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New cubic phase of lithium nitride to 200 GPa

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We present a new cubic (γ) Li_3N phase discovered above $40(\pm 5)$ GPa. Structure and electronic bands are examined at high pressure with synchrotron x-ray diffraction and inelastic x-ray scattering in a diamond anvil cell, and also with first-principles calculations. We observe a dramatic band-gap widening and volume collapse at the phase transition. γ - Li_3N remains extremely stable and ionic to 200 GPa, with predicted metallization near 8 TPa. The high structural stability, wide band-gap and simple electronic structure of γ - Li_3N are analogous to that of such lower valence closed-shell solids as NaCl, MgO and Ne, meriting its use as a low-Z internal pressure standard.

Most elements in the periodic table form thermodynamically stable nitride compounds, with chemical bonding ranging from covalent to ionic to semimetallic. Lithium nitride is the only known alkali metal nitride and is one of the most ionic of all known nitrides. At ambient pressure, the nitrogen exists in an anomalous multiply charged (nearly N^{3-}) state [1, 2] which is stable only because of its crystal environment - a hexagonal bipyramid of Li^+ ions. This layered structure (α - Li_3N , P6/mmm) consists of Li_2N layers, widely separated and connected by one lithium atom per unit cell occupying a site between the nitrogen atoms in adjacent layers [3, 4]. This material is a superionic conductor via vacancy-induced Li^+ diffusion in the Li_2N layers. [5–7] Its potential for use as an electrolyte in lithium batteries [4], a hydrogen storage medium [8–11] and a component in the synthesis of GaN [12] has prompted several studies including an investigation into its behavior at high pressure [13].

At ~ 0.5 GPa, α - Li_3N transforms into a second layered hexagonal structure (β , P6₃/mmc) with honeycomb-like LiN layers [14]. In this structure, each nitrogen binds an additional lithium atom above and below the plane and, unlike the Li_2N layers in α - Li_3N , adjacent LiN layers are shifted relative to one another. β - Li_3N is metastable at ambient pressure and is typically found mixed with the α -phase. It remains stable up to 35 GPa – the high-pressure limit of experiments on Li_3N to date. A second phase transition to a cubic structure - P-43m at 37.9 GPa [13] or Fm3m at 27.6 GPa [15] - has been predicted. If it exists, the similarity of this phase to those of other simple ionic cubic solids such as NaCl makes it an interesting study, particularly in light of its higher ionicity. Understanding the behavior of the unstable and highly charged N^{3-} ions under large compression will be particularly important.

In this paper, we present the first concrete experimental evidence that β - Li_3N indeed transforms to a cubic structure (γ - Li_3N) in the pressure range of 36–45 GPa.

This transformation represents an increase in structural and bonding strength and isotropy, and is accompanied by a relatively large volume collapse and a significant widening of the electronic band gap. γ - Li_3N is uniquely stable and quite compressible in this pressure regime and up to at least 200 GPa, making it a good candidate for an internal pressure standard.

Polycrystalline lithium nitride powder (99.5 % purity, CERAC, Inc) was loaded into a membrane diamond-anvil cell of LLNL design. Several diamond sizes were used to obtain an extended range of pressure up to 200 GPa. In the lower pressure experiments, argon was used as a pressure medium and internal pressure standard. For the high-pressure experiments, no pressure medium was used, and copper or ruby ($\text{Al}_2\text{O}_3:\text{Cr}^{3+}$) were included in the sample chamber as pressure indicators. Under non-hydrostatic conditions, we found that the equation of state fitting parameters differed from those obtained under quasi-hydrostatic conditions by 8.7%, 0.8% and

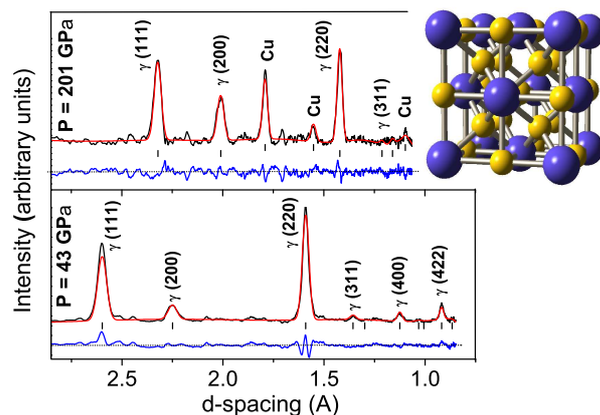


FIG. 1: ADXD diagrams of γ - Li_3N at 43 and 201 GPa, with refined and difference patterns. Miller indices are based on the cubic (Fm3m) structure shown, with large atoms representing the highly negative nitrogen ions, small representing lithium.

TABLE I: Volume per formula unit V_o , bulk modulus B_o , its pressure derivative B_o' , volume change at the $\beta \rightarrow \gamma$ transition and transition pressure as obtained in experimental(*) and theoretical work in present and other studies. Experimental errors are primarily a result of non-hydrostaticity in the DAC. The γ -phase predicted in reference [13] is space group P-43m.

		V_o ($\frac{\text{\AA}^3}{f.u.}$)	B_o (GPa)	B_o'	$\frac{\Delta V}{V_o}$	P(GPa)
Exp.*	β	34.4(.8)	71(19)	3.9(.9)	8(2)%	40(5)
	γ	30.8(.8)	78(13)	4.2(.2)		
Th.	β	34.44(.08)	68(3)	3.6(.1)	6.7%	40.4
	γ	31.16(.08)	73.1(.8)	3.85(.01)		
[13]*	β	35.04	74(6)	3.7(.7)		>35
[13]	β	30.88	78.2	3.77	8%	37.9
	γ	28.08	82.8	3.84		
[15]	β	33.36				28(5)
	γ	30.44				
[14]*	β	34.48				>10

13.2% for B_o , V_o and B_o' , respectively. Samples were loaded in an argon environment as Li_3N is hygroscopic. High-pressure behavior of was investigated by angle-dispersive powder x-ray diffraction (ADXRD) at 16IDB and inelastic x-ray Raman scattering (XRS) at 16IDD of the High-Pressure Collaborative Access Team (HP-CAT) beamlines at the Advanced Photon Source (APS). For the ADXD experiments, we used intense monochromatic x-rays ($\lambda = 0.3683$ or 0.4126 \AA) microfocused to ~ 10 μm at the sample using a pair of piezo-crystal controlled bimorphic mirrors. X-ray diffraction patterns were recorded on a high-resolution image plate detector (MAR 350) and analyzed with the FIT2D, XRDA and GSAS programs. For the XRS experiments, we used monochromatic x-rays (9.687 keV) focused to $\sim 20 \times 50$ μm at the sample through an x-ray translucent Be gasket by a pair of 1 m-long Kirkpatrick-Baez focusing mirrors. Six spherically bent Si(660) single crystal analyzers (50 mm in diameter) were vertically mounted on a 870 mm Rowland circle to refocus inelastically scattered x-ray photons onto a Si detector (Amp Tek) at a scattering angle of 25° in a nearly back scattering geometry (Bragg angle of 88.6°). This configuration corresponds to a momentum transfer of $q \sim 2.2$ \AA $^{-1}$. The overall system provides an energy resolution of ~ 1 eV.

At low pressures our x-ray data indicate a coexistence of α and β phase, consistent with previous observations [13]. The mixture converts to single phase β - Li_3N near 0.5 GPa, which appears opaque. Between 35 and 45 GPa, we observed β -phase transform to a new transparent phase, γ - Li_3N , which remains stable up to 201 GPa, the maximum pressure achieved in the present study. A possible beginning of this transition was reported in [13] and [14]. An additional transition to I- Na_3N structure predicted at 168 GPa [15] is not seen. Figure 1 shows the measured and refined diffraction patterns of γ - Li_3N

at 43 GPa and 201 GPa. The refinement was performed based on the Li_3Bi structure (Fm3m) with one formula unit per primitive fcc cell consisting of one lithium and one nitrogen ion occupying m-3m sites, and the two additional lithium ions in the -43m sites, each tetrahedrally coordinated with 4 nitrogen ions. Slight changes in relative intensities of the diffraction peaks appear to be due to increasing occupancy of the Li -43m sites with pressure (92.2% at 43 GPa and 99.9% at 201 GPa).

The pressure-volume data for the β and γ phases and their 3rd order Birch-Murnaghan equation of state (BM-EOS) fits are shown in Figure 2, with fitting parameters summarized in Table I. The $\beta \rightarrow \gamma$ transition is accompanied by an 8% volume collapse and an increase in the coordination number for every atom. In the β phase ($a \sim 3.20$ \AA, $c \sim 5.71$ \AA, at the transition), each N^{3-} ion is surrounded by 11 Li^+ ions (three in the hexagonal planes at 1.85 \AA, two above and below the plane at 1.90 \AA and six in trigonal prismatic coordination at 2.1 \AA). In the γ phase ($a \sim 4.5$ \AA), 14 Li^+ ions surround N^{3-} , eight tetrahedrally coordinated with N at 1.95 \AA and six octahedrally at 2.25 \AA. Across the phase transition there is no discontinuity in the nearest-neighbor N distance (~ 3.23 \AA), and the nearest N-Li distance even increases slightly. The significant increase in packing without a decrease in distance between highly charged N ions makes the γ phase highly preferred at high pressures. The more populated and symmetric distribution of Li ions serves to effectively shield the highly charged N ions from one another and even potentially to compress their ionic radii [16], stabilizing the cubic structure up to very large lattice constant reduction.

The compressibility of γ - Li_3N rivals other common and highly compressible closed-shell cubic solids as seen in the inset of Figure 2. Values for bulk modulus were obtained as a function of pressure by fitting to the pressure-volume data with a BM-EOS formulated relative to some non-ambient-pressure reference volume within the cubic

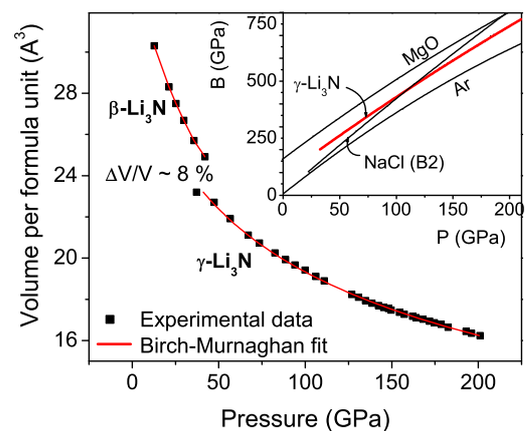


FIG. 2: Equation of state of β - and γ - Li_3N . In the inset, the high pressure bulk modulus of γ - Li_3N is compared to other common highly compressible materials [17].

phase. [17]. The fitting parameters were then used to calculate values for bulk modulus over a wide-range of pressures. The results clearly show that γ -Li₃N is harder than argon, but softer than MgO and even NaCl above 100 GPa. The distinct lack of broadening in the measured ADXD (seen at 201 GPa in Figure 1) even in the absence of a pressure medium suggests that this material has very low shear strength, which is consistent with a high compressibility.

The differences in crystal structure of γ -Li₃N from the α - and β - phases suggest that there will also be distinct differences in electronic structure. To validate this conjecture, we have carried out XRS experiments at the nitrogen K absorption edge. The most distinct features of the spectrum (shown in black in Figure 3) are a narrow near edge peak near 397 eV and a main edge near 403 eV, both of which change under pressure. The near edge peak decreases in intensity as the structure becomes more isotropic, leading us to believe that the two are somehow correlated. Because of the structural anisotropy in the hexagonal phases, there are large empty spaces in the crystal lattice and therefore larger distances between ions in the *c* direction and weaker coulombic interactions. This is evident from the larger compressibility parallel to the *c* axis which was observed for the β phase in this study and, to an even greater extent, for the α phase in [2]. The N³⁻ ions would, therefore, be highly polarizable (as is also predicted in [18, 19]), and the electrons more loosely bound in the hexagonal phases. Thus, the lower energy near-edge peak could reflect more accessible conduction states which arise because of the more weakly interacting ions perpendicular to the *a*-*b* planes. The main edge would then describe the ion interactions within the plane. The higher energy of these states indicates that they are less accessible due to the relatively smaller interstitial space in which to house itinerant electron states within the planes. The shift to higher energy

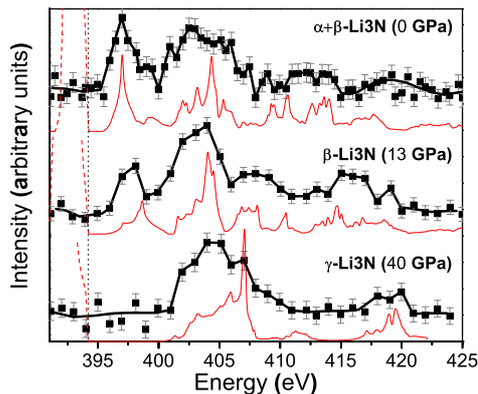


FIG. 3: XRS of α -, β -, and γ -Li₃N shown in black and error bars estimated as the square root of the counts. Nitrogen 2p projected DOS are shown in red beneath experimental curves, offset by 394.2 eV with occupied valence states as dashed lines, and vertical line representing the Fermi energy.

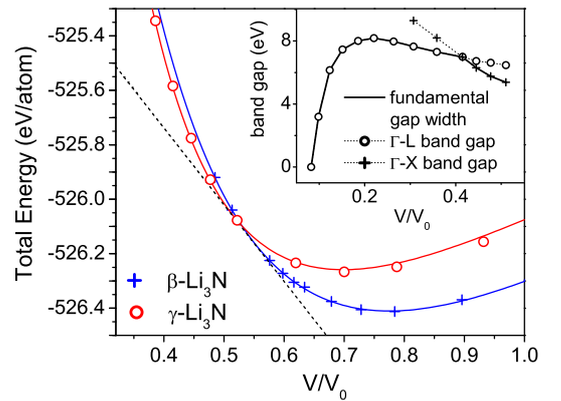


FIG. 4: Calculated EOS of β - and γ -Li₃N. The slope of the dotted line represents the transition pressure. The collapse of the electronic band gap under pressure is shown in the inset. V_0 refers to the volume at ambient pressure in the α phase.

of all peaks in each subsequent phase transition reflects a further decrease in interstitial space in all directions.

In order to explain the observed structural and spectral changes, we have performed first-principles electronic structure calculations. Because of the large six-fold compression carried out in these calculations, we used two methods for comparison: full-potential linearized augmented plane-waves (LAPW) as implemented in WIEN2k code [20] within the Generalized Gradient Approximation [22] and a full-potential nonorthogonal local-orbital minimum basis bandstructure scheme [21], within the local spin-density approximation (LSDA) [23]. We found good agreement between these two results. Figure 4 summarizes total energy calculations on the β and γ phases at various compressions and their BM-EOS fits (lines). The calculated equilibrium volumes, bulk moduli, and transition pressure are all in good agreement with experimental values (see Table I), giving us confidence in the theoretical models employed.

Using this model, we have calculated the projected nitrogen p density of states (pDOS) to compare with the XRS data. (Figure 3) A linear shift in energy is applied to the pDOS to align the leading edge peak with experiment. Within the dipole approximation, the relative intensities of the XRS spectrum are proportional to the pDOS and the squared dipole transition-matrix element. If the matrix element does not have a sharp energy dependence, the XRS and pDOS should be rather closely related, which is clearly seen in Figure 3. This agreement reveals that the electronic states observed in the XRS study can be well understood as $1s \rightarrow 2p$ transitions, and also that core-hole-electron interactions (excitons) have a lesser effect on these spectra than often observed [24–26]. It confirms that the near edge peak is indeed a reflection of low energy conduction bands which result in a smaller electronic band-gap in the hexagonal phases than in the

cubic phase. Experimentally, such a band-gap difference is evident from an optical change from opaque β to transparent γ phase. If these low-lying conduction states are coming from more loosely bound electrons in the direction of weaker ion interactions, one may expect to see some amount of free charge in the interstitial regions outside of the planes, which has indeed been indicated [1]. In the cubic phase, then, we expect more localized and symmetric charge distributions around the nitrogen atoms, making them even closer to the ideal ionic state (N^{3-}) and explaining the insulating character and high stability of this phase. In spite of a similarity between this transition and the $1s$ to π^* and σ^* transitions in graphite-diamond and boron nitride, an examination of the p_x , p_y and p_z character in Li_3N reveals that the near edge peak has a significant $p_{x,y}$ component in addition to p_z , and the main edge composition (primarily Li and N p character) does not suggest any significant hybridization between Li s and N p orbitals as is the case between B s and N p states in boron nitride.

Our calculated electronic structures further reveal that, upon decreasing unit cell volume, the (indirect) band-gap continues to increase up to a calculated pressure of ~ 760 GPa ($V/V_o = 0.22$) before beginning to collapse (see Figure 4, inset). Assuming no further phase transitions, band-gap closure does not occur until ~ 8 TPa ($V/V_o = 0.08$), which will be a lower limit because of the tendency of the local density approximation to underestimate the band gap. This high metallization pressure places γ - Li_3N in the same family as other closed-shell, cubic insulating solids such as Ne, MgO and NaCl, which are predicted to metallize near 134 TPa, 21 TPa and 0.5 TPa, respectively [27–29]. An examination of metallization in these materials reveals a trend that the states with smaller ℓ (orbital character) increase in energy with respect to larger ℓ because they are more extended and hence more greatly affected by pressure, so metallization occurs as a result of overlap between valence sp states and conduction d states [30]. Neon is isoelectronic to N^{3-} , and its metallization pressure is remarkably high because there is no 2d band, so that the overlap must occur between 2sp valence states and the much higher energy 3d conduction states. Li_3N is more complex, however, as the low-lying conduction bands have primarily Li(2p) character, with a negligibly small amount of N d character at band closure. Therefore, the phenomenon in Li_3N is more of an interspecies metallization which perhaps explains its 20-times lower metallization pressure than predicted for the intraspecies metallization in neon.

In summary, we have provided a coherent picture of the structural and spectral changes associated with a graphite-diamond-like phase transition in Li_3N , understood here for the first time. The high-pressure cubic phase has several interesting and unique properties including unusually high phase stability to pressures exceeding 200 GPa, high compressibility rivaling commonly

used pressure media such as NaCl, and also an ultra-high metallization pressure that makes it one of the most difficult materials to metallize that we know of.

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