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Article

# A Thermodynamic Landscape of Hydrogen Cyanide-Derived Molecules and Polymers

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KEYWORDS: prebiotic chemistry, iminoacetonitrile, diaminomaleonitrile, tholins, adenine, Titan

# INTRODUCTION

dazole and adenine.

In this work, we use quantum chemistry to map the thermodynamic landscape of hydrogen cyanide (HCN)derived molecules and polymers. HCN is a common molecule in the universe, having been identified on various planets,<sup>1</sup> dwarf planets,<sup>2</sup> Saturn's moon Titan,<sup>3</sup> comets,<sup>4</sup> and in the interstellar medium.<sup>5</sup> HCN was likely present on the early Earth.<sup>6</sup> The molecule is also quite reactive. In experiments, biologically relevant molecules such as amino acids, nucleobases, and pterins<sup>7</sup> have all been extracted from the products of HCN polymerization experiments (see ref 8 for a review).

The prevalence of HCN in many astrochemical environments, and its reactive nature, places the molecule, and its possible reaction products, at the center stage of prebiotic chemistry and astrobiology.<sup>9,10</sup> However, a clear picture of HCN reactivity, both in the laboratory and in astrochemical environments, remains elusive.

HCN-derived polymers are complex materials that can form under a variety of laboratory conditions.<sup>11</sup> These product mixtures are typically heterogeneous and insoluble in most common solvents.<sup>12</sup> Various forms of characterization have been attempted to elucidate which chemical structures can form from HCN. For example, mass spectrometry, infrared spectroscopy, and nuclear magnetic resonance (NMR) spectroscopy have been used to infer a wide variety of functional groups in polymerization product mixtures.<sup>13–16</sup> A plethora of different polymerization pathways have been proposed to explain this complexity (see ref 11 for a review). Due to their diverse nature, some HCN-derived polymers are functional materials with proposed uses in catalysis,<sup>10,17</sup> as adhesives and as coatings.<sup>18,19</sup>

**The Complex Space of HCN-Derived Chemistry.** For this analysis, we have collected HCN-based materials that appear as suggestions, hypotheses, or confirmed detections in the literature. The limitations of our selection are motivated by practicality. The full structural space of possible HCN-derived polymers and molecules is immense. For example, one automated search for reactions that result in HCN tetramers identified 678 compounds.<sup>20</sup> Several other structures have also been computationally predicted.<sup>21</sup> We here consider a selection of pure HCN-based materials, i.e., structures where the stoichiometries of H, C, and N are 1:1:1 (and no additional elements). Other HCN-based materials can incorporate oxygen,<sup>11</sup> additional nitrogen, and hydrogen<sup>22</sup> or form through condensation reactions, e.g., through the formation of NH<sub>3</sub>.<sup>11,11,23,24</sup> Such materials, which may also

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be important, are beyond the scope of this work. In the following, we will first introduce the materials we study and then proceed to evaluate their relative thermodynamic stability.

**HCN-Derived Molecules.** All HCN-derived molecules that we investigated are shown in Figure 1. In addition to HCN (1), several of these molecules have been proposed to act as monomers in polymerization:<sup>11,13,15,25-33</sup> the HCN dimers iminoacetonitrile (2) and aminocyanocarbene (3), the trimer aminomalononitrile (4), and the tetramer diaminomaleonitrile (5) (Figure 1). Out of these molecules, only 5 has been directly observed during polymerization of HCN.<sup>31,34</sup> Speculations that 2 and 4 might form are based on indirect evidence.<sup>34</sup> The proposed formation of 3 is controversial, as we will return to discuss. Aminoimidazole carbonitrile (6) and adenine (7) (Figure 1) are less likely to take part in polymerization but are included in our study because of their relevance to prebiotic chemistry.<sup>11,25</sup>

To aid our discussion, we outline a collection of pathways, which connect some of the molecules in Figure 1 to different proposed polymer structures. Sometimes, these pathways will be mentioned in the context of mechanistic details (e.g., baseor radical-catalyzed). However, we mostly use pathway to refer to the causal connections between reactant, intermediates, and products.

**Suggested Products of HCN Polymerization.** Two of the simplest HCN-derived polymers are polyimine (8) (Figure 2, pathway 1) and head-to-tail polymer (9) (Figure 2, pathway 2). Structures 8 and 9 have been suggested to form through different mechanisms. He et al.<sup>15</sup> have suggested that the formation of 8 is base-catalyzed, proceeding through the successive additions of cyanide anions. Mozhaev et al.,<sup>32</sup> who studied radiation-induced HCN polymerization, and Mamajanov and Herzfeld, who studied polymerization in the presence of radical initiators,<sup>31</sup> have both suggested 9 as a plausible product. Mamajanov and Herzfeld have further proposed that 9 can subsequently transform into two-dimensional polytriazine sheets (10, Figure 2). Both structures 9 and 10 have been computationally predicted to form when solid HCN is subjected to pressures above 72 GPa.<sup>35</sup>

Figure 3 shows 11, the only three-dimensionally connected polymer in this study. This material can be viewed as an NN  $\leftrightarrow$  HCN substituted analogue of cubic gauche nitrogen—a metastable allotrope of nitrogen synthesized above 42 GPa.<sup>36</sup> Structure 11 has been predicted to be dynamically stable (a minimum on the potential energy surface).<sup>37</sup>

**Polymerization of Iminoacetonitrile (2).** Several polymers have been proposed to form from the polymerization of 2. One such structure is Völker's ladder polymer polyaminocyanomethylene (12) (Figure 4, pathway 3a). In Völker's model, 12 converts to a cyclic "ladder" structure (13).<sup>13</sup> Umemoto and colleagues have suggested an alternative "single-



Figure 1. Different molecules linked to HCN-derived polymers.



**Figure 2.** Examples of HCN polymerization pathways catalyzed in different ways. Top: base-catalyzed formation of polyimine (8).<sup>15</sup> Bottom: radical-catalyzed<sup>31,32,35</sup> and/or pressure-induced<sup>35</sup> formation of one- and two-dimensional materials.



Figure 3. Three-dimensional HCN-derived polymer (11).<sup>36</sup> Left: color representations: carbon—brown, hydrogen—pink, and nitro-gen—gray.



**Figure 4.** Products of iminoacetonitrile (2) polymerization proposed by Völker<sup>13</sup> and Umemoto<sup>33</sup> (pathways 3a and 3b), and Ruiz-Bermejo and colleagues<sup>11</sup> (pathways 4a and 4b).

ladder" structure (14) (pathway 3b).<sup>33</sup> Ruiz-Bermejo and colleagues have suggested that 2 may react to form structures 16 and 17 (pathways 4a and 4b).<sup>11</sup>

**Polymerization of Diaminomaleonitrile (5).** Molecule **5** features prominently in HCN reaction hypotheses because it is

readily produced in polymerization experiments.<sup>16,31,34</sup> One suggested role of **5** is as an intermediate in the hypothesized formation of 7.<sup>25,34</sup> The structure **5** is known to undergo photochemical isomerization to 6.<sup>8,38</sup> It has been suggested<sup>15,17,39</sup> that the *cis*- and *trans*-forms of **5** are possible starting points for the Völker<sup>13</sup> and Umemoto<sup>33</sup> models (**12**, **13**, and **14**) shown in Figure 4.<sup>15,30</sup> Other structures proposed to form from **5** include the linear polymers **21** and **22** (Figure 5, pathway 6).<sup>11</sup>

**Polyaminomalononitrile (23).** Figure 6 shows the two last pathways that we consider. Interest in polyaminomalononitrile (23) is motivated by a suggestion by Matthews and Moser: that 23 may be a precursor to polypeptides.<sup>29</sup> This proposal has been criticized for lack of experimental evidence.<sup>40,41</sup> 3, 4, and 5 have been proposed as precursors to 23 (Figure 6, pathways 7 and 8).<sup>11,27–29,42</sup>

To facilitate comparison with laboratory experiments, predicted thermodynamics of reaction ( $\Delta G_{298K}^o$ ) are provided under standard conditions of temperature (298 K) and concentration (1 M). Relevant experiments have been conducted in a wide range of temperatures: HCN (195–363 K),<sup>16</sup> 2 (>233 K),<sup>43</sup> 4 (~298 K),<sup>18,44</sup> and 5 (298–483 K).<sup>16,24</sup> We note that astrochemical environments rich in HCN, such as Saturn's moon Titan, can be considerably colder than the temperature of our estimates and in most experiments. Although the main conclusions drawn from our predictions are largely unaffected by changes in temperature, there are some exceptions that we will return to discuss. Estimating the temperature dependence of thermodynamic stability for our entire collection of materials is beyond the scope of this work.

# RESULTS AND DISCUSSION

In this work, we compute the free energy of all of the abovementioned structures as well as a few additional ones. Our study makes use of a sampling algorithm to identify candidates to the lowest-energy ground states in the vast configurational space of the HCN-derived molecules and polymers (see the Methods section for details).<sup>45</sup> To limit computational costs, polymers have been represented as oligomers of approximately 60 atoms. Figure 7 exemplifies predicted stable conformers for three such structures evaluated with the implicit consideration of water solvation. Most experimental literature studies have been performed in aqueous solution (see ref 11 for a review) or in neat liquid HCN.<sup>15,31,32</sup> HCN and water are both polar and hydrogen bonding solvents and can be expected to cause comparable trends in the relative solvation energies.



**Figure 5.** Two pathways for polymerization of diaminomaleonitrile (5) proposed in the literature.<sup>11,31</sup>



<u>Pathway 8</u>



Figure 6. Proposed pathways to form polyaminomalononitrile (23).<sup>11,27-29,42</sup>



Figure 7. Examples of predicted HCN oligomer structures. (A) Headto-tail polymer (9), (B) cyclic polymer (16), (C) copolymer (26) of repeating units belonging to 19 and 8. Color representations: carbon—brown, hydrogen—pink, and nitrogen—gray.

Figure 8 summarizes our best estimates for the relative Gibbs free energy of all considered structures. Corresponding relative energies in the absence of solvation are available in the Supporting Information.

The energy content per HCN in the studied structures varies between +7.5 and -16.1 kcal/mol (Figure 8). In other words, note that to arrive at a reaction energy, the values shown in Figure 8 need to be multiplied with the number of HCN in each structure. For example, for adenine, this multiplication factor is 5, while for a polymer, it is *n*. Most investigated structures are clearly thermodynamically downhill from HCN in aqueous solution at 298 K. The overall thermodynamic favorability of HCN self-reactions is thus one part of the explanation for the commonly observed chemical complexity



Figure 8. Thermodynamic landscape of HCN-derived structures. Gibbs energies ( $\Delta G_{298K}^o$ ) are provided per mole of HCN and relative to a reference chain of HCN molecules. Polymeric materials are organized by the number of HCN molecules in their monomer unit.

of reaction mixtures. The only materials predicted to be unstable relative to HCN are **3**, **9**, **12**, and **14**, effectively ruling these out as plausible reaction products under the studied conditions.

Our calculations infer new insight into the viability of Völker's suggestion that 2 is the monomer unit of 12 and 13 (Figure 4, pathway 3a). Our data predict that the formation of 13 from 2 is thermodynamically favored. At the same time, we can rule out 12 as a major product since it is unstable relative to 2, as well as 8 + 1 (and most other structures). We can also rule out Umemoto's single-ladder model (14) (Figure 4, pathway 3b) as a plausible polymerization product on the grounds of unfavorable thermodynamics.

Figure 8 shows that 5 is more stable than most of the considered structures. This prediction has several important consequences. For example, structures 12, 13, and 14 are unlikely to be the products of polymerization of 5 because such transformations are predicted to be thermodynamically unfavored. Other similar examples include structures 23 and 20, predicted to lie ~1.7 and 5.8 kcal/mol of HCN above 5, respectively. Establishing that the formation of 23 from 5 is not thermodynamically spontaneous is important as the former is a proposed precursor to heteropolypeptides.<sup>27,40</sup> However, we note that our predictions do not exclude the possibility of 23 forming through other higher-energy intermediates, such as the HCN trimer 4 ( $\Delta G_{298 \text{ K}}^2 = -1.7 \text{ kcal/mol HCN}$ ).

We have found only a limited number of HCN-derived polymers and molecules to be thermodynamically downhill relative to 5. For example, and in agreement with previous theoretical studies,<sup>46</sup> the formation of 7 from 5 is thermodynamically allowed. The linear polymer 21 is another

example that could feasibly form from 5. Structures 19 and 33 are predicted to be the most stable polymers out of those considered, with only molecule 7 being more stable. Formation of 19 and 33 from 5 is predicted to be spontaneous by -6.0 and -6.3 kcal/mol of HCN, respectively. The corresponding polymerization enthalpy ( $\Delta H_{298 \text{ K}}^o$ ) is calculated as -9.9 and -9.7 kcal/mol of HCN (Supporting Information, Table S3). These energies are in fair agreement with estimates derived from differential scanning calorimetry thermograms of solid state and melt polymerization of 5.<sup>17</sup> That study, by Ruiz-Bermejo and colleagues, estimated heat releases of 3-5 kcal/mol HCN at 423-463 K.<sup>17</sup>

We furthermore predict that the HCN dimer **3** has an energy content of +7.5 kcal/mol HCN, i.e., its formation from HCN is uphill by 15 kcal/mol. This value is 8.4 kcal/mol lower than the corresponding value in gas phase (Supporting Information, Table S1), in turn comparable to other computational results.<sup>43,47</sup> We note that **3** has a similar energy content to hydrogen isocyanide in solution.<sup>48</sup> Nevertheless, **3** is high in energy and can be ruled out as a major component of reaction mixtures.

**Energy of Two- and Three-Dimensional Materials.** Our methodology used for calculating the data shown in Figure 8 is limited to molecules and one-dimensional polymers. For polymers with higher-dimensional connectivity, we have relied on DFT calculations with periodic boundary conditions (see the Methods section). Figure 9 shows our stability estimates for 2D and 3D polymers 10 and 11. Both systems are composed of tetrahedrally coordinated C and N atoms in a graphane-like configuration (Figure 9). Phonon band structures of 10 (including the isomers 10A and 10B)



**Figure 9.** Structure and predicted energies of **10** and **11**. Gibbs energies  $(\Delta G_{298 \text{ K}}^{0})$  in kcal/mol are provided relative to an infinite chain of HCN molecules in vacuum. Color representations: carbon—brown, hydrogen—pink, and nitrogen—gray.

and 11, provided in the Supporting Information (Figures S2–S4), confirm that these materials are dynamically stable. In other words, these structures all constitute local minima on the potential energy surface, which allows us to account for temperature effects in our relative stability estimates. Although structures 10A and 10B are calculated as only slightly thermodynamically favored (-1.9-0.4) relative to our reference of an infinite chain of HCN molecules at ambient conditions (Figure S1), 11 is instead predicted to be unstable by +0.5 kcal/mol.

Approximate Stability Prediction of HCN-Based Polymers. The repeating units of several structures in Figure 8 can be considered combinations of simpler monomer units. For example, structure 23 can be viewed as a combination of 8 and 15 (Figure 8). Figure 10 illustrates how our calculated energies of smaller polymer units can be used to estimate the relative energies of more complex HCN-based structures through a weighted averaging approach (see the Methods section for details). Such linear extrapolation is approximate and can be expected to fail, e.g., in the presence of extensive conjugation. Nonetheless, this approach constitutes a rapid route to exploring the thermodynamics of a larger structural space of HCN-derived materials.

**Effect of Temperature.** Before speculating on the consequences of our predictions on different astrochemical environments, we remind that relative Gibbs energies depend on temperature. To understand such dependencies, we recall that the Gibbs energy can be expressed as

$$\Delta G = \Delta H - T \Delta S \tag{1}$$

where H is the enthalpy, T is the temperature, and S is the entropy. The energies we discuss (Figures 8 and 9) include



**Figure 10.** Quantum mechanically calculated stability (c.f. Figure 8) plotted against a weighted average extrapolation of known energies of smaller subunits (see the Methods section). The mean absolute error is 3.3 kcal/mol.

corrections for enthalpic and entropic contributions at 298 K. This choice was made to approximately align with experiments, typically performed under near Earth ambient conditions. Although the broad strokes of the presented thermodynamic map will persist at lower temperatures, changes in the order of a few kcal/mol can be expected. In general, relative enthalpies ( $\Delta H$ ) are not substantially affected by temperature and Tables S1–S4 exemplify how  $\Delta H_{298 \text{ K}}$  is approximately equal to  $\Delta H_{T\to0 \text{ K}}$  ( $\Delta E$  including zero-point energy) for a selection of studied reactions.

The largest temperature effect on our Gibbs energy estimates derives from changing entropy, the  $T\Delta S$ -term of eq 1. The combination of HCN molecules with progressively larger structures is resisted by decreasing entropy ( $\Delta S < 0$ ), a change dominated by the removal of rotational and translational degrees of freedom. In low-temperature environments, such as those on Titan, the resulting entropic penalty for making larger molecules and polymers ( $-T\Delta S$ ) decreases. Figure 11 exemplifies how such entropy-driven temperature effects can disfavor the formation of smaller molecules, relative to polymerization. In other words, thermodynamic changes brought by temperatures dropping, e.g., from Earth to Titan surface conditions, can change the outcome of prebiotic reaction chemistry.

# CONCLUSIONS

Our study shows that several hypotheses regarding HCN's reactions to molecules and polymers can be ruled out on the grounds of thermodynamics. For example, diaminomaleonitrile (5), a prominently suggested reaction intermediate in HCN-based prebiotic chemistry, is predicted to be lower in energy compared with many proposed polymer products. Our study also infers a new perspective on adenine (7), a nucleobase,



Figure 11. Example of how temperature can affect the relative stability of HCN-derived materials. As the temperature drops below the eutectic point of HCN:water mixtures, polyimine (8) becomes favored over the HCN dimer iminoacetonitrile (2). Such stability inversions may have consequences for chemical evolution in cold environments, e.g., in the atmosphere and surface of Saturn's moon Titan.

present in all life as we currently know it (Figure 8). Formation of 7 is predicted to be thermodynamically downhill from *all* structures considered herein under ambient conditions. The low experimental yields ( $<0.01\%^8$ ) of 7 from HCN in typical condensed phase experiments therefore means such reactions must be kinetically controlled. One class of materials that closely competes with the thermodynamic favorability of adenine is polyaminoimidazoles (19, Figure 5, pathway 5). In other words, under the assumption that reaction kinetics are permissive, structures such as 19 and 33 should be expected as products of HCN self-reaction chemistry.

The presented thermodynamic landscape of HCN-derived materials has several consequences for interpreting HCN's reactivity in various astrochemical environments, in and beyond the solar system. Most (but not all) previously proposed HCN-derived polymers and molecules can form spontaneously ( $\Delta G_r < 0$ ) close to Earth ambient conditions. In low-temperature environments, such as those on Titan, the entropic penalty for making larger molecules and polymers ( $-T\Delta S$ ) naturally decreases. In other words, the chances for thermodynamic spontaneity of macromolecular formation increase in cold environments.

We end by emphasizing that thermodynamics is only one side of the chemical reactivity coin. At the same time, as more structures can become thermodynamically viable at lower temperatures, such chemistry will naturally be more limited by reaction kinetics.<sup>49</sup> Enhanced kinetic control of HCN-based prebiotic chemistry means a greater ability of colder environments to generate specific metastable products from an increased pool of thermodynamically allowed options. We can therefore speculate on the potential of colder worlds, such as Titan, in generating kinetically selected products that may otherwise not form at higher temperatures. Some such materials could, hypothetically, be crucial templates for the origin of life.<sup>10</sup> We report on reaction kinetics for the formation of 2, 4, 5, and 8 in refs 49 and 50, and will return in future work with a detailed study of the formation of adenine (7).

# METHODS

**Conformational Search.** The Conformer-Rotamer Ensemble Sampling Tool (CREST)<sup>45</sup> was used in automated searches of lowest-lying conformers of molecules and oligomers. The energy evaluations in CREST make use of the semiempirical xtb program, version 6.4.0.<sup>51</sup> Conformational structure searches in aqueous solution were performed with the analytical linearized Poisson–Boltzmann (ALPB) model.<sup>52</sup> Structural differences between predicted larger and smaller oligomers risk influencing the computed energy content, e.g., through different end-effects. We therefore additionally relied on visual inspection to ensure models of oligomers of different lengths were similar in appearance/ folding.

**Electronic Structure Calculations.** With the exception for two and three-dimensional networks, all structures were optimized using the B3LYP<sup>53</sup> functional, as implemented in Gaussian 16, revision B.01.<sup>54</sup> Missing treatment of dispersion effects in B3LYP was corrected for using Grimme's method combined with Becke-Johnson damping (D3BJ).<sup>55</sup> The standard polarizable continuum (PCM) model of Gaussian was used to model interactions with water solvent.<sup>56</sup> Our best estimate to total Gibbs free energies,  $G_{\rm fp}$  was calculated as

$$G_{\rm f} = E_{\rm f} + G_{\rm corr} \tag{2}$$

where  $G_{corr}$  denotes thermal corrections obtained following optimization and frequency analyses using the cc-pVDZ basis set. Final electronic energies,  $E_{tr}$  in eq 2 were calculated as single-point calculations at the B3LYP-D3BJ/aug-cc-pVTZ level of theory. Method validations were performed against the M06-2X<sup>57</sup> meta-GGA functional, and close agreement was found (Tables S2 and S4). The average absolute deviation between B3LYP-D3 and M06-2X results in PCM-water is 0.65 kcal/mol, with a maximum of 3.2 kcal/mol for compound 9.

**Calculations of Extended Structures.** Periodic DFT calculations were carried out using the Vienna Ab Initio Simulation Package (VASP)<sup>58</sup> version 5.4.4. The B3LYP-D3BJ level of theory was used for both structure optimizations and calculations of phonon spectra. These calculations relied on projected augmented wave (PAW) potentials<sup>59</sup> used together with a plane wave kinetic energy cutoff of 800 eV.  $\Gamma$ -Centered *k*-point mesh with a resolution <  $2\pi$  0.066 Å<sup>-1</sup> was used for integration over reciprocal space. Two-dimensional polymers were separated by 30 Å vacuum layers. Energies and forces were converged to less than 1 meV/atom and 1 meV/(Å × atom), respectively. Thermal corrections were calculated using PHONOPY<sup>60</sup> version 2.11.0 using supercell dimensions of 2 × 2 × 2, 2 × 2 × 1, and 4 × 1 × 2 for the **11, 10A**, and **10B** polymers, respectively.

**Estimating the Stability of Molecules.** The average energy per unit of HCN is here used when comparing different materials

$$\Delta G_{298\,\mathrm{K}}^{0} = \frac{G_{\mathrm{molecule}}}{k} - G_{\mathrm{HCN}} \tag{3}$$

where  $G_{\text{molecule}}$  is the total free energy of the molecule, k is the number of HCN in the molecule, and  $G_{\text{HCN}}$  is the average energy of HCN inside a one-dimensional H-bonded chain.

**Estimating the Stability of Polymers.** The average Gibbs energy per unit of HCN inside polymeric materials is here evaluated as

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$$\Delta G_{298 \text{ K}}^{0} = \frac{G_{\text{f,Long}} - G_{\text{f,Short}}}{k} - G_{\text{HCN}}$$
(4)

where  $G_{f,Long}$  and  $G_{f,Short}$  are the total free energies of the (n)long and (n-k)-short polymer models, respectively, n is the number of HCN in the long oligomer model, and k is the number of HCN in a monomer unit. The sizes of our oligomer models were chosen to have at least 4 repeating polymer units, and n is always near 20. Each polymer model was capped by a cyanide group at one end and a hydrogen atom at the other end.

**Computing the Energies of 10 and 11.** Thermodynamic stabilities of **10A**, **10B**, and **11** were estimated by dividing the calculated total energy of each material by the number of HCN in its unit cell, and by subtracting the average energy of HCN in an infinite hydrogen-bonded chain. Our molecular and periodic calculations are of comparable levels of theory, also evidenced by the hydrogen bond distances between HCN differing by less than 0.025 Å.

**Energy Estimates of Composite Polymers.** The Gibbs free energy content of HCN in composite polymers,  $G_{cp}$ , can, as a first approximation, be estimated from a weighted average of our quantum mechanically computed free energies  $G_{f,i}$ ,

$$G_{\rm cp} = \frac{\sum_{i=1}^{l} k_i G_{{\rm f},i}}{\sum_{i=1}^{l} k_i}$$
(5)

where  $k_i$  is the number of HCN in the polymer units of polymers *i* and *l* is the number of monomer units in the combined polymer.

# ASSOCIATED CONTENT

#### **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsearthspace-chem.4c00088.

Relative free energy, enthalpy, and zero-point energies in aqueous solution; relative free energy, enthalpy and zero-point energies in vacuum; and phonon bands of structures **10A**, **10B**, and **11** (PDF)

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# **Author Contributions**

Conceptualization: M.R. Investigation: H.S., F.I.-R., M.C., R.D., S.S., C.B. Methodology: H.S., M.R., F.I.-R. Writing—original draft: H.S., M.R., F.I.-R.

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# Notes

The authors declare no competing financial interest.

# ABBREVIATIONS

HCN, hydrogen cyanide; DFT, density functional theory; CREST, Conformer-Rotamer Ensemble Sampling Tool; ALPB, analytical linearized Poisson-Boltzmann; VASP, Vienna Ab initio Simulation Package; PAW, projected augmented wave; B3LYP, Becke, 3-parameter, Lee-Yang-Parr

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