Carbon Nitride Nanothread Crystals Derived from Pyridine

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**Supporting Information**

**ABSTRACT:** Carbon nanothreads are a new one-dimensional sp³ carbon nanomaterial. They assemble into hexagonal crystals in a room temperature, nontopocemical solid-state reaction induced by slow compression of benzene to 23 GPa. Here we show that pyridine also reacts under compression to form a well-ordered sp³ product: C₅NH₅ carbon nitride nanothreads. Solid pyridine has a different crystal structure from solid benzene, so the nontopocemical formation of low-dimensional crystalline solids by slow compression of small aromatics may be a general phenomenon that enables chemical design of properties. The nitrogen in the carbon nitride nanothreads may improve processability, alters photoluminescence, and is predicted to reduce the bandgap.

Relative to the vast corpus of organic molecules, we have few examples of extended carbon networks: diamond, graphite, graphene, and carbon nanotubes. Nevertheless, they have attracted much interest due to their superlative thermal, electronic, and mechanical properties and symmetric geometries. More than 500 new carbon networks have been predicted, but this diversity remains largely unrealized. Recently, we showed that polycrystalline benzene transforms under pressure into sp³-bonded and one-dimensional single-crystalline packings of carbon nanothreads. Fully saturated degree-6 nanothreads could exhibit a unique combination of strength, flexibility, and resilience, whereas partially saturated degree-4 threads may act as novel organic conductors. Moreover, the transformation to nanothread carbon cannot be topochemical because it involves large changes in volume and symmetry as multiple short, strong carbon–carbon bonds form between each benzene pair. Such nontopocemical reactions usually disrupt single-crystalline order and often yield amorphous solid products, but the benzene nanothread reaction instead creates single-crystalline order. The requirement for commensuration between reactant and product structures for topochemical reactions severely constrains the number of suitable monomers; freedom from this requirement could allow for chemical design through solid-state organic synthesis of crystalline extended networks from new types of monomers.

Here we show that slow compression and decompression of polycrystalline pyridine allows for the formation of single-crystal packings of carbon nitride nanothreads (Figure 1a,b) with empirical formula C₅NH₅. Chemical substitution of heteroatoms such as nitrogen into extended carbon networks is generally difficult because nitrogen substituting carbon is not isoelectronic; carbon nitride nanothreads, in contrast, are isoelectronic to carbon nanothreads. Carbon nitrides are often thought of as compounds that have more N than C (e.g., C₅N₄) to distinguish them from nitrogen-doped carbons. Nonetheless, any solid state compound such as the sp³ nanothreads described here in which nitrogen is intrinsic to the unit cell can be considered a carbon nitride.

The isomeric possibilities and nomenclature for substituted nanothreads with chemical formula ((CH)N)ₓ (X = heteroatom or -CR substituent) have been explored systematically. For the isomers of carbon nitride nanothreads with chemical formula ((CH)N)ₓ we adopt the naming convention of the enumerated carbon nanothreads by an integer string that indicates the positions of nitrogen atoms in every six-membered progenitor (i.e., molecular) ring. For example, isomer tube (3,0)₁₂₃₄₅₆ can be generated from the tube (3,0) carbon nanothread where the C–H pairs on positions 1, 2, 3, 4, 5, and 6 of six successive rings in the stack are substituted with N atoms, as shown in Figure 1a,b (see Figure S1 for numbering). Degree-4 unsaturated nanothreads have one double bond per pyridine formula (Figure 1c) whereas degree-2 threads have two.

Liquid pyridine was loaded into a double-stage membrane diamond anvil cell. As we slowly compressed the pyridine
over 8–10 h, it first froze to solid pyridine at 1–2 GPa\(^{15,16}\) and then reacted at \(\sim 18\) GPa (Figure S2). A crystalline reaction product has previously been reported, but does not exhibit the diffraction pattern of nanothreads.\(^{17}\) After holding the sample at 23 GPa for 1 h, the pressure was released to ambient over 8–10 h. The recovered sample is a yellow/orange translucent solid. The diamond cell induces an anisotropic (uniaxial) stress parallel to the force applied by the diamonds.\(^{18}\)

Synchrotron X-ray diffraction patterns collected on the recovered solid (Figure 2a, left) with the beam parallel to the applied uniaxial stress axis are characteristic of a hexagonal single crystal with its \(c\)-axis parallel to the X-ray beam, suggesting a role for uniaxial stress in guiding the growth of the crystal.\(^{13}\) There is good agreement with the predicted 6-fold single-crystal diffraction pattern of a representative tube (3,0)\(_{123456}\) carbon nitride nanothread with structural parameters determined by modeling (Figure 2a, right). Additional much weaker \(\{hk0\}\) reflections in the [001] zone are evident at smaller interplanar spacings (Figure S3). Again, as for carbon nanothreads, the diffraction patterns for carbon nitride nanothreads are pseudohexagonal,\(^6\) i.e., with small deviations from perfect hexagonal symmetry. Observation of a pseudohexagonal X-ray diffraction “signature” similar to that observed for carbon nanothreads\(^6\) provides compelling evidence for the synthesis of carbon nitride nanotubes of similar diameter (6.5 Å). Nevertheless, these single-crystal diffraction patterns do not yet constrain which of several possible pyridine nanothread atomic structures\(^{13}\) are formed. The ability of uniaxial stress to align nanothreads is remarkable (see also Figure S4). In contrast, other than ref 17, previous investigations of the reaction products from the compression of pyridine reported them to be amorphous,\(^{15,16}\) possibly because faster compression/decompression rates led to more highly branched reaction pathways for intermolecular bonding.

The interplanar spacings for the three “pseudohexagonal” Friedel pairs of arcs from experiment can be compared with

![Figure 1](image1.png)

Figure 1. Carbon nitride nanothread structures: (a) view down the hexagonal \(c\) axis and (b) view perpendicular to the \(c\)-axis of a degree-6 carbon nitride nanothread tube (3,0)\(_{123456}\). (c) Degree-4 carbon nitride nanothread structure (IV-7) with C–N double bonds in red.

![Figure 2](image2.png)

Figure 2. Diffraction experiment and modeling. (a) 6-Fold synchrotron diffraction pattern (left) collected down the hexagonal \(c\) axis of the nanothread crystal. Simulated diffraction pattern (right) of carbon nitride tube (3,0)\(_{123456}\). (b) Experimental and simulated {100} interplanar spacings for the 3 Friedel pairs of carbon nitride nanothread structures. The measured spacings are adjusted for thermal expansion.

![Figure 3](image3.png)

Figure 3. Infrared spectrum. The major radial breathing mode (top left) and the inter-ring C–N stretch mode (bottom right) in tube (3,0)\(_{123456}\) are shown. Peaks with asterisks may be associated with pyridinic derivatives either as amorphous carbon or as substituted pyridine linking nanothreads. The intensities of these peaks are significantly lower than in the IR spectra of amorphous recovered samples.\(^{15,16}\)

![Figure 4](image4.png)

Figure 4. NMR spectra. Solid-state (a) \(^{15}\)N and (b) \(^{13}\)C NMR spectra of nanothreads made from \(^{15}\)N-enriched pyridine. Peak areas are nearly quantitative. Selective spectra of NH and of nonprotonated C are shown by colored thin lines in panels a and b, respectively. The original signal positions of pyridine are marked by dashed vertical blue lines. The natural-abundance \(^{13}\)C spectrum shows the characteristic chemical-shift increases due to N-bonding of nearly half of all C.
those from simulations (Figure 2b, see also Figure S1 and Supporting Information (SI)) performed at 0 K for six optimized pseudohexagonally packed carbon nitride nanothreads, each of which is energetically most stable among their isomeric structures. After shrinking the experimental spacings by 1−3% to account for thermal contraction to 0 K, there is reasonable agreement with the predicted spacings for tube (3,0)123456. Polytwistane_13_3_25 and zipper polymer_24_13 are also consistent with experiment, but with a slightly denser packing along one dimension (see SI for ideal carbon nitride nanotread structures). Polymer I_3_3_25 and polymer I_4_2_36_13 seem less likely candidates, with significant differences of spacings when ordered packed, but azimuthal and/or axial packing disorder will narrow the differences among the interplanar spacings.

Amorphous carbon nitride (α-C:H:N) films deposited by chemical vapor deposition have broad infrared (IR) absorption spectra.19 In contrast, carbon nitride nanotreads exhibit more and narrower absorption features (Figure 3) suggestive of a 1D thread or tube structure along the thread axis (see SI). We calculated the IR absorption spectra of the six candidate structures mentioned above. Two notable characteristic vibrations of a 1D thread or tube are radial breathing and axial elongation-compression modes.1 The former is Raman (but not IR) active in carbon nanotreads and is observed experimentally at 805 cm−1,4 much higher in frequency than corresponding modes for (larger-diameter) sp2 carbon nanotubes.1 Symmetry breaking due to the nitrogen atoms in carbon nitride threads splits this mode and makes it IR active. In all of the simulated spectra, there are one or several absorption peaks from 575 to 650 cm−1 associated with radial breathing. In contrast to the corresponding higher-symmetry pure-carbon threads, mode eigenvectors for the radial breathing modes of carbon nitride threads include nonradial (axial, circumferential) motions, the magnitudes of which depend on the precise nitrogen placements on successive rings.

Considering the complexity of the vibrational modes in this region and the possibility of more than one type of nanotread forming during reaction, these calculated modes are consistent with the broad peaks observed in experiment at 580 and 645 cm−1.14 Axial elongation and compression in carbon nitride nanotreads include stretching of both C−C and C−N single bonds formed between pyridine rings upon reaction. A peak in the IR spectrum at 1117 cm−1 is identified as inter-ring C−N stretching because all the calculated spectra show this mode near 1120 cm−1 with moderate-to-strong intensity. Unlike the peak at 1178 cm−1 that is associated with intra-ring C−C and C−N stretching, neither the 1117 cm−1 inter-ring peak nor the radial breathing modes near 600 cm−1 appear in prior samples derived from pyridine at high pressure (Figure S5),15 i.e., they seem to be diagnostic of a thread-like structure, although the current IR spectra cannot yet uniquely determine which thread structure is formed.13 The SI provides detailed discussion of assignments, comparison with other candidate structures, modeling methods and animated vibrations.

We prepared larger (mg) samples in a Paris-Edinburgh press for chemical analysis (Materials and Methods, Figure S4). Only C, N, and O were found in X-ray photoelectron spectroscopy scans (Figure S6). The observed C:N ratio, 5.08:1, is very close to that of pyridine, which contrasts with chemical vapor deposition approaches for sp2 systems, which form low nitrogen content (x = 0.01−0.02) CNx nanotubes when using pyridine as a precursor.20 Bulk combustion analysis gave a reasonably close composition, 5.5:1, whereas nuclear magnetic resonance (NMR) analysis gave 4.1 ± 0.9:1 (see SI for details).

15N and 13C solid-state NMR spectra collected on 15N-enriched carbon nitride nanotreads (Figure 4) reveal nearly identical sp2/sp3 ratios for nitrogen and carbon (22−23% sp2, 77−78% sp3), indicating significant conversion from sp2 to sp3 bonding during polymerization and similar outcomes for C and N. About 45% of sp3 carbons have chemical shifts between 50 and 95 ppm, indicating that they are bonded to N. This exceeds the C−N bond fraction of 2/5 = 40% in pyridine and thus indicates C−N bond formation between pyridine rings, which is consistent with the observation of inter-ring C−N stretching in IR. While the 13N chemical shift of −310 ppm is similar for pyridine and imine N=CH, the pronounced CH resonances near 170 ppm in the 13C spectrum, distinct from pyridine signals at ≤152 ppm, document the presence of imine N=CH as found in degree-4 nanotreads.

Furthermore, the relative 13C NMR peak intensities indicate that in degree-4 threads, N=CH is strongly favored over N−CH=CH, which would produce a CH peak near 106 ppm that is not significantly observed in Figure 4b. The estimated fraction of degree-6 carbon nitride nanotreads is ca. 40%, after taking into account the sp3-hybridized carbons associated with degree-4 nanotreads. These two different types of threads still appear to pack into ordered crystals (Figure 2a). Figure 4a and Figures S8 and S9 also show some amides and N−H moieties, which may be due to reaction with O2 and/or H2O from air.

Synthesis of nanotreads from benzene16 and pyridine with distinct progenitor molecular crystal structures21 suggests that general principles govern nanotread formation. Under compression, an efficient molecular packing forms stacks of parallel aligned flat molecules. Such stacks provide a natural direction for a polymerization that yields 1D structures, i.e., nanotreads. The small size of the rings impedes π overlap between molecules in adjacent stacks, disfavoring cross-link polymerization to form 2D networks. The uniaxial component of the applied stress defines the thread axis, and the enthalpic driving force for efficient packing under pressure strongly favors a dense near-hexagonal packing of the threads. Low (i.e., room) reaction temperature favors the single lowest-barrier kinetic pathway toward thread formation and requires slow compression and decompression to afford the reaction sufficient time to proceed.
Nanotreads exhibit photoluminescence, likely due to defects or sp² carbon functions along their length (Figure 5a,b) or bandgap emission.22,23 The blue, green, and red emission properties of carbon nitride nanotreads differ from those of carbon nanotreads (Figure 5a,b, Figures S10 and S11). The shift to longer emission wavelengths for carbon nitride nanotreads is likely to be associated with the decrease in bandgap predicted for carbon nanotreads24 or nitrogen-related defects. Thus, with further synthesis work nanotreads might be chemically designed for specific biological imaging applications.25

ASSOCIATED CONTENT

Supporting Information
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Degree-6 carbon nitride nanotread structure: Polymer 1_3-3-2_5 (CIF)
Degree-6 carbon nitride nanotread structure: Polymer 1_4-2-36 (CIF)
Degree-6 carbon nitride nanotread structure: Polytwistane_153 (CIF)
Degree-6 carbon nitride nanotread structure: Tube (3,0)_{1245} (CIF)
Degree-6 carbon nitride nanotread structure: Tube (3,0)_{123456} (CIF)
Degree-6 carbon nitride nanotread structure: Zipper polymer_24 (CIF)
Degree-4 carbon nitride nanotread structure: IV-7 (CIF)
Vibrational mode animations (ZIP)
Materials and Methods, Additional Discussion (PDF)

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Notes
The authors declare no competing financial interest.

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