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OPEN Synthesis of diamond-like phase **from graphite by ultrafast laser driven dynamical compression**

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Rapid variations of the environmental energy caused by ultrashort laser pulses have induced phase transitions in carbon allotropes, therefore bringing the promise of revealing new carbon phases. Here, by exposing polycrystalline graphite to 25 fs laser pulses at 4J/cm2 fluence under standard air atmosphere, we demonstrated the synthesis of translucent micrometer-sized structures carrying diamond-like and onion-like carbon phases. Texturized domains of the diamond phase were also identified. Concerning different synthesized carbon forms, pulse superposition and singularities of the thermodynamical process, we pinpoint the synthesis mechanism by the laser-induced subsequent products energetically evolving to attain the diamond-like phase.

In recent years carbon allotropes, such as nanodiamonds, have shown promising new applications in many fields due to its physical, chemical and surface characteristics. Their high electron mobility, field electron emission and magnetic properties^{[1](#page-4-0)} make them important players in carbon based electronics^{2,[3](#page-4-2)}. Their tribological and mechanical^{[1](#page-4-0)[,4](#page-4-3),5} properties give rise to harder coatings^{1,5}, which are biocompatible and can provide improved biological prosthesis joints^{[6](#page-4-5)} with decreased wear. This biocompatibility allied to their biosensing, optical and nanoprobing functionalities^{[7,](#page-4-6)[8](#page-4-7)} provide drug delivery and cellular labeling capacity[7](#page-4-6) . This wide range of novel applications has fostered the active search for new and more efficient synthesis and production methods of carbon allotropes.

In this pursuit, it has been shown that dynamical compression is a way to steer carbon allotropes, through rapid energy variations, to phase transitions. As examples, transition from graphite to diamond headed by shock compression experiments occurs through the fast martensitic mechanism⁹; in a setup designed for quenching shock-compressed samples at a fast cooling rate, the n-diamond phase¹⁰ has been created, and in regions of shock impacted meteorites nanodiamonds have been formed¹¹. These observations brought to light the importance of carbon phase transformations in non-equilibrium states, spurring studies in a faster time scale using ultrashort pulses, which trigger shockwaves[12,](#page-4-11)[13](#page-4-12) carrying extreme temperature and pressure in the matter. Experiments employing high peak power ultrashort laser pulses with durations around 100 femtoseconds irradiating Highly Oriented Pyrolytic Graphite (HOPG), usually under vacuum¹⁴⁻¹⁶ or in the graphite/liquid interface¹⁷, have originated sp³-bounded lattices on the graphite^{[14,](#page-4-13)18} and provided preliminary evidences of diamond formation on the substrates^{14-17[,19](#page-4-16),[20](#page-4-17)}. Ultrashort laser shots raise promises for synthesizing known²¹ and still theoretically predicted^{22,23} sp³ carbon structures. Here, we have taken advantage of moderate energy ultrashort laser pulses to generate shockwaves and induce the formation of a diamond-like phase from the sp² carbon polycrystalline graphite precursor.

In the present work, laser shots irradiating graphite under air dramatically modified the surface and created micrometer scale translucent structures where diamond-like crystallites were found coexisting with onion-like phases and quasi-amorphous nanometer sized graphite. Such materials were recovered

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after the laser irradiation process and analyzed by micro Raman spectroscopy, Scanning Electron Microscopy (SEM) and High Resolution Electron Microscopy (HREM). Considering the ultrafast optical excitation, ablation regime and the distinct laser-synthesized carbon structures, we propose an indirect mechanism in which subsequent laser-induced transformations lead to the synthesis of the diamond-like phase.

Results and Discussion

The Raman spectra and the SEM micrographs shown in [Fig. 1](#page-1-0) present the evolution of the induced modifications in the ultrashort laser pulses irradiated polycrystalline graphite sample through assessing the stages from the precursor ([Fig. 1a](#page-1-0),b), the intermediary stage found at the irradiated surface [\(Fig. 1c](#page-1-0),d), to the final diamond-like structure [\(Fig. 1e](#page-1-0),f).

In the [Fig. 1a](#page-1-0) the pristine graphite exhibits its characteristic Raman modes at 1343 and 1615 cm[−]¹ (related to structural defects bands D1 and D2, respectively), and at 1579 cm[−]¹ originating from the stretching of the sp²-bounded carbon atoms forming the well-known hexagonal structure (G-band)^{[24](#page-5-3)}. After irradiation, notable changes were observed in the Raman spectrum of the laser modified surface, shown in [Fig. 1c](#page-1-0), with respect to the precursor [\(Fig. 1a](#page-1-0)). While the D1 band appears at 1340 cm⁻¹, the displacement of the G band to 1595 cm⁻¹ [\(Fig. 1c](#page-1-0)) confirms the laser-induced pressure, since similar shifts to higher wavenumbers were verified for HOPG upon room-temperature compression²⁵. Additionally, Raman resonances, which do not manifest in the precursor [\(Fig. 1a](#page-1-0)), emerge at 1087, 1245, 1425 and 1552 cm[−]¹ . Analogous spectra have been reported for an experiment that quenched Carbon Black from high-pressure high-temperature environment²⁶ (15GPa and 1700 °C for 15min) and also in shocked meteorites²⁷. In the former work, the new vibrations were declared unexplained Raman modes, while in the latter they were attributed to a new carbon phase. Considering previous works $26-28$ and our experimental observations, we suggest that these vibrations are evidence of the creation of sp³ carbon phases by the ultrafast laser excitation, as theoretical models²⁹ have predicted sp³ lattices with rather specific Raman resonances.

The Raman spectrum of the laser-created structure transferred to a Cu grating [\(Fig. 1e\)](#page-1-0) shows an expressively pronounced G-band at 1580 cm⁻¹. While this band nearly dominates the spectrum, less intense vibrational resonances similar to the ones for the laser modified surface are also present. The dominance of the G-band closely resembles the Raman spectrum of HOPG²⁶, therefore, indicating that the laser shockwaves induced reconstruction of the disordered arrangement of the pristine graphite into a more ordered graphitic phase. However, Selected Area of Diffraction (SAD) analysis of crystallites at multiple regions of this laser created structure [\(Fig. 1f\)](#page-1-0) turned out being rather different from graphite, as can be noted in [Fig. 2a](#page-2-0) (laser created particle) and 2**b** (polycrystalline graphite precursor). By assessing the related diffraction peaks plotted as a function of the inverse of the d-spacing [\(Fig. 2d\)](#page-2-0), the laser induced changes are corroborated by the clear absence of the 0.338nm interlayer distance – a signature of graphitic phase – in the created structure. Moreover, the electron diffraction pattern of this laser syn-thesized carbon form reasonably matches that of zinc blend diamond phase^{[10](#page-4-9)[,30](#page-5-8)} and, accordingly, lead us to argue that the laser excitation synthesized a diamond-like phase. [Figure 2c](#page-2-0) exhibits a High Resolution Image corresponding to a small region of the diffraction pattern area shown in the [Fig. 2a](#page-2-0). The whole image area $(16 \times 16 \text{ nm}^2)$ of the [Fig. 2c](#page-2-0) presents planes with 0.205 nm distance (diamond characteristic plane distance) and an amorphous background, which is also noticed in [Fig. 2a](#page-2-0).

The dark field image shown in [Fig. 3b](#page-3-0) was obtained using an objective aperture to select the diffraction hales corresponding to the two more intense peaks of the diamond-like phase electron diffraction pattern ([Fig. 3a\)](#page-3-0). Texturized micrometer domains of the latter phase ([Fig. 3b](#page-3-0)) can be observed with typical sizes around 50×25 nm². In their vicinities, onion-like structures were also encountered ([Fig. 3c](#page-3-0),d).

Figure 3. Texturized domains of the diamond-like phase and onions-like carbon coexisting at the laser created particle. (**a**) SAD of the diamond-like phase from a small area on the region of the laser created structure displayed in (**b**). The highlighted red spot in the frame (**a**) represent the selected diffraction peaks for construction the patterned domains shown in (**b**). Onion-like phases also found at the laser created structure are presented in the frames (**c**) and (**d**).

Here we propose that the mechanism for the transformation from graphite to the diamond-like phase follows an indirect pathway that is underpinned on the morphology of the starting material, specific thermodynamical events dictated by the fluence of overlapping ultrashort laser pulses¹⁸ and the forma-tion of natural catalysts, as the onion-like structures³¹ and the laser driven nanometer sized graphite^{[26](#page-5-5)}. Initially, the high densities of free electrons accumulating at the boundaries of the precursor defective graphitic flakes favored the absorption of the ultrashort pulses energy steering to the creation of superexcited states, and ultimately to ablation³². As a consequence of the ultrafast excitation above the high fluence ablation threshold and the following explosive ablation, nonthermal shockwaves first propagate into the material after the electronic relaxation, and then heating and thermal equilibrium take place. This thermodynamical process, which happens at each laser shot, produces cumulative incremental lattice distortions^{14,33} leading to more ordered carbon forms³⁴. Considering the exposure time with pulse superposition, these latter structures behave as transient states whose energetic barrier to phase transition lowers while the crystallinity gradually increases at each shockwave shot. Therefore, the formation of the diamond-like phase is assisted by those intermediary carbon structures which assume the role of nucleation sites more prone to phase transitions 35 .

Conclusion

In summary, we have demonstrated, for the first time to our knowledge, the synthesis of a diamond-like phase from blasting polycrystalline graphite with spatially overlapping 25 fs ultrashort laser pulses under air atmosphere, relaxing the experimental conditions to create nanodiamonds. The diamond-like phase was found coexisting with onion-like carbon structures, in a micrometer sized translucent laser created particle that was investigated by micro Raman spectroscopy, SEM and HREM. Based on results of previous literature work and on our observations, we proposed a mechanism for the formation of the diamond-like phase from the graphite precursor. The deduced pathway of synthesis correlates the dynamics of excitation, temperature and pressure with phenomenological observations to yield relevant insights into the phase transformation of carbon allotropes by ultrafast extreme conditions.

Methods

Samples were prepared by cutting 99.99% pure polycrystalline graphite bars (Alpha Aesar) into rectangular blocks ($15 \times 7 \times 4$ mm³) whose larger surfaces were manually polished with fine sandpaper up to 10μm (RMS) roughness. Micro Raman spectroscopy (Xplora Plus microscope from Horiba with laser excitation at 532nm), Scanning Electron Microscopy (SEM, FEI Inspect 50F), and High Resolution Electron Microscopy (HREM, JEOL JEM 2100F) were used to characterize the precursor sample and the subsequent products of the laser irradiation. To accomplish the ablation experiments, the samples were irradiated by ultrashort pulses from a Ti:Sapphire amplified laser system (Femtopower Compact Pro HR/ HP from Femtolasers). This system generated 25 fs pulses centered at 780 nm, with 558 μ J of energy, in a 4 kHz pulse train. The beam was focused in air by a 75mm focal distance achromatic doublet, and the sample was placed 1mm before the focal plane, with the laser beam impinging on its polished face. In order to create a proper surface for the analyses, the sample was moved transversely to the beam, at 10mm/s, in such a way to etch a line across its width; subsequently, 400 parallel lines, displaced by $10 \mu m$, were etched and a 7×4 mm² irradiated area was formed. At the sample surface, the calculated beam spot size (radius) was 67 μ m, resulting in 3.96 J/cm² fluence per pulse, which corresponds to a 1.58×10^{14} W/cm² peak intensity. This applied fluence is three times higher than the 1.3 J/cm² single shot high fluence ablation threshold that we measured for this material³⁶. Moreover, we estimate that the pulses superposition were close to 300 pulses per spot, at which high fluence ablation threshold is lowered to 0.8 J/cm^{2,36}. As a comparison, ultrashort laser pulses with fluencies as low as 60 mJ/cm² are known to cause changes in the interplanar distance of the graphite¹⁴, a process that starts the laser ablation. After irradiation the sample was annealed (at 450° C for 5 hours) in ambient atmosphere to eliminate debris³⁷. Upon visual inspection of the ablated surface through the $10\times$ magnification objective of the Raman microscope, the white illumination highlighted translucent micrometer-sized structures with different shapes, sizes ranging from $10 \mu m$ to $50 \mu m$ (Figures S1 to S3 of the Supplementary Information) and pronounced photoluminescence (Figure S2 of the Supplementary Information). Electrostatic attraction between those structures, which were found detached from the substrate probably due to the ablation process, and the tip of an Atomic Force Microscope, allowed their transfer to a Cu grating for the HREM analysis.

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Author Contributions

R.E.S. and N.D.V. Jr. made the ablation experiments. F.C.B.M. performed Raman measurements. R.O.F. steered the experiments with the Atomic Force Microscope. J.B. accomplished electronic micrographs and diffraction experiments. N.M.S.-N. proposed the experiments and throughout coordinated the work. All authors took part in the discussion and effectively contributed to the conclusions.

Additional Information

Supplementary information accompanies this paper at <http://www.nature.com/srep>

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