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Nanosecond formation of diamond and lonsdaleite by shock compression of graphite

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The shock-induced transition from graphite to diamond has been of great scientific and technological interest since the discovery of microscopic diamonds in remnants of explosively driven graphite. Furthermore, shock synthesis of diamond and lonsdaleite, a speculative hexagonal carbon polymorph with unique hardness, is expected to happen during violent meteor impacts. Here, we show unprecedented *in situ* X-ray diffraction measurements of diamond formation on nanosecond timescales by shock compression of pyrolytic as well as polycrystalline graphite to pressures from 19 GPa up to 228 GPa. While we observe the transition to diamond starting at 50 GPa for both pyrolytic and polycrystalline graphite, we also record the direct formation of lonsdaleite above 170 GPa for pyrolytic samples only. Our experiment provides new insights into the processes of the shock-induced transition from graphite to diamond and uniquely resolves the dynamics that explain the main natural occurrence of the lonsdaleite crystal structure being close to meteor impact sites.

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he formation of diamond after applying pressure and heat on graphite is highly relevant to the artificial synthesis of diamond¹⁻⁴, and also for a general understanding of high pressure phase transitions in environments comparable to meteor impacts or planetary interiors^{5,6}. In 1967, a hexagonal form of diamond⁷, later called lonsdaleite⁸, in its pure form supposedly harder than diamond⁹, was identified for the first time inside fragments of the Canvon Diablo meteorite, the asteroid which created the Barringer Crater with an impact velocity of $\sim 10 \,\mathrm{km \, s^{-1}}$ generating pressures exceeding 200 GPa (ref. 5). Since then, occurrences of lonsdaleite and also nanometer-sized diamonds have been suggested to serve as marker for meteor impacts all over the world, for example, the Tunguska explosion in the early twentieth century¹⁰, the Ries crater¹¹, the Younger Dryas event¹² or the impact which has been correlated to the extinction of the dinosaurs¹³. However, due to difficulties in creating a pure lonsdaleite phase under static compression conditions, serious doubts on the overall existence of lonsdaleite were published recently^{2,14}. Yet, static compression is not well suited to mimic the fast processes during impact events.

Dynamic shock compression experiments, driven by explo-sives^{15,16}, gas guns^{4,17} or pulsed high-energy lasers¹⁸, are much closer to such conditions despite the smaller dimensions and thus shorter timescales (nanoseconds to microseconds) compared with meteor impacts (seconds). For this reason, laboratory shock experiments are limited to detect phase transitions happening on corresponding timescales and cannot explore some complex structural changes, which are thought to require nucleation times up to seconds¹⁹. The shock-induced transition from graphite to diamond is thought to be a fast martensitic transition⁴, but up to now, shock compression experiments on graphite have lacked a direct in situ measurement of the microscopic crystal structure. In fact, all conclusions regarding structural transitions were based on studies from the material that was recovered after applying the shock drive or dynamic measurement of macroscopic quantities, such as density and pressure. As a consequence, there is a speculation in understanding the dynamic processes for shock synthesis of diamond and lonsdaleite $^{20-27}$ and conclusions regarding meteor impacts^{2,5}. Particularly, the different concepts involve nucleation²⁴, plane-sliding and buckling²², wave-buckling and slipping²⁷ or the intermediate formation of so-called layered diamond²⁰, which is a mixture of hexagonal and cubic diamond²³. These models give very different results under which conditions, or if at all, diamond or lonsdaleite are formed from graphite. Moreover, while a fast martensitic process on nanosecond timescale has been accepted for compressing highly oriented graphite samples along the c axis^{4,17}, much slower diffusion-based mechanisms are thought to be more effective for highly defective graphite samples²⁸.

In the following, we address these questions in an unprecedented way by applying ultrafast X-ray diffraction for *in situ* structure measurements of the shock-induced transition from graphite to diamond. Our results show nanosecond formation of the diamond structure after shock compressing pyrolytic as well as porous polycrystalline graphite samples. At the highest pressures, we find evidence for a relatively pure lonsdaleite structure when compressing pyrolytic graphite, whereas the porous polycrystalline samples are transferred to a complex liquid in this regime.

Results

Experimental method. The experiments reported here were performed at the Matter at Extreme Conditions endstation of the Linac Coherent Light Source of Stanford National Accelerator Laboratory²⁹. Here, the newly available combination of high-energy lasers ($\gtrsim 10$ J per pulse) with an X-ray free electron laser

allows for femtosecond X-ray diffraction to directly access the kinetics of the shock-induced transition. A schematic of the experiment is shown in Fig. 1.

The recorded diffraction angle θ is directly connected to the lattice spacing *d* via the Bragg relation

$$\sin(\theta) = \frac{n\lambda}{2d},\tag{1}$$

where *n* is the diffraction order, and λ the wavelength of the incident light source. For the symmetric cubic diamond crystal structure, the lattice spacing given by the (111) diffraction peak directly determines the density $\rho \propto d^{-3}$. Knowing the density of the shocked state ρ_1 as well as the shock transit time, and thus the shock velocity $v_{\rm S}$, allows for inferring the pressure *p* using the Rankine–Hugoniot relations, which simply apply mass, momentum and energy conservation³⁰

$$p = v_{\rm S}^2 \rho_0 \frac{\rho_1 - \rho_0}{\rho_1}.$$
 (2)

Here, ρ_0 is the initial density of the graphite sample.

Pyrolytic graphite. Figure 2 shows X-ray diffraction results for oriented pyrolytic graphite ($\rho_0 = 2.21 \text{ g cm}^{-3}$) driven parallel to the graphite c axis with different drive pressures. For $\sim 20 \text{ GPa}$, we observe the (002) Bragg reflection shifting to larger diffraction angles, thus providing a signature of compressed graphite. For pressures above ~ 55 GPa, the graphite (002) signature has completely vanished, whereas a broad peak at ~ 60 degrees appears. This new feature is in agreement with the (111) Bragg reflection of diamond. The large width of the diffraction peak at this pressure most probably documents remaining disorder (that is, very small diamond crystallites) and thus implies that the transition from graphite to diamond is not fully completed within the shock transit time of ~ 9 ns. Higher drive pressures result in faster transition times, reducing the width of the (111) diamond diffraction peak due to less disorder, that is, creation of larger diamond crystallites. For drive pressures above 100 GPa, we observe the formation of diamond on timescales less than 1 ns (inset in Fig. 2d). For these pyrolytic samples at shock pressures higher than 170 GPa, we see two diffraction ring fragments



Figure 1 | Schematic of the experimental setup at the Matter at Extreme Conditions endstation of the Linac Coherent Light Source. Two highenergy laser beams drive shock waves into graphite samples generating pressures from 20 to 230 GPa. The shock transit times of few nanoseconds are recorded by a VISAR system, which detects the shock-induced reflectivity drop of a 100-nm thick aluminum coating when the shock exits on the target rear side. The microscopic state is probed by a single X-ray pulse with 6 keV photon energy and 50 fs pulse duration. X-ray diffraction is recorded by a large area X-ray detector.



Figure 2 | Transition from pyrolytic graphite to diamond. X-ray diffraction data of (**a**) cold and (**b**-**d**) compressed pyrolytic graphite samples, driven parallel to the graphite *c* axis; t_0 is denoting the start of the drive laser pulse. For 19 GPa, diffraction shows graphite, which is mainly compressed along the *c* axis, together with some cold material from the rear side of the sample not reached by the shock. For 60 GPa at exactly the moment when VISAR records the shock having traversed the whole sample, every signature from graphite vanishes. In this case, only a broad cubic diamond (111) diffraction peak remains. For higher pressures (for example, 122 GPa), the width of the peak decreases due to the reduced transition time and we observe the formation of a sharp Bragg reflection within 1 ns.

emerging that are clearly caused by different crystal planes showing some degree of texture (see Fig. 3), conserving the preferred crystallite orientation of the pyrolytic samples (see Supplementary Figs 1 and 2). The inferred spacing is compatible with diffraction from the (010) and (002) planes of lonsdaleite. For this structure, we obtain an average c/a lattice constant ratio of ~1.69, which is slightly above the ideal value of $\sqrt{8/3}$ for a wurtzite lattice with equidistant atoms. This reasonably follows the trend given by reported values for ambient conditions⁸ (c/a = 1.64) and simulations for 20 GPa (ref. 23; c/a = 1.66). Density functional theory coupled with molecular dynamics (DFT-MD) simulations of the lonsdaleite structure at 4.4 g cm⁻³ and 6,000 K, resulting in a pressure of 205 GPa, show a very good agreement with our diffraction measurements.

The observed diffraction pattern is only compatible with lonsdaleite and not with any other structure that has been predicted for this pressure regime. Faulted and twinned cubic diamond, as shown by Nemeth *et al.*², would result in a cubic diamond main peak with small side peaks due to the defects. Furthermore, a mix of cubic and hexagonal diamond, the



Figure 3 | **Diffraction from lonsdaleite.** When compressing pyrolytic graphite to pressures above 170 GPa, we observe two strong diffraction peaks consistent with lonsdaleite (010) and (002) planes. The grey-dashed line shows a DFT-MD simulation of the lonsdaleite phase at $4.4 \,\mathrm{g\,cm^{-3}}$ and 6,000 K, resulting in a pressure of 205 GPa, which is in qualitative agreement with the observed diffraction pattern. As the range of detectable diffraction angles ends at $2\theta = 68$ degrees in our experiments, the (011) peak at $2\theta = 70.5$ degrees is not recorded. The occurrence of partial diffraction rings shows a preferred orientation of the crystallites that is compatible with the preferred orientation of the initial pyrolytic graphite sample.

so-called layered diamond struture, would not result in the observed clear doublet of peaks²⁰. Other more complicated polymorphs show many more diffraction maxima²³ and can certainly be excluded by our measurements. Hexagonal lonsdaleite being favoured at higher drive pressures is conceivable since the direct transition path to cubic diamond requires more lattice distortions^{22,24,26}. Recent MD calculations²⁷ demonstrate that plane sliding of the graphite basal planes to rhombohedral graphite, which in turn would lead to the formation of diamond, is prevented by the emergence of few interplanar sp³ bonds, when the spacing between the graphite basal planes falls below a certain distance. For this case, these simulations predict the formation of lonsdaleite²⁷. Our experiments indicate that this limit is reached by dynamic shock compression of pyrolytic graphite above 170 GPa.

Figure 4 combines the recorded microscopic crystal structure with the macroscopic quantities density and pressure. Applying the simple relation between Bragg angle and lattice spacing, and thus density, in combination with equation (2), we can create an experimental pressure-density diagram, giving a line of states achievable by shock compression, the shock Hugoniot curve (all experimental values including error estimates can be found in Supplementary Tables 1 and 2). Classical Hugoniot measurements usually determine only macroscopic quantities like density and pressure via velocity measurements. As such, the shock Hugoniot is one of few experimentally accessible quantities capable of benchmarking high-pressure equation of state theories, from which, for example, the temperature of the created states can be deduced. Only based on these models, it is then possible to draw conclusions on microscopic properties. In contrast, our measurements directly record the microscopic structure of the sample in addition to determining density and pressure. Thus, they give a more complete picture joining both microscopic and macroscopic observations to an absolute equation of state measurement.

For compressed pyrolytic graphite, our results connect and overlap with established literature data at lower pressures^{15–17,31,32} as well as first principles simulations of the carbon phase diagram³³ (see Fig. 4a). At higher pressures, our data extend to a regime which has not been well characterized before. For states which remain in the graphite phase, it needs to be considered that graphite mainly compresses along the *c* axis of the unit cell³⁴.



Figure 4 | Summary of the experimental results. (a) Recorded pressure-density diagram for compressed pyrolytic graphite ($\rho_0 = 2.21 \text{ g cm}^{-3}$) compared with literature data without structure information and a first principles phase diagram³³. At lower pressures, there is very good agreement, whereas at higher pressures, due to the formation of lonsdaleite, we observe higher densities than predicted by a proposed shock Hugoniot³¹, which instead suggests a transition to the liquid. (b) Recorded pressure-density diagram for porous polycrystalline graphite ($\rho_0 = 1.84 \text{ g cm}^{-3}$) compared with experiments without structure information^{16,32} and a first principles phase diagram. Comparable to the pyrolytic samples, the diamond formation is not fully completed within few nanoseconds at lower pressures, resulting in a broad peak which sharpens up to ~100 GPa. At higher pressures, however, the increasing temperature leads to melting of the diamond structure, resulting in broader and fainter diffraction peaks in agreement with a bonded liquid¹⁸. No signature of lonsdaleite is observed when compressing porous graphite.

The (002) diffraction peak position directly determines the lattice spacing d_c along the c axis and using the relation $\rho \propto d_c^{1.14}$ from static compression experiments³⁵, we can infer the density of the compressed graphite states. The rearrangement in the microscopic structure, that is, the graphite-diamond transition, is clearly visible due to a density jump at the transition pressure of \sim 50 GPa and a lower compressibility of the diamond states. We do not observe a two-wave structure as, for example, Erskine and Nellis¹⁷ did for this transition after shock compression of pyrolytic graphite samples with a higher grade of orientation. These experiments achieved the transition already at 27 GPa, while the two-wave structure was overdriven at 50 GPa. In agreement with our experiments, Erskine and Nellis did not record a two-wave structure for graphite samples with lower grade of orientation. At higher pressures, we observe a second smaller jump in density at \sim 170 GPa, pointing at the transition to lonsdaleite consistent with a slightly increased energy that is required for the structural transition compared with diamond³⁶. In turn, we obtain higher densities than predicted by a proposed shock Hugoniot³¹ that does not take the formation of lonsdaleite into account in this regime. However, this phase exhibits even higher stiffness, supporting predictions that the hardness of cubic diamond is exceeded by its hexagonal polymorph⁹ and possibly leading to a re-connection with the proposed Hugoniot line in the liquid phase. Using a very recent multi-phase equation of state model for carbon³⁷, we expect the temperatures of the states, which remain in the graphite phase in the order of 4,000 K, 4,000-6,000 K for the diamond data and 6,000-7,000 K for the measurements, which show the lonsdaleite structure.

Porous polycrystalline graphite. For porous polycrystalline samples, we obtain slightly different results (see Fig. 4b). When compressing porous samples, the voids close instantly nearly without resistance, resulting in an entropy increase. Further compression then leads to states of higher entropy and thus temperature compared with shock compression of the higher density pyrolytic samples to the same pressure³⁰. For porous

polycrystalline graphite, we record nanosecond formation of diamond with a transition threshold at slightly lower pressure compared with the pyrolytic samples. The width of the diffraction peak is marginally larger than for pyrolytic samples. This indicates that only slightly smaller diamond crystallites are formed for the polycrystalline samples. Starting from ~ 100 GPa, the diffraction again broadens and weakens due to melting and, instead of reaching the hexagonal phase, a bonded liquid is formed, which is in good agreement with the first principles phase diagram. It has been shown that short-time bonding remains active in liquid carbon, probably up to temperatures of more than 20,000 K (ref. 38). As the liquid structure is diamond-like, the diffraction signatures do not immediately vanish or shift above the melting line. Again using a recent multiphase equation of state model³⁷, we expect the temperatures of the states which remain in the graphite phase in the order of 4,000-5,000 K, 5,000-8,000 K for the diamond data and temperatures exceeding 8,000 K for the data points, which show the transition to the liquid.

Discussion

The data of dynamic diamond formation presented here resolve controversies on the shock-induced graphite to diamond transition: a fast transition from graphite to diamond on nanosecond timescale does not require highly oriented graphite samples and shock loading perpendicular to the c axis. Furthermore, recording the first clear diffraction signal of the meta-stable hexagonal diamond polymorph, that is not obscured by dominant cubic diamond or graphite, demonstrates that this state exists. Finally, observing lonsdaleite only for pyrolytic graphite samples compressed along the c axis to very high pressures suggests that sliding of the graphite basal planes and thus diamond formation can be prevented under these conditions. Hence, the picture that lonsdaleite synthesis is highly difficult applying static pressure but can be formed in violent impact events is strongly supported.

Methods

Laser-driven shock compression. The graphite samples (pyrolytic graphite, GoodFellow C000720, $\rho_0 = 2.21$ g cm⁻³, mosaicity ~10 degrees and porous polycrystalline graphite, SGL Carbon R6650, $\rho_0 = 1.84 \,\mathrm{g \, cm^{-3}}$, 10% volume porosity, 5 µm average grain size) were compressed using two pulsed high-energy lasers (527 nm, up to 16 J per beam, 10 ns pulse duration, 150–200 µm focal spot diameter, smoothed with random phase plates). The samples were laser-cut out of larger graphite pieces and then polished to achieve a surface roughness of 0.1 µm. The sample thickness was varied between 60 and 100 µm and had individually been characterized for each sample with an accuracy of $\sim 0.5 \,\mu\text{m}$. The shock transit time was recorded with a VISAR system, which monitored the drop in reflectivity of a 100 nm aluminium coating on the sample rear side at the moment of shock release. The VISAR system at MEC has two streak cameras and the transit time was determined as the average of the two cameras. The arrival time of the drive laser was determined by the same streak cameras as used for the VISAR system by scattering light of the drive laser beams from the exact position of sample interaction into the VISAR system. Typical transit times varied between 6 and 12 ns, depending on sample thickness and drive pressure. Recording transit times for different sample thickness and using the spatial resolution of the VISAR, sufficient shock steadiness as well as shock planarity could be guaranteed for the region that was irradiated by the X-ray beam. This was confirmed by measurements using graphite samples with a LiF window on the rear side, where a very steady interface velocity could be monitored. Using this interface velocity and the shock velocity in graphite, as well as assuming the LiF equation of state to be well known, density and pressure in the shocked graphite sample can be inferred in principle. These results are in agreement with the X-ray diffraction method, but by far not as accurate for our experiments (particularly in terms of density) and cannot provide additional insights.

Ultrafast X-ray diffraction. The graphite samples were probed using the LCLS XFEL beam in SASE mode (5.9–6.1 keV photon energy, 0.3% spectral bandwidth, 50 fs pulse duration, 20 µm spot size, ~3 mJ per pulse corresponding to 3×10^{12} photons per pulse). The input spectrum was recorded for each run with a high-resolution silicon crystal spectrometer. The timing of the X-ray pulse relative to the drive laser was chosen to probe the sample slightly before or exactly at the moment when the shock wave had traversed the whole sample. X-ray diffraction was measured by an 8×8 cm² Cornell-Stanford Pixel Array detector (CSPAD) at 11 cm distance to the sample covering diffraction angles from 27 to 68 degrees on top of the X-FaL beam axis. In this way, interfering effects due to the horizontal polarization of the X-ray beam could be avoided. Angular calibration for the recorded 2θ range of 20–68 degrees was obtained by diffraction of a LaB₆ powder sample.

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Author contributions

D.K., R.W.F., B.N., H.J.L., T.D., S.H.G., G.S., D.O.G., J.V., G.G., P.N. and M.R. were involved in the project planning. D.K., A.R., M.G., S.F., J.H., L.B.F., B.N., B. Barbrel, B. Bachmann, E.J.G., S.G., E.G., H.J.L., W.S. and T.D. carried out the experiment. G.S., J.H., S.F., M.R. and D.K. designed and built the samples. Experimental data were analysed and discussed by D.K., S.H.G., A.R., M.G., D.O.G., J.V. and T.D. The manuscript was written by D.K., S.H.G. and D.O.G.

Additional information

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