standard ILL “Orange” cryostat [54, 55]. This is the same kind of vertical cryostat used in the neutron scattering experiment. The inner capillary of the stick was connected through a valve to the end of the manifold labeled as “sample” in Figure S1. Using the known value of V_R all the needed volumes are determined by expansion of He loads. In particular, the volume comprising from “sample” valve (closed) down to the sample can (not shown in the Figure) was $20.64 \pm 0.15 \text{ cm}^3$. Once the system volumes were determined, 0.521 g of the MWCNT sample were loaded in the sample can, the sample stick was introduced in the cryostat and the temperature set to 20 K. From repeated D_2 loads in V_{inj} and subsequent expansions, the adsorbed moles were determined [29]. The sample temperature during the adsorption/desorption isotherm was $T = \langle T_S^i \rangle = 20.23 \pm 0.03 \text{ K}$.

Appendix B.6. Neutron scattering experimental setup

The neutron scattering instrument of choice was D20 at the ILL in Grenoble, a reactor-based neutron facility (continuous neutron beam). The most characteristic feature of D20 is its very high neutron flux, which allows for real-time studies of many processes. The setup was chosen to maximize the neutron flux corresponding to a neutron wavelength of 1.30 Å. D20 is a powder diffractometer with a circular detection zone 1.47 m of radius, 4 m long and 15 cm high with the sample holder at the center of the circle (see [56] for details about D20). For the setup chosen the resolution remains below 0.02 \AA^{-1} until $Q = 5.5 \text{ \AA}^{-1}$, reaching 0.1 \AA^{-1} at $Q = 9 \text{ \AA}^{-1}$. Such resolution values are high enough to avoid resolution integrals in the data treatment. The gas handling manifold, described in Fig. S2, was located outside the neutron beam safety zone connected to the cryostat insertion stick through a 5-m-long 1-mm-diameter steel capillary. The injection volume, V_{inj} , used during the 20 K loads remained open during the neutron scattering measurements as a safety expansion volume. The sample can was a standard sample holder for the D20 instrument, namely, a 0.1 mm thick vanadium cylinder of 60 mm high and 8 mm of inner diameter. The neutron beam size was 8 mm width and 4 cm high. It was centered at the axis of the vanadium cylinder and the lower vertically edge of the beam located at the bottom part of the vanadium can. All the neutron irradiated volume was filled with MWCNTs sample (2 g). The pressure was monitored with a MKS 640A manometer (1000 Torr range) and a MKS 627B manometer (15000 Torr range). Since V_{inj} was open, the moles of D_2 irradiated by the neutron beam increased by a factor of around 1.4 from the equilibrium after load at 20 K to zero pressure solid at 2 K.

Appendix B.7. Neutron scattering data reduction

The raw data were corrected from detector efficiency and transformed to scattering vector reciprocal units using the LAMP software package freely available at the ILL facility [57, 58]. Self-shielding and neutron

absorption correction were estimated using the DAVE software package [59, 60]. As expected, given the small D₂ quantities and the cylindrical geometry of the sample can, self-shielding and neutron absorption amounts to a small and essentially Q -independent correction. Likewise, multiple scattering corrections are again Q -independent [61]. Both can be ignored for our purposes, which do not depend on absolute scattering values. An important point to take into the account is the variation in the number of scatterers irradiated by the beam. During a thermal cycle, due to the varying thermodynamic conditions (thermal expansion/contraction, D₂ condensation and diffusion, etc), some grains of the MWCNT powder move in and out the irradiated volume. In absolute terms the effect is small but not negligible with respect to the smaller D₂ loads. To compensate the effect the $I(Q)$ s of each run are divided by their corresponding total neutron count, which is proportional to the number of scatterers. In order to improve the statistics, all the runs corresponding to the pristine MWCNTs were added together, while for the low temperature $I(Q)$ s of the loaded MWCNTs, only those below 11.5 K (well below solidification temperature) were considered.

With respect to the subtraction of the MWCNT signal, since self-shielding and neutron absorption can be considered Q -independent, $I_C(Q)$ is just given by an attenuation transmission factor applied to the $I(Q)$ corresponding to the empty MWCNTs. The porous nature of the D₂ distribution within the MWCNT sample does not allow for a theoretical calculation of the effective transmissions. Instead, we have just heuristically adjusted the corresponding values, namely, 0.994 at 62 hPa, 0.9675 at 175 hPa, 0.940 at 225 hPa and 0.930 at 284 hPa.

Appendix B.8. Numerical methods. Molecular Dynamics

Molecular Dynamics (MD) simulations used in our nonlinear fitting analysis have been carried out using Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) code [62]. Taking into account the quantum delocalization of the nuclear rotational degrees of freedom we have considered for the D₂–D₂ interaction an isotropic effective potential [12] adapted to the LAMMPS code. For the C–C interaction we have relied in the Adaptive Intermolecular Reactive Empirical Bond Order (AIREBO) potential described in [63] as it is included in the LAMMPS package. In the case of the D₂–C interaction, we have relied in the anisotropic pair potential developed in [25] out of accurate *ab initio* calculations using Density Functional Theory (DFT) based on Symmetry Adapted Perturbation Theory (SAPT(DFT)), again adapted to LAMMPS. For details, including the parameterization used, see Supplementary Material in [25]. Thermalization has been achieved through the Berendsen thermostat [64], which presents a large thermal stability suitable for small samples. The chosen time step is one fs. The specifics of the procedure in the present calculations are given in the Supplementary Information.

Appendix B.9. Numerical methods. DFT

For the DFT calculations we have relied on the Perdew, Burke, and Ernzerhof (PBE) approach [65] as implemented in the CASTEP code [66]. The pseudopotentials are those corresponding to the 00PBE_OP.recpot

475 family as given by the Materials Studio package. The dispersive interactions are treated using the semi-
empirical dispersion correction scheme (DFT-SEDC) [67] with Tkatchenko and Scheffler (TS) correction [68]
(hydrogen atoms corrected to deuterium mass). The energy cutoff of the plane wave basis set was 900 eV,
only one k point was used and the maximum force tolerance per atom was set to 0.006 eV/Å. The CNT
containing the D₂ molecules is a (10,5) SWCNT with a length of 22.6211727 Å and 280 carbon atoms. The
480 specifics of the procedure are given in the Supplementary Material and in Table SIII therein.

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Formation of one-dimensional quantum crystals of molecular deuterium inside carbon nanotubes: Supplementary Material

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I. NONLINEAR FITTING ANALYSIS

We have performed MD simulations of the D_2 molecules inside two types of MWCNTs. The inner diameters are chosen so that the cross section of the D_2 1D system accommodates at least two molecules, given that a strict 1D chain is not compatible with the measurements. Specifically, the smallest MWCNT is $(1,13)@(4,20)@(31,0)@(5,37)@(8,44)$ where $(a,b)@(c,d)$ means tube (a,b) within (c,d) . We denote it as MWI(1,13), from “Multi-Walled Inner” with a $(1,13)$ inner nanotube. The largest is a $(18,0)@(10,20)@(35,0)@(44,0)@(5,50)$ MWCNT, that is, a MWI(18,0). In the latter case, the inner nanotube allows for the formation of a D_2 shell adsorbed at the nanotube wall and a 1D chain at the center. The D_2 arrangements obtained by MD inside the MWCNTs are in agreement with previously published results on H_2 in single-walled carbon nanotubes [1–3], and they are shown in Fig. S4. For the larger diameter, MWI(18,0), two relaxed D_2 structures are depicted, one consisting of just an adsorbed shell and the other consisting of a shell plus a 1D central chain. As mentioned in the main text, owing to the strong influence of the adsorption potential, the shells can be seen as a rolled portion of a 2D D_2 triangular lattice with a nearest-neighbor distance consistent with the ND patterns ($d = 3.605 \text{ \AA}$). Remarkably, even the compact structure inside the MWI(1,13) is well reproduced with such approach. Following this insight, we have explored an ample series of diameters and lengths. As in the MD simulations shown in the Fig. S4, none of the obtained $I_D(Q)$ s are able to reproduce adequately the experimental ND

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signals.

Confronted with these results, we turned to the opposite perspective, i.e., to study structures preserving the bulk crystalline molecular arrangements. We have chosen cylindrical cuts of the HCP and FCC crystals which maximize the number of molecules in the minimum of the adsorption potential. Fig. S5 and Fig. S6 clearly show many structures with $I_D(Q)$ s compatible with both the 62 hPa and 175 hPa ND measurements. Therefore, we performed a nonlinear fitting analysis (using the NonlinearModelFit procedure of the Mathematica[®] package) of the various structures assuming a log-normal distribution of lengths for each given type. In such a case, the model for the ND signal reads (see Equation (1) in main text):

$$I_D(Q) = C_{1D} \left(\sum_{l=1}^{l=40} \frac{e^{-\frac{(\ln(l/m)-\sigma^2)^2}{2\sigma^2}}}{\sqrt{2\pi\sigma^2}l} 4a_d^2 j_0(Qd_{D_2}/2)^2 (S_{1D}^M(Q, u_{1D}, l) - 1) + F_{inc}(Q, u_{1D}) \right) + C_{3D} \left(F_{inc}(Q, u_{3D}) - 4a_d^2 j_0(Qd_{D_2}/2)^2 \exp \left[-\frac{(Q u_{3D})^2}{3} \right] \right) + bg_0 + bg_1 Q^2 + bg_2 Q^4,$$

where the fitting parameters are the scaling factor of the scattering coming from the chosen type of 1D crystals, C_{1D} ; m and σ , which define a log-normal distribution of lengths l of such 1D crystals, being l the number of 1D crystal layers, m its mode and σ is the dispersion of $\ln(l)$; u_{1D} is the corresponding rms of the molecular center of mass (COM) displacement; C_{3D} is the scaling factor of the external small crystals (or crystal); the bg_i 's constants define the background up to second order in Q^2 . Since u_{3D} is the rms COM molecular displacement of external small crystals not confined within the MWCNTs, it was fixed to 0.46 Å, the most recent experimental value corresponding to the bulk case [4]. With respect to the ortho concentration a value of 2/3, i.e., the value corresponding to room temperature [5], was needed in order to obtain successful fits.

In Figures S5 and S6 the fits are shown as green solid lines superimposed to the data. Focusing in the 62 hPa load, Fig. S5, the best fit corresponds to the FCC structure labeled as B. However, this fit, as that of the HCP structure C, is not physical since it requires negative $S(Q)$ values (those below the dashed line in the left). Imposing restrictions to force positivity leads to low quality fits with unnatural length distributions. Remarkably, the structure A, in spite of being structurally near, yields a fit in excellent agreement with the measurements. It corresponds to the TCP 1D crystal discussed in the main text. We notice that despite the great quality of the fit, the reduced χ_ν^2 value is around 4 instead of ≈ 1 ,

as expected for good fits [6]. This implies an underestimation of the data dispersion that, in fact, can be visually gauged in the data figures which show fluctuations incompatible with the lengths of the error bars. The error bars account only for the counting statistics and therefore some other fluctuations must be present. We have traced the extra noise as a consequence of fluctuations in the density of buffer gas (He; needed for a proper thermal conductivity) around the sample cell within the cryostat. The effect, most notorious at the lowest working point of the cryostat, is approximately reproduced by doubling the data error bars. Doubling the error bars would divide the χ_ν^2 value by 4, but otherwise it would not alter neither the quality of the fits nor the utility of χ_ν^2 values to compare the goodness between fits.

Considering the 175 hPa load, which corresponds to larger 1D crystal diameters (see Fig. S6), the best fit is for the structure C. This is just the THCP 1D crystal, as denoted in the main text, which once more comes from the HCP bulk crystal. Notice that for this load the χ_ν^2 values are larger since the two Bragg peak precursors coming from the external nanocrystals are not included in the model. At any rate, the quality of the fit is again excellent not only for the C structure, but also for the A and E, all derived from the HCP bulk crystal (see Fig. S6). However, when including the interference term, $I_{DC}(Q)$, the cases A and C are compatible with the diffraction patterns whereas the E case is not so. Otherwise, although the C structure (a denser version of A) is the most probable, from the fits the presence of some proportion of the A structure cannot be disregarded.

In Table SI and Table SII we present the fitting parameters along with their standard deviation for the TCP and THCP 1D crystals as yielded by the NonlinearModelFit procedure in Mathematica[®].

II. NUMERICAL CALCULATIONS PROCEDURE DETAILS: MOLECULAR DYNAMICS

For the MD results finally shown in this work, the initial configurations of the D₂ 1D crystals inside the MWCNTs were those obtained from the nonlinear fitting analysis of the ND results. In a first step, the D₂ 1D crystals were kept fixed inside the corresponding MWCNT. In order to relax the MWCNT structure, it was submitted to an annealing pro-

cedure by cooling down from 100 K to 0.01 K in a stepwise manner, leaving the structure to thermalize at each temperature. Once the MWCNT are relaxed to the energy minimum, the D_2 molecules are released, now fixing all the CNTs except the most inner one during 0.5 ns. This procedure was applied to the MWI(1,13) and MWI(18,0) nanotubes chosen as explained in the previous section.

In the former, simulation runs with 1D crystal seeds corresponding to those shown in Fig. S5A, B and C composed of 87 molecules were performed using a cell length of 171.667 Å. In the latter, crystal seeds corresponding to Fig. S6C and D with 279 D_2 molecules and a cell length of 304.560 Å were addressed. These lengths yield supercells ensuring that the D_2 configurations do not interact with themselves. In all the simulation runs, the initial 1D crystal arrangements destabilize in around 50 ps in favor of structures adsorbed onto the CNT wall. In the case of the initial seeds corresponding to the TCP and HCP 1D crystals we performed a further annealing, thermalizing first at 10 K and cooling down to 0.1 K in 0.99 ns, keeping the system at 0.1 K during 2.33 ns. As a result of this procedure, the TCP seed gives the structure shown in Fig. S4A, while the HCP breaks in two independent structures displayed in Fig. S4 B and C.

III. NUMERICAL CALCULATIONS PROCEDURE DETAILS: DFT

For the DFT calculations, three initial 1D crystals composed of 11 molecules were used to find the minimum energy configuration, namely, those shown in Fig. S5A (TCP) and B (FCC bulk parent structure) as well as the stable configuration obtained through MD (see Fig. S4A). In regarding the optimization of the electronic density, we start with a rough criterium for the energy minimization, namely, a convergence tolerance for the total energy per atom of 0.25×10^{-8} eV. The minimization criteria sought for the geometry minimization, made using the BFGS algorithm [7], are a change in energy per atom $|\Delta E| < 0.02$ meV, a displacement per atom $|\Delta R| < 0.001$ Å and a force per atom $|F_{max}| < 6$ meV/Å for all the atoms involved. Notice that the maximum force per atom is set here far more stringent than the CASTEP default. We do so because the minimum of the potential with respect to the molecular orientation is very shallow. The minimization starts by leaving the COMs of the molecules and carbon atoms fixed with an initial orientation of the molecules along the

SWCNT axis. The rotational degrees of freedom of the D_2 molecules were first minimized. Then, the carbon degrees of freedom are released and, finally, all the system is minimized without any restriction. For the three initial configurations CASTEP reaches a minimum for the given energy tolerance corresponding to final structures near the initial ones, but fails for both, the $|\Delta R|$ and the $|F_{max}|$ criteria. From these configurations we restart optimization runs without any restriction requiring a total energy convergence tolerance of 0.9×10^{-9} eV. The initial configuration corresponding to the MD yielded the global minimum of total energy, E , fulfilled the criterium for $|\Delta R|$ while $|F_{max}|$ was marginally above the threshold (see Table SIII). In order to check the quality of such a minimum we performed a calculation of the density of vibrational modes, obtaining 8 low energy unstable modes out of the total of 66, a reasonable result that serves also to gauge the difficulties associated to the shallowness of the orientational interaction potential. The final configuration is shown in Figure 5A of the main text. The initial structure corresponding to the FCC parent configuration failed again both criteria, in particular yielding a high residual force per atom. Finally, the TCP initial configuration gave as final (meta)stable geometry that shown in Figure 5B, this time fulfilling all three criteria (see Table SIII). In this case the density of vibrational modes showed 7 low energy unstable modes.

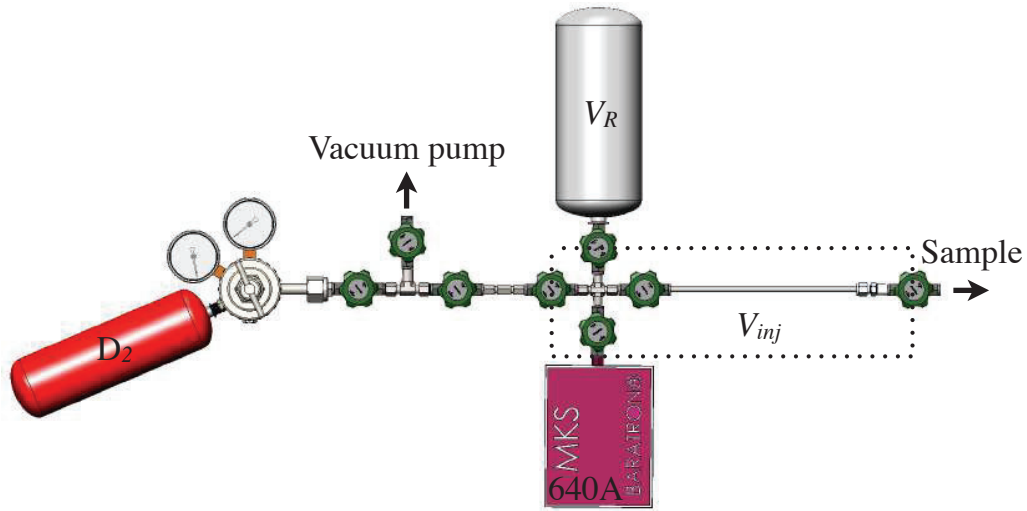


Figure S1: Layout (top view) of the home-made gas handling manifold employed to measure the adsorption isotherm. The dotted line defines the volume V_{inj} : valves enclosed by the line are open, valves crossed by the line are closed. The volume V_R is defined with its own valve closed (not shown). Their corresponding values are $V_R = 1045.7 \pm 2.3 \text{ cm}^3$, and $V_{inj} = 55.21 \pm 0.11 \text{ cm}^3$.

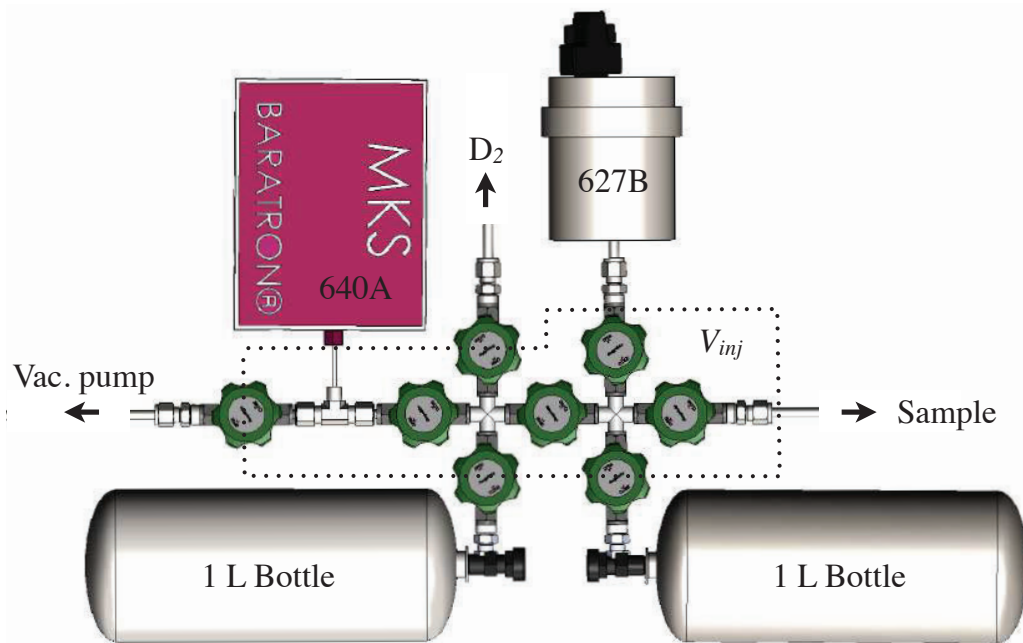


Figure S2: Layout (top view) of the home-made gas handling manifold employed in the neutron scattering experiment. The dotted line defines the volume V_{inj} : valves enclosed by the line are open, valves crossed by the line are closed ($V_{inj} = 30 \text{ cm}^3$).

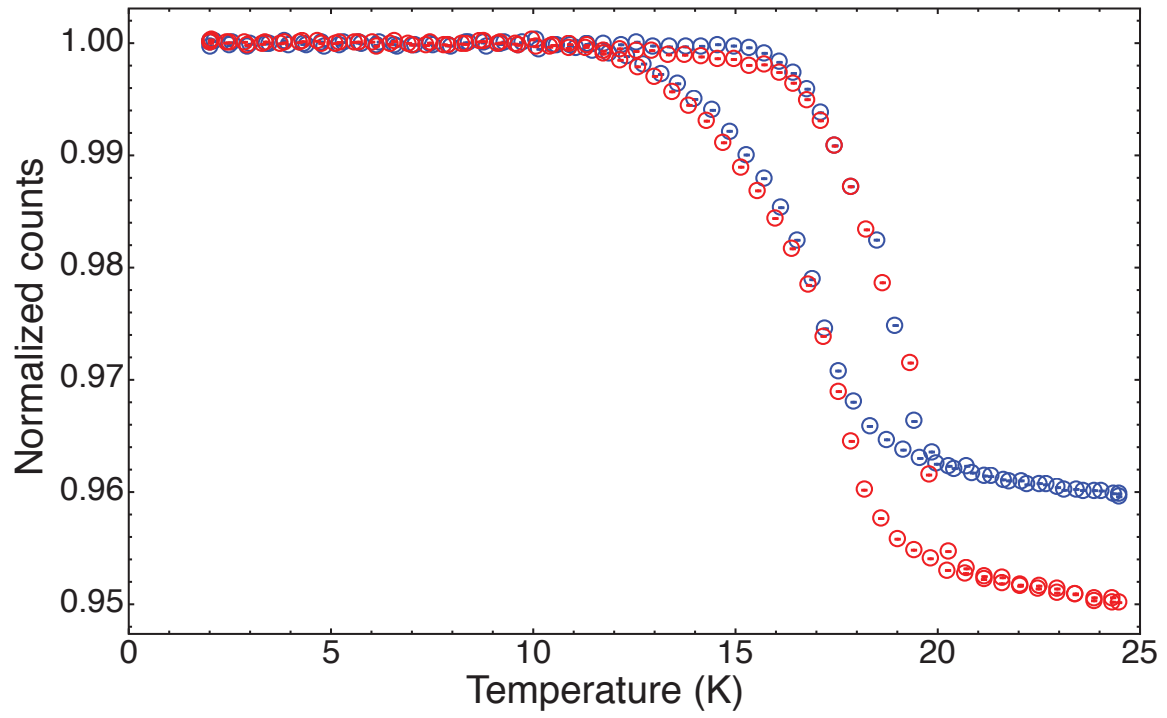


Figure S3: The evolution during a thermal cycle of the normalized area (one at 2 K) under $I(Q)$ within the red rectangle shown in Figure 2 (main article) for the 225 hPa load (blue circles) and for the 284 hPa case (red circles).

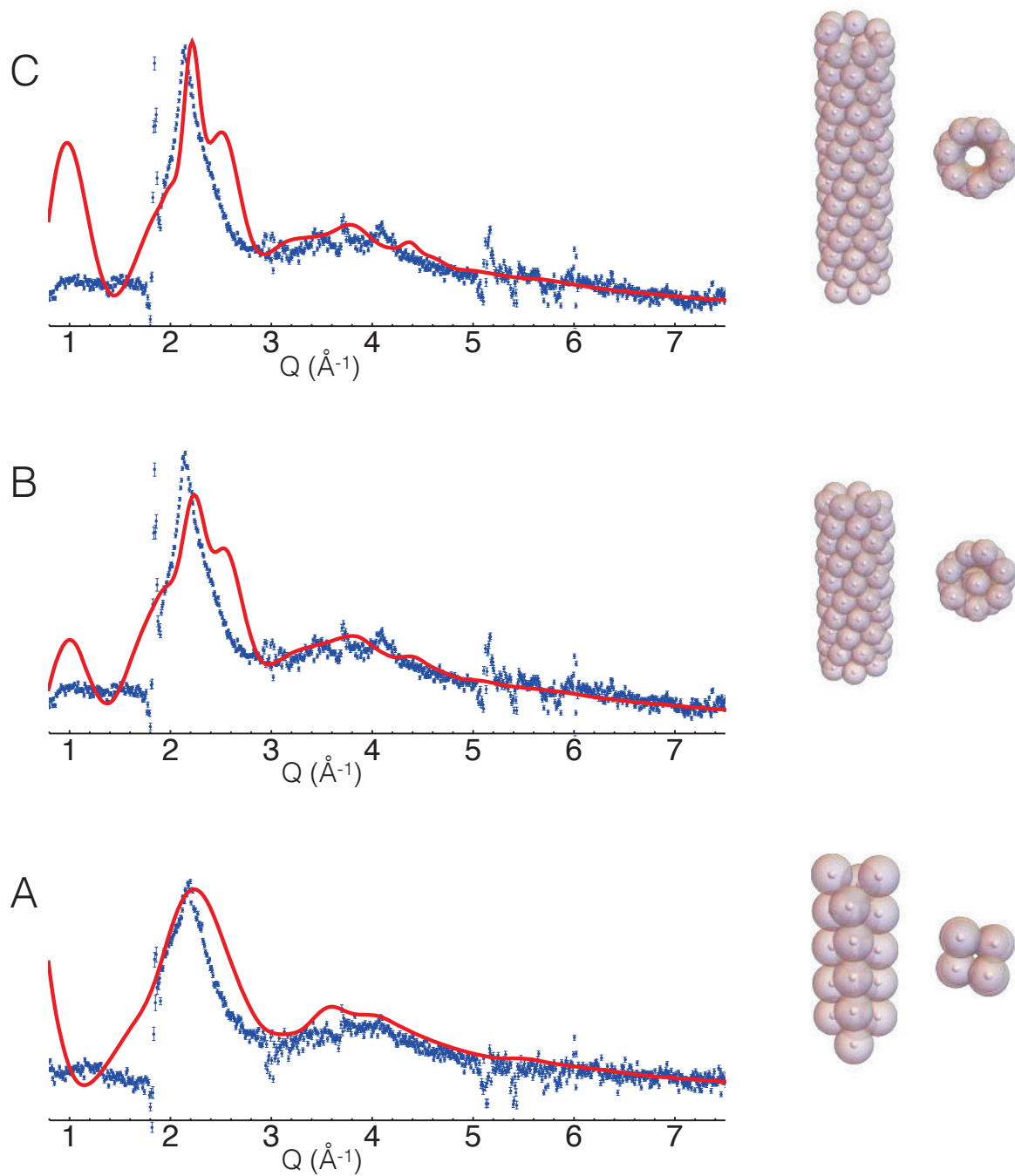


Figure S4: 1D D_2 structures obtained from molecular dynamics simulations: (A) Within the MWI(1,13) nanotube; (B) and (C) within the MWI(18,0) nanotube. For each structure, $I_D(Q)$ s are shown superimposed to the ND pattern (blue data) at (A) 62 hPa and (B) and (C) at 175 hPa.

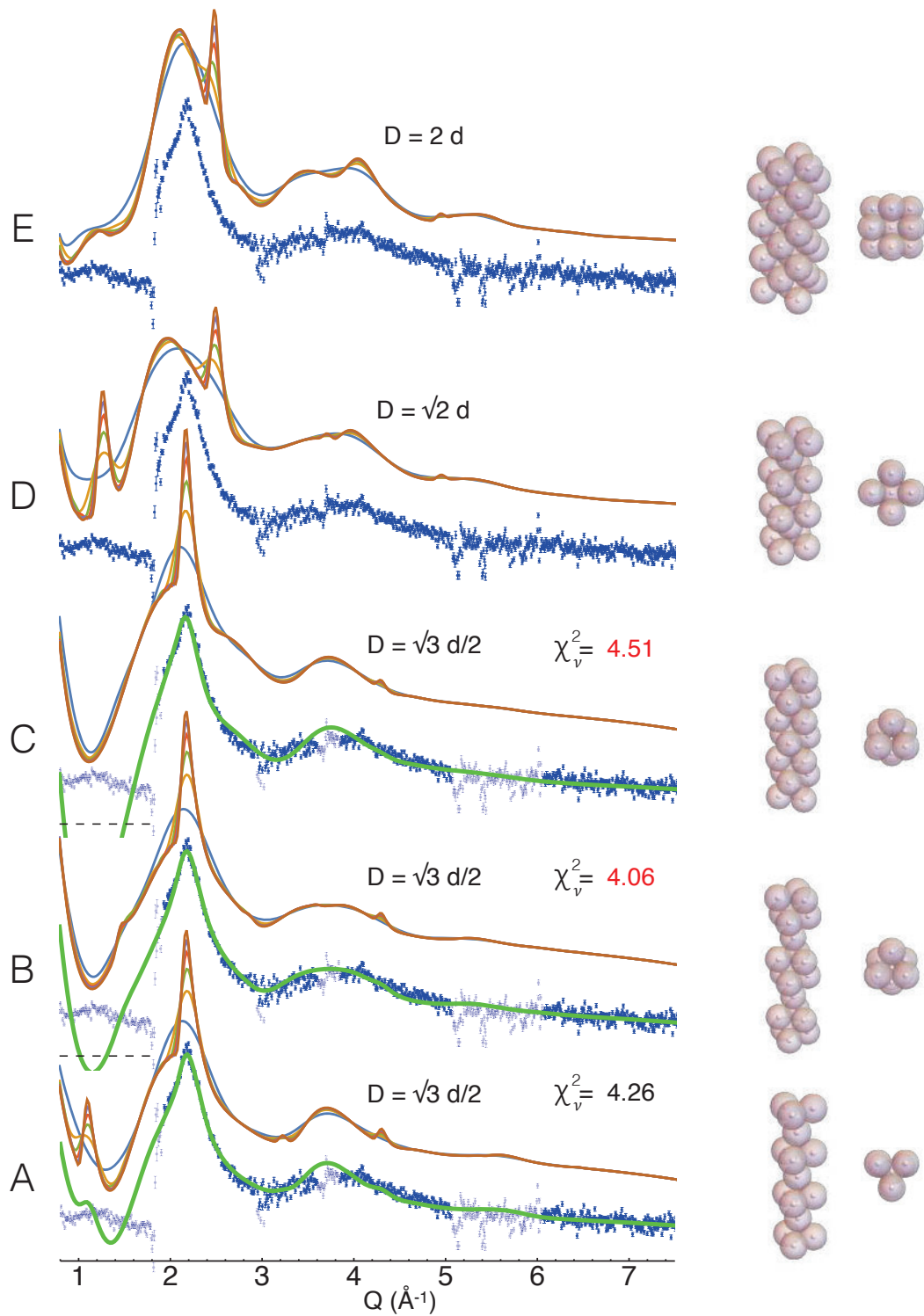


Figure S5: 1D D_2 structures compatible with the HCP (A, and C) and FCC (B, D, and E) bulk crystals ordered in increasing diameters, D , given in terms of the nearest-neighbor distance, d , along with their $I_D(Q)$ s. For each structure, $I_D(Q)$ s corresponding to 4, 8, 12, 16, and 24 layers are shown superimposed in different colors. The ND data at 62 hPa are also shown (blue data). When it makes sense a fit corresponding to a log-normal distribution of lengths is also shown as a solid green curve on top of the ND data. In the fits only the dark blue data are actually used. The cases B and C (χ_v^2 in red) are unphysical since they reach negative values (zero corresponds to the dashed lines in the left).

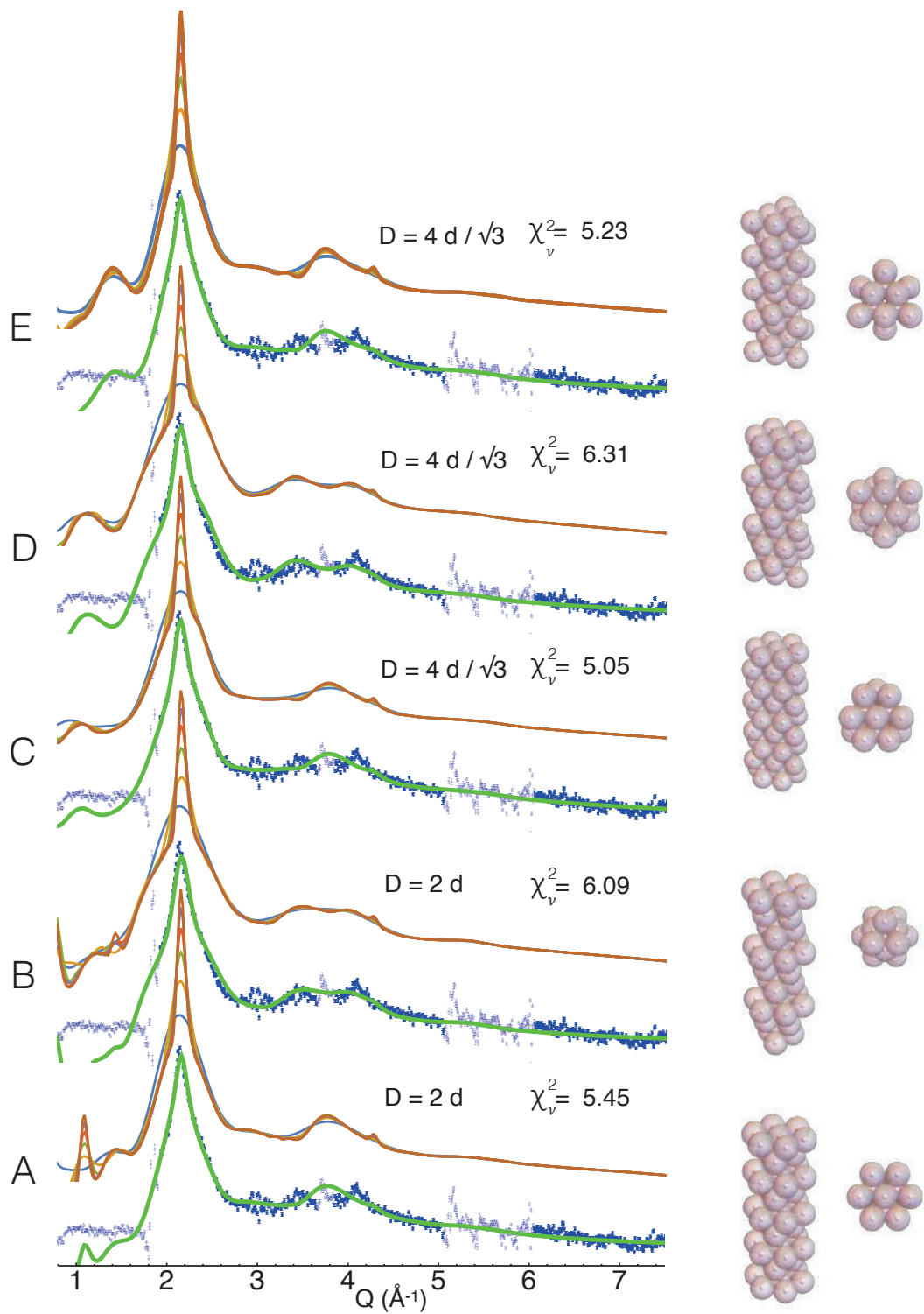


Figure S6: Continuation of 1D D_2 structures compatible with the HCP (A, C, and E) and FCC (B, D) bulk crystals ordered in increasing diameters, D , given in terms of the nearest-neighbor distance, d , along with their $I_D(Q)$ s. For each structure, $I_D(Q)$ s corresponding to 4, 8, 12, 16, and 24 layers are shown superimposed in different colors. The ND data at 175 hPa are also shown (blue data). Fits corresponding to a log-normal distribution of lengths is also shown as a solid green curve on top of the ND data. In the fits only the dark blue data are actually used.

Table SI: Fitting parameters at 62 hPa load (TCP 1D crystal).

C_{1D}	m	σ	u_{1D}
0.036 ± 0.001	5.4 ± 0.4	0.49 ± 0.05	0.33 ± 0.02
C_{3D}	bg_0	bg_1	bg_2
0.010 ± 0.003	-3.21 ± 0.45	0.09 ± 0.009	-0.00071 ± 0.00006

Table SII: Fitting parameters at 175 hPa load (THCP 1D crystal).

C_{1D}	m	σ	u_{1D}
0.0520 ± 0.0007	8.2 ± 0.2	0.39 ± 0.02	0.45 ± 0.02
C_{3D}	bg_0	bg_1	bg_2
0.026 ± 0.002	2.9 ± 0.3	0.090 ± 0.006	-0.00057 ± 0.00004

Table SIII: DFT results

	TCP	FCC parent	MD
E	-43703.99311226 eV	-43704.09768858 eV	-43704.21699157 eV
$ \Delta E $	2.655354×10^{-7} eV	4.461872×10^{-7} eV	2.010411×10^{-8} eV
$ F_{max} $	5.50953 meV/Å	18.14 meV/Å	8.6932 meV/Å
$ \Delta R $	1.321504×10^{-4} Å	1.866585×10^{-2} Å	2.88899×10^{-4} Å

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