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1 **A reactive molecular dynamics study of the surface catalysis and**
2 **ablation behavior of graphene sheets colliding by hyperthermal**
3 **atomic oxygens**

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9
10 **Abstract**

11 Carbon-based composite materials, whose mechanisms of surface catalysis and
12 ablation during the impact of oxygen atoms remain unclear, are widely used in the
13 aerospace field due to their light weight, ideal physical and chemical properties. In this
14 study, the surface catalysis and ablation behaviors during the erosion process of
15 hyperthermal atomic oxygens was investigated by molecular dynamics (MD)
16 simulation method with ReaxFF potential. Seven models were established to study the
17 impact of the energy flux density for the energetic oxygen atoms, the presence of
18 multiple layers beneath the graphene sheet, and the effect of three different graphite
19 surfaces, namely graphite basal plane, armchair edge surface and zigzag edge surface,
20 respectively. The simulation results show that for all the seven models, the adsorption
21 of oxygen atoms dominates at the beginning for all models, generating O₂ molecules
22 through the surface catalytic recombination effect. The ablation rate of graphene surface
23 accelerates significantly as oxygen energy flux density increases due to the higher
24 system temperature. The multi-layered graphite slab can be etched by hyperthermal
25 atomic oxygen with a ‘layer-by-layer’ phenomenon. The more graphene stacks
26 appearing below the top layer, the slower the ablation rate due to the fact that additional
27 graphene layers can act as a heat sink to slow down the temperature rise by conduction
28 due to oxygen-atom collisions. Graphite surfaces with AC surfaces show the largest

1 etching rate and basal presenting the lowest etching rage, revealing the fact that the
2 binding energies of the AC edge tends to be weaker than those of the ZZ edge, causing
3 that the AC surface model is difficult to form stable functional group structures to resist
4 the etching of high-energy oxygen atoms than the ZZ surface. This research advances
5 our understanding of the detail surface catalysis and ablation mechanisms during the
6 erosion process of graphene sheets from the prospective of atomistic scale.

7 **Keywords**

8 Reactive Molecular dynamics (RMD) simulation, hyperthermal atomic oxygens,
9 surface catalysis and ablation, graphene sheet

11 **1 Introduction**

12 Carbon-based materials have been applied as heat shields in spacecraft
13 applications due to their desirable physical and chemical properties [1, 2]. At low-earth-
14 orbit (LEO) altitude, the dominant component of this rarefied atmosphere environment
15 is atomic oxygen with a number density of around 10^{15} atoms/m³. This atomic oxygen
16 density, for instance, with a combination of the orbital velocity of 8 km/s for the
17 spacecraft would yield an oxygen flux of 10^{19} atoms/(m²·s) at a mean collision energy
18 of 5 eV [3]. Thermal protection materials would undergo complex physical and
19 chemical response to such conditions including surface catalysis and ablation reactions
20 [4-7]. Aiming to identify the material damage and characterize the aerodynamic heating
21 due to the collisions of hyperthermal atomic oxygens accurately, a fundamental
22 understanding of the surface catalysis and ablation reaction mechanisms for carbon-
23 based materials is essential for thermal protection system design of with space based
24 applications.

25 Recent experimental works using multi-scale imaging techniques have provided
26 valuable information for such a high-energy collision process with a focus on carbon-
27 based material surfaces. Murray et al. [8, 9] conducted hyperthermal scattering
28 experiments of both atomic oxygen (O) and molecular oxygen (O₂) collisions on High

1 Orientation Pyrolytic Graphite (HOPG) surfaces to reveal the mechanisms of carbon
2 oxidation over a wide temperature range. Nicolson et al. [10] further reported the effect
3 of temperature on corrosion rate and surface roughness. These experimental results
4 indicated that the ablation rate of graphite was more dependent on the graphite surface
5 temperature, which was controversial with other experimental conclusions [11-13]
6 implying the exposure impacts of atomic oxygens. From the perspective of
7 experimental design, many of key process parameters are closely interrelated, such as
8 temperature, pressure and energy flux of incoming atomic oxygens, making it
9 challenging to change one only while keeping others unchanged, which in turn makes
10 an identification of the key mechanism difficult. In parallel with these experimental
11 research, theoretical investigations aimed at understanding the surface catalysis and
12 ablation process of carbon-based materials upon hyperthermal oxygen atom collision
13 have also been carried out in the past. Arrhenius type expressions have been excessively
14 applied to fit with the experimental reaction rates, while the possible physiochemical
15 mechanism insights are still hard to achieve aiming to serve as a fundamental guide to
16 design materials.

17 Density Functional Theory (DFT) based modelling has been shown its capability
18 of providing such detailed physical and chemical information during the interactions
19 between carbon element and energetic atomic/molecular oxygen. Studies on the
20 interaction between graphite surface and high-energy oxygen atoms [14-17] reported
21 that oxygen molecules were generated during the collisions due to the surface catalysis
22 effect through the Eley-Rideal (E-R) mechanism, which was much more important than
23 the Langmuir-Hinshelwood (L-H) mechanism. The formation mechanism of CO and
24 CO₂ was further investigated with DFT method by Paci et al. [18-21], reporting that
25 oxygen atoms were adsorbed on the graphite surface to form epoxy functional groups
26 firstly, which were then diffused to form ketone groups or carbonyl groups, resulting
27 different formation of CO and CO₂. It can be found that these ab initio simulations
28 based on DFT can provide accurate information during the physical interaction and

1 chemical reaction processes, but their associated computational cost limits their
2 applicability to systems with sizes of only a few angstroms and time scales of a few
3 picoseconds.

4 With rapid advances in high-performance computing systems in the last few
5 decades, the classical Molecular Dynamics (MD) simulations have shown their abilities
6 to simultaneously model large system sizes for a long time. Thus MD method has been
7 recently employed to study the mechanical and thermal properties for graphite and
8 graphene materials [22-24] and their complicated physical interactions with other
9 species [25, 26] at the atomistic level. These MD simulations, however, have not
10 elucidated the chemical reaction mechanisms. With the continuous development of
11 force field potentials, an empirical reactive potential ReaxFF was recently proposed by
12 van Duin et al. [27] to simulate the physics, chemistry and dynamics of various
13 processes for graphitic materials [28-30]. Poovathingal et al. [31] studied the oxidation
14 of hyper-thermal oxygen atom beams with HOPG based on reactive molecular
15 dynamics (RMD). The simulation predicted that the composite reaction occurred on the
16 graphene sheet, but the carbon removal reaction only occurred on the edge of the
17 corrosion pit. Srinivasan et al. [32] simulated the ablation process of high-energy
18 oxygen atoms colliding with the diamond surface in a low-earth orbit (LEO)
19 environment, observing that a variety of functional groups such as ethers, peroxides,
20 oxygen radicals and dioxins were formed on the diamond surface. Ermakov et al. [33]
21 later performed a simulation process of exposing three layers of graphene to atomic
22 oxygen, and reported that the graphene gasified uniformly layer by layer. Majumder et
23 al. [34] studied the energy transfer when O₂ collided with the graphite surface. It is
24 found that the average energy transferred to the graphite surface and the average energy
25 retained in the O₂ translation were linearly related to the initial translation energy. Liu
26 et al. [35] simulated the bombardment of single-layer graphene films by different
27 energetic atoms. The results showed that the incident energy played an important role
28 in the evolution and final shape of graphene defects. Bai et al. [36] systematically

1 explored the effects of incident angle and incident energy on defects and vacancies by
2 using molecular dynamics with reactive force fields. It is found that oxygen ions with
3 an incident angle of 70° showed the highest ion replacement rate. Qiao et al. [37] fine-
4 tuned the theoretical graphene oxide (GO) model based on RMD simulations and
5 experimental characteristics. The scattering process of N_2 impinging on graphite was
6 also simulated to investigate the effects of angular distribution of N_2 , the average
7 translational energy and rotational energy, and the number of rebounds [38, 39]. It also
8 reported the simulation of other materials impacting graphite, exploring the effects of
9 nickel pellets on graphene at the supersonic velocity, and the dynamics of H_2O , CO_2 ,
10 and glycine (GLY) colliding with graphite [40-42]. These RMD simulations suggest
11 that ReaxFF potential allows for the simulation of chemical reactions on large systems
12 while retaining most of the accuracy of DFT, which can be used as an effective tool to
13 screen and characterize materials for applications in extreme environments.

14 Though previous RMD studies have advanced our understanding of the collisions
15 between atomic oxygen and graphitic materials, the surface catalysis and ablation
16 mechanism still remains unclear due to its complex physiochemistry nature during this
17 gas-solid interaction process. In this work, RMD simulation method with ReaxFF
18 reactive force field has been employed to gain an atomically detailed understanding
19 through seven independent models with a twofold purpose, firstly using Model I to
20 demonstrate the suitability of the choice of ReaxFF force field and calculation set-up,
21 and secondly to further apply this approach to obtain a deeper understanding of the
22 catalytic and ablative reaction process when the graphene is exposed to hyperthermal
23 atomic oxygen collisions: The impact of the energy flux density for the energetic
24 oxygen atoms is investigated; In addition, a study of the effect of the presence of
25 multiple layers beneath the graphene sheet is carried out; Finally, the effect of three
26 different graphite surfaces is examined, namely graphite basal plane, armchair edge
27 surface and zigzag edge surface, respectively. A detailed description of the
28 computational methods and model constructions is presented in section 2, followed by

1 a discussion of the results in section 3 and conclusions in section 4.

2

3 **2. Molecular Dynamics Simulation Details**

4 **2.1 ReaxFF Reactive Force Field Method**

5 The ReaxFF reactive force field [27] were applied to investigate the hyperthermal
6 collisions of atomic oxygen with graphene in the present work. ReaxFF is a general
7 transferrable bond-order (BO) based empirical potential that uses relationships between
8 bond distance and bond order on the one hand, and relationships between bond order
9 and bond energy E_{bond} on the other to describe bond formation and bond dissociation
10 properly. The system energy E_{sys} in ReaxFF is calculated as the sum of a number of
11 energy terms according to

$$12 \quad E_{\text{sys}} = E_{\text{bond}} + E_{\text{val}} + E_{\text{tors}} + E_{\text{vdWaaals}} + E_{\text{Coulomb}} + E_{\text{lp}} \quad (1)$$
$$13 \quad \quad \quad + E_{\text{over}} + E_{\text{under}} + E_{\text{pen}} + E_{\text{conj}}$$

13 where the valence angle energy E_{val} and torsion angle energy E_{tors} are made functions
14 of bond order so that their energy contributions go to zero smoothly upon bond
15 dissociation; Non-bonded interactions, including intermolecular van der Waals
16 potential E_{vdWaaals} and electrostatic interaction potential energy E_{Coulomb} , are calculated
17 between every atom pair, irrespective of the connectivity; And other correction terms,
18 including energy compensation terms related to lone pair electrons E_{lp} , over-
19 coordination energy E_{over} , under-coordination energy E_{under} , compensation function E_{pen} ,
20 and four-body conjugation energy E_{conj} , are used to describe the fracture and formation
21 of chemical bonds in different chemical environments properly. Excessively short range
22 interactions are avoided by using a shielding term in the energy expression for the
23 nonbonded interactions. Instead of Ewald summation to calculate long-range Coulomb
24 interactions, ReaxFF uses a seventh-order taper function with an outer cutoff radius of
25 10 Å.

26 The force-field parameters used in the present simulations were extended by
27 Chenoweth et al. [43] as ReaxFF_{C/H/O}, which was parameterized against an extensive

1 training set consisting of atomic charges, bond lengths, valence and torsion angle
2 energies, heats of formation and various hydrocarbon reaction energies. A detailed
3 description of the data included in the parameterization of ReaxFF_{C/H/O} formulation can
4 be found in the original work. ReaxFF_{C/H/O} has been successfully used to study the
5 oxidations of graphene, graphite and diamond subject to high energy oxygen atom
6 bombardment, which results show a good agreement with earlier experiments and first-
7 principle-based calculations [44].

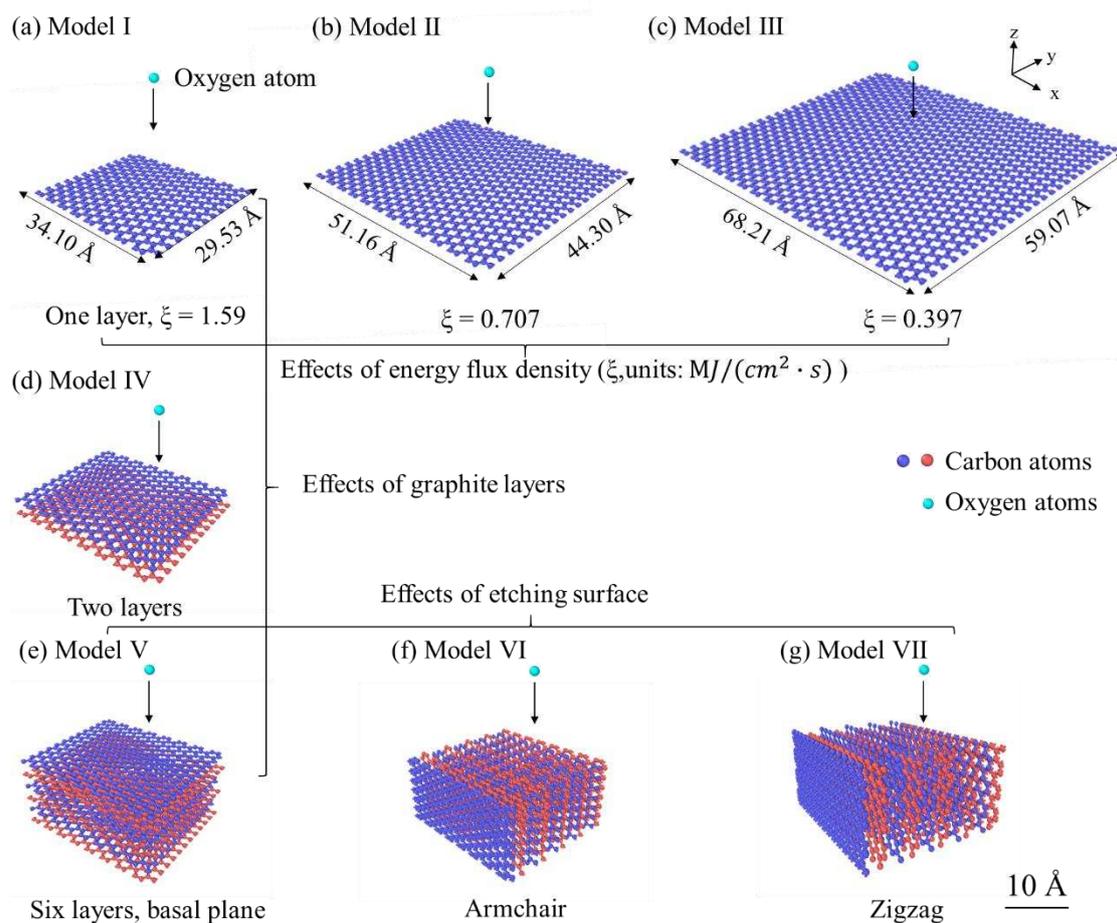
8 **2.2 Model Construction**

9 Firstly, to verify our simulations, a careful benchmarking of the approach was
10 established aiming to reproduce the results of Srinivasan et al. [44], as a reference
11 Model I shown in **Figure 1(a)**. Model I had a graphene sheet with a total of 384 carbon
12 atoms, where the dimensions of the graphene basal plane were $34.10 \text{ \AA} \times 29.53 \text{ \AA}$ in
13 the x - and y -axis directions. The impinging atomic oxygen was initially generated every
14 1.0 ps at a vertical separation of 5.0 \AA above the graphene basal plane, whereas the in-
15 plane coordinates were chosen randomly. The graphene sheet was subjected to collide
16 by these oxygen atoms with their translational energies of 5 eV, which value was also
17 considered as the mean energy when oxygen atoms hit the spacecraft in low-Earth orbit
18 [3]. To ensure that the leaving species after collision did not interfere the surface
19 chemistry, the z dimension of the unit cell was chosen to be 240 \AA . Periodical boundary
20 conditions were used in the directions parallel to the graphene basal plane, while the
21 reflective boundary condition was used in the z -axis direction. To avoid the downward
22 displacement of the graphene sheet due to the continuous bombardment by
23 hyperthermal atomic oxygens, the positions of carbon atoms at each corner of the
24 graphene sheet were fixed to ensure that the center of mass of the graphene in the z -
25 axis direction was remaining stable.

26 Three sets of ReaxFF-based MD simulations, comprised of seven independent
27 models, were carried out, as presented in **Figure 1**. The first set of simulations
28 investigated the effect of energy flux density for the impingement of the hyperthermal

1 atomic oxygens on the reactive events: Through keeping the same impinging energy of
2 the hyperthermal oxygen atoms as Model I, the particle flux in Model II and III was
3 decreased by expanding the graphene sheet structure along each of the coordinate
4 directions of the basal plane by 1.5 and 2 times, respectively. Thus, three different
5 energy flux densities ζ of the impinging atomic oxygens were studied in these
6 simulations, which were calculated as a result of 1.590, 0.707 and 0.397 MW/cm²,
7 respectively. **Figure 1(b)** and **1(c)** shows the expanded sheets, having a total of 864 and
8 1536 carbon atoms. The second set of simulations involved the effect of the presence
9 of multiple layers beneath the graphene sheet on the surface catalysis and ablation
10 process. Two-layer and six-layer graphene stacks in the AB arrangement with an
11 interlayer distance of 3.2 Å [44] were established as shown in **Figure 1(d)** and **1(e)**,
12 denoted as Model IV and V. The carbon atoms are colored in blue and red to distinguish
13 them among different layers. The third set of simulations investigated the continuous
14 bombardment outcome of graphite with different edges by hyperthermal atomic oxygen.
15 Through advancing Model V as a standard graphite basal plane, Model VI and Model
16 VII were built up where the graphite armchair edge and zigzag edge of a model graphite
17 were exposed to hyperthermal oxygen atom collisions normal to the surface, as
18 presented in **Figure 1(e, d, f)**.

19 All the systems were energy-minimized before being used for MD simulations
20 with the LAMMPS package. A time step of 0.10 fs was used for all simulations with
21 the microcanonical (*NVE*) ensemble, because a local increase in temperature along the
22 time due to the collision event is essential in driving the surface catalysis and ablation
23 reactions [44]. A total of 500 ps simulations was conducted for all the models, except
24 for Model I with only 200 ps calculations, as the graphene sheet was shown to get
25 broken into fragments by then. The trajectories of each atomic species for each system
26 were collected every 0.10 ps for all the post-processing process. More information of
27 model construction details in this work can be found in the **Supplementary Material**.



1
2 **Figure 1** The schematic illustrations of initial configurations for the simulated systems

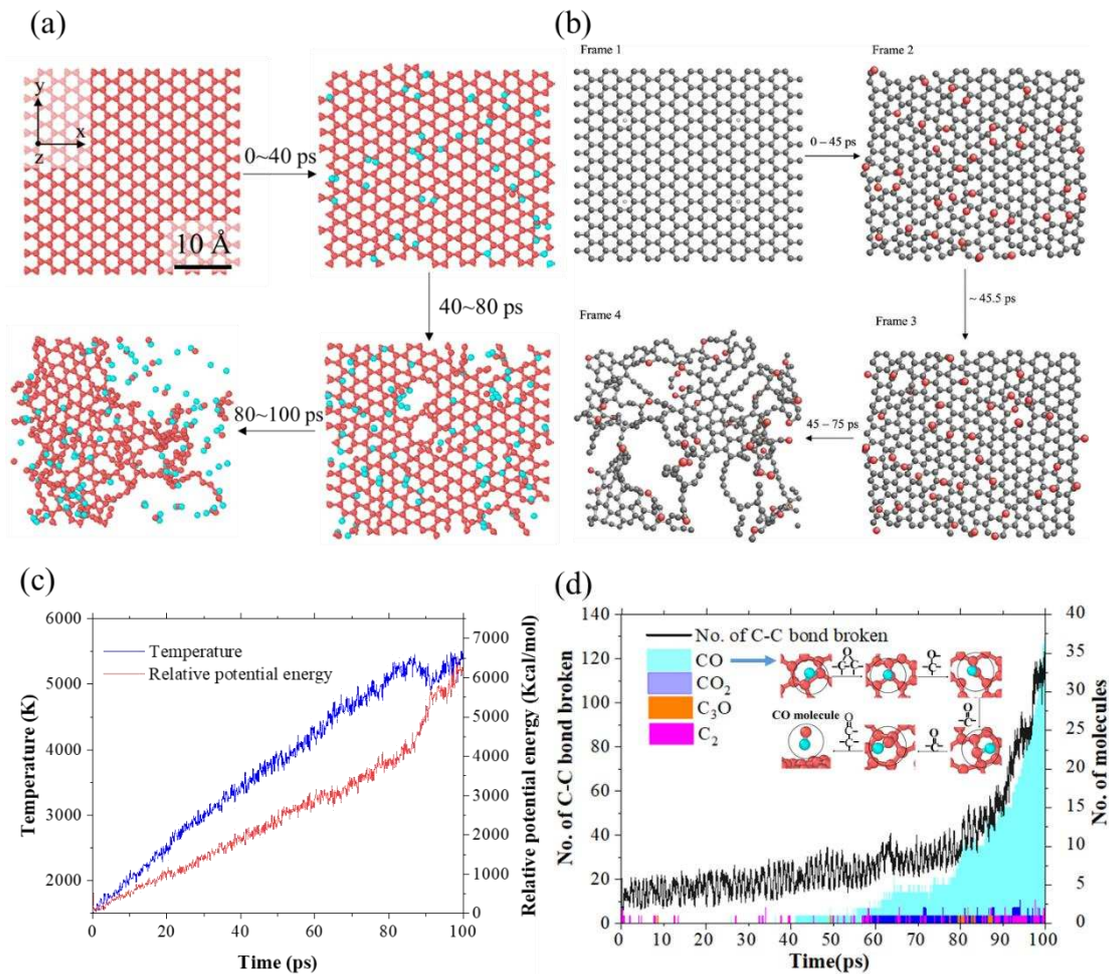
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4 **3. Results and Discussion**

5 Section 3.1 discusses the validity of the choice of ReaxFF_{C/H/O} force field and MD
6 calculation set-up through a benchmark case investigation of Model I. Section 3.2, 3.3
7 and 3.4 report the physical/chemical interactions between the hyperthermal atomic
8 oxygens and the solid surface with the effects of the oxygen energy flux density ξ with
9 1.590, 0.707 and 0.397 MW/cm², the presence of multi layers beneath the graphene
10 sheet surface (a total of 1/2/6 layers) and the graphite surfaces with different edges
11 (graphite basal plane surface, armchair edge surface and zigzag edge surface).

12 **3.1 A benchmark case for validation**

13 As mentioned previously, the validation of our simulations is demonstrated by a
14 careful benchmarking of the approach on Model I aiming to reproduce the results of
15 Srinivasan et al. [44]. The evolution of reaction events leading to the breakup of the

1 graphene sheet is illustrated in **Figure 2(a)**, which is compared with that of Srinivasan
 2 et al. [44]. It can be observed that in the time period from 0 to 40 ps, the impinging
 3 oxygen atoms are adsorbed and bind onto the surface in the forms of epoxide. The hole-
 4 shape defects in the sheet can be visualized at a snapshot of 80 ps from the simulation
 5 results. With rapid development of the defect formation and growth, the graphene sheet
 6 is broken into fragments by 100 ps. This process shows a good agreement with that of
 7 Srinivasan et al. [44] presented in **Figure 2(b)**, indicating the validity and stability of
 8 our calculation setup.



9
 10 **Figure 2** The detail simulation results of Model I. ((a) The breakup process of the
 11 graphene sheet due to successive atomic oxygen impacts (Carbon atoms are colored red
 12 and oxygen atoms are colored cyan) with (b) a comparison of the result from Srinivasan
 13 et al. [44]; (c) Variations of system temperature and potential energy for Model I; (d)
 14 Number of C-C bond broken and the product analysis as a function of time with an
 15 illustration of the formation process of a CO molecule)

16 During the continuous impingement of atomic oxygens, both the temperature and

1 the relative potential energy (relative to that of the pristine graphene sheet) increase, as
2 depicted from **Figure 2(c)**. It should be noted that the system temperature remains
3 approximated stable from 80 to 100 ps, accompanied with a sharp rise in the potential
4 energy. This indicate that most of the energy input to the system through the addition
5 of successive oxygen atom impacts is consumed by breaking the carbon-carbon (C-C)
6 bonds of the graphene sheet. The evidence can be quantitatively observed from the
7 significant increase of the C-C bond broken number variation as a function of time in
8 **Figure 2(d)**, where an interatomic distance of less than 1.8 Å between two carbon atoms
9 is applied as a criterion for the C-C bond breakage for the graphene film structure [44].
10 The statistics of all the gas species leaving from the graphene sheet are obtained as also
11 shown in **Figure 2(d)**. It can be found that the dominant products during the reaction
12 process are CO molecules, where the generation and release process of a CO molecule
13 is illustrated in **Figure 2(d)**. The production of CO is monitored to occur due to the
14 epoxide formation, migration and ring-opening reactions. A small number of CO₂
15 molecules are observed after around 57 ps, with some C₂ and C₃O as intermediate
16 products.

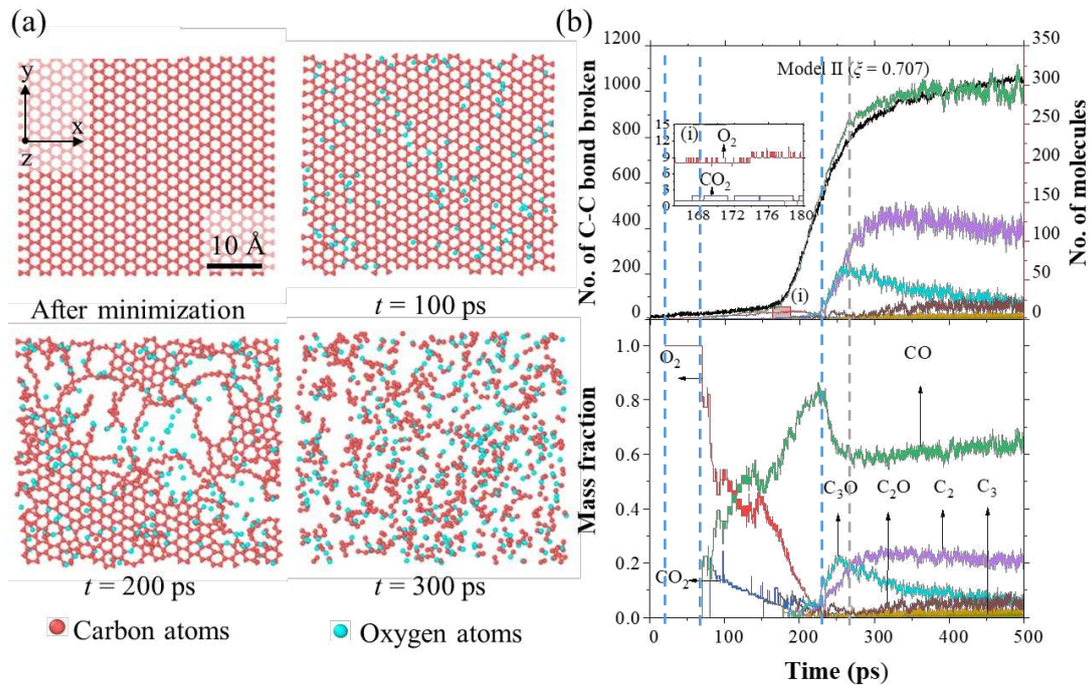
18 **3.2 The effect of impinging energy flux density on the graphene erosion**

19 The impinging energy flux density effect of hyperthermal atomic oxygens on the
20 erosion process of a graphene sheet is reported in this section, which is 1.590, 0.707
21 and 0.397 MW/cm², respectively. This is implemented through expanding the graphene
22 sheet structure in Model I by 1.5 and 2 times, respectively, along each of the coordinate
23 directions of the basal plane to decrease the particle flux, while keeping the same
24 impinging energy of the hyperthermal oxygen atoms as Model I. Thus, three different
25 energy flux densities ξ of the impinging atomic oxygens were studied in these
26 simulations, which were calculated as a result of 0.397 MW/cm² for Model III, 0.707
27 MW/cm² for Model II and 1.590 MW/cm² for Model I, respectively.

28 **Figure 3(a)** illustrates the snapshots of Model II with evolutions of time from 0 to

1 300 ps. It can be observed that, as the impinging energy flux density decreasing from
 2 1.590 MW/cm^2 to 0.707 MW/cm^2 , the breakup rate of Model II seems to be delayed
 3 but still following the same erosion procedure with Model I: The adsorption behavior
 4 of atomic oxygens onto the graphene sheet is dominant with the formation of epoxides
 5 from 0 to 100 ps; Some hole-shape voids/defects in the sheet can be visualized at a
 6 snapshot of 100 ps; The graphene sheet is rapidly broken into fragments by 200 ps with
 7 the development of defect formation and growth, following by completely destroyed
 8 by 300 ps.

9



10

11 **Figure 3** The erosion process of graphene sheet with energy flux densities $\xi=0.707$
 12 MW/cm^2 of the impinging atomic oxygens. (a) the snapshot illustrations of graphene
 13 sheet with the evolution of time (b) the gas species analysis during the erosion process

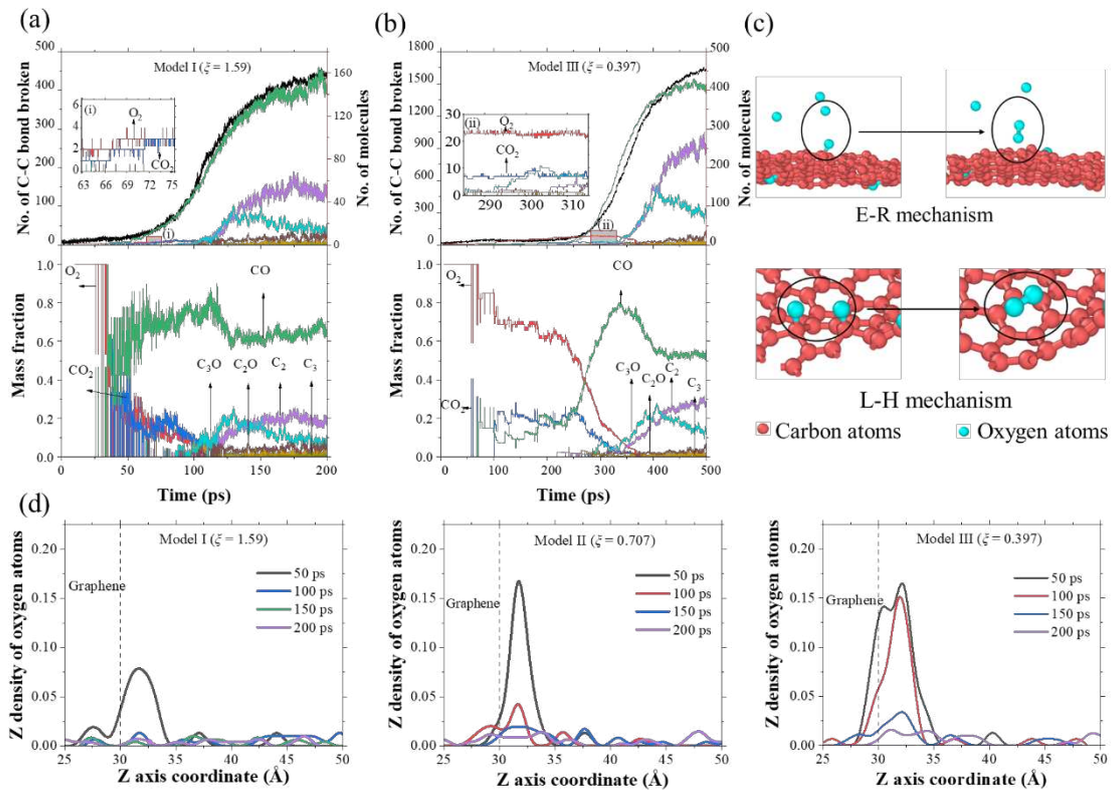
14

15 To further analyze the erosion process of Model II with the incoming energy flux
 16 density of 0.7 MW/cm^2 , the C-C bond breaking analysis, the number of generated
 17 gaseous species and their corresponding mass fractions are quantified as plotted in
 18 **Figure 3(b)**. The formations of the gaseous species are summarized more than those
 19 similar compounds of CO , CO_2 , C_2 , and C_3O observed by Srinivasan et al. [44].
 20 Dominant O_2 molecules amongst all the gaseous species are observed incipiently from

1 around 20 ps: (1) At Stage 1, though a few C-C bonds on the graphene sheet are broken,
2 there is no gaseous species releasing from the graphene sheet before around 20 ps; (2)
3 At the second stage, only O₂ molecules are released approximately from 20 to 70 ps,
4 resulting in whose corresponding mass fraction of 1 (100%); (3) Both CO and CO₂
5 molecules are formed (thereby creating vacancy defects on the graphene sheet) during
6 this third stage of this erosion process, which is also accompanied by rapid consumption
7 of O₂ molecules. The number of the broken C-C bonds for the graphene sheet trends to
8 increase exponentially from around 170 ps. Moreover, almost all the O₂ molecules are
9 consumed at around 220 ps, along with few CO₂ oxidation products are generated
10 afterwards. At this point, the CO molecule becomes the dominant gaseous production
11 with its corresponding mass fraction plot reaches to its peak value of 0.86 (86%). An
12 instance of C₂, C₃, C₂O and C₃O formations can also be monitored at this stage; (4) At
13 Stage 4, the CO yield is monitored to be dominant due to the epoxide formation,
14 migration and ring-opening reactions, where with C₃O and C₂ molecules to a less extent
15 by 270 ps. (5) Finally, with further successive oxygen impingement, a 60% mass
16 fraction of CO molecules and 20% of C₂ molecules are becoming the main composition,
17 accompanied with a remaining small number of C₃, C₂O and C₃O formations.

18 The quantitative analyses of generated gaseous species number and their
19 corresponding mass fractions with evolutions of time for Model I and III with the
20 incoming energy flux density of 1.590 and 0.397 MW/cm² are summarized and plotted
21 in **Figure 4(a, b)**. Compared with Model II in **Figure 3(b)**, the trends of the number of
22 C-C bonds broken, the number of each production molecules and their mass fraction
23 for Model I, III are consistent, while the whole erosion process occurs faster for Model
24 I and slower for Model III. The formation of O₂ molecules are also monitored
25 incipiently for both Model I and III. Through capturing the trajectories of O₂ formation
26 procedures, it can be found that the formation of O₂ molecules are resulted from the
27 surface catalytic recombination effect with both E-R and L-H mechanisms. As
28 illustrated with the snapshots in **Figure 4(c)**, some O₂ molecules are formed by one

1 adsorbed oxygen atom on the graphene surface recombining with another atomic
 2 oxygen from the incoming particle flux due to the collisions, as known as E-R
 3 mechanism; there are also some O₂ molecules formed by the recombination of two
 4 adsorbed atomic oxygens binding onto the graphene surface due to the diffusions, called
 5 L-H mechanisms. It can be noticed that the formation of O₂ molecules due to the surface
 6 catalysis effect is closely related with the incipient adsorption of atomic oxygens onto
 7 the graphene sheet surface. Therefore, the z-density plots of atomic oxygens as a
 8 function of time for Model I ~ III are summarized in **Figure 4(d)**. It can be observed
 9 that with the decrease of the impinging energy flux density from 1.590 to 0.397
 10 MW/cm², the adsorption period of the atomic oxygens onto the graphene surface is
 11 extended obviously. For instance, when the impinging energy flux density decreases to
 12 0.397 MW/cm² (Model III), the number of adsorbed atomic oxygens remains relatively
 13 stable from 50 to 100 ps, following by a trend of descent from 100 to 150 ps due to the
 14 surface catalysis recombination effect and then releasing from the surface.
 15



16

17 **Figure 4** The gas component and z-density analysis during the oxygen impingement

1 for models with different energy flux densities of the atomic oxygens.

2

3 A comprehensive comparison for the effect of incoming energy flux density on the
4 surface catalytic and ablation processes for Model I, II, and III is shown in **Figure 5**.

5 The system temperature profile and the fraction of C-C bond broken profile for each
6 graphene sheet are presented in **Figure 5(a)**. Within the procedure of the NVE

7 simulation of the graphene sheet, the system temperature profiles for all the three

8 models with different incoming energy flux density of atomic oxygens are ascending

9 continuously, which is attributed to these incoming hyperthermal atomic oxygens input

10 to the system. As a result, at the same impingement time, the graphene temperature is

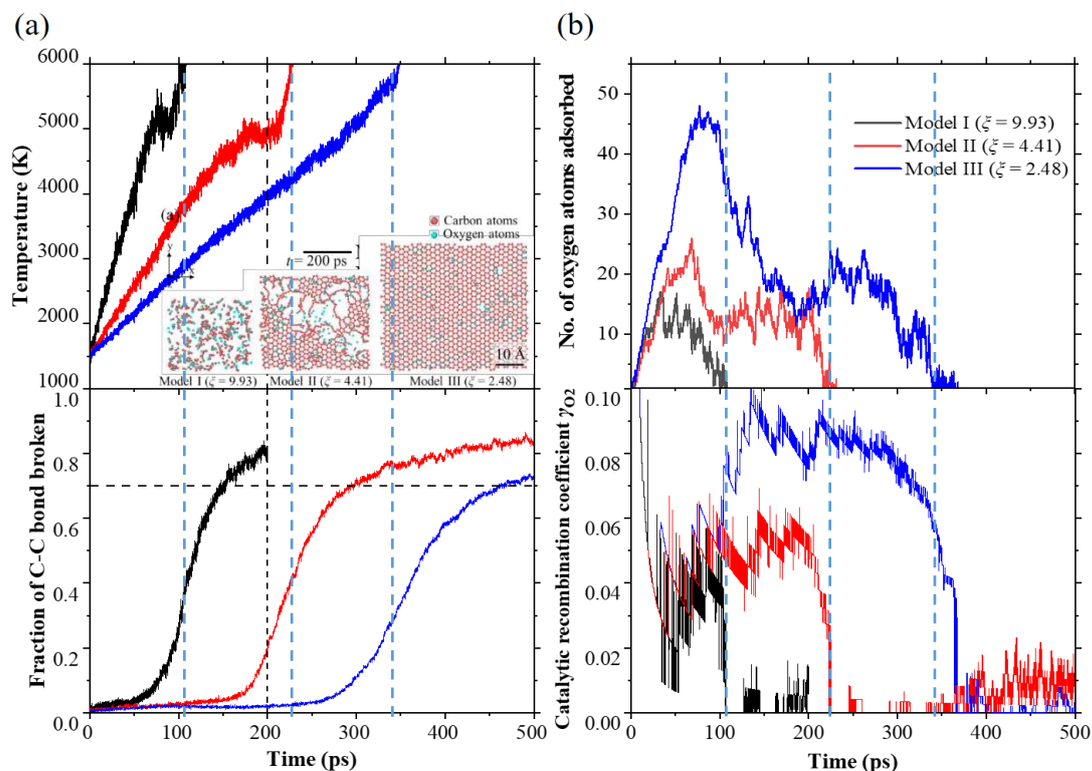
11 much higher for the case with greater energy flux density of the impinging oxygen

12 atoms, which leads to a greater erosion rate, as shown in the C-C bond breaking fraction

13 profiles. The inset snapshots of the graphene surface at the time of 200 ps for Model I

14 ~ III shown in **Figure 5(a)** indicate that a severer erosion process can be achieved with

15 greater energy flux density of oxygen atoms colliding with the graphene sheet.



16

17 **Figure 5** A comprehensive comparison for the effect of incoming energy flux density
18 on the surface catalytic and ablation processes for Model I, II, and III. (a) The system

1 temperature with the snapshot illustrations for Model I, II, III at 200 ps, along with the
2 Fraction of C-C bond broken as a function of time. (b) the number of adsorbed oxygen
3 atoms on the graphene surface and corresponding catalytic recombination coefficient
4 as a function of time.

5

6 To further identify the surface catalysis process for the O₂ formations, the number
7 of oxygen atoms adsorbed onto the graphene sheet and the corresponding
8 recombination coefficient profiles are shown in **Figure 5(b)**. The recombination
9 coefficient γ_{O_2} is defined to quantify the surface catalysis process as the fraction of
10 impinging atoms that recombining to O₂ molecules at the graphene surface. The results
11 indicate that the recombination coefficient is highly dependent on the energy flux
12 density of incoming oxygen atoms, which is closely related to the surface temperature.

13 The surface catalytic recombination coefficient γ_{O_2} for Model III with the
14 smallest energy flux density of 0.397 MW/cm² is the shown to be largest with an
15 average value of 0.081, which results is controversial to a traditional linear-like
16 Arrhenius fit with the monotonically increasing correlation model between surface
17 temperature and surface catalytic recombination coefficient. Our result can indicate that
18 the traditional Arrhenius expression for the surface catalytic recombination coefficient
19 as a function of temperature may only be valid within a certain temperature range. Since
20 the system temperature range to evaluate the surface catalysis can be as great as 6000
21 K for all the three cases in this study, the average surface catalytic recombination
22 coefficient is found to decrease when the energy flux density is increased from 0.4 to
23 1.6 MW/cm² due to the effect of severe ablation concurrence. To further address this
24 point, another Model 1S with the energy flux density of 0.07 MW/cm² has been
25 simulated with more detail results providing in the **Supplementary Material**, where
26 the average surface catalytic recombination coefficient for Model 1S is calculated to be
27 a value of 0.041.

28

3.3 The erosion process of multilayer graphene films

To investigate the effect of multilayer graphene films on the erosion process, the bombardments of graphene stack with 2 and 6 layers (or graphite in general) in the AB arrangement with hyperthermal atomic oxygens are examined in this section through Model IV and V in comparison with the one-layer graphene sheet of Model I.

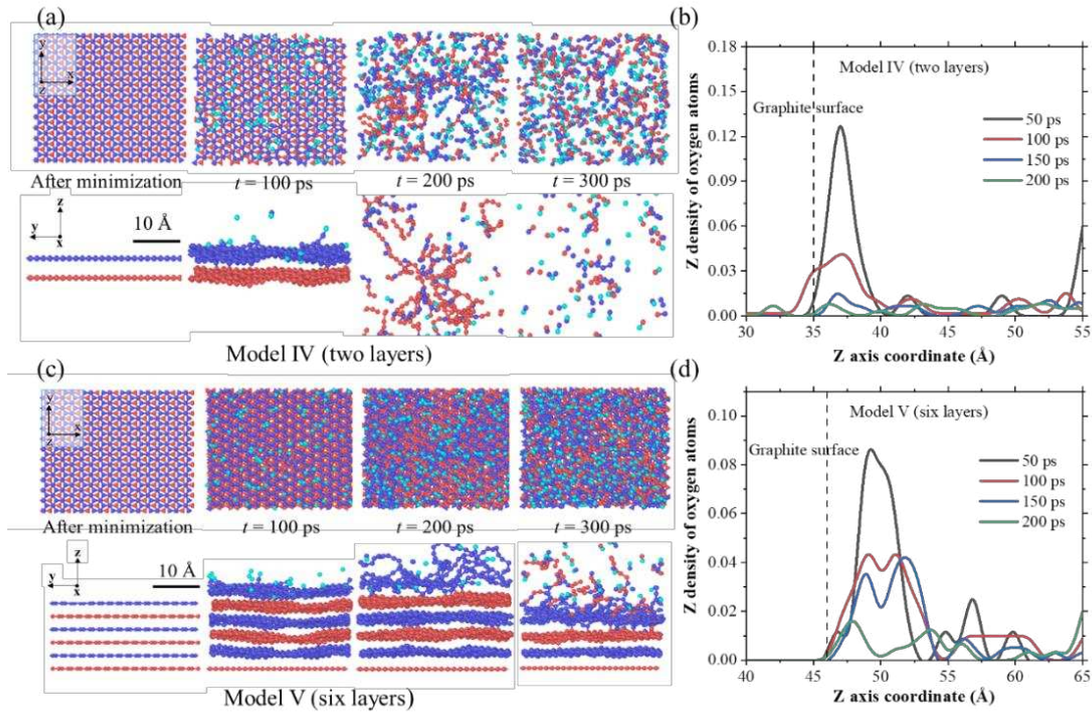
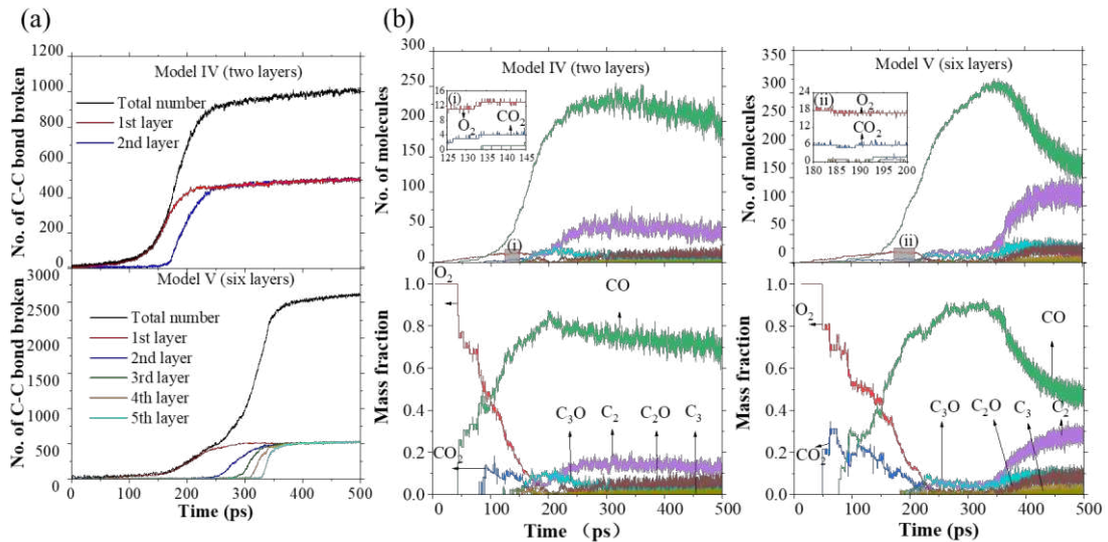


Figure 6 The snapshot illustrations of Model IV and V with oxygen z-density analysis with the evolution of time.

Figure 6 illustrates the snapshots of the erosion processes for Model IV and Model V from both top and side views with the evolution of time from 0 to 300 ps. For both models, graphite erosion proceeds firstly through atomic oxygens adsorb on the graphene stack surface with the formation of epoxides, as shown in the peaks of oxygen z-density profiles in the early stage as presented in **Figure 6(b, d)**. Then, The growth of vacancy defects on the top layer surface, followed by the same procedure on the subsequent layer below. The breakup of graphite for both models is observed to be a layer-by-layer event with the growth rate of defects much larger along the basal plane directions compared to the axial direction. Our result is consistent with that of Srinivasan et al. [44], reporting that the second layer began to be etched when the

1 carbon atoms in the top layer were almost completely consumed. This phenomenon is
 2 proved by the qualitative trends of the C-C bond broken number plot as a function of
 3 time, as shown in **Figure 7(a)**. Eventually, the number of broken C-C bonds for all the
 4 layers reach the same maximum value, indicating that the graphene layer is completely
 5 broken at this time. The leaving population and mass fraction for each gas composition
 6 during the reaction for both models are obtained as shown in **Figure 7(b)**, which trends
 7 are similar with the single-layer models (Model I~III) as shown in **Figure 3(b)** and
 8 **Figure 4(a, b)**. The gas phase are populated initially with O₂ and CO₂ molecules due
 9 to the surface catalysis and oxidation reactions, respectively. Later, as the system
 10 temperature increasing, CO and C₂ dominates among all the leaving gas species. The
 11 recombination coefficients and fraction of C-C bond broken for multilayer graphene
 12 models are derived from these and presented in **Figure 8(b)**.

13



14

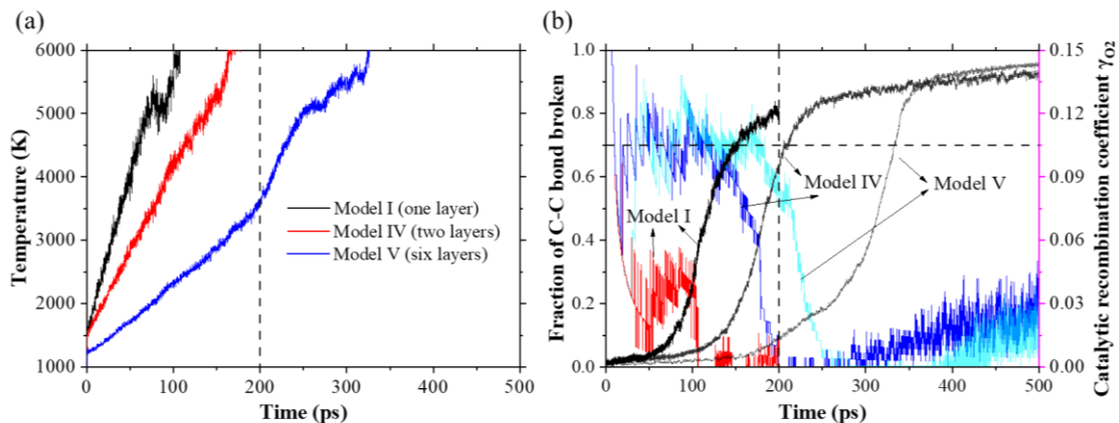
15 **Figure 7** The C-C bond broken profiles and gasified component analysis as a function
 16 of time for Model IV and V over time.

17

18 Additionally, it should be noted from **Figure 7(a)** that, the C-C bond broken of the
 19 top layer (1st layer) for Model V is slower than that of Model IV. In the meanwhile, the
 20 ablation rate for Model V with six graphene layers is the slowest as shown in **Figure**
 21 **8(b)**, compared with the single-layer Model I and two-layer Model IV. This is because
 22 that, compared with Model IV, more graphene stacks appearing below the top layer in

1 Model V, acting as a heat sink that slows down the temperature rise by conduction due
2 to oxygen-atom collisions. The temperatures of Model I and IV both exceed 6000 K at
3 200ps with Model I almost completely broken (~80% C-C bond broken) and Model IV
4 partially broken (~60% C-C bond broken), while the temperature of Model V is around
5 3500 K with the model slightly ablated (~10% C-C bond broken) as shown in **Figure**
6 **8 (a, b)**.

7



8

9 **Figure 8** The system temperature profiles and the surface catalysis / ablation analysis
10 as a function of time with the comparisons of Model I, IV and V.

11

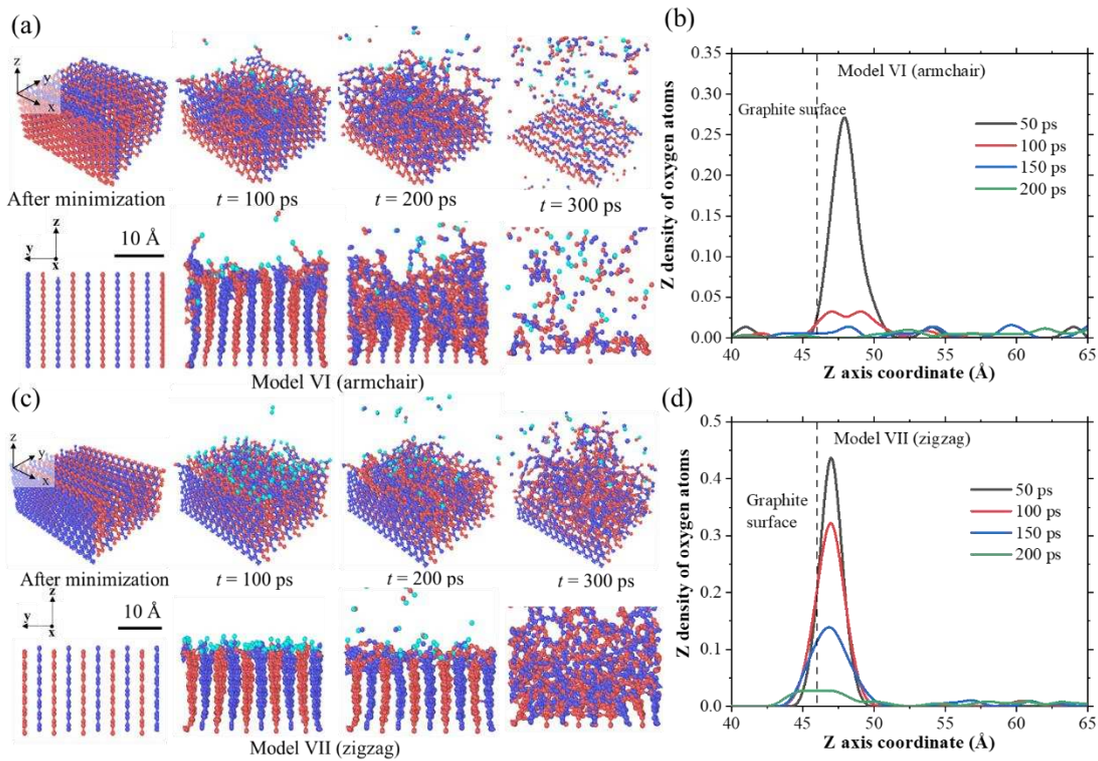
12 **3.4 Graphite surface effects on the atomic oxygen erosion process**

13 To quantifying the effect of the graphite orientation on the surface catalysis and
14 oxidative erosion rates, the simulation results of Model V, VI, VII are compared in this
15 session with graphite basal plane, armchair (AC) edge surface and zigzag (ZZ) edge
16 surface exposing to energetic oxygen atom collisions normal to the surfaces.

17 A sequence of states through which graphite surfaces with armchair (Model VI)
18 and zigzag (Model VII) conformations proceeding upon hyperthermal atomic oxygen
19 collision is illustrated in **Figure 9**, associating with the z-density profiles of oxygen
20 atoms as a function of time. Combining with the graphite basal plane of Model V,
21 successive oxygen atom collisions indicate that all the three surfaces can be etched by
22 hyperthermal atomic oxygens. The breakup of the graphene stack happens through a
23 layer-wise sequential erosion process. Taking Model VI with armchair surface as an

1 example, from the initial state to 100 ps, the z-density peaks of oxygen atoms near the
 2 graphite surface in **Figure 9(b)** indicate the dominant adsorption behavior of oxygen
 3 atoms on the graphene surface. Moreover, z-density of oxygen atoms in Model VII is
 4 greater than that in Model VI from 0 ps to 200 ps, as shown in **Figure 9(b, d)**. Compared
 5 with Model VI with AC surface, Model VII with ZZ surface is ablated more slowly
 6 compared with Model VI and its structure at around 300 ps becomes chaotic and
 7 disorderly. This can be proved by the fact that the binding energies of the AC edge
 8 tended to be weaker than those of the ZZ edge [45]. Therefore, it is more difficult for
 9 the AC surface to form stable functional group structures to resist the etching of high-
 10 energy oxygen atoms than the ZZ surface.

11



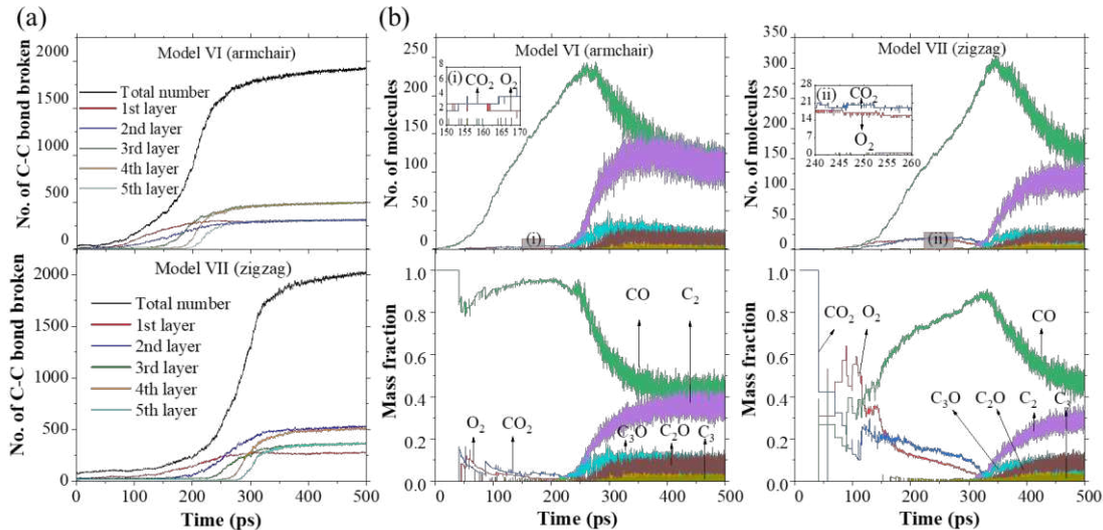
12

13 **Figure 9** The snapshot illustrations of Model VI and VII with oxygen z-density analysis
 14 with the evolution of time.

15

16 **Figure 10(a)** shows the number of C-C bond broken for different layers with the
 17 evolution of time. The breaks of the C-C bond from the 1st layer to the 5th layer for

1 both Model VI and VII are observed to be gradually delayed and reach their maximums,
 2 proving the oxidative erosion of the basal planes being a ‘layer by layer’ phenomenon.
 3 The total number of broken C-C bond for Model VI grows exponentially from about
 4 100 ps while that of Model VII grows exponentially from about 200 ps. The sequence
 5 of erosion events leads to the loss of C atoms from the graphite slabs, forming gas
 6 molecules as shown in **Figure 10(b)**. It is found that the starting time of gas generation
 7 for graphite with AC surface is earlier than that of graphite with ZZ surface. The mass
 8 fractions of O₂ and CO₂ for both Model VI and VII decrease gradually, while the mass
 9 fraction of CO increases rapidly during the catalytic ablation reaction. Later, the mass
 10 fraction of CO reaches its peak, while the mass fractions of O₂ and CO₂ molecules
 11 approach zero. Subsequently, the number of CO molecules starts to decrease while the
 12 number of C₂ molecules begins to increase rapidly. In the meanwhile, the mass fractions
 13 of C₃ and C₂O molecules increase slightly.
 14

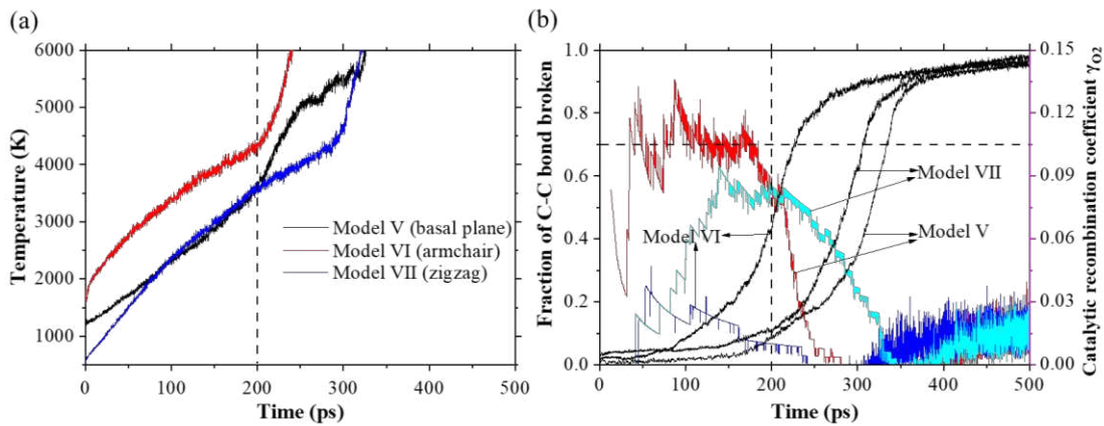


15
 16 **Figure 10** The C-C bond broken profiles and gasified component analysis as a function
 17 of time for Model VI and VII over time.

18
 19 In order to compare the surface catalysis and ablation rate of the graphite slabs
 20 with different orientations, the catalysis recombination coefficients and the fraction of
 21 graphite C-C bond broken are obtained as shown in **Figure 11(b)**. Z-density of oxygen

1 atoms for Model VI decreases rapidly from 100 ps afterwards in **Figure 9(b)**, resulting
 2 in the catalytic recombination coefficient γ of Model VI smaller than that of Model VII,
 3 as shown in the **Figure 11(b)**. Besides, though all the surfaces can be etched by
 4 hyperthermal atomic oxygen with a layer by layer phenomenon, the armchair surfaces
 5 show the largest etching rate and basal presenting the lowest etching rage. It is clear
 6 from **Figure 11(a)** that an increase in system temperature causes a speedup of the
 7 graphite ablation process: Model VI with AC edge surface is ablated fastest with the
 8 largest C-C bond broken rate and the smallest catalytic recombination coefficient.
 9 Graphite with basal plane reveals better resistant to energetic oxygen atom etching
 10 compared to the graphite with AC edge surfaces and that with ZZ edge surfaces.

11



12

13

14 **Figure 11** The system temperature profiles and the surface catalysis / ablation analysis
 15 as a function of time with the comparisons of Model V, VI and VII.

16

17 **4. Conclusions**

18 The surface catalysis and ablation processes of hyperthermal atomic oxygens
 19 colliding with graphite are simulated by MD method using ReaxFF reactive force field.
 20 The physical/chemical interactions between the hyperthermal atomic oxygens and the
 21 solid surface were investigated with seven models to study the effects of the oxygen
 22 energy flux density, the presence of multi layers beneath the graphene sheet surface and
 23 the graphite surfaces with different edges (graphite basal plane surface, AC edge surface

1 and ZZ edge surface). The simulation results can be concluded as followed:

2 (1) For all the seven models, the system temperature increases gradually due to the
3 energy input to the system through the addition of successive oxygen atom impacts.
4 The adsorption of oxygen atoms dominates at the beginning for all models when the
5 temperature is relative low, generating O₂ molecules through the surface catalytic
6 recombination of oxygen atoms. The ablation of graphene sheet accelerates while
7 system temperature further going forward, along with the number of O₂ and CO₂ gas
8 species declining gradually and forming the main incomplete oxidation production of
9 CO gas molecules.

10 (2) The ablation rate of graphene surface accelerates significantly as oxygen
11 energy flux density increases due to the higher system temperature resulting from the
12 denser impingement of oxygen atoms. The traditional Arrhenius expression to describe
13 the temperature effect on the surface catalysis process is inefficient due to the
14 simultaneous ablation behavior.

15 (3) With the presence of multiple layers beneath the graphene sheet surface, the
16 graphite slab can be etched by hyperthermal atomic oxygen with a layer by layer
17 phenomenon. The more graphene stacks appearing below the top layer, the slower the
18 ablation rate. This is due to the fact that additional graphene layers can act as a heat
19 sink, slowing down the temperature rise by conduction due to oxygen-atom collisions.

20 (4) For graphite surfaces with different edges, the AC surfaces show the largest
21 etching rate and basal presenting the lowest etching rage, revealing the fact that the
22 binding energies of the AC edge tends to be weaker than those of the ZZ edge. Therefore
23 it is more difficult for the AC surface to form stable functional group structures to resist
24 the etching of high-energy oxygen atoms than the ZZ surface.

25 In the future, we will broaden our research through further exploring the
26 temperature and pressure effects of both gas/solid phases on the surface catalysis and
27 ablation behaviors with multiple carbon-based materials.

28

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4

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