Melting of Partially Fluorinated Graphene: From Detachment of Fluorine Atoms to Large Defects and Random Coils

Sandeep Kumar Singh,† S. Costamagna,‡ M. Neek-Amal,*† and F. M. Peeters†

Universiteit Antwerpen, Department of Physics, Groenenborgerlaan 171, BE-2020 Antwerpen, Belgium., and Facultad de Ciencias Exactas Ingeniería y Agrimensura, Universidad Nacional de Rosario and Instituto de Física Rosario, Bv. 27 de Febrero 210 bis, 2000 Rosario, Argentina.

E-mail: mehdi.neekamal@gmail.com

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*To whom correspondence should be addressed
†Universiteit Antwerpen, Department of Physics, Groenenborgerlaan 171, BE-2020 Antwerpen, Belgium.
‡Facultad de Ciencias Exactas Ingeniería y Agrimensura, Universidad Nacional de Rosario and Instituto de Física Rosario, Bv. 27 de Febrero 210 bis, 2000 Rosario, Argentina.
Abstract

The melting of fluorographene is very unusual and depends strongly on the degree of fluorination. For temperatures below 1000 K, fully fluorinated graphene (FFG) is thermomechanically more stable than graphene but at \( T_m \approx 2800 \) K FFG transits to random coils which is almost twice lower than the melting temperature of graphene, i.e. 5300 K. For fluorinated graphene (PFG) up to 30% ripples causes detachment of individual F-atoms around 2000 K while for 40-60% fluorination, large defects are formed beyond 1500 K and beyond 60% of fluorination F-atoms remain bonded to graphene until melting. The results agree with recent experiments on the dependence of the reversibility of the fluorination process on the percentage of fluorination.

Introduction

Several distinct atomic arrangements of adatoms (fluor, hydrogen, chlorine, etc) have been proposed for tuning the electronic properties of graphene (GE).\(^1\)\(^2\)\(^3\) The main advantage of using fluor is that C-F bonds are energetically more stable than e.g. C-H ones\(^4\)\(^5\) because F-atoms possess larger binding and desorption energies to C than H-atoms.\(^1\) Furthermore, fluorination is easier to control via temperature and by reactant gases leading to reproducibly precise C/F stoichiometries\(^6\)\(^7\)\(^8\) In the presence of F adatoms C-bonds in graphene transit from \( sp^2 \) to \( sp^3 \) hybridization, which turn the conjugated, graphitic C-C bonds into single C-C bonds. The lattice structure results in an Angstrom scale out-of-plane buckled shaped membrane known as chair configuration\(^4\) that influences the high temperature stability of FG. Owing to the properties mentioned above, a complete understanding of the thermal behavior of the FG sheet is hence very important.

Previous studies have shown that, different from graphene, fully fluorinated and also hydrogenated\(^9\)\(^10\) graphene, are more rigid for temperatures up to 1000 K. The situation beyond this temperature and up to melting is not yet fully understood. Raman spectroscopy experiments revealed that the fluorination time and the \( N_C/N_F \) ratio (where \( N_C \) and \( N_F \) are the number of carbon and fluor atoms, respectively) are two important key factors in the preparation of FG.\(^11\)\(^2\) It was
found that the process can be reversed for low coverage and during the fluorination process large membrane holes could appear due to losses of C atoms even at room temperature.\(^{11,12}\) Despite this macroscopic information, the microscopical features and temperature stability is not understood. The aim of this letter is to provide such a detailed microscopic understanding for different level of F coverage and to explain recent experiments. We investigate the importance of the \(N_C/N_F\) ratio and the modifications induced by vacancy defects on the stability of FG and the melting process.

**Simulation Method**

Molecular dynamics (MD) simulations using reactive force fields (ReaxFF\(^{15-17}\)) present in the large-scale atomic/molecular massively parallel simulator (LAMMPS) code\(^{18}\) are used. ReaxFF is a bond-order-dependent potential that describes bond formation and dissociation. Many body interactions such as the valence angle and torsional interactions are formulated as function of bond order. Non bonded interactions, e.g. Coulomb and van der Waals interactions, are included for all pair atoms which are not well treated usually by quantum mechanical methods such as density functional theory (DFT).\(^{19}\) Excessively close range interactions are avoided by shielding. To account for the van der Waals interaction, a distance-corrected Morse-potential is used. ReaxFF uses the geometry dependent charge calculation scheme (EEM scheme) of Mortier et al.\(^{20}\) Intratomic contributions of atomic charges which is required to polarize the atoms, are included in the energy scheme, which allows to apply the force fields to ionic compounds.\(^{21}\) The system energy in ReaxFF consists of a sum of terms:

\[
E_{sys} = E_{bond} + E_{under} + E_{over} + E_{tp} + E_{val} + E_{pen} + E_{tors} + E_{conj} + E_{vdW} + E_{Coulomb}
\]

A detailed general description of each of these terms and their functional forms can be found in the original work\(^{15}\) and for fluorinated graphene in our previous paper.\(^{9}\) ReaxFF is able to predict very precisely the equilibrium C-F bond length and the C-F bond dissociation energy, close
to DFT based energies and geometries for a number of molecules and reactions. We considered square shaped computational unit cells having $N_C = 1008$ C-atoms in the graphene sheet with partial fluorination up to fully fluorinated graphene (FFG). The simulations were performed in the NPT (P=0) ensemble with periodic boundary conditions. Temperature was maintained by the Nosé-Hoover thermostat and the MD time-step was taken to be 0.1 fs. The partial covered samples (PFG) were designed by adding randomly F atoms equally distributed on both sides of an initially flat graphene lattice such that each carbon gets maximum one fluorine atom. The dependence of the averaged lattice parameter with increasing F concentration was found to be in agreement with recent results.

To account for the melting transition we analyzed the variation of the total potential energy $E_T$ per atom with temperature identifying partial contributions from C-atoms ($E_C$) and F-atoms ($E_F$). The Lindemann criterion was used to characterize the ordered state by considering the modified parameter $\gamma$, used previously for 2D systems.

**Results**

We first analyze the case of FFG. Figures 1(a,b) show the variation of the potential energy per atom of carbon and fluor atoms, i.e. $E_C$ and $E_F$ respectively, with time at 2800 K and 2900 K. The sharp increase (decrease) in $E_C$, which is about 4.5% ($E_F$ about 10%), is a signature of melting at 2900 K. During melting, (10 ps) the number of six-membered rings (R6) of the crystalline phase is reduced (Fig. 1(c)) and chains composed by single C-atoms bonded to F-atoms are formed (Fig. 1(d)). The melting temperature $T_m = 2800$ K is further confirmed by the Lindemann parameter $\gamma$ (Fig. 1(e)). Due to the strong covalent nearest-neighbor C-C interaction $\gamma$ increases linearly up to close the melting temperature where it diverges.

After melting the C-atoms in the single chains remain bonded to the F-atoms, i.e. a spaghetti of 3D-polymers constructed from C-F monomers (a snap shot of the molten FFG will be shown in Fig. 1(c)). For large simulation time the molten structure is composed of C-chains which form
an entangled three-dimensional network which looks more like a polymer gel than a simple liquid.\[13\] The larger reduction in $E_F$ suggests that F atoms prefer to be bonded to carbon atoms of rings/chains rather than the carbons of GE. This indicates that in experiments during fluorination F atoms first prefer to be bonded to defective regions, where the environment is more similar to a C-chain. The radial distribution function (rdf) indicates that the C-C distance in chains is shorter and after melting (double bonds appear) only one significant peak remains (see Fig. \[1\]e)). However the C-F rdf in Fig. \[1\]e) shows that after melting there are two significant peaks which correspond to the appearance of -C-F$_2$ and -C-F bonds. In the case of PFG, conjugated C-C double bonds in the non-fluorinated parts coexist at low temperatures with covalent C-F bonds in corrugated fluorocarbon regions.\[26,27\] However the behavior at higher temperature is more complicated. We found three different melting processes depending on the fluor percentage:

**Low fluorination**

(10-30%). In Figs. \[2\] (a,b) we display $E_C$ and $E_F$ against temperature. The main feature here is evaporation of F-atoms and the formation of large rings, e.g. R10, R14. While $E_C$ increases linearly against temperature due to a smooth expansion of the C-C inter-atomic distance (small deviations are due to the formation of large rings), because of C-F bond breaking, $E_F$ instead behaves non-linear (and obtains higher energies: F atoms are in the gas phase, see Fig. \[4\]a)) with Temperature (indicated by the arrows). Detachment starts around 2000 K and continues until almost all the F-atoms are removed from the membrane.\[29\] With further increase in temperature, the Lindemann criterion indicates that melting occurs at 5100 K, 4500 K and 3500 K for 10, 20 and 30% fluorination, respectively and F-atoms are less likely to be bonded to the GE. Thus, the linear increase in $E_C$, Fig. \[2\]a), shows that the PFGs with $N_F/N_C \leq 30\%$ turns into pristine graphene (with a few number of rings) around 3000 K.
Intermediate fluorination

(40-60%). In this regime at about $T \approx 1500$ K, $E_C$ shows small deviations from the linear behavior due to the formation of large defective rings (Fig. 2(c)). In contrast to the case of low $N_F/N_C$ ratio, $E_F$ shown in Fig. 2(d), becomes more negative since F-atoms instead of being detached from the sheet, now break the original local bonds and are transferred to the edges of the rings (a snapshot of this configuration is shown in Fig. 4(b)). Then, at the melting temperature clusters of C-F are formed near the defects and they detach from the sheet. The microscopic configurations consist of a mixture of 2D and 3D phases.

High fluorination

(70-90%). Finally, in the high-fluorination regime with increasing temperature large ripples develop while F atoms remain bonded to graphene until melting occurs. Now, $E_C$ receives positive contributions due to the breaking of C-C bonds at melting (Fig. 2(e)) and since the C-F bond becomes stronger $E_F$ is reduced (Fig. 2(f)). The microscopical features of melting in this case posses similar characteristics as FFG. Notice that arrows in Fig. 2 are not pointing necessarily to the melting transitions, instead they indicate mostly evaporation (b) and defect formation thresholds Figs. 2(d,f). The reduction in $E_F$ indicates the appearance of polymers consisting of C-F monomers. The average melting temperatures $T_m$ were estimated from the behavior of the Linde-mann parameter. Starting from 60%, when we increase further the F-content $T_m$ is seen to pass a minimum around 70% and then increases due to the suppression of long wave-length ripples.9

We depict the energy variation during melting for a typical case of 90% fluorination in Figs. 3(a,b). Here, the number of hexagons approach zero and many single chains are formed as shown in Figs. 3(c,d). From the slope of $E_C$ versus time we obtain the rate at which the hexagonal crystal structure of the sample becomes lost (e.g. dashed blue curve in Fig. 3(a) shows $\delta = dE_C/dt$). Values of $\delta$ are shown in Fig. 3(e) together with $T_m$. The main result is that the larger the ratio $N_F/N_C$, the higher the melting temperature and the faster that melting takes place.
**Effect of vacancies on the melting of FFG**

It is important to study the effect of the presence of atomic vacancies in FFG on the melting temperature. We performed several additional simulations for FFG with \( p \) number of vacancies which were randomly distributed over FFG, i.e. \( N_C \rightarrow N_C - p \) and \( N_F \rightarrow N_F - p \). The presence of atomic defects in the GE sheet makes it less stiff and consequently results in a lowering of the melting temperature, see Fig. 3(f). The melting temperature is fitted by the red line \( T_m(N)=a+bN \), where \( a=(2780 \pm 75) \) K, \( b=(-60 \pm 6) \) K and \( N \) is the number of vacancies. These results clearly indicate that \( T_m \) decreases sharply as the number of defects in the FFG increases.

**Discussion**

FFG is an insulator with a large band gap while graphene shows metallic properties, thus one naturally expects the transition from GE to FFG by increasing the ratio \( N_F/N_C \). This gives the appropriate melting trend for a typical metal to an insulator. However, the melting of GE and FFG is different because of their 2D-nature. The existence of flexural phonons makes GE and FFG also different from typical 3D crystals and the bending rigidity is a temperature dependence parameter. Although at low temperature FFG is thermo-mechanically more stable than GE, by increasing temperature, due to the excitation of the vibrational C-F mode, in FFG the entropy term increases faster with \( T \) which is being responsible for the observed lower melting temperature.

In PFG the distribution of masses through the system is non-uniform, hence the vibrational frequencies are not well defined and are position dependent. This randomness in the system produces very large out-of-plane fluctuations even at low temperature. This broadening in the frequency range brings the system closer to the melting transition point.

Moreover, for a low ratio \( N_F/N_C \) when we heat the system F-atoms are evaporated in order to reduce the total energy. Then, the system behaves like pristine graphene and evaporated F atoms have no chance to be re-bonded to the system. For intermediate ratio \( N_F/N_C \), the concentration of F atoms in some random domains make the system unstable due to the growth in the mean square
value of the height fluctuations $\langle h^2 \rangle$ resulting in the formation of ring-defects with increasing temperature and the melting of this new defected FG is more complex. Finally, for high fluorination we deduce that the melting temperature is proportional to the coverage percentage, e.g. decreasing the coverage percentage from 100% to 90% (80%) decreases the melting temperature with about 16% (30%).

Let's now compare our results with recent experiments.\[1,2\] Raman spectra of graphene after exposure to atomic F shows dramatic changes induced by fluorination: the D peak emerges at early times of fluorination and the 2D peak is suppressed which indicates that F atoms are bonded to C atoms and atomic defects appear. The process of fluorination has been shown to be reversible only when the exposure time is relatively short (< 20 hours) and the concentration of F atoms is presumably low. FFG is obtained only for longer exposure time, beyond 30 hours, where the evaporation of F atoms and the restoring to pristine graphene is impossible. In our simulations, we found that for low coverage percentage (less than 40%) increasing the temperature causes evaporation of F atoms and almost pristine graphene can be recovered in agreement with experiment. However, percentage beyond 40% are related to long time fluorination (> 20 hours) where the process was reported to be irreversible. The presence of vacancies decrease dramatically the melting temperature. Indeed, in experimental PFG samples containing a few vacancies it was found that when they were heated up to much lower temperatures (~500-700 K) it was possible to restore to not-perfect pristine graphene.\[\text{1}\]

**Conclusion**

The phase diagram displayed in Fig.\[4\] summarizes our results: green squares refer to the transition temperature and blue circles indicate the starting point of evaporation of F-atoms. Two insets (b) and (c) show a top views of the system after the transition points, inset (a) shows top view of a snap shot of a system which lost fluor and inset (d) shows a side view of a typical PFG at low temperature. The PFG above the blue symbols for $N_F/N_C < 40\%$ looses fluor and become pristine.
graphene with some defects and above the green line it transits to the liquid phase, however for $N_F/N_C \sim 40-60\%$ above the green line and below ($\sim 5000$ K) PFG does not loose F and very slowly transits to the liquid phase so that below 5000 K the liquid and the solid phases coexist. For larger $N_F/N_C \sim 60-100\%$ above the green curve PFGs rapidly transits to a 3D liquid phase. The minimum melting temperature is found to be around 70\% fluorination. The vertical dashed lines separate three different regions for different percentage of F atoms. Our findings are therefore consistent with the experimental reversibility of the fluorination process in single layer graphene.

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Figure 1: (Color online) Fully fluorinated graphene. (a) $E_C$, (b) $E_F$, (c) number of six-membered rings $R_6$, and (d) chains (nc) with more than three connected twofold-coordinated atoms, as a function of time for $T=2800$ K and at $T=2900$ K where melting occurs between these temperature range. (e) Modified Lindemann parameter $\gamma$ versus temperature. (f) C-C and (g) C-F radial distribution function.
Figure 2: (Color online) Dependence of $E_C$ (upper panels) and $E_F$ (lower panels) versus temperature for the three regimes of melting indicated. Curves were shifted for a better comparison (i) for $E_C$: 30, 40 and 60% fluorination, curves are shifted by 0.05, 0.45 and -0.5 eV/atom, respectively. (ii) for $E_F$: for 20, 30, 70, 80, 90, and 100% fluorination, curves are shifted by 0.3, 0.4, 0.65, 0.7, 0.75, and 0.8 eV/atom, respectively). Notice that $E_F$ increases in (b) while it decreases in (d) and (f) which is an indication of the evaporation of fluor in “low” concentration regime and ring/defect formation and their saturation by F in “intermediate" and “high" concentration regimes".
Figure 3: (Color online) Partially fluorinated graphene 90%. (a) Total energy of C-atoms, (b) total energy of F-atoms, (c) number of six-membered rings $R_6$, (d) number of chains ($n_c$) with more than three connected twofold-coordinated atoms, as a function of time at $T=2300$ K (below melting) and at $T=2400$ K where melting occurs, (e) $\delta$ and $T_m$ versus percentage of F atoms, and (f) variation of the melting temperature $T_m$ against the number of defects in fully fluorinated graphene.
Figure 4: (Color online) Melting phase diagram for fluorinated graphene. Circular symbols refer to the evaporation of F atoms (blue circles). The insets show the top view of the simulated FG before (a) and after melting (b,c). The inset (d) is a side view of the simulated PFG with $N_F/N_C = 80\%$. 