Structural transformations of Li$_2$C$_2$ at high pressures

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Structural changes of Li$_2$C$_2$ under pressure were studied by synchrotron x-ray diffraction in a diamond anvil cell under hydrostatic conditions and by using evolutionary search methodology for crystal structure prediction. We show that the high-pressure polymorph of Li$_2$C$_2$, which forms from the Immm ground-state structure ($Z = 2$) at around 15 GPa, adopts an orthorhombic Pnma structure with $Z = 4$. Acetylide C$_2$ dumbbells characteristic of Immm Li$_2$C$_2$ are retained in Pnma Li$_2$C$_2$. The structure of Pnma Li$_2$C$_2$ relates closely to the anticotunnite-type structure. C$_2$ dumbbell units are coordinated by nine Li atoms, as compared to eight in the antifluorite structure of Immm Li$_2$C$_2$. First-principles calculations predict a transition of Pnma Li$_2$C$_2$ at 32 GPa to a topologically identical phase with a higher Cmcm symmetry. The coordination of C$_2$ dumbbell units by Li atoms is increased to 11. The structure of Cmcm Li$_2$C$_2$ relates closely to the Ni$_2$ In-type structure. It is calculated that Cmcm Li$_2$C$_2$ becomes metallic at pressures above 40 GPa. In experiments, however, Pnma Li$_2$C$_2$ is susceptible to irreversible amorphization.

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I. INTRODUCTION

Carbides of alkali and alkaline-earth metals typically occur as saltlike acetylides which consist of C$_2^-$ dumbbell anions isoelectronic to dinitrogen [1]. Recent theoretical studies suggested that acetylide carbides should transform to modifications with polymeric carbon structures at moderate pressures (below 10 GPa) [2–6]. The predicted “polycarbides” consist of carbon polyanions with chain, ribbon, or layer structures which are stabilized by electrostatic interactions with surrounding cations. Such polyanions occur typically in Zintl phases and are well known for, e.g., silicon and germanium. For carbon they represent a hitherto unknown chemical and structural feature. Polycarbides display distinct electronic structures and are predicted to be superconductors [3–5].

Yet the computational predictions deviate notably from results of experimental high-pressure studies. Hitherto investigated Li$_2$C$_2$, CaC$_2$, and BaC$_2$ have in common that acetylide C$_2$ dumbbells are retained until irreversible amorphization occurs at pressures far higher than the calculated transition pressures for polymeric carbide formation [7–9]. The discrepancy has been attributed to kinetic hindrance [3]. Prior to amorphization BaC$_2$ and Li$_2$C$_2$ undergo structural transformations at around 4 and 15 GPa, respectively, in room temperature experiments [7–9]. These transformations correspond to a “conventional” increase of coordination numbers with pressure, leading to denser packings of cations and dumbbells. In the ambient-pressure structure of BaC$_2$ Ba$^{2+}$ and C$_2^-$ ions are six-coordinated and arranged as in the NaCl structure. The rhombohedral high-pressure modification relates to the CsCl structure, with both types of ions attaining an eightfold coordination [7]. For Li$_2$C$_2$ the structure of the high-pressure form has not been conclusively characterized [8,9].

Here we present the elucidation of the high-pressure behavior of Li$_2$C$_2$ from combined synchrotron diffraction experiments and crystal structure prediction methodology. To prevent the generation of enthalpically more favorable polymeric carbides in the computations, a constrained evolutionary algorithm was employed that enforced retention of C$_2$ dumbbell units at high pressures [10]. We further show that if amorphization of Li$_2$C$_2$ were suppressed, a high-pressure form predicted here would approach metallic behavior at pressures above 40 GPa.

II. METHODS

A. Experiments

All steps of sample preparation were performed in an Ar-filled glove box (H$_2$O and O$_2$ concentration <1 ppm). Starting materials for Li$_2$C$_2$ synthesis were lithium (ABCR, 99.99%) and graphite powder (Sigma-Aldrich, 99.9998%), which was degassed at 800 °C under dynamic vacuum for 24 h prior to use. Stoichiometric amounts of lithium and graphite were transferred into a purified Ta ampoule. Afterwards the ampoule was sealed in He atmosphere (800 mbar) and was placed inside a quartz ampoule, which was sealed under vacuum. The quartz ampoule was heated for 24 h at 1073 K in air (tube furnace), after which the sample was allowed to cool by turning off the furnace. An air- and moisture-sensitive fine powder with a light-gray color was obtained. The phase purity of the sample was checked by powder x-ray diffraction (PXRD, Huber G670, Cu Kα$_\alpha$ radiation, capillary). Apart from a small amount of unreacted graphite, no impurities were detected.

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In-situ high-pressure monochromatic PXRD experiments were performed with a membrane-driven diamond anvil cell (DAC) using a culet size of 400 microns. Powdered samples were loaded under inert gas atmosphere into a 150-μm-sized hole drilled in a stainless steel gasket. The pressure-transmitting medium (PTM) was helium. Diffraction data were collected at room temperature at the ID09 beamline of the ESRF using a MAR555 flat panel detector. The x-ray wavelength was λ = 0.41558 Å, and the beam diameter on the sample was set to 30 μm. In order to improve powder averaging, the DAC was rocked by ±3 deg. The pressure was monitored by the ruby luminescence method [11]. The two-dimensional diffraction data were integrated using the software FIT2D [12].

All diffractograms were inspected using the STOE WINXPOW software system [13]. DICVOL [14] within WINXPOW was used for indexing and ENDEAVOUR [15] for an ab-initio structural solution using a direct-space approach. Rietveld refinements were performed with GSAS [16]. More details of the structure solution and refinement are given in Sec. III.

B. Computations

Structure searches were carried out using the evolutionary algorithm USPEX [17–19]. The search over configurational space was constrained to structures containing C2 acetylide units. C-C bond connectivities were enforced using the Z-matrix representation [20] available in the ab-initio code SIESTA [21]. However, computationally demanding SIESTA was only used in the initial phase of a search as a means to quickly optimize the structure by constraining the molecular geometry and degrees of freedom of the C2 acetylide units. These calculations employed the Perdew-Burke-Ernzerhof (PBE) exchange correlation [22] as well as the single-ζ basis set. The plane-wave cutoff was set at 100 Ry, and a Monkhorst-Pack grid defined at a cutoff of 10 Å was used. The pseudopotentials used were Troullier and Martins norm-conserving pseudopotentials [23]. The final stages of a search were performed using the Vienna Ab Initio Simulation Package (VASP) [24]. The target pressure for searches was chosen to be 20 GPa. All populations contained 30 structures, and the initial population’s structures were randomly generated. All structures contained 16 atoms constrained to the chemical composition of Li2C2 (i.e., Z = 4).

Enthalpy versus pressure relations of Li2C2 phases were calculated using the first-principles all-electron projector augmented waves (PAW) method [25] as implemented in VASP. Exchange-correlation effects were treated within the generalized gradient approximation (GGA) using the PBE parametrization [22]. The structures were relaxed with respect to pressure, lattice parameters, and atomic positions. Forces were converged to better than 1 × 10−3 eV/Å. The integration over the Brillouin zone (BZ) was done on a grid of special k points of size 6 × 6 × 6, determined according to the Monkhorst-Pack scheme and using Gaussian smearing to determine the partial occupancies for each wave function [26]. The kinetic energy cutoff was set to 675 eV. To obtain the band structure and enthalpies, the tetrahedron method with Blöchl correction was employed for BZ integration [27]. Structure relaxations and phonon calculations were performed at pressures ranging from 0 to 40 GPa. Once a structure was relaxed at a target pressure, zone-centered phonon calculations were executed using VASP’s density functional perturbation theory approach.

III. RESULTS AND DISCUSSION

A. Experimental observations

The ground-state structure of Li2C2, Immm Li2C2, relates to the antifluorite structure. Li atoms are coordinated by four dumbbell units and each dumbbell unit by eight Li ions. When recording Raman spectra of Li2C2 in a DAC, it was consistently observed that Immm Li2C2 transforms reversibly at around 15 GPa into a high-pressure modification. This is shown in Fig. 1.

The retention of the dumbbell units is evidenced by the persistence of the acetylide C-C stretching vibration (A g). The stretching mode frequency drops discontinuously by about 20 cm−1 at the transition. At higher pressures the Raman spectra became featureless and remained featureless upon decompression. This phenomenon is attributed to irreversible amorphization of Li2C2 at high pressures [9]. In the Raman experiments no PTM was used in order to avoid any background scattering from possible surface contaminations. The nonhydrostatic pressure conditions do not appear to influence the transition into the high-pressure modification. However, pressures at which irreversible amorphization occurs varied between 17 and 25 GPa.

Figure 2 shows synchrotron PXRD patterns of Li2C2 across the phase transition and up to 24.7 GPa. Different from the Raman studies, pressure conditions here were hydrostatic. Below 16 GPa patterns correspond to Immm Li2C2. At 16.5 GPa additional reflections appear. The onset pressure of

FIG. 1. (a) Raman spectra of polycrystalline Li2C2 at different pressures, (R) = decompression, and (b) observed mode frequencies as a function of pressure from two experiments (black and white circles, respectively). The broken vertical line marks the transition pressure for a reversible structural transformation. No PTM was used in order to avoid scattering by sample surface contaminations. Li2C2 amorphizes irreversibly at pressures between 17 GPa (black circle experiment) and 24 GPa (white circle experiment, according to Ref. [9]).
To aid the structure elucidation, crystal structure searches by USPEX were performed at a target pressure of 20 GPa, well above the experimental transition pressure and below possible amorphization under hydrostatic conditions, respectively. Previous efforts using crystal-structure prediction methodology in the structure search for high-pressure Li$_2$C$_2$ have been restricted to simulation cells containing two formula units (i.e., eight atoms) [9]. This resulted in an energetically favorable structure (with Cmc$_{21}$ symmetry) for pressures above 15 GPa. However, calculated frequencies of Raman active modes for Cmc$_{21}$ Li$_2$C$_2$ deviated considerably from experiment. When extending the simulation cells to contain four formula units (16 atoms), as suggested by the diffraction experiments, the search indeed yielded a structure with Pnma symmetry. Figure 3 shows the enthalpy differences (with respect to the Immm ground-state structure) as a function of pressure for Pnma Li$_2$C$_2$ and earlier predicted Cmc$_{21}$ Li$_2$C$_2$. At pressures around 13 GPa the enthalpy of Pnma Li$_2$C$_2$ becomes lower than the ground state. This value for the transition pressure is slightly lower than the experimental observation. The minor discrepancy may be attributed to the negligence of zero-point-energy contributions and temperature effects in our calculations. Importantly, Pnma Li$_2$C$_2$ is dynamically stable in the pressure range 10–30 GPa (see Supplemental Material [29], Fig. S1). The structural parameters at 20 GPa are compiled in Table I. Additionally, Ref. [29] contains parameters for the relaxed structures of Immm and Pnma Li$_2$C$_2$ for the complete investigated pressure range 0–40 GPa (Tables S1 and S2).

For Rietveld refinement the structural parameters of the model obtained with USPEX were used as starting parameters. The refined parameters for Pnma Li$_2$C$_2$ at 18.7 GPa are given in Table II. Details of the measurement and the refinement
are summarized in Table S3 [29]. In Table III interatomic distances are compared with those of the computed structure at 20 GPa. The refinement of the Li atom positions was quite unstable and led to a few short Li-Li distances. This can be attributed to the modest data quality and the strong overlap of reflections from coexisting Immm Li2C2. The reduced data quality could be a consequence of Li disorder, connected to the occurrence of an intermediate phase between Immm and Pnma Li2C2. Such an intermediate phase has been identified for the high-pressure-phase transition of Li2S [30] which, as we will discuss later, relates closely to that of Li2C2. Also, an intermediate phase with varying Li disorder might explain from our diffraction data. The final fit of the 18.7 GPa data is shown in Fig. 4. Differences between the calculated and measured profiles (in particular, extra sharp reflections) can mainly be attributed to ruby and solid helium. Attempts to improve the fit by applying parameters for stress, strain, or anisotropic peak broadening gave unstable refinements and did not lead to physically meaningful results. Only the refinement of preferred orientation parameters (March-Dollase) gave a significant improvement of the fit. In Table S4 we also present the results from Rietveld refinements of the data at 7.2, 18.1, 18.7, and 19.3 GPa, respectively [29].

Figure 5 shows the pressure-volume (p-V) relations of Immm and Pnma Li2C2. Unit-cell parameters as a function of pressure from diffraction data are given in Tables S5 and S6 [29]. Both experimental and calculated p-V data were fitted to a three-parameter Birch-Murnaghan equation of state (EOS) expression [31]. Generally there is good agreement between calculated and experimentally determined p-V data. Computed volumes are somewhat underestimated, by 2%–2.5%. The first-order phase transition from Immm to Pnma Li2C2 is accompanied by a 7% volume reduction. The fitted EOS parameters are presented in Table IV. For Immm Li2C2 computed and experimental p-V data give virtually identical parameters. The ambient-pressure bulk modulus of this phase is around 40 GPa. For Pnma Li2C2 the bulk modulus extracted from the experimental data is 112 GPa at the reference pressure \( p_r = 16.5 \) GPa (\( V_r = 34.5 \) Å\(^3\)).

C. Cmcm Li2C2 and structural relationships

The high-pressure-phase Pnma Li2C2 amorphizes irreversibly in room temperature Raman experiments at \(~17\) GPa (nonhydrostatic) but persists up to at least 25 GPa under hydrostatic conditions. Computationally, Pnma Li2C2 may be further compressed. Interestingly, as shown in Fig. 6, at around 32 GPa this yields a transition into another structure. The new structure is topologically equivalent to Pnma Li2C2 but adopts the higher-symmetry space group Cmcm. The structural parameters for Cmcm Li2C2 and their variation with pressure are compiled in Table S7 in Ref. [29]; EOS parameters are included in Table IV.

Figure 7 depicts the structural relations between ground-state Immm Li2C2 and the Pnma and Cmcm high-pressure

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**TABLE I. Structural parameters of Pnma Li2C2 at 20 GPa (DFT optimized).**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Exp. structure (18.7 GPa)</th>
<th>Comp. structure (20 GPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li2–Li</td>
<td>1.86–2.68 Å (4×)</td>
<td>2.39–2.62 Å (4×)</td>
</tr>
<tr>
<td>Li2–C</td>
<td>1.97 Å (2×), 2.14 Å (2×)</td>
<td>2.05 Å (2), 2.15 Å (2×)</td>
</tr>
<tr>
<td>C–C</td>
<td>2.62 Å (2×), 2.33 Å (2×)</td>
<td>2.19 Å (2×), 2.22 Å (2×)</td>
</tr>
<tr>
<td>C–Li</td>
<td>1.97–2.55 (7×)</td>
<td>1.96–2.52 (7×)</td>
</tr>
</tbody>
</table>

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**TABLE II. Structural parameters of Pnma Li2C2 at 18.7 GPa (Rietveld refinement).**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>a (Å)</th>
<th>b (Å)</th>
<th>c (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li2 (4c)</td>
<td>5.098(2)</td>
<td>4.505(1)</td>
<td>5.909(2)</td>
</tr>
<tr>
<td>Li2 (4c)</td>
<td>0.144(4)</td>
<td>0.25</td>
<td>0.938(3)</td>
</tr>
<tr>
<td>Li2 (4c)</td>
<td>0.999(5)</td>
<td>0.25</td>
<td>0.227(5)</td>
</tr>
<tr>
<td>C (8d)</td>
<td>0.742(1)</td>
<td>0.1163(3)</td>
<td>0.9100(6)</td>
</tr>
</tbody>
</table>
phases. As mentioned earlier, the *Immm* structure relates to the antifluorite type: $C_2$ dumbbells are arranged as a quasi-cubic close packing in which Li atoms occupy the tetrahedral voids. Consequently, each $C_2$ dumbbell is surrounded by eight Li atoms and each Li atom by four dumbbell units. Alternatively, the *Immm* structure can be viewed as a stacking of layers consisting of planar, rectangular nets formed by the Li ions which are stuffed by $C_2$ dumbbells oriented perpendicularly. Layers are stacked along the $b$ direction (which is the elongation direction of dumbbells) and consecutive layers $A$ and $B$ are related by the $I_2$ centering.

Also within *Pnma* $Li_2C_2$ Li ions form planar nets (parallel to the $ac$ plane) that consist of triangle ribbons running along the $a$ direction. Interatomic distances within triangles are short compared to distances in between (2.5 Å vs 3.1 Å at 20 GPa). Planar Li nets are completed to layers by perpendicularly oriented $C_2$ dumbbells interspersed between triangle ribbons. In the *Pnma* structure consecutive layers $A$ and $B$ are stacked in a way that $C_2$ dumbbells (e.g., in a layer $A$) attain a trigonal prismatic coordination by two triangles from adjacent layers above and below (layers $B$). A dumbbell is coordinated additionally by three Li ions which are situated in the same layer and cap the rectangular faces of the trigonal prism.

**TABLE IV.** Equation-of-state parameters for phases of $Li_2C_2$. Note that the experimental results for *Pnma* $Li_2C_2$ refer to a reference pressure of 16.5 GPa, not zero pressure.

<table>
<thead>
<tr>
<th>$Li_2C_2$</th>
<th>$V_0$ (Å$^3$)</th>
<th>$K_0$ (GPa)</th>
<th>$K_0'$</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Immm</em> exp</td>
<td>47.9</td>
<td>39(1)</td>
<td>3.9(2)</td>
</tr>
<tr>
<td><em>Pnma</em> exp</td>
<td>$V_0 = 34.5$</td>
<td>$K_0 = 112(5)$</td>
<td>4 (fixed)</td>
</tr>
<tr>
<td><em>Immm</em> calc</td>
<td>46.7</td>
<td>40.8</td>
<td>3.9</td>
</tr>
<tr>
<td><em>Pnma</em> calc</td>
<td>44.13</td>
<td>34.7</td>
<td>4.9</td>
</tr>
<tr>
<td><em>Cmcm</em> calc</td>
<td>42.71</td>
<td>38.7</td>
<td>4.3</td>
</tr>
</tbody>
</table>

Consequently, compared to *Immm* $Li_2C_2$ the coordination of a dumbbell by Li ions is increased to 9.

As *Immm* $Li_2C_2$ relates to the antifluorite type so does the *Pnma* structure to the anticotunnite type. *Pnma* $Li_2C_2$ is isostuctural to the recently discovered ternary carbides CsK$C_2$ and CsRb$C_2$ [32], and antifluorite–anticotunnite transitions are frequently observed for alkali metal chalcogenides $A_2B$ at high pressures. In particular, $Li_2O$ and $Li_2S$ display this transition at around 45 and 12 GPa, respectively [30,33]. For $Na_2S$ the antifluorite ground-state structure transforms to the anticotunnite structure at even lower pressures, at around 7 GPa. At about 16 GPa another transition takes place which results in a phase with the Ni$_2$In-type structure [34].

Interestingly, the sequence antifluorite $\rightarrow$ anticotunnite $\rightarrow$ Ni$_2$In type is also shown by $Li_2C_2$ as *Cmcm* $Li_2C_2$ relates to the hexagonal Ni$_2$In structure. The topology of planar Li-ion nets is identical in *Pnma* and *Cmcm* $Li_2C_2$. However, in the higher-symmetry *Cmcm* structure ribbons are straightened into distinct zigzag chains in which triangles are strictly oriented up and down [cf. Fig. 7(b)]. These chains run along the $c$ direction. The orientation of triangles from adjacent chains yields five-membered rings which are centered by the dumbbell units. Because the trigonal prismatic environment of a dumbbell by Li triangles situated in layers above and below is maintained, its total coordination by Li ions is increased to 11 with respect to the *Pnma* structure. The coordination polyhedron corresponds to an Edshummar polyhedron, which is the signature of the Ni$_2$In structure type [35].

To conclude the discussion of structural relationships, we address the evolution of interatomic distances with pressure [referring to the density-functional theory (DFT) optimized structures]. The C-C distance within dumbbell units is only slightly compressible. Within the *Immm* structure this distance reduces from 1.256 Å at ambient pressure to 1.239 Å at 40 GPa. This is similar for the high-pressure forms. Here this
FIG. 7. (Color online) (a) Crystal structures of Immm, Pnma, and Cmcm Li$_2$C$_2$ represented as layers consisting of planar nets formed by Li ions, which are centered by perpendicularly oriented dumbbell units. Li ions are shown as light gray circles and C atoms as red circles. Layers are stacked with an AB sequence in the dumbbell direction, as described in the text. (b) View of the structures along the layer stacking direction. A-type layers are distinguished by bold lines. B-type layers by thin lines and pale colors. (c) Coordination of C$_2$ dumbbells within the three phases. The numbers indicate the distances between the dumbbell center and surrounding Li ions in Å (referring to DFT optimized structures at 20 GPa).

distance decreases from 1.254 Å at 10 GPa to 1.244 Å at 40 GPa. The Li-Li distances defining the coordination around C$_2$ dumbbells are 2.55, 2.81, and 3.02 Å for the Immm structure at ambient pressure. They reduce to 2.35, 2.58, and 2.84 Å at 14 GPa, which is close to the calculated transition pressure. At this pressure the corresponding Li-Li distances in the Pnma structure are between 2.46 and 3.17 Å.

D. Electronic structure changes with pressure

The band structures of Immm Li$_2$C$_2$ and Pnma/Cmcm Li$_2$C$_2$ are shown in Fig. 8. At pressures below 10 GPa both the ambient- and high-pressure forms exhibit insulating properties. At ambient pressure Immm Li$_2$C$_2$ has an indirect band gap of 3.3 eV, with the bottom of the conduction band at Γ and the top of the valence band at T. The valence bands mirror the molecular orbital (MO) diagram of the acetylide anion. Their topology for Li$_2$C$_2$ is similar to CaC$_2$, whose electronic structure has been studied earlier [36,37]. The weakly dispersed band centered at −12 eV below the Fermi level corresponds to the sp$_{σ}$ bonding MO. Bands corresponding to the two lone-pair states (sp$_{σ_u}$ and sp$_{σ_g}$) have dispersions of about 2 eV and are located in the range −4 to −1 eV below the Fermi level. The two π-bonding bands constitute the top of the valence band. It is clearly seen that pressure increases especially in the lone-pair–Li interactions because the dispersion of lone-pair–based bands increases most. The pressure dependence of the DFT-GGA–computed band gap is shown in Fig. 9. It decreases linearly, but Immm Li$_2$C$_2$ obviously stays insulating.

At low pressure (below 10 GPa) Pnma Li$_2$C$_2$ exhibits an indirect band gap of <2.5 eV with the bottom of the
STRUCTURAL TRANSFORMATIONS OF Li$_2$C$_2$ AT . . . PHYSICAL REVIEW B 92, 064111 (2015)

conduction band at $\Gamma$ and the top of the valence band lying along $T$-$Y$. The band gap of Pnma Li$_2$C$_2$ diminishes faster with pressure compared to the Immm structure. In high-pressure Raman experiments a darkening of the sample is observed after the Immm-to-Pnma phase transition [9]. This possibly relates to the considerably decreased band gap of Pnma Li$_2$C$_2$. At 35 GPa the Pnma structure merged into the Cmcm structure. At this pressure the calculated band gap dropped below 0.5 eV. Above 40 GPa the band gap of Cmcm Li$_2$C$_2$ has closed. The comparatively low pressure for (hypothetical) metallization of an ionic structure is remarkable. The changed pressure dependence of the band gap for the high-pressure phases of an ionic structure is remarkable. The changed pressure compared to the rock salt type for (hypothetical) metallization it would transform at around 32 GPa to a higher-symmetry Cmcm structure that is closely related to the Ni$_2$In type. Cmcm-Li$_2$C$_2$ would metallize at pressures above 40 GPa as a result of indirect band overlap. The sequence antifluorite $\rightarrow$ anticotunnite $\rightarrow$ Ni$_2$In-type mirrors a common trend of high-pressure-phase transitions in A$_2$B compounds toward higher coordination.

We point out that the high-pressure behavior of the acetylide carbides Li$_2$C$_2$ and MC$_2$ ($M = \text{Ca, Sr, Ba}$) appears strikingly similar to the corresponding sulfides. Experimental and/or calculated transition pressures for the sequences antifluorite $\rightarrow$ anticotunnite $\rightarrow$ Ni$_2$In-type (referring to Li$_2$C$_2$/Li$_2$S) and rock salt $\rightarrow$ CsCl-type (referring to MC$_2$/MS) are remarkably close [38–43]. This may be attributed to a similar polarizability of the C$_2^2-$ and S$^2-$ anions. However, unlike sulfides, acetylides will undergo amorphization with pressure and expected phase transitions may not be observed. The origin of the pressure-induced amorphization of acetylides is uncertain, and different scenarios can be envisioned. For example, amorphization could indicate compositional instability and phase segregation, which appears to be the case for BaC$_2$ [5]. Furthermore, it could connect with a pressure limit for stability of multiple bonded light element entities, as suggested in Ref. [7]. However, the enthalpic pressure limit for the stability of C$_2^2-$ units is rather low, as computations show clearly that with pressure carbides with polymeric anions become rapidly favored over acetylides. Specifically, for Li$_2$C$_2$ a phase with the CrB structure becomes more stable than Immm Li$_2$C$_2$ at already 5 GPa [2,3] This is far below the experimentally observed Immm-to-Pnma phase transition (see also Fig. S2 in Ref. [29]) and it has been concluded that acetylides are distinguished by a high kinetic stability [3]. The elucidation of the origin of the kinetic stability and pressure-induced amorphization of acetylides will require the analysis of the composition and local structure of the amorphous carbides by, e.g., synchrotron extended x-ray absorption fine structure (EXAFS) and/or total scattering experiments, preferably in combination with molecular dynamics simulations.

IV. CONCLUSIONS

In summary, we have employed a combination of synchrotron x-ray diffraction experiments and computational evolutionary search methodology to elucidate the high-pressure crystal structure of the acetylide carbide Li$_2$C$_2$. The observed high-pressure phase has Pnma symmetry and relates to the anticotunnite structure (Z = 4). In hydrostatic experiments Pnma Li$_2$C$_2$ does not amorphize under pressures up to 25 GPa. We find that if Pnma Li$_2$C$_2$ were prevented from amorphization it would transform at around 32 GPa to a higher-symmetry Cmcm structure that is closely related to the Ni$_2$In type. Cmcm-Li$_2$C$_2$ would metallize at pressures above 40 GPa as a result of indirect band overlap. The sequence antifluorite $\rightarrow$ anticotunnite $\rightarrow$ Ni$_2$In-type mirrors a common trend of high-pressure-phase transitions in A$_2$B compounds toward higher coordination.

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FIG. 9. (Color online) Band gap–pressure relations of the ground-state and high-pressure phases of Li$_2$C$_2$. Figure text: acetylide lone pairs are coordinated by triangles of Li ions. With pressure this coordination will develop into a covalent bonding interaction between C and Li, formally corresponding to electron donation from the dumbbell C$_2^2-$ to Li$^+$, and eventually leading to metallization.