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# Direct *in situ* observation of toughening mechanisms in nanocomposites of silicon nitride and reduced graphene-oxide



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#### ABSTRACT

In situ observations are performed, for the first time, of stable crack propagation in nanocomposites of silicon nitride  $(Si_3N_4)$  and reduced graphene-oxide (rGO) inside a scanning-electron microscope. Two different specimen geometries (wedge-splitting and double-cantilever beam) are used to observe crack interactions with rGO stacks in two different orientations (cross-section and in-plane). These observations provide new insights into the unique, effective pull-out of crack-bridging rGO stacks, which appears to be responsible for the extraordinary toughness in the  $Si_3N_4$ /rGO nanocomposites. These insights could be used to design and create future ceramic/ rGO nanocomposites with superior mechanical properties.

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There has been great interest in using carbon nanostructures, especially one-dimensional (1-D) carbon nanotubes (CNTs), to toughen brittle ceramics (see recent reviews [1,2]). This is because CNTs have a set of exceptional structural features and properties that are not found in conventional ceramic reinforcements such as platelets, whiskers, or fibers. Accordingly, the toughening mechanisms in ceramic/CNTs nanocomposites are also unique [3,4]. With the advent of 2-D graphene and its derivatives, which also have a set of unprecedented structural features and properties, it is a natural progression to use them as reinforcements in ceramic nanocomposites. (The term 'graphene' refers to flexible fewlayer graphene stacks, in addition to single-sheet graphene, but not stiff graphite 'platelets.') We argue that the 2-D nature of both the graphene reinforcements, and the crack that these reinforcements are expected to bridge, is likely to make graphene a more effective reinforcement compared to the 1-D CNTs. Early reports showed significant toughening in alumina  $(Al_2O_3)$  [5] and silicon nitride  $(Si_3N_4)$  [6] ceramics with the introduction of graphene reinforcements in the form of reduced graphene oxide (rGO) stacks. Since then, there have been numerous reports showing toughening in a wide variety of ceramic/graphene nanocomposites [7,8]. There is, however, large variation in toughness gains, which can be attributed to large variation in the processing and the microstructures. Generally, rGO stacks are found to be more effective because of the better dispersion of hydrophilic GO containing surface

\* Corresponding author. E-mail address: nitin\_padture@brown.edu (N.P. Padture). functional groups (O<sup>-</sup>, OH<sup>-</sup>, COOH<sup>-</sup>) in the initial GO/ceramic powder mixture, as opposed to non-functionalized hydrophobic graphene [9]. While GO typically has inferior mechanical properties such as elastic modulus and strength, graphene-like behavior can be recovered by reducing the GO (rGO) during heat-treatments as part of the composite densification processes such as spark-plasma sintering (SPS) [10]. Furthermore, the functional groups on the starting GO surfaces can promote better interfacial adhesion between the outer layers of the rGO stacks and the ceramic surfaces [11].

While most of the relevant studies report toughness values, measured using either bend or indentation tests, some also provide micrographs of static cracks that show bridging by rGO stacks in the wake of the crack tip and/or on fracture surfaces [6-8,12]. Toughening mechanisms are then speculatively proposed based on those observations. In this context, it is well established that in situ observations of crack propagation provides significantly deeper insight into the toughening mechanisms in other materials [4,13,14]. To that end, direct in situ investigation of crack propagation in fully-dense SPSed Si<sub>3</sub>N<sub>4</sub>/rGO nanocomposites were performed for the first time, to provide new insights into the unique toughening mechanisms that operate in these types of materials. Note that Si<sub>3</sub>N<sub>4</sub> is perhaps one of the most widely studied ceramic matrix in the context of incorporating graphene fillers because of the high toughness levels achieved in the resulting nanocomposites [8,17]. Two different specimen geometries (Fig. 1A and B) are used to enable stable crack growth during loading in a micro-test device, inside of a scanning electron microscope (SEM): (i) wedge-splitting [15], and





**Fig. 1.** *In situ* test specimen geometries used to obtain stable crack propagation inside the SEM: (A) wedge-splitting (a = 10 mm, b = 3 mm, d = 5 mm) and (B) DCB in compression (a = 4.8 mm for in-plane, 3 mm for cross-section, b = 1 mm, d = 1.9 mm, h = 0.6 mm, w = 1.2 mm). (C) SEM micrograph of fracture surface of Si<sub>3</sub>N<sub>4</sub>/rGO nanocomposite (arrow indicates SPS pressing direction). Schematic depiction of the rGO-stacks orientation with respect to the main crack (arrows indicate SPS pressing direction): (D) cross-section and (E) in-plane.

(ii) double-cantilever beam (DCB) in compression [16]. The SPSed nanocomposite pellets are unintentionally orthotropic, where most of the rGO stacks naturally align in planes perpendicular to the SPS pressure axis. Fig. 1C is a cross-sectional SEM micrograph of Si<sub>3</sub>N<sub>4</sub>/rGO nanocomposite showing this alignment of rGO stacks. This presents a unique opportunity to study crack propagation in two different orientations: 'cross-section' (Fig. 1D) and 'in-plane' (Fig. 1E), where the rGO stacks are 'normal' and 'edge-on', respectively, with respect to the planar crack front.

Fully-dense SPSed Si<sub>3</sub>N<sub>4</sub>/rGO nanocomposite pellets (3 mm thickness, 20 mm diameter) from a previous study [17] are used here. They contain 4.3 vol% rGO (Fig. 1C), with an impressive toughness of ~10 MPa  $\cdot$  m<sup>0.5</sup> in the 'cross-section' orientation, as measured using the reliable surface-crack in flexure (SCF) toughness test [18]. This represents over two-fold increase in the toughness compared to monolithic Si<sub>3</sub>N<sub>4</sub> ceramic (~4.5 MPa  $\cdot$  m<sup>0.5</sup>) without the rGO. (See Refs. [10,17] for

details regarding processing, microstructures, and mechanical properties.) Both types of specimens were cut from the same pellets in the two orientations using a low-speed diamond saw, followed by surface-polishing. A piezo-driven micro-test device (MT10141, Deben, Woolpit, UK) with specially fabricated fixtures, equipped with a load cell (2 kN capacity), was used to load the test specimens in compression (Fig. 1A and B) [4]. In both types of specimens, sharp 'V' notches were cut using a razorblade with a 1-µm diamond suspension. Compression was applied to propagate a sharp, stable crack ( $\Delta c$ ) at the root of the 'V' notch. The load was then increased and held, allowing us to photograph the crack using secondary-electron imaging in the SEM. This process was repeated at successively higher applied loads. There are pros and cons to the two specimen geometries. While the crack growth is more stable in the DCB geometry, it requires much larger loads (~2 kN), limiting the dimensions of DCB specimens. As mentioned earlier, the specimens are oriented in two different ways. Both specimen geometries (wedge-splitting and DCB) are used in the in-plane orientation (Fig. 1E), but only wedge-splitting specimen geometry is used for cross-section orientation (Fig. 1D) due to the limitation on the thickness (3 mm) of the available pellets.

Fig. 2A–D are a sequence of SEM micrographs taken from a fixed location at different loads as the main crack tip propagates (downward) using the DCB geometry (Fig. 1B) in the cross-section orientation (see Fig. 1D; horizontal rGO stacks are perpendicular to the plane of the image). In Fig. 2A it can be seen that thinner rGO-stack bridges (upper) have already pulled-out and fractured. The thicker rGO-stack bridge is still intact (lower). With increasing horizontal crack-wall separation (downward main-crack propagation), the rGO stack appears to stretch thin and split along the weak rGO layers, resulting in sliding between the layers (Fig. 2B and C). The rGO stack thins further, bends, and eventually fractures (Fig. 2D). (Note that there is always some vertical displacement of the crack walls during crack propagation, resulting in the rGO bending.) However, in all of the SEM images (Fig. 2A-D), there is no relative motion between the Si<sub>3</sub>N<sub>4</sub> matrix and the embedded rGO stack within the composite on either side of the crack. Instead, the rGO stack appears to be anchored strongly within the Si<sub>3</sub>N<sub>4</sub> matrix. This implies the absence of classical pull-out mechanism observed in toughened ceramics, where the bridging reinforcement (e.g. platelet, whisker, fiber) slides out frictionally, leaving behind an empty socket [13]. This type of non-classical pull-out behavior is found to be pervasive in the Si<sub>3</sub>N<sub>4</sub>/rGO nanocomposite in the cross-section orientation. Another example of a sequence of in situ SEM micrographs in cross-section orientation (wedge-splitting geometry) is presented as a movie in Supplementary Information (SI) showing very similar behavior. Thus, these results show that the unique 2-D layered structure of the rGO stacks results in pull-out via a fundamentally different mechanism.

Now consider the in-plane orientation (see Fig. 1E), where rGO stacks are in the plane of the image. Figs. 2E-H show a sequence of SEM micrographs taken from a fixed location as the crack walls separate using the wedge-splitting geometry. The main crack tip (front) intersects the large rGO stack edge-on when it first encounters it in this orientation. Here, it appears that the rGO stack is tough enough to resist fracture as the crack walls separate and the main crack tip propagates downward through the matrix. The stretching and narrowing of the rGO-stack width is dramatic going from Fig. 2E to G, which confirms that the rGO stack is anchored strongly within the Si<sub>3</sub>N<sub>4</sub> matrix. As the cracks walls separate, the rGO stack narrows further and eventually fractures, most likely from an edge flaw (arrow). Here also, the rGO stack does not appear to pull-out via the classical mechanism. (See SI for a video of this sequence of SEM micrographs.) Another sequence of in situ SEM micrographs is shown in Fig. S1A-D (in-plane orientation, wedge-splitting geometry) in SI. Once again, stretching of a well-anchored rGO stack and its eventual fracture is observed.

In contrast, smaller rGO stacks appear to pull-out in a more classical manner, as seen in Fig. 2I–K (in-plane orientation, DCB geometry). The rGO stack does not appear to stretch with increasing crack-wall

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