Formation of the –N(NO)N(NO)– polymer at high pressure and stabilization at ambient conditions

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Contributed by William A. Goddard III, December 31, 2012 (sent for review November 12, 2012)

A number of exotic structures have been formed through high-pressure chemistry, but applications have been hindered by difficulties in recovering the high-pressure phase to ambient conditions (i.e., one atmosphere and 300 K). Here we use dispersion-corrected density functional theory (DFT-uldg) (Perdew-Burke-Ernzerhof flavor of DFT with the universal low gradient correction for long range London dispersion) to predict that above 60 gigapascal (GPa) the most stable form of N2O (the laughing gas in its molecular form) is a one-dimensional polymer with an all-nitrogen backbone analogous to cis-polyacetylene in which alternate N are bonded (ionic covalent) to O. The analogous trans-polymer is only 0.03–0.10 eV/molecular unit less stable. Upon relaxation to ambient conditions, both polymers relax below 14 GPa to the same stable nonplanar trans-polymer. The predicted phonon spectrum and dissociation kinetics validates the stability of this trans-poly-NNO at ambient conditions, which has potential applications as a type of conducting nonlinear optical polymer with all-nitrogen chains and as a high-energy oxidizer for rocket propulsion. This work illustrates in silico materials discovery particularly in the realm of extreme conditions (very high pressure or temperature).

DFT | high pressure physics and chemistry | prediction of novel materials

With strong interplay between experiment and theory, such as N2 (1, 2), CO2 (3, 4), CO (5, 6), NH3 (7), and benzene (8) have been transformed into extended solids (covalent and ionic bonded networks) under high pressures. These studies have enhanced our understanding of chemical bonds under compression and provide opportunities to seek additional novel materials; however, it has been difficult to retain these remarkable structures at the ambient conditions needed for most applications (9). For CO2, a 3D covalent network was synthesized (3) at high pressure (40 GPa) and temperature (1,800 K) that is isomorphic to the β-cristobalite phase of SiO2 (10), with each carbon atom bonded tetrahedrally to four oxygen atoms. This phase of CO2 was proposed to have potential applications as superhard (initial experiments estimated a bulk modulus of 365 GPa (11), but theory and experiment later found it to be 136 GPa (10, 12)), nonlinear optical, and high-energy density material, so efforts were made to quench this phase down to 1 atm and 300 K (3); however, it reverted back to the molecular phase at pressures lower than 1 GPa.

Because it is isoelectronic to CO2 but polar, attempts were made to form an extended solid from N2O using compression (above 20 GPa) and laser heating (above 1,000 K) in a diamond anvil (13). However, instead it decomposed into a mixture of an ionic crystal NO+NO− and compressed N2 molecules. No covalent extended framework similar to the polymeric CO2 phase was found. Indeed, because the nitrogen atom forms one less covalent bond than the carbon atom, it is not obvious that it would be possible to construct a dense extended solid phase of N2O other than the ionic form. Even so, we decided to use first principle method to explore a large number of space groups and bonding patterns at high pressures. Because we start with molecular crystals of nitrous oxide (NNO) in which London dispersion attractions (van der Waals attraction) is dominant, we used the PBE-uldg flavor of density functional theory (DFT-uldg) in which corrections accounting for London dispersion attractions are included (14). We searched for high-pressure structures with two independent strategies.

In the first approach, we started with 41 known AB2-type crystal structures, replaced A with O and B with N, and used DFT to determine the optimum atom positions and packings using a fixed density of 3.915 g/cm3 (a relative compression of 0.384 and the same as polymeric CO2 at 41 GPa) (10). Then we took the new configurations from the lowest energy structures, built appropriate extended cells, and refined further with DFT to obtain the optimum stacking.

In the second approach, we used the USPEX (Universal Structure Predictor: Evolutionary Xtallography) code (15-17) based on an evolutionary algorithm developed by Romanov, Glass, and Lyakhov (15) and featuring local optimization, real-space representation, and flexible physically motivated variation operators. Here we considered a 12-atom unit cell at 40, 60, and 80 GPa.

Results

Both methods led to the same two final high-pressure structures: both corresponding to a 1D N2O polymer but with either cis- (Fig. 1B) and trans- (Fig. 1C) conformations (cis is lower), predicted to be the most stable species above 60 GPa. The valence bond descriptions of these structures has an alternating N and N+ backbone polyacetylene like zigzag chain with each N+ forming a covalent bond to O2. The valence bond description would suggest alternating single and double bonds (1,448 Å and 1,309 Å), which we find for cis; however, for trans we find all NN bonds to have the same bond distances (1,390 Å at 1 atm), indicating full resonance along the chain. Fig. 1 shows the enthalpies of various relevant phases of N2O as a function of pressure, taking the ground state molecular crystal with space group Pn3 (α-nitrogen structure) as the reference. For the two molecular phases, Pn3 and space group Cmc a, experiments find head-to-tail orientation disorder (18), so we considered all possible conformations in the unit cell to determine the lowest energy. The calculated dependence of crystal volume on pressure is comparable with experiment (Fig. S1), leading to a calculated transition pressure between the two molecular phases of 5.8 GPa at 0 K, compared with the experimental value of 4.8 GPa at room temperature (18).

In the range of 10–55 GPa and 1,000–3,400 K, the diamond anvil experiments on NNO observed a mixture of the ionic NO+N2O3− phase in equilibrium with an N2 molecule phase, formed from the dissociation of N2O at high pressure and temperature (13, 19, 20). We also calculated these phases at the pressures considered here. However, the structure of NO+N2O3− is not well-established experimentally: experiments reported the formation of monoclinic P21/m phase ionic salt from the high-pressure


The authors declare no conflict of interest.

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This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10.1073/pnas.1222890110/-/DCSupplemental.
(2 GPa) reaction of an N₂ and O₂ mixture (21), whereas the original experiments on dissociation of N₂O under laser heating in a diamond anvil found an orthorhombic phase related to the aragonite to form after cooling (13, 22). Therefore, for the DFT study, we constructed the NO(NO₃) crystal by cell relaxation starting with the aragonite structure. This led to an orthorhombic Pna2₁ phase (Fig. S2 and Table S1), which we found to be 0.03 eV per formula more stable than the monoclinic phase. We carried out DFT calculations of N₂ molecular crystals, starting with the known phases (α, γ, ε, ζ) at low temperature (23), and at each pressure, the energy calculated to be most stable was used in the enthalpy calculations. Surprisingly, this combination of NO(NO₃)⁻ and N₂ is more stable than the NNO molecular phases for all pressures down to 0 GPa. This is plausible, as the heat of formation of N₂O molecule is 82 kJ/mol (1 kcal/mol) above N₂ and O₂, which were shown to form NO(NO₃)⁻ at 2 GPa using 10.2 keV synchrotron X-ray radiation (21).

Fig. 1 shows that the cis and trans 1D polymeric phases remain energetically close (cis more stable by 0.03–0.10 eV/molecular unit) and become more stable than both the molecular and the mixed ionic phases at pressures above 60 GPa. Including zero point energy (ZPE) and entropy from phonons has little effect (Fig. S3):

- including ZPE to obtain the 0 K equation of state (EOS), we find that the critical pressure for the transition from the mixed ionic phase to the cis-polymer increases from 58.5 GPa to 61 GPa,
- while introducing entropy and enthalpy corrections to 300 K leads to a further increase to 62 GPa at 300 K (64 GPa at 500 K).

The dominant factor in determining the enthalpy at high pressure is the compressibility. For the mixed phase, the ionic NO(NO₃)⁻ component is a dense solid, but the global compressibility of the mixed phase is dominated by the molecular phase of N₂, which polymerizes only above 110 GPa (2). In contrast, the two new NNO polymeric phases each forms a pure extended solid with mixed covalent and ionic frameworks. Thus, these phases dominate at high pressures (where the PV term in the enthalpy becomes dominant) over all of the other phases of N₂O containing molecular forms.

The major difference between these cis and trans forms is that the planar cis-NNO polymer possesses alternating single and double bonds as in polyenes (polyacetylene) with an even number of carbons, whereas the planar trans-conformation of NNO has equal NN bond lengths (strong resonance) along the chain, perhaps due to strong electrostatic repulsion between adjacent negatively charged oxygen atoms.

We further investigated the stability of the two NNO polymers by phonon analysis. Indeed, for both crystals we find no imaginary phonon modes under high pressures, as shown in the phonon spectra in Fig. 2 (at 80 GPa), which confirms their existence as stable species (local minima). However, as the pressure is released, both crystals develop imaginary phonon modes at ~14 GPa, and both relax to one single nonplanar trans-conformation (Dataset S1) at zero pressure, with a phonon spectrum that attests its stability. Under high pressures, the planar conformations with their higher compressibilities are more stable, but they lead to larger electrostatic repulsion between negatively charged oxygen atoms and lone-pair on nitrogen in the cis-case and between adjacent negatively charged oxygen atoms in the trans-case. It is the competition between compressibility and electrostatics that leads to the transition to the nonplanar polymer at low pressure.

Discussion

Besides accommodating electrostatics, the transition between planar and nonplanar trans-conformations is also accompanied by an abrupt change from uniform bonding to alternating single and double bonds, as shown in Fig. 3, implying synergistic Peierls distortion, which is relevant to the electronic properties. Therefore, we calculated band gaps of all polymeric structures (opti-
fied with PBE-ulg flavor of DFT) and plotted against pressures in Fig. 3. Interestingly, PBE-ulg predicts that both planar NNO polymers are 1D metals, with transitions to insulators below 72 GPa for cis-polymer and 25 GPa for trans-polymer, the latter with
simultaneous conformation transition from planar to nonplanar. Fig. 3 also shows the bond length differences between neighboring \( N-N \) bonds for the various phases. We find a trend from uniform bonding (exact for \( \text{trans} \)-polymer, quasi-uniform for \( \text{cis} \)-polymer with 0.03 Å difference) at high pressures to alternating single and double bonds when pressure is released. Combining this with the trend in band gaps indicates a first-order Peierls distortion as in polyacetylene. However, it is well documented (24, 25) that density functionals, such as PBE that are based only on local density approximation and generalized gradient approximation, significantly underestimate band gaps for insulators, due to intrinsic delocalization error (26) from including the self-interaction in the Coulomb energy. To verify the metallic properties of these polymers under high pressures, we reinvestigated all electronic structures with the hybrid functional B3PW91 (27), which we showed to predict accurate band gaps for a wide range of semiconductors (28). In contrast to PBE, B3PW91 predicts insulating states for all three polymers throughout the whole range of pressures considered, as shown in Fig. 3. Here the density of states (Fig. S4) shows that both planar \( \text{cis} \)- and \( \text{trans} \)-polymers at high pressures with (quasi-) uniform bonding are charge-transfer insulators, with the valence band dominated by negatively charged oxygen and bridging nitrogen atoms, whereas the conduction band is dominated by the positively charged nitrogen atom. Accordingly, B3PW91 results

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**Fig. 2.** Phonon spectra for polymer phases of \( \text{N}_2\text{O} \) predicted at 80 and 0 GPa. Both planar \( \text{cis} \)- and \( \text{trans} \)-polymer phases develop imaginary phonon modes, due to electrostatic repulsions and/or second-order Peierls distortion, and transform into the same nonplanar \( \text{trans} \)-polymer phase, the phonon spectrum of which at 0 GPa shows no imaginary modes.

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**Fig. 3.** Band gap (\( A \)) and neighboring \( N-N \) bond length difference (\( B \)) predicted as functions of pressure for all three polymer phases of \( \text{N}_2\text{O} \) at ground states. The dashed lines with arrows indicate where the transition from planar to nonplanar conformation happens.
infer the transition to be second-order Peierls distortion as in polyacene. At this point we cannot be sure whether to trust the B3PW91 or the PBE description, so it would be most valuable to carry out experiments on the conductivity or electronic spectra of these new phases.

To further evaluate the stability of nonplanar trans-polymer, we investigated the dissociation kinetics of the polymer, using DFT for a model oligomer containing 8 N₂O units and terminated with methyl groups at both ends. We stretched this finite chain until it fractured, locating the transition state (TS) shown in Fig. 4. This leads to an activation energy of 20.6 kcal/mol. At the TS, the partially dissociated N₂O has N-N distances of 2.062 and 1.830 Å to its two neighbors, much longer than corresponding equilibrium distances of 1.465 and 1.447 Å. This indicates that a large free space is required to activate the dissociation process, making the dissociation less favorable in the solid. This high-activation barrier of 20.6 kcal/mol from the finite model corresponds to a surface energy of 1.56 × 10⁻³ erg/cm², considering the packing in the crystal. Thus, we consider that it is likely that the nonplanar trans-polymer of N₂O will be stable at 1 atm pressure and 300 K.

After passing the TS, the oligomer would start releasing N₂O one by one from each end, accompanied with significant heat release of 40.6 kcal/mol per N₂O molecule, which might lead to catastrophic decomposition. Thus, the nonplanar trans-NNO polymer is a high-energy content structural material. Indeed, we calculated (PBE-ulg) the nonplanar N₂O trans-polymer to provide an internal energy release of 3.5 kJ/g when dissociated into N₂ and O₂, which is comparable to the energy release of trinitrotoluene (TNT) (4.2 kJ/g). Thus, nonplanar trans-NNO is a potential high-energy oxidizer for new explosive composites and rocket propellants.

Since the N₂O polymer chain is composed of alternating single and double bonds, similar to polyacetylene, albeit with an all-nitrogen backbone, it might form the basis for a unique type of conducting polymer, through appropriate doping or structure modification. Similarly we expect that it might have strong nonlinear polarizabilities for nonlinear optical applications.

Summarizing, we used DFT to predict that the NNO molecular crystal can be transformed into novel polymeric phases at high pressures (beyond decomposition into mixture phase of ionic compound NO⁺NO⁻ and N₂ gas previously observed in experiments). The two most stable 1D N₂O polymers with planar cis- and trans-conformations were identified to be energetically favorable at pressures above 60 GPa. More importantly, when the pressure is released, these polymers transform into the same nonplanar trans-conformation, stable at ambient pressure and temperature. This was substantiated by analysis of the phonon spectrum and by calculating the dissociation kinetics. This unique poly-NNO material might be an excellent high-energy oxidizer for a polymer composite in which nonpolar NNO forms the matrix.

The PBE calculations suggest metallic property in the high-pressure polymer resulting from the strong resonance in these systems, which seems to disappear in the low-pressure nonplanar polymer due to changes in bonding, leading to Peierls distortion, together with electrostatic repulsions. Thus, with doping these NNO polymers, such as I, which works successfully in polyacetylene, may give rise to a new type of conducting polymer based on all-nitrogen chains. This whole work serves to illustrate the unique in silico process of discovering materials by theoretical modeling particularly under extreme conditions.

**Methods**

PBE-ulg calculations were performed using Vienna Ab-initio Simulation Package (VASP) package (29–31), modified to describe PBE-ulg and using the projector augmented wave method (32) to account for core-valence interactions. The kinetic energy cutoff for plane wave expansions was set to 500 eV, and the reciprocal space was sampled by Γ-centered Monkhorst-Pack scheme with a fine resolution of 2 × 1/60 Å⁻¹.

In the search of structures using USPEX (15–17), the kinetic energy cutoff was lowered to 400 eV, with reciprocal grids of 2 × 0.06 Å⁻¹. The convergence criteria were set to 1 × 10⁻¹ eV energy difference for solving for the electronic wavefunction and 1 × 10⁻⁶ eV/Å force for geometry optimization.

To obtain the phonon spectra, very tight convergence criteria were used, with 1 × 10⁻⁸ eV energy difference and 1 × 10⁻⁶ eV/Å force thresholds. To calculate force constants, we used the supercell approach with finite displacements, as implemented in the Phonopy code (33).

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**Fig. 4.** Dissociation path calculated for dissociation of the model 8-mer. The oligomer contains 8 N₂O units and is terminated with methyl groups at both ends.
B3PW91 calculations were conducted using CRYSTAL09 package (34) and 6–311G* triple-zeta quality basis sets for both nitrogen and oxygen. Similar k-space grids and convergence criteria as in fine calculations with VASP were set for consistency.

For finite model calculations, the geometry optimization, Hessian calculation, and TS search were carried out at UB3LYP/–311G* level using Jaguar (35). The Hessian was used to provide the vibrational frequencies for ZPE and thermo-corrections to enthalpy. The TS was shown to have exactly one imaginary vibrational mode by following the minimum energy path scan to connect reactant and product. Enthalpies are reported at 298.15 K and 1 atm.

ACKNOWLEDGMENTS. This work was supported by the Office of Naval Research (N00014-12-1-0538; program manager, Cliff Bedford) and the Defense Advanced Research Planning Agency (program manager, Judah Goldwasser).

3. Iota V, Yoo CS, Cynn H (1999) Quartzlike carbon dioxide: An optically nonlinear ex-


Supporting Information

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Fig. S1. Comparison between PBE-ulg calculated (green filled symbols) and experimental (1, 2) (black open symbols) equation of states for molecular phases of N\textsubscript{2}O. Error bars are shown for all experimental data, and the agreement is excellent for high pressures (> 2 GPa), with the largest deviation of 1.8% for volume (0.6% for length dimension), which is most likely the limitation of parent PBE functional (3). However, at low pressure of 1 atm (0.001 GPa), PBE-ulg gives overestimation of 6.6% for volume (2.2% for length dimension), which may be again originated from PBE's description of molecular bonding: for N\textsubscript{2}O molecule, PBE gives N-N and N-O bond lengths of 1.143 Å and 1.198 Å, overestimated by 1.3% and 1.2% with respect to experimental data of 1.128 Å and 1.184 Å. This error gets amplified through dipole in evaluating volume of the molecular crystal. At high pressure, molecular crystals resemble more as extended solids, for which PBE is better from theoretical construction. Indeed, as shown in this figure, PBE-ulg calculated volume agrees better and better when pressure is increased.

Fig. S2. Illustration of the orthorhombic crystal structure of ionic NO\(^+\)NO\(_3^−\) with space group Pna2\(_1\). The structure was derived from aragonite, with lower energy than the monoclinic phase reported, and cell coordinates are reported in Table S1.
Fig. S3. The effect of temperature (particularly entropy) and quantum effects (particularly zero point energy, ZPE) on the transition pressure (TP). A shows relative enthalpy (without ZPE) curves, which predict TP = 58.5 GPa; B shows relative Gibbs free energy (at 0 K; i.e., no entropy contribution, but ZPE is included) curves, rendering TP = 61 GPa; C shows relative Gibbs free energy (at 300 K) curves, increasing TP to 62 GPa; D shows relative Gibbs free energy (at 500 K) curves, further boosting TP to 64 GPa. Since including both ZPE and entropy at 300 K increases TP by only 3.5 GPa, we consider that high temperature is not essential for these systems.
Fig. S4. Total and partial density of electronic states (DOS) calculated with the B3PW91 flavor of DFT for planar cis-polymer (A) and planar trans-polymer (B) at 80 GPa. The different colors indicate atoms on which the total DOS is projected. The valence band maximum (highest occupied molecular orbital) is set as the energy zero for this ground state structure.

Table S1. Fractional cell coordinates for ionic NO$^+$NO$_3^-$ with orthorhombic space group $Pn_a2_1$ at 0 GPa

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<th>$y/b$</th>
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The lattice parameters are $a = 6.3603$ Å, $b = 8.0200$ Å, and $c = 6.9563$ Å.

Other Supporting Information Files

Dataset S1 (TXT)