Critical Length Scales and Strain Localization Govern the Mechanical Performance of Multi-layer Graphene Assemblies

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Critical Length Scales and Strain Localization Govern the Mechanical Performance of Multi-layer Graphene Assemblies

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Abstract

Multi-layer graphene assemblies (MLGs) with a staggered architecture exhibit high toughness and failure strain that surpass those of the constituent single sheets. However, how the architectural parameters such as sheet overlap length affect these mechanical properties remains
unknown due in part to limitations of mechanical continuum models. By exploring the mechanics of MLG assemblies under tensile deformation using our established coarse-grained molecular modeling framework, we identify three different critical interlayer overlap lengths controlling the strength, plastic stress, and toughness of MLGs, respectively. The shortest critical length scale $L_c^S$ governs the strength of the assembly as predicted by the shear-lag model. The intermediate critical length $L_c^P$ is associated to a dynamic frictional process that governs the strain localization propensity of the assembly, and hence the failure strain. The largest critical length scale $L_c^T$ corresponds to the overlap length necessary to achieve 90% of the maximum theoretical toughness of the material. Our analyses provide general guidelines for tuning constitutive properties and toughness of multilayer 2D nanomaterials using elasticity, interlayer interaction energy and geometry as molecular design parameters.

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Graphene, one of the strongest materials known,$^{1-3}$ is particularly suited for advanced structural and mechanical applications.$^{4,5}$ In practice, its range of application remains limited due to the difficulty of harnessing the mechanical properties at larger length scales than those limited by the processing and synthesis of individual graphene sheets (in the order of nanometers to a few micrometers). In addition, its low extensibility and brittle failure behavior further restricts the usage in electronics, energy storage devices, and other applications where toughness and ductility are critical.

Drawing inspiration from biological architectures such as nacre, one possible way to simultaneously increase its extensibility and harness the properties of graphene in a scalable manner is to stack multiple sheets in a staggered fashion forming a so-called multi-layer
graphene assembly (MLG). In materials with multi-layer staggered architecture the tensile load is transferred through shear at the interfaces and the deformation occurs by relative sliding between the sheets in different layers. For small deformations, where sheet and interface behave linearly elastic to a good approximation, the continuum shear-lag model can be used to adequately predict the mechanical properties. However, as the deformation increases and more complex deformational mechanisms, such as strain localization, are activated, the shear-lag model breaks down. Understanding how architectural parameters, in particular the overlap length between sheets in different layers, impact the mechanical behavior of the assembly is of critical importance for the rational design of optimal MLGs and other engineered materials with analogous architecture. Here, we use coarse-grained molecular dynamics (CG-MD) of MLGs under uniaxial tensile strain to identify the critical overlap length scales that control the strength, toughness and failure strain.

The CG molecular model of graphene used here (Fig. 1(a)) has been shown to accurately reproduce the elastic and fracture properties of single crystal graphene, as well as its interlayer shear behavior. Details of the CG force-field can be found in the original publication but have also been included in the SI, together with a detailed presentation of the simulation parameters and protocols used in this work. We have simulated multiple MLG systems with varying number of layers ($n_t = 2, 3, 5, \text{ and } 10$) and one sheet per layer ($n_s = 1$), which constitute a representative volume element (RVE) (Fig. 1(b)). The simulated MLGs in the RVE have overlap lengths $L_o$ ranging from ~ 3 to 780 nm. The sheet’s width is set to be 6.4 nm for all the cases and our preliminary results indicate that the width does not affect our findings. Each graphene sheet in the assembly is a single crystal without defects. We do not consider grain boundary effects as the typical grain size of graphene (~10 μm) is much larger than the sizes of our system. To verify
that our findings are applicable to multiple sheets per layer, we also perform simulations for the 
\( n_s = 3 \) case. The results have been included in the Supporting Information (SI) and do not alter 
any of the conclusions presented. It is worth noting that the bonds of the CG model are breakable 
and that no further constraints, other than the imposed uniaxial strain, are used, i.e., we do not 
impose any a-priori failure or deformational mechanisms on the system.

The typical stress-strain response of MLGs exhibits a short linear elastic regime that ends with 
a sudden drop in the stress. A plastic regime that ends in a rapid decay of the stress until failure 
follows the stress drop (Fig. 1(c)). At the microscopic level, the fundamental deformational 
process of interlayer sliding is not continuum and consists of discrete transitions between 
commensurate stacking configurations of graphene sheets in neighboring layers (i.e. stick-slip 
mechanism). The elastic regime in this microscopic picture corresponds to the deformational 
period before the system undergoes the first slip event, which occurs within the range from \(~1\% 
to 4\% strain depending on \( L_o \) for the all the cases studied (inset of Fig. 1(c)). For small 
deformations, the van der Waals-governed interfaces and the sheets behave linearly elastic, and 
the tensile stress of MLGs is resulted from the static friction between layers. Under these 
conditions, the Young’s modulus and the maximum stress \( \sigma_m \), which we also refer as the tensile 
strength here, can be predicted by the continuum shear-lag model:\(^{14,15}\)

\[
\sigma_m = \frac{\sinh(L_o / l) \gamma_s' E \mu h}{2(1 + \cosh(L_o / l))(l)}
\]  

(1)

where \( l = \sqrt{\frac{E_g h^2}{4G}} \) (\( \approx 5.2 \) nm) is a parameter that represents the length scale over which most of 
the stress is transferred in the interface, \( E_g (\sim 950 \) GPa) is the Young’s modulus of graphene, 
\( G (\sim 1 \) GPa) is the shear modulus of the interface, \( h (= 3.35 \) Å) is the equilibrium interlayer 
distance between the two layers, and \( \gamma_s' (\sim 0.35) \) is the critical interlayer shear strain. The values
of these fundamental parameters are set by the properties of the CG model.\textsuperscript{18} The prediction of the MLG strength by the shear-lag model as a function of $L_o$ is in very good agreement with the simulation results (Fig. 1(d)). If we define the critical length scale governing the strength of MLGs as the $L_o$ at which 90\% of the maximum strength is achieved, we obtain $L_c^\gamma \sim 17$ nm, which is consistent with previous theoretical calculations that predicted $L_c^\gamma \sim 3 \ell$ for an analogous system.\textsuperscript{17} Although the maximum strength of MLGs ($\sim 10$ GPa) is an order of magnitude lower than that of monocrystalline graphene ($\sim 100$ GPa),\textsuperscript{1} its strength still surpasses most of structural engineered materials.

At the end of the elastic regime we observe a sudden drop in stress. We associate this drop to the transition from a static (before the first slip event) to a dynamic frictional situation \textsuperscript{20, 21} (referred here as the plastic regime because the deformations are irreversible). This drop after the peak stress could originate from the irregular stress distributions within the sheet, which will not strictly follow the shear-lag prediction. The plastic stress, $\sigma_p$, is defined as the average stress in the plateau plastic regime, which is indicated by the horizontal dashed lines in Fig 1(c). Interestingly, the dependence of the post-peak plastic stress $\sigma_p$ on $L_o$ reveals that shear-lag scaling still holds (Fig. 1(d)). In fact, the trend can be quantitatively captured by Eq. (1) if $\ell$ ($\sim 16.3$ nm) and $\gamma_{cr}$ ($\sim 0.67$) are used as fitting parameters. This resulting weaker interface (i.e. softer and more ductile) in the plastic regime compared to the interface properties at small deformations is compatible with the static-to-dynamic frictional transition proposition. Similar to the critical length scale identified for strength, we can define a second critical length scale associated with the saturation of the plastic stress to a near maximum value around $L_c^\beta \approx 50$ nm.

The physical meaning of $L_c^\beta$ becomes clear when looking at the mechanical response of a single interface. Fig. 2(a) illustrates the bilayer model with a single interface used to characterize
the constitutive force-displacement \((f-u)\) response. To simplify our analysis, we apply the bilinear curve to characterize the \(f-u\) response (Fig. 2(b)). It can be observed that the nearly constant force regime lasts until the displacement \(u\) reaches \(\sim (L_o - L_c^p)\), after which the force decays approximately linearly up to failure by complete loss of overlap between the sheets. This analysis suggests that \(L_c^p\) is the critical overlap length after which the shear force at the interface does not depend on the overlap length. This observation can also be inferred from the shear-lag theory in which the strength \(\sigma_s\) and plastic stress \(\sigma_p\) are nearly independent of \(L_o\) as \(L_o\) beyond \(L_c^s\) and \(L_c^p\), respectively.

To be able to generalize these findings to other 2D materials, we need to understand the dependence of \(L_c^p\) on the constituent material properties, in particular the Young’s modulus of and interlayer adhesion energy. According to the shear-lag model\(^{22}\), the critical length scale at which the tensile strength starts to saturate follows the scaling, \(L_c^s \sim \sqrt{\frac{E_g}{G}}\). In the stick-slip microscopic deformational mechanism the shear rigidity depends linearly on the adhesion energy \(\gamma\) between the two sheets (\(G \sim \gamma\)). Therefore, from a theoretical standpoint, we expect \(L_c^p\) to follow a similar scaling, \(L_c^p \sim \sqrt{\frac{E}{\gamma}}\). In fact, we find that the best fits to the simulation results \(L_c^p \sim (E/E_g)^{0.41}\) and \(L_c^p \sim 1/(\gamma/\gamma_g)^{0.52}\) (Fig. 3(a) and 3(b)), are in close agreement with the shear-lag exponent of 0.5. Although \(L_c^p\) may also depend on other factors, such as the pulling velocity as in the case of atomic friction\(^{21,23}\), we expect the scaling on constituent material properties to hold at a given rate.

Visualization of the simulation trajectories reveals that the deformation is uniformly distributed over all of the interfaces up to a certain point (the intermediate state), after which the strain localizes in a certain region along the length of the MLG (Fig. 4(a) and Movie S1). The
system ends up failing by deoverlapping in the region where the strain localizes. The deformation process identified from our simulations is schematically illustrated in Fig. 4(b). We characterize the intermediate state or equivalently the onset of localization by the displacement of each interface \( u_{int} \) at that moment. After the intermediate the strain localizes only on one side that keeps deoverlapping until failure. Using simple geometric arguments, one can show that if localization occurs irreversibly at \( u_{int} \) the failure strain \( \varepsilon_f \) is directly related to \( u_{int} \) via:

\[
\varepsilon_f = \frac{u_{int}}{2L_o} + \frac{1}{2}
\]  

or more generally for \( n_s \) sheets per layer,

\[
\varepsilon_f = \frac{(2n_s - 1)u_{int}}{2n_s L_o} + \frac{1}{2n_s}
\]  

In order to predict the failure strain of the MLG, we need to know when strain localization occurs, or equivalently the value of \( u_{int} \). Shearing of an interface proceeds through a series of slips between equilibrium states where the lattices are in commensurate positions. Thus, we can think of interlayer sliding as a series of thermally activated jumps over energy barriers.\(^{24,25} \) and use a kinetic model based on Bell’s theory\(^ {26-28} \) to describe the shear behavior of each interface. The lifetime, \( \tau \), of a single jump under an applied force is given by \( \tau = \exp \left( \frac{E_b - x_b f}{k_B T} \right) / \omega_0 \), where \( \omega_0 (\sim 1 \times 10^{13} \text{ } \text{1/s}) \) is the vibrational frequency of the interface in the energy well\(^ {26} \), \( k_B T \) is the thermal energy, and \( x_b \) is the distance from equilibrium to the transition state, which depends on the lattice spacing. The probability \( P_{over} \) of overcoming an energy barrier and advancing the system to the next equilibrium state (i.e. slipping) within a time interval \( \Delta t \) can therefore be approximated by:

\[
P_{over} = 1 - \exp \left( -\frac{\Delta t}{\tau} \right)
\]  
Using this theoretical scheme, we can numerically simulate the deformation of two interfaces in mechanical equilibrium and measure \( u_{int} \), which
would correspond to the displacement at which one of the interfaces gets trapped in a local minimum while the other keeps sliding under the applied force. A detailed description of this kinetic model is given in the SI. The results of this model reveal that strain localization always initiates when the displacement is $u_{int} = L_o - L_c^o$, for $L_o > L_c^o$. This finding suggests that $L_c^o$ is a critical length also associated to strain localization, and hence failure strain. Substituting $u_{int} = L_o - L_c^o$ into Eq. (2), $\varepsilon_f$ can be directly expressed as:

$$\varepsilon_f = 1 - \frac{L_c^o}{2L_o}$$

(4)

This equation accurately captures the $\varepsilon_f$ obtained from the kinetic model analysis (Fig. 4(c)). When $L_o < L_c^o$, the kinetic model predicts that localization initiates from the start, which leads to $\varepsilon_f \sim 0.5$. It is worth noting that the minimum and maximum theoretical failure strain in this system ($n_s = 1$) is 0.5 and 1 respectively. The predictions of the kinetic model agree with what is observed in the CG-MD simulations for $L_o > L_c^o$ (Fig. 4(d)), suggesting that the origin of the strain localization is kinetic trapping of one of the interfaces in a local minimum. For $L_o < L_c^o$, the CG-MD simulations show slightly greater failure strains than the kinetic model predictions, which we attribute to the fact that at a finite rate and short $L_o$, some homogeneous deformation can still carry on before strain localization occurs. It should be noted that changing the $x_o$ of the CG model to the value of atomistic graphene will not change the kinetic model predictions.

Once we are able to theoretically predict the plastic stress and the failure strain of MLGs, we can attempt to predict the toughness of the system, or equivalently its energy dissipation capacity, defined as the area under the stress-strain curve. If we follow the definition of toughness and simplify the stress-strain response as a constant plateau at $\sigma_p$ followed by a linear decay after strain localization, $T$ can be directly approximated by a simple expression:
The quantitative agreement between this relationship and the toughness from the simulations is remarkable considering the simplicity of the model (Fig. 5(a)). Notably, Eq. (5) implies that the toughness of MLG saturates at large $L_o$. Defining the critical length as before following the 90% criteria, we get $L_c^T \approx 400$ nm or $\sim 8L_c^P$ for $n_s = 1$. Further increase in $L_o$ beyond this critical length will not enhance the toughness significantly. This critical overlap length $L_c^T$ is an extremely relevant design guideline for the design of tough MLG papers. Our simulations show that the theoretical maximum toughness ($\sim 7 \times 10^3$ MJ/m$^3$) or specific energy dissipation ($E_p^* \sim 3$ MJ/kg) that can be achieved by the MLG is roughly an order of magnitude higher comparing to other engineering materials, such as steel or Kevlar armor.$^{19,29,30}$ Our simulations also suggest that the theoretical maximum toughness values are much higher than those recently measured by supersonic ballistic experiments on MLGs ($E_p^* \sim 1.5$ MJ/kg).$^{19}$

It is important to notice that the energy of the intralayer interactions (i.e. interactions between different sheets in the same layer) is negligible compared to the surface energy between sheets in different layers. This is because within each layer, the side-by-side interaction energy of the graphene sheets is much smaller than the strength of the covalent bonds within each sheet, so its contribution to features such as modulus and strength are negligible. Most of the load transfer occurs through shear transfer across layers rather than through serial force transfer between sheets in the same layer. Accordingly we have not included these edge effects in the analysis presented above. Also, although we considered only monocrystalline graphene for our sheets, the presence of grain boundaries will most likely not change our results, as the energy to break those bonds will still be much larger than the shear barriers imposed by the weak van der Waals
interactions, maintaining the failure mode by deoverlapping unchanged. These assumptions will need to be revisited if the system under study includes cross-links that strengthen the intralayer interactions or the interfaces.

Our findings provide several strategies for improving toughness. First, given that $L_c^T$ is directly related to $L_c^P$, the failure strain and toughness can be controlled by tuning the stiffness $E$ of the constituent sheets and the interfacial adhesion energy $\gamma$ between the sheets by functionalizing the surface. The prediction of $L_c^T$ as a function of $E$ and $\gamma$ is shown in Fig. 5(b). There are various pathways through which these material properties could be tailored. For example, by oxidizing the carbon atoms at different functionalization levels (the fraction of oxidized carbon atoms), forming the so-called graphene oxide (GO) paper, the enhanced interfacial adhesion energy and reduced modulus compared to pristine graphene can be utilized to design these length scale parameters and thus materials toughness.$^{4,31}$ Second, the interlayer shear response of graphene, such as shear modulus, interaction energy, and frictional force, depends on the stacking orientation due to the hexagonal lattice feature,$^{18,32,33}$ and thus it would be expected that $L_c^P$ and $L_c^T$ also varies for different stacking order (e.g. Bernal vs. turbostratic). The scaling of $L_c^P$ relations for dependence on factors such as critical shear strain and adhesion energy can be readily used to predict $L_c^T$ for different stacking orientations. Such predictions may enable the design of tougher protection materials, since recent projectile penetration experiments have already shown multilayer graphene to be excellent for this purpose due to its superior energy absorption capacity.$^{19}$ Our findings on mechanical behaviors of MLG with staggered architecture can be applied to design materials with dramatically enhanced toughness and energy adsorption. Moreover, our results of the size-dependent mechanical response and failure mechanism of MLG can be adapted to other 2D materials whose interfacial properties are governed by vDW
interactions as well, such as phosphorene, MoS	extsubscript{2} and hBN.\textsuperscript{34-37} For instance, for the case of phosphorene which has a rougher potential energy landscape and a lower sheet stiffness compared to graphene, its $L^C$ of a multilayer system will be smaller than that of graphene based on our predictions as shown in Fig. 5(b). Future work on theoretical description of plastic stress associated with the dynamic friction process will be a key to quantitatively predicting the mechanical responses of multi-layer assemblies of 2D nanomaterials at large deformation regime.

**Conclusions**

In summary, we have performed CG-MD simulations to provide unprecedented quantitative understanding of the dependence of the mechanical properties of MLG on interlayer overlap length $L_o$, including the elusive large deformation and failure regimes where continuum models often break down. We have developed analytical expressions for a hierarchy of critical overlap lengths that govern the strength ($L^S_c \approx 17$ nm), failure strain and plastic stress ($L^P_c \approx 50$ nm), and the toughness ($L^T_c \approx 400$ nm) of MLGs. Particularly interesting, we found that the toughness of MLGs, and therefore its ability to dissipate energy, does not significantly increase with overlap length for very large $L_o$ beyond $L^T_c$. In fact, 90% of the maximum theoretical toughness can already be achieved for overlap lengths $\sim L^T_c$, which can be considered as an extremely important design parameter to design tough multilayer graphene systems. More importantly, we have elucidated the functional dependence of these critical length scales with the mechanical properties of the constituent materials, thus providing transferrable guidelines to assemblies with similar multi-layered staggered architectures based on 2D materials with different interfacial or elastic properties, such as phosphorene, MoS	extsubscript{2} and hBN.
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References

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Figure 1. (a) Mapping scheme from the atomistic lattice (gray) to the coarse-grained structure (blue). (b) Schematic of staggered multi-layer graphene (MLG) with an overlap length of $L_o$. (c) Typical stress-strain response of MLG with $L_o \approx 155$ nm under uniaxial tension. Inset highlights the linear elastic response at small strain. (d) The tensile strength $\sigma_m$ and plastic stress $\sigma_p$ of the MLG as a function of $L_o$. The dashed curves are the “shear-lag” model fits to the data. $L_c^e$ and $L_c^p$ are the critical overlap lengths for $\sigma_m$ and $\sigma_p$, respectively, beyond which the $\sigma_m$ and $\sigma_p$ start to saturate.
Figure 2. (a) Schematic of simplified bilayer model used to quantify the constitutive behavior of graphene with single interface. (b) Force-displacement ($f$-$u$) curve obtained from the pulling test. The resultant simplified constitutive law consists of a plateau force $f_p$ regime and a linear decay regime as the remaining contact length is less than $L_c^p$, which is denoted by the bilinear solid lines.
Figure 3. Dependence of the critical length $L_c$ on (a) sheet stiffness $E$ and (b) interlayer adhesion energy $\gamma$ determined from the pulling test of the bilayer model.
Figure 4. (a) MD snapshots of MLG deformation process. (b) Schematic of sequence of different states during the deformation process. (c) Kinetic model prediction and (d) MD simulation results of failure strain $\varepsilon_f$ as a function of overlap length $L_o$. The inset of (c) shows the schematic of energy landscape used to predict strain localization in the simplest bilayer system. The solid curves in (c) and (d) show the scaling of $n$ by assuming the localization occurs when the remaining contact length of each part reduces to $L_c^P$. 
Figure 5. (a) Normalized toughness $T/\sigma_p$ as a function of overlap length. The solid curve shows the prediction from Eq. (5) for $n_s = 1$. (b) The predicted critical length $L_c^T$ for the toughness as a function of sheet stiffness $E$ and interlayer adhesion energy $\gamma$. 
Three critical length scales govern the deformation mechanisms and constitutive response of multilayer graphene.