Graphitization of Glassy Carbon after Compression at Room Temperature

T. B. Shiell,1,* D. G. McCulloch,2,3 D. R. McKenzie,4 M. R. Field,3 B. Haberl,5 R. Boehler,5,6 B. A. Cook,5 C. de Tomas,7 I. Suarez-Martinez,7 N. A. Marks,7 and J. E. Bradby1

1Department of Electronic Materials Engineering, Research School of Physics and Engineering, The Australian National University, Canberra, Australian Capital Territory 2601, Australia
2Physics, School of Science, RMIT University, Melbourne, Victoria 3001, Australia
3RMIT Microscopy and Microanalysis Facility, RMIT University, Melbourne, Victoria 3001, Australia
4School of Physics, The University of Sydney, New South Wales 2006, Australia
5Neutron Scattering Division, Neutron Science Directorate, Oak Ridge National Laboratory, Oak Ridge, Tennessee 37831, USA
6Geophysical Laboratory, Carnegie Institution of Washington, 5251 Branch Road, Northwest Washington, D.C. 20015, USA
7Department of Physics and Astronomy, Curtin University, Perth, Western Australia 6845, Australia

(Received 17 July 2017; revised manuscript received 5 April 2018; published 23 May 2018)

Glassy carbon is a technologically important material with isotropic properties that is nongraphitizing up to ~3000 °C and displays complete or “superelastic” recovery from large compression. The pressure limit of these properties is not yet known. Here we use experiments and modeling to show permanent densification, and preferred orientation occurs in glassy carbon loaded to 45 GPa and above, where 45 GPa represents the limit to the superelastic and nongraphitizing properties of the material. The changes are explained by a transformation from its sp² rich starting structure to a sp³ rich phase that reverts to fully sp² bonded oriented graphite during pressure release.

DOI: 10.1103/PhysRevLett.120.215701

The flexibility of the carbon atom in forming covalent bonds with different hybridization states leads to carbon solids that exhibit a range of useful properties from the extreme hardness of diamond to the extreme “slipperiness” of graphite [1–3]. Glassy carbon (GC) is a predominately sp² bonded carbon that has exceptional superelastic mechanical properties in which it completely recovers its shape after large compressive strains [4]. The material is a noncrystalline solid that is formed by the high temperature decomposition of cross-linked polymers [5]. GC has a low macroscopic density (~1.5 g/cm³) when compared to graphite (2.27 g/cm³) and is by definition a nongraphitizing carbon (i.e., it resists the development of graphite crystals) even when heated to temperatures up to 3000 °C [6,7]. It is superior to graphite for applications such as noncontaminating crucibles and electrodes in electrochemical devices. GC is also being investigated for use in tribological applications, including medical prosthetics [8] and wear resistant surfaces under extreme conditions (e.g., aircraft brakes) [9,10]. Transmission electron microscope (TEM) images reveal that GC has a characteristic nanostructure that has been interpreted as either a tangle of graphitic ribbons [5] or an assemblage of sheetlike structures containing fullerene-like elements [11]. Its novel nanostructure also makes it a useful precursor material for the synthesis of new carbon polymorphs such as nanocrystalline hexagonal diamond [12].

The structural origin of the resistance of GC to graphitization remains a topic of discussion in the literature [5,13,14]. GC can be classified into two “types,” where type 1 has a heat treatment temperature less than 2000 °C, and type 2 has a heat treatment temperature greater than 2000 °C. Only subtle differences in the high pressure behavior between the two types have been reported [15]. Some authors have proposed that GC contains a small fraction of sp³ bonds that act as cross-links between graphitic sheets or ribbons [11]. The structural origin of the superelastic property also remains unknown. Some authors propose that it is associated with a shape memory effect embodied within the sheetlike nanostructure [4,15], with the graphene sheet as a structural element retaining the memory of its initial shape. When compressed, the sheets develop some bonding interactions that do not change the original topology of the sheets, enabling the structure to return to its original configuration after decompression. Any sp³ bonds acting as cross-links in GC contribute to this superelastic behavior, governing the time response of the shape recovery.

Some high pressure studies have been used to show that GC undergoes a reversible bonding transformation from sp² rich to sp³ rich at pressures beyond 40 GPa at room temperature [16], while other work reports that a substantial content of sp² bonding remains while the material is held under pressures of up to 60 GPa [17]. It has also been reported that GC becomes transparent when compressed above 33 GPa [18], suggesting that the majority of the material becomes sp³ bonded. Another study found a permanent increase in macroscopic density of 10%.
following compression to 35 GPa [15]. However, none of the room temperature, high pressure studies of GC provide clear evidence of \( sp^3 \) bonds in the recovered samples at ambient pressure. Several simulations do show that compressing graphitic materials induces the formation of \( sp^3 \) bonding, leading to many different hybrid \( sp^2/sp^3 \) structures with diamondlike densities, but only while the pressure is maintained [19–23]. Above room temperature, the high pressure results are different again. Recent reports have shown hexagonal diamond formed at 400 °C [12], increased \( sp^3 \) bonding by heating up to 1200 °C [24], and the recovery of a highly disordered, almost 100% \( sp^3 \) bonded material after laser heating [25].

An unanswered question regarding the high pressure behavior of GC is the maximum pressure to which GC can be subjected before it loses its distinctive nongraphitelike properties and isotropic tangled sheetlike nanostructure. This question is of direct relevance to tribological performance of GC, since the pressure threshold determines how much of a GC wear surface is converted to a graphitic structure, with the remainder retained as a superelastic 3D network solid. In this study, we address this question by subjecting GC to compression in diamond anvil cells (DACs) and characterizing the recovered samples using Raman spectroscopy and TEM. To assist in the interpretation of the experiments, the compression and decompression processes are simulated using molecular dynamics.

The GC sample used in this Letter has a macroscopic density of 1.42 g/cm³ and was purchased in the form of a solid plate from Hochtemperatur Werkstoffe (Sigradur-G). It was manufactured from a phenolic resin heated to 3000 °C. Small chips (≈80 × 80 × 50 μm) were loaded without a pressure medium into a Bohler Almax plate-DAC and raised to maximum pressures of 4, 10, 25, 35, 45, and 54 GPa using diamonds with 400 μm diameter culets. The absence of a pressure medium results in a significant uniaxial component in the supplied stress field. For all experiments, stainless-steel gaskets were used with sample chambers initially 200 μm in diameter and 55 μm deep. Pressures were determined in situ by the shift of the R1 ruby fluorescence line and the shift of the main diamond Raman peak in the center of the gasket hole [26]. Figure 1 shows a schematic of the experimental procedure for preparing TEM specimens, which involved using a focused ion beam (FEI Scios Dualbeam) to retrieve samples of GC following compression in a DAC.

Figure 2(a) is a TEM image (taken using a JEOL 2100F TEM operating at 200 kV) of uncompressed GC, showing the tangled isotropic nanostructure consisting of curved graphitic stacks of up to ~10 layers. A TEM image from the recovered sample following compression to 35 GPa is shown in Fig. 2(c). This image shows a similar structure to that observed in the uncompressed case, with the entanglement of graphitic sheets still present. After compression to 35 GPa, the characteristic GC nanostructure remains intact, consistent with previous observations [15]. However, a significant change in the nanostructure is evident in the TEM image from the recovered sample following compression to 45 GPa, as shown in Fig. 2(e). This image shows that some of the curved graphitic sheets are destroyed and are fully replaced with small graphitic crystallites with significant short-range order.

The observed changes in the nanostructure were further analyzed by electron diffraction. Diffraction patterns are shown in Fig. 2(b) for uncompressed GC, Fig. 2(d) for GC after compression to 35 GPa, and Fig. 2(f) for GC after compression to 45 GPa. The diffraction pattern of uncompressed GC shows the rings typical of nanocrystalline graphite, exhibiting strong \{002\}, \{100\}, and \{110\} reflections. It should be noted that, for the uncompressed GC sample, each diffraction ring has a uniform distribution of intensity, proving that the nanostructure has random orientation. Figure 2(d) shows that, after compression to 35 GPa, the diffraction rings still have uniform intensity, although there is some evidence for increased structural order from the sharpening of the diffraction rings. However, the diffraction pattern of the sample after compression to 45 GPa shows a highly nonuniform distribution of intensity. Therefore, the sample undergoes a significant permanent structural change. The graphitic planes are now
preferentially oriented and aligned perpendicular to the direction of compression. This new preferred orientation minimizes the elastic strain energy as a result of the strongly anisotropic elastic moduli of graphite [27,28]. A similar phenomenon has previously been described for pure graphite and is used to create highly oriented pyrolytic graphite under uniaxial stress conditions and high temperatures [29]. The development of nanocrystalline graphite following compression to 45 GPa is indicative of a loss of the superelastic property of GC, as the isotropic topology of the original sheet nanostructure has been destroyed and replaced with graphite that does not exhibit the superelastic property [30,31].

Direct evidence for the loss of the superelastic properties comes from electron energy loss spectroscopy (EELS) density measurements (Gatan Tridiem imaging filter fitted to a JEOL 2100F TEM operating at 80 kV), which are known to be a sensitive measure of both the bonding and density [32]. The position of the plasmon peak in the low loss EELS spectra [Supplemental Material Fig. S1(a) [33]] is proportional to the density of free electrons and therefore can be used to estimate the local microscopic density [34] of the uncompressed GC, giving a value of 1.62 g/cm³. This value is lower than the density of graphite (2.27 g/cm³), but it is higher than the manufacturer’s specified macroscopic density of 1.42 g/cm³. The macroscopic density includes the contribution from voids and thus differs from the EELS density [35]. The density of the recovered GC samples as a function of pressure is shown in Fig. 3. It reveals no significant change in microscopic density for samples compressed up to 35 GPa, but a significant permanent increase in microscopic density to 1.86 g/cm³ occurs for samples compressed to 45 GPa and above. A nanostructure consisting of stacked graphitic regions [Figs. 2(e) and 2(f)] is more efficiently packed than the tangled GC nanostructure leading to this increase in microscopic density.

To investigate bonding changes in the material, the carbon K-edge EELS of the uncompressed and recovered GC materials was measured [Supplemental Material Fig. S1(b)]. All spectra exhibit a strong 1s to π* peak at 285 eV, consistent with a dominance of \( sp^2 \) bonding [32]. The \( sp^2 \) fractions were calculated from the carbon K-edge spectra using a method similar to that described by Berger et al. [36]. Care must be taken when calculating \( sp^2 \) fractions of the recovered samples, as preferred orientation is known to strongly influence the intensity of the 1s to π* peak [37]. This influence of preferred orientation can be accounted for by performing EELS measurements at the so-called magic angle conditions, where the intensity of the 1s to π* peak does not vary with orientation of the graphitic crystallites relative to the incident beam [38]. This magic angle EELS (performed using specific electron beam convergence and spectrometer collection angles) enables the \( sp^2 - sp^3 \) bonding ratio to be

![FIG. 2. TEM images and selected area diffraction patterns (indexed to graphite) of (a),(b) uncompressed GC, (c),(d) the sample recovered from 35 GPa, and (e),(f) the sample recovered from 45 GPa. (Inset) The compression axis of the DAC is indicated by the blue arrows.](image)

![FIG. 3. Local microscopic density (black squares), \( sp^2 \) fraction (red circles) obtained from EELS, and average in-plane crystallite size \( L_{a\parallel} \) (blue crosses) obtained from Raman spectroscopy. Interestingly, the Raman spectra were better fitted with two peaks in the \( f \)-band region. However, the physical origin of these peaks remains unknown.](image)
measured, even in the highly oriented sample recovered after compression to 45 GPa. At least four measurements were taken on each sample and the results are plotted in Fig. 3. The results show that the uncompressed GC has a $sp^2$ bonding fraction of 95 ± 2% and a small fraction of $sp^3$ bonds. The fraction of $sp^2$–$sp^3$ bonding does not change for samples compressed up to 35 GPa. For samples recovered after compression to 45 GPa and above, there is an increase in the $sp^3$ bonding fraction to ~100%. This result suggests that, when raised to the threshold pressure of 45 GPa, any preexisting $sp^3$ bonds are destroyed either on loading or on pressure release.

Raman spectra were collected (using a Renishaw InVia micro-Raman spectrometer, equipped with a 532 nm excitation laser) from each of the recovered GC samples (Supplemental Material Fig. S2 [33]). The Raman peak positions do not change substantially from reported peak positions [39] after compression; however, relative integrated intensities do show a sharp discontinuity between 35 and 45 GPa (fitted peak parameters in Supplemental Material Table S1). The relative integrated intensities can be used to determine the average in-plane graphitic crystallite sizes $L_a$ [40]. Figure 3 shows that $L_a$ of the uncompressed GC is 10.3 nm and decreases to 8.1 nm between 35 and 45 GPa, supporting our observations of significant changes in nanostructure observed in the TEM results.

To assist in the interpretation of our experimental data, atomistic simulations of uniaxial compression and decompression were performed using the LAMMPS molecular dynamics package [41], with interactions described by the environment-dependent interatomic potential (EDIP) for carbon [42]. The starting point of the simulations was a 1.5 g/cm$^3$ structure [Fig. 4(a)] containing 32768 atoms, generated using a liquid quench and annealing methodology [43]. The starting structure contains graphene layers with different orientations, characteristic of GC, and is highly $sp^3$ bonded. Upon compression, the graphene layers gradually align perpendicular to the applied stress [Fig. 4(b)]. At higher pressures, cross-linking $sp^3$ bonds form between layers, increasing the $sp^3$ fraction substantially [Fig. 4(d)]. Structures compressed to 10, 25, 35, 45, and 56 GPa were fully decompressed using the same strain rate as the compression, in combination with annealing of the decompressed structure at 3500 K to activate atomic rearrangement. At low maximum compression (e.g., to 10 GPa), the structure recovers its original bonding architecture upon decompression, although the details of the bonding have been altered in some regions [Fig. 4(c)]. These simulations are consistent with our experimental data for pressures less than 35 GPa and are consistent with the superelastic behavior of GC. At higher pressure (e.g., 45 GPa), the decompressed structure has a similar $sp^2$ fraction to the original structure; however, the graphitic layers have been preferentially realigned perpendicular to the applied stress direction [Figs. 4(d) and 4(e)]. The development of preferred orientation of the graphitic layers is consistent with our experimental observations at similar pressures. Additionally, the structures that have been compressed to pressures above ~35 GPa do not recover their original density upon decompression [black circles in Fig. 4(f)], but have a permanently increased density as seen experimentally (Fig. 3). Figure 4(f) also shows the $sp^3$ fraction of the compressed phase (blue squares), displaying a transition at around 35 GPa, to a structure with a high $sp^3$ fraction.
fraction. This high $sp^3$ structure is unstable and reverts to a high $sp^2$ fraction structure upon pressure release, as shown in the Figs. 4(d) and 4(e).

Following recovery after compression up to 35 GPa, both experiment and simulation show that the isotropic nanostructure of GC is retained. This demonstrates that the GC nanostructure, which consists of extremely strong and tangled graphene sheets, is highly resilient to compression. The superelastic property of GC is the manifestation of this resilience.

At pressures of 45 GPa and above, our simulations show that a majority $sp^3$ phase is formed that is unstable when pressure is released. Several other theoretical studies have also predicted the formation of exotic unstable $sp^3$ rich phases at high pressures [19–23]. These findings from theory are consistent with the experimental work of Lin et al. that showed the $sp^3$ bonds formed by loading GC to pressures up to 44 GPa reverted to $sp^2$ bonds on pressure release [16]. Moreover, it has recently been reported that an exotic $sp^3$ phase, known as $M$-carbon, has been experimentally observed during the room temperature compression of graphite at $\sim$ 40 GPa [44], which spontaneously transformed back to $sp^2$ bonded graphite on pressure release. We propose that a similar high pressure $sp^3$ rich phase is formed in our GC at pressures of 45 GPa and above. This $sp^3$ structure is unstable at ambient. At high pressure, the tangled GC nanostructure, including any fullerenelike elements, is destroyed as the dominant bonding changes from $sp^2$ to $sp^3$. The transformation to a $sp^3$ rich phase and then down to a fully $sp^2$ bonded phase would explain why, on pressure release, the sample does not recover its original isotropic tangled sheetlike structure but instead shows permanent preferred orientation of the newly formed $sp^2$ bonded graphite. The graphite is oriented [Fig. 2(f)] reflecting its formation in a stress field with a strong uniaxial component. Since the formation of extensive preferred orientation of the graphitic layers is essentially graphitization, we propose that compression of GC to 45 GPa and above causes a loss of its superelastic properties, since graphite is not superelastic.

In summary, we have used TEM imaging, electron diffraction, EELS, Raman spectroscopy, and atomistic modeling to understand the superelastic and nongraphitizing characteristics of GC as it is compressed at room temperature. After loading up to 35 GPa, the recovered material is found to retain its tangled nanostructure, including its minority content of $sp^3$ bonding ($\sim$5%). In contrast, the samples recovered after compression to over 45 GPa contain a negligible $sp^3$ bonding content ($\sim$0%). Pressures above this threshold cause the tangled sheet structure to align as the sheets come closer together, and the modeling shows that new $sp^2$ bonds form but are unstable at ambient. Experimental results also show permanent densification and the formation of oriented graphite layers in GC subjected to such pressures. Accordingly, this 45 GPa threshold represents the ultimate limit to the nongraphitizing and superelastic properties of GC and this work provides valuable insight into the transition pathway of pressure-induced structural and bonding changes in GC.

J. E. B. would like to acknowledge the Australian Research Council (ARC) (FT130101355) and J. E. B. and D.G.M. acknowledge funding under the ARC Discovery Project scheme (DP140102331). This work was also supported by the Australian Research Council (DP150103487). Computational resources are provided by the Pawsey Supercomputing Centre with funding from the Australian Government and the Government of Western Australia. B. H. gratefully acknowledges funding through the ORNL Neutron Scattering User Facilities supported by the U.S. Department of Energy, Office of Sciences, Basic Energy Sciences. ORNL is funded under DOE-BES Award No. DE-AC05-00OR22725.

This manuscript has been authored by UT-Battelle, LLC under Award No. DE-AC05-00OR22725 with the U.S. Department of Energy. The United States Government retains and the publisher, by accepting the article for publication, acknowledges that the United States Government retains a nonexclusive, paid-up, irrevocable, world-wide license to publish or reproduce the published form of this manuscript, or allow others to do so, for United States Government purposes. The Department of Energy will provide public access to these results of federally sponsored research in accordance with the DOE Public Access Plan. Work by R. B. was supported by the Energy Frontier Research in Extreme Environments (EFrEE) Center, an Energy Frontier Research Center funded by the U.S. Department of Energy (DOE), Office of Science, Basic Energy Sciences. ORNL is funded under DOE-BES Award No. DE-SC0001057.

*Corresponding author. tom.shiell@anu.edu.au