Platinum Clusters on Vacancy-Type Defects of Nanometer-Sized Graphene Patches

Takashi Yumura 1,*, Tatsuya Awano 2, Hisayoshi Kobayashi 2 and Tokio Yamabe 2

1 Department of Chemistry and Materials Technology, Kyoto Institute of Technology, Matsugasaki, Sakyoku, Kyoto 606-8585, Japan
2 Nagasaki Institute of Applied Science, 536 Aba-machi, Nagasaki 851-0193, Japan

* Author to whom correspondence should be addressed; E-Mail: yumura@chem.kit.ac.jp; Tel.: +81-75-724-7571; Fax: +81-75-724-7580.

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Abstract: Density functional theory calculations found that spin density distributions of platinum clusters adsorbed on nanometer-size defective graphene patches with zigzag edges deviate strongly from those in the corresponding bare clusters, due to strong Pt-C interactions. In contrast, platinum clusters on the pristine patch have spin density distributions similar to the bare cases. The different spin density distributions come from whether underlying carbon atoms have radical characters or not. In the pristine patch, center carbon atoms do not have spin densities, and they cannot influence radical characters of the absorbed cluster. In contrast, radical characters appear on the defective sites, and thus spin density distributions of the adsorbed clusters are modulated by the Pt-C interactions. Consequently, characters of platinum clusters adsorbed on the sp² surface can be changed by introducing vacancy-type defects.

Keywords: density functional theory; graphene; cluster; catalyst; spin state

1. Introduction

Graphitic carbon materials serve as a support material [1] for anode catalysts such as platinum clusters in proton exchange membrane (PEM) fuel cells [2–9]. The supported Pt clusters catalyze the activation of hydrogen molecules to form protons and electrons on the anode of fuel cells. Consequently, the size of Pt clusters is a crucial parameter in determining their catalytic activity.
Actually, clusters of less than 3 nm are more effective for catalyzing the H\textsubscript{2} dissociation [9]. During the catalytic reactions, adjacent clusters tend to coalescence, forming larger clusters. Accordingly, their catalytic activity decreases as the reaction proceeds. To retain the catalytic activity of supported Pt catalysts, a plausible approach is to strengthen interactions between Pt clusters and underlying carbon sp\textsuperscript{2} surface.

To come up with a strategy for constructing carbon supports suitable for Pt catalysts, computational simulations are becoming a powerful tool [9–31]. Several computational studies suggest that disrupting the sp\textsuperscript{2} surface by introducing defects (vacancy-type and Stone-Wales type) [9,18,21,25,26,29–31], dopants (nitrogen or boron impurities) [14–16], and mechanical strain [25] can enhance the interactions with Pt clusters. With respect to the formation of the vacancy-type defects, recent high-resolution transmission electron microscopy (TEM) studies [32–35] show that electron irradiation of graphene creates vacancy-type defects by removing a few carbon atoms from the surface. After these events, unsaturated carbon atoms are generated. Some of the unsaturated atoms make a covalent bond with an adjacent atom to form a five-membered ring, whereas the others remain two-coordinated. These carbon atoms, which cannot be seen in pristine graphene, are more chemically reactive, and thus they serve as sites for strong adsorption of Pt clusters. Reactive carbon atoms can be also found on edges of zigzag-nanoribbons and zigzag-graphene patches, because their frontier orbital coefficients are located on edge carbon atoms [36–42]. Thus, one can utilize such reactive edge atoms to trap well Pt clusters [22,27]. Previously, we investigated by means of density functional theory (DFT) calculations how a Pt cluster is bound to the nanometer-size rhombic sp\textsuperscript{2} patch with zigzag-edges (C\textsubscript{96}H\textsubscript{26}) (Figure 1) [22]. Such H-terminated sp\textsuperscript{2} patches are contained in activated carbons as condensed-aromatic-ring fractions [43]. The DFT calculations found that a Pt\textsubscript{6} cluster preferentially binds into edge atoms of C\textsubscript{96}H\textsubscript{26} rather than into center atoms. In fact, the Pt\textsubscript{6} additions to edge atoms were about 50 kcal/mol more stable than those to center atoms [22].

**Figure 1.** Optimized structure for C\textsubscript{96}H\textsubscript{26} graphene patch.

Another interesting feature of nanometer-size graphenes is that they have radical character in the ground state, depending on their shape and size [44–51]. Thus, we assume that interactions between a Pt cluster and a radical sp\textsuperscript{2} patch can modulate the catalytic activity of the supported cluster due to the onset of unpaired electrons on Pt atoms. Based on the assumption, the current study will focus on whether spin states of C\textsubscript{96}H\textsubscript{26} support have a power to influence the properties of the adsorbed
Pt clusters. Furthermore, we are interested in how introduction of vacancy-type defects on the radical C_{96}H_{26} support changes the electronic properties of the surface. These changes would have an impact on determining the properties of Pt clusters adsorbed on the sp² support. To increase our understanding of the interactions between a radical sp² support and a Pt cluster, we performed density functional theory (DFT) calculations. The main aim in the current DFT study is to clarify how different electronic properties of C_{96}H_{26} support with or without vacancy-type defects influence the interactions with Pt clusters, and concomitantly the properties of the absorbed clusters.

2. Results and Discussion

2.1. Platinum Clusters on C_{96}H_{26} in the Triplet State

To obtain a basic insight on how different spin states of C_{96}H_{26} patch affect the interactions with Pt clusters, we investigated how Pt₆ clusters bind into the sp² surface. Following the previous study [22], two types of Pt₆ cluster were considered, denoted by (i) and (ii) in Figure 2. The DFT calculations found that their triplet states are energetically stable relative to the corresponding singlet states. The stability of spin-polarized states in Pt clusters was also reported by other groups [52–54]. We obtained three optimized geometries for Pt₆(i) or Pt₆(ii) clusters adsorbed on C_{96}H_{26} (Pt₆-C_{96}H_{26}) in the singlet and triplet states. Within the three optimized geometries in Figure 3, one is that the Pt₆(i) cluster makes four Pt-C bonds with the sp² surface, and the other two are distinguished by whether the number of Pt-C bonds formed between the Pt₆(ii) cluster and the surface is 2 or 4. In these geometries, optimized lengths of Pt-C bonds range from 2.288 to 2.334 Å.

We estimated in Table 1 the energy difference between the triplet and singlet spin states in each configuration, ΔE_{state}(Pt₆-C_{96}H_{26}), defined as [E_{total}(triplet state) − E_{total}(singlet state)] where E_{total}(triplet state) or E_{total}(singlet state) is the total energy in each state. As shown in Table 1, the three configurations have negative ΔE_{state}(Pt₆-C_{96}H_{26}) values. These negative ΔE_{state}(Pt₆-C_{96}H_{26}) values indicate that the triplet state of a Pt₆-C_{96}H_{26} configuration is energetically favorable relative to the singlet state, irrespective of the cluster shapes. Furthermore, we see from Table 1 more significant ΔE_{state} values in the Pt₆-C_{96}H_{26} configurations than those in the bare Pt₆ clusters. Thus, the Pt-C interactions influence relative stability of the triplet to singlet states of the Pt₆ clusters.

Spin density distributions in the triplet Pt₆-C_{96}H_{26} structures are also displayed in Figure 3, where isosurface α- and β-spins are given by pink and blue, respectively. As shown in Figure 3, the Pt-C
interactions induce spin densities on C\textsubscript{96}H\textsubscript{26}, although the stable triplet state of pristine C\textsubscript{96}H\textsubscript{26} has radical characters only on edge carbon atoms. Likewise, we see spin densities on the adsorbed Pt cluster in the configurations. Basically their spin density distributions are similar to those in the bare Pt\textsubscript{6} clusters (Figure 2), but spin densities slightly decrease on Pt atoms that participates the Pt-C bond formation. The similarity between Pt clusters with and without the carbon support is understandable, because underlying carbon atoms do not have radical characters in pristine C\textsubscript{96}H\textsubscript{26} in the stable triplet state, and thus they cannot perturb the spin density distributions of Pt clusters even though they interact substantially.

**Figure 3.** Spin density distributions of optimized C\textsubscript{96}H\textsubscript{26} and Pt\textsubscript{6}-C\textsubscript{96}H\textsubscript{26} configurations in the triplet state. Parts of the optimized geometries, corresponding to the region surrounded by pink hashed lines in Figure 1, are given. Isosurface \(\alpha\)- and \(\beta\)-spins are given by pink and blue, respectively. Optimized bond lengths are in Å.

**Table 1.** Energy difference between the singlet and triplet states in Pt\textsubscript{6}-C\textsubscript{96}H\textsubscript{26} (\(\Delta E_{\text{state}}\) in kcal/mol)\textsuperscript{a}.

|                  | Pt\textsubscript{6}(i) | Pt\textsubscript{6}(ii)-(1) | Pt\textsubscript{6}(ii)-(2) |
|------------------|------------------------|-----------------------------|-----------------------------|
| Bare clusters    | –14.5                  | –5.4                        | –5.4                        |
| Clusters on C\textsubscript{96}H\textsubscript{26} | –31.4                  | –23.6                       | –22.1                       |

\(\Delta E_{\text{state}}(\text{Pt}_6-C_96H_{26}) = E_{\text{total}}(\text{triplet state}) - E_{\text{total}}(\text{singlet state})\). Negative \(\Delta E_{\text{state}}\) values indicate that the triplet state of a Pt\textsubscript{6}-C\textsubscript{96}H\textsubscript{26} configuration is energetically stable relative to the singlet state.

2.2. Vacancy-Type Defects Formed by Removing Carbon Atoms from C\textsubscript{96}H\textsubscript{26}

Prior to discussing Pt clusters adsorbed on the sp\textsuperscript{2} surface with vacancy-type defects, we look at how introduction of a vacancy-type defect on C\textsubscript{96}H\textsubscript{26} changes its electronic structures. In this study, we considered the number \(n\) of carbon atoms removed from C\textsubscript{96}H\textsubscript{26}, ranging from 1 to 3. Removed
carbon atoms are colored in Figure 4. First, we constructed mono-, di-, and tri-vacancy defects by removing the green atom, the green and blue atoms, and the three colored atoms, respectively. The vacancy-type defects will be denoted by $C_{96-n}H_{26}$. Using the initial geometries, we obtained optimized structures for the vacancy-type defects in the triple and singlet states. Then, the energy difference between the two spin states was evaluated in each vacancy-type defect, given as $\Delta E_{\text{state}}(C_{96-n}H_{26})$ in Table 2. We can see from Table 2 negative $\Delta E_{\text{state}}(C_{96-n}H_{26})$ values irrespective of the number of carbon atoms removed from $C_{96}H_{26}$. The negative $\Delta E_{\text{state}}(C_{96-n}H_{26})$ values indicate that each $C_{96-n}H_{26}$ has energetically stable triplet state.

Figure 4. Vacancy type defects ($C_{96-n}H_{26}$), constructed by removing a few carbon atoms from $C_{96}H_{26}$ where $n$ ranges from 1 to 3. Parts of the optimized geometries, corresponding to the region surrounded by pink hashed lines in Figure 1, are given. Optimized bond lengths are given in Å. Their spin density distributions in the triplet state are also given. Isosurface $\alpha$- and $\beta$-spins are given by pink and blue, respectively.

| $N$ | $\Delta E_{\text{state}}$ (kcal/mol) |
|-----|------------------------------------|
| 0   | -12.8                              |
| 1   | -14.0                              |
| 2   | -14.8                              |
| 3   | -31.4                              |

$\Delta E_{\text{state}}(C_{96-n}H_{26}) = E_{\text{total}}(\text{triplet state}) - E_{\text{total}}(\text{singlet state})$. Negative $\Delta E_{\text{state}}$ values indicate that the triplet state of a $C_{96-n}H_{26}$ configuration is energetically stable relative to the singlet state.

Figure 4 also displays their optimized structures in the triplet state as well as corresponding spin density distributions. As shown in Figure 4, all optimized structures for the vacancy-type defects have some five-membered rings formed by connecting two orange atoms. There is one five-member ring in the mono-vacancy defect ($C_{95}H_{26}$), while there are two five-member rings in the other vacancy-defects ($C_{94}H_{26}$ and $C_{93}H_{26}$). Besides, removing odd-numbered carbon atoms from $C_{96}H_{26}$ generates one coordinatively unsaturated carbon atom, given by red in Figure 4. In fact, they are bound to only two neighboring atoms. The presence of vacancy-type defects perturbs significantly spin density
distributions of pristine C\textsubscript{96}H\textsubscript{26}. As displayed in Figure 4, we can see radical carbon atoms around defective sites. In particular, significant spin densities were found in the tri-vacancy defect site. More interestingly, we found that the structural features of the defects have a correlation with how spin densities are distributed. On the unsaturated (red) atoms in the mono- and tri-vacancy-defects, spin densities are distributed on the carbon plane, which come from non-bonding orbitals. In contrast, spin densities on the orange atoms, which are a part of five-membered rings, are found perpendicular to the plane. The onset of radical carbon atoms at the center of the patch differentiates the defective surfaces from the pristine in terms of the interactions with Pt clusters, as will be mentioned below.

2.3. Platinum Clusters on Vacancy-Type Defects (C\textsubscript{96}H\textsubscript{26}) Depending on Spin States

2.3.1. Singlet State

Despite the preferences of the triplet state of C\textsubscript{96-n}H\textsubscript{26} over the singlet state, let us first use the singlet state to increase our understanding of how a Pt\textsubscript{k} cluster interacts with a defective site on the sp\textsuperscript{2} surface. Following the previous study on interactions between a Pt\textsubscript{6} cluster and pristine C\textsubscript{96}H\textsubscript{26}, we have a special interest on how the presence of a vacancy-type defect of the patch affects the interactions with a Pt\textsubscript{6} cluster. In addition, we will discuss dependences of the interaction energies on size of clusters whose number of contained Pt atoms (k) being smaller than 6. Figures 5–7 show optimized structures for a Pt\textsubscript{6} cluster adsorbed on the mono-, di-, and tri-vacancy-type defects, respectively.

Several modes for the cluster bindings were considered. For example, we obtained six optimized geometries for a Pt\textsubscript{6} cluster binding into the mono-vacancy-type defect in Figure 5. The four Pt\textsubscript{6}-C\textsubscript{95}H\textsubscript{26} structures displayed in Figure 5 are relatively stable in energy. In general, stable Pt\textsubscript{k}-C\textsubscript{95}H\textsubscript{26} structures have a Pt\textsubscript{k-1} moiety contained in stable Pt\textsubscript{k-1}-C\textsubscript{95}H\textsubscript{26} structures. Of course, there are other possibilities for Pt\textsubscript{k} binding modes. However, our computational resource is limited, reluctantly we did not obtain other optimized geometries. We evaluated the binding energy in each configuration defined as $E_{\text{bind}} = E_{\text{total}}(\text{Pt}\textsubscript{k}-\text{C}\textsubscript{96-n}H\textsubscript{26}) - E_{\text{total}}(\text{C}\textsubscript{96-n}H\textsubscript{26}) - E_{\text{total}}(\text{Pt}\textsubscript{k})$, where k ranges from 1 to 6 (Tables 3–8).

**Figure 5.** Optimized geometries for Pt\textsubscript{6} cluster on the mono-vacancy-type defect in the singlet state (Pt\textsubscript{6}-C\textsubscript{95}H\textsubscript{26}). Optimized bond lengths are given in Table 3.
Table 3. Key parameters of Pt\textsubscript{k} on mono-vacancy defect (C\textsubscript{95}H\textsubscript{26}) (k is 1 or 6) in Figure 5. Separations of a Pt atom from orange atoms (Pt-C(orange)) and those from the red atom (Pt-C(red)). Separations of carbon atoms from a Pt atom except for the nearest Pt atom (other Pt-C), and those between the two orange atoms (C-C). Bond lengths are in Å. The $E_{\text{bind}}$ and $\Delta E(Pt_k)$ values are given in kcal/mol. Their definition was given in the text.

|       | $E_{\text{bind}}$ | Pt-C(orange) | Pt-C(red) | other Pt-C | C-C Bond | $\Delta E(Pt_k)$ |
|-------|-------------------|--------------|-----------|------------|----------|-----------------|
| Pt\textsubscript{1} | -145.7            | 1.953, 1.954 | 1.942     | —          | 2.764    | —               |
| Pt\textsubscript{6(i)-(A)} | -152.0            | 1.986, 1.988 | 1.968     | —          | 2.745    | 4.2             |
| Pt\textsubscript{6(i)-(B)} | -146.6            | 1.947, 1.968 | 1.958     | 2.215      | 2.768    | 16.8            |
| Pt\textsubscript{6(ii)-(C)} | -151.6            | 1.980, 1.981 | 1.991     | 2.074      | 2.728    | 17.4            |
| Pt\textsubscript{6(ii)-(D)} | -157.1            | 1.983, 1.983 | 2.010     | 2.114      | 2.729    | 8.1             |
| Pt\textsubscript{6(ii)-(E)} | -136.7            | 1.978, 1.976 | 1.991     | 2.200, 2.083, 2.252 | 2.803 | 15.3 |
| Pt\textsubscript{6(ii)-(F)} | -143.4            | 1.986, 1.977 | 1.982     | 2.113      | 2.740    | 6.3             |

Table 4. Key parameters of Pt\textsubscript{k} on di-vacancy defect (C\textsubscript{94}H\textsubscript{26}) (k is 1 or 6) in Figure 6. Separations of a Pt atom from orange atoms (Pt-C(orange)), those of carbon atoms from a Pt atom except for the nearest Pt atom (other Pt-C), and those between the two orange atoms (C-C). Bond lengths are in Å. The $E_{\text{bind}}$ and $\Delta E(Pt_k)$ values are given in kcal/mol.

|       | $E_{\text{bind}}$ | Pt-C(orange) | other Pt-C | C-C Bond | $\Delta E(Pt_k)$ |
|-------|-------------------|--------------|------------|----------|-----------------|
| Pt\textsubscript{1} | -106.1            | 1.999, 1.985, 1.999, 1.985 | —          | 2.846, 2.846 | —               |
| Pt\textsubscript{6(i)-(A)} | -106.7            | 2.010, 2.001, 2.091, 2.117 | 2.088, 2.033 | 2.810, 2.940 | 12.7            |
| Pt\textsubscript{6(i)-(B)} | -106.3            | 2.014, 2.101, 2.109, 2.008 | 2.087, 2.108 | 2.918, 2.921 | 14.8            |
| Pt\textsubscript{6(ii)-(C)} | -94.5             | 2.014, 2.119, 2.088, 2.005 | 2.206, 2.314, 2.084, 2.119 | 2.928, 2.921 | 17.3            |
| Pt\textsubscript{6(ii)-(D)} | -85.1             | 1.994, 2.021, 2.036, 2.114 | 2.040      | 2.836, 2.941 | 17.9            |
**Figure 7.** Optimized geometries for Ptₖ cluster on the tri-vacancy-type defect in the triplet state (Pt₆-C₉₃H₂₆). Optimized bond lengths are given in Table 5.

![Figure 7](image)

**Table 5.** Key parameters of Ptₖ on tri-vacancy defect (C₉₃H₂₆) (k is 1 or 6) in Figure 7. Separations of a Pt atom from orange atoms (Pt-C(orange)) and those from the red atom (Pt-C(red)). Separations of carbon atoms from a Pt atom except for the nearest Pt atom (other Pt-C), and those between the two orange atoms (C-C). Bond lengths are in Å. The $E_{\text{bind}}$ and $\Delta E$($Pt_k$) values are given in kcal/mol.

|       | $E_{\text{bind}}$ | Pt-C(orange)          | Pt-C(red) | other Pt-C | C-C Bond | $\Delta E$ (Ptₖ) |
|-------|-------------------|-----------------------|-----------|------------|----------|------------------|
| Pt₁   | –162.6            | 2.114, 2.333, 2.097, 2.641 | 2.059     | —          | 2.637, 2.641 | —                |
| Pt₆(i)-(A) | –198.2          | 1.992, 1.959, 1.971, 1.969 | 2.105     | 2.054      | 2.753, 2.830 | 12.2             |
| Pt₆(ii)-(B) | –193.4          | 2.006, 1.988, 1.974, 1.938 | 2.144     | 2.130      | 2.825, 2.746 | 5.9              |
| Pt₆(ii)-(C) | –188.4          | 1.971, 2.081, 1.970, 1.969 | 2.046     | 2.378, 2.217, 2.107 | 2.924, 2.922 | 20.0             |
| Pt₆(ii)-(D) | –142.0          | 2.006, 2.015           | 2.021     | 2.130, 2.231 | 2.710 | 17.9             |

Corresponding optimized structures for the smaller Ptₖ clusters adsorbed (k = 2, 3, 4, and 5) are also displayed in Figures 8–10.

**Figure 8.** Optimized geometries for Ptₖ cluster (k = 2~5) on the mono-vacancy-type defect in the singlet state (Ptₖ-C₉₅H₂₆). Optimized bond lengths are given in Table 6.
Figure 8. Cont.

Table 6. Key parameters of Pt\textsubscript{k} on mono-vacancy defect (C\textsubscript{95}H\textsubscript{26}) (\(k = 2 \sim 5\)) in Figure 8.

| \(E_{\text{bind}}\)  | Pt-C(orange) | Pt-C(red) | other Pt-C | C-C Bond | \(\Delta E(\text{Pt}_k)\) |
|----------------------|-------------|-----------|------------|----------|-----------------|
| Pt\textsubscript{2}-(A) | \(-131.7\)  | 1.964, 2.006 | 1.969 | —       | 2.755 | 29.6      |
| Pt\textsubscript{2}-(B) | \(-146.1\)  | 1.965, 1.965 | 1.942 | —       | 2.743 | 9.0       |
| Pt\textsubscript{3}-(A) | \(-132.4\)  | 1.971, 1.972 | 1.971 | —       | 2.715 | 0.9       |
| Pt\textsubscript{3}-(B) | \(-126.9\)  | 1.952, 1.976 | 1.962 | —       | 2.790 | 2.5       |
| Pt\textsubscript{4(ii)}-(A) | \(-139.7\) | 1.983, 1.984 | 1.967 | —       | 2.759 | 7.4       |
| Pt\textsubscript{4(ii)}-(B) | \(-140.0\) | 1.981, 1.987 | 1.987 | 2.054   | 2.743 | 7.4       |
| Pt\textsubscript{4(i)}    | \(-128.3\)  | 1.953, 1.966 | 1.960 | 2.227   | 2.757 | 19.8      |
| Pt\textsubscript{5(ii)}-(A) | \(-140.5\) | 1.976, 1.976 | 1.980 | —       | 2.753 | 4.3       |
| Pt\textsubscript{5(ii)}-(B) | \(-147.8\) | 1.982, 1.982 | 2.002 | 2.105   | 2.741 | 5.4       |
| Pt\textsubscript{5(i)}    | \(-132.4\)  | 1.947, 1.966 | 1.972 | 2.211, 2.275 | 2.790 | 4.6 |

Figure 9. Optimized geometries for Pt\textsubscript{k} cluster (\(k = 2 \sim 5\)) on the di-vacancy-type defect in the singlet state (Pt\textsubscript{k}-C\textsubscript{94}H\textsubscript{26}). Optimized bond lengths are given in Table 7.
Table 7. Key parameters of Pt on di-vacancy defect (C_{94}H_{26}) \((k = 2 ~ 5)\) in Figure 9.

| \(E_{\text{bind}}\) | \(\text{Pt-C(orange)}\) | \(\text{other Pt-C}\) | \(\text{C-C Bond} \downarrow\) | \(\Delta E(\text{Pt}_k)\) |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| Pt\_2-(A)       | -91.2           | 1.999, 2.028, 2.038, 2.066 | -           | 2.831, 2.904 | 10.6 |
| Pt\_2-(B)       | -59.2           | 1.934, 1.935, 1.997, 1.997 | -           | 2.786, 2.740 | 7.1  |
| Pt\_3-(A)       | -70.9           | 2.025, 2.118, 2.025, 2.119 | 2.045, 2.046 | 2.893, 2.893 | 9.6  |
| Pt\_3-(B)       | -63.2           | 1.990, 2.013, 2.125, 2.199 | 2.081, 2.082 | 2.791, 2.976 | 7.0  |
| Pt\_4-(A)       | -57.6           | 1.972, 1.987, 1.971, 1.986 | -           | 2.690, 2.690 | 1.1  |
| Pt\_4-(B)       | -84.8           | 2.034, 2.035, 2.101, 2.100 | 2.053, 2.054 | 2.852, 2.853 | 6.4  |
| Pt\_5-(A)       | -75.8           | 2.004, 2.005, 2.064, 2.134 | 2.067, 2.066 | 2.786, 2.896 | 4.7  |
| Pt\_5-(B)       | -93.4           | 2.005, 2.006, 2.085, 2.099 | 2.122, 2.138 | 2.917, 2.923 | 12.3 |
| Pt\_5-(C)       | -81.3           | 1.991, 2.001, 2.113, 2.123 | 2.053, 2.082 | 2.806, 2.932 | 13.4 |

Figure 10. Optimized geometries for Pt\_k cluster \((k = 2~5)\) on the tri-vacancy-type defect in the singlet state \((\text{Pt}_k\text{-C}_{93}H_{26})\). Optimized bond lengths are given in Table 8.

Table 8. Key parameters of Pt\_k on tri-vacancy defect \((\text{C}_{93}H_{26})\) \((k = 2 ~ 5)\) in Figure 10. Separations of a Pt atom from orange atoms (Pt-C(orange)) and those from the red atom (Pt-C(red)). Separations of carbon atoms from a Pt atom except for the nearest Pt atom (other Pt-C), and those between the two orange atoms (C-C). Bond lengths are in Å. The \(E_{\text{bind}}\) and \(\Delta E(\text{Pt}_k)\) values are given in kcal/mol. Their definition was given in the text.

| \(E_{\text{bind}}\) | \(\text{Pt-C(orange)}\) | \(\text{Pt-C(red)}\) | \(\text{other Pt-C}\) | \(\text{C-C Bond}\) | \(\Delta E(\text{Pt}_k)\) |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Pt\_2-(A)       | -172.3          | 1.944, 1.968, 1.988, 1.950 | 2.038           | -               | 2.839, 2.852 | 22.0 |
| Pt\_2-(B)       | -132.0          | 2.002, 1.985    | 2.009           | 2.021           | 2.874, 1.630 | 8.3  |
| Pt\_3           | -167.5          | 1.948, 1.964, 1.989, 2.124 | 2.021           | 2.067           | 2.999, 2.832 | 9.8  |
| Pt\_4           | -180.3          | 1.978, 2.068, 1.973, 1.981 | 2.046           | 2.050           | 2.897, 2.704 | 3.6  |
| Pt\_5           | -173.0          | 1.950, 1.976, 1.934, 1.997 | 2.099           | -               | 2.840, 2.750 | 13.9 |

Here \(E_{\text{total}}(\text{Pt}_k\text{-C}_{96-n}H_{26})\) is the total energy of an optimized \(C_{96-n}H_{26}\) geometry, \(E_{\text{total}}(C_{96-n}H_{26})\) is that of the optimized \(C_{96-n}H_{26}\) geometry, and \(E_{\text{total}}(\text{Pt}_k)\) is that of the optimized \(\text{Pt}_k\) cluster. For the DFT calculations of the binding energies, a counterpoise correction for basis set superposition error (BSSE) was included [55]. When an \(E_{\text{bind}}\) value has a negative sign, the binding of a Pt cluster or the Pt atom
into C\textsubscript{96–n}H\textsubscript{26} is energetically preferable. As shown in Tables 3–5, the calculated \(E_{\text{bind}}\) values in the single Pt addition are similar to those reported in [35]. These similarities verify the reliability of our DFT results.

Looking at the \(E_{\text{bind}}\) values, the bindings of a Pt\textsubscript{k} cluster into the sp\textsuperscript{2} surface are strongly facilitated by introducing vacancy-type defects. In fact, their stabilizing energies (\(-E_{\text{bind}}\)) in stable Pt\textsubscript{k}-C\textsubscript{95}H\textsubscript{26} structures are around 150 kcal/mol. These values are much larger than the pristine cases (about 50 kcal/mol [22]). Similar enhancement in the stabilization energies was also found in the Pt\textsubscript{1}-C\textsubscript{94}H\textsubscript{26} and Pt\textsubscript{1}-C\textsubscript{93}H\textsubscript{26} structures. Judging from the \(E_{\text{bind}}\) values, reactivity of vacancy-type defects toward Pt clusters declines in the order: tri-vacancy > mono-vacancy > di-vacancy. These results suggest that the tri-vacancy defect is more suitable for binding of Pt clusters into carbon surface rather than the mono- and di-vacancy defects.

From Tables 3–8, we see different behaviors between the three types of defect in terms of dependences of the \(E_{\text{bind}}\) values on Pt cluster size. Most stable Pt\textsubscript{k}-C\textsubscript{95}H\textsubscript{26} structures except for Pt\textsubscript{3}-C\textsubscript{93}H\textsubscript{26} have \(E_{\text{bind}}\) values similar to that in Pt\textsubscript{1}-C\textsubscript{95}H\textsubscript{26}. In contrast, the \(E_{\text{bind}}\) values in Pt\textsubscript{k}-C\textsubscript{94}H\textsubscript{26} and Pt\textsubscript{k}-C\textsubscript{93}H\textsubscript{26} are deviated from those in Pt\textsubscript{1}-C\textsubscript{94}H\textsubscript{26} and Pt\textsubscript{1}-C\textsubscript{93}H\textsubscript{26}. When a Pt\textsubscript{k} cluster binds into the di-vacancy-defect, their \(E_{\text{bind}}\) values are smaller than the Pt\textsubscript{1}-C\textsubscript{94}H\textsubscript{26} value. However, these absolute values increase gradually with an increase of the cluster size, and seem to converge to the Pt\textsubscript{1}-C\textsubscript{93}H\textsubscript{26} value at \(k = 6\). In the Pt\textsubscript{k}-C\textsubscript{93}H\textsubscript{26} cases, the \(E_{\text{bind}}\) values, being around 180 kcal/mol, are always larger than the Pt\textsubscript{1}-C\textsubscript{93}H\textsubscript{26} value.

To understand the energetics in the optimized Pt\textsubscript{k}-C\textsubscript{96–n}H\textsubscript{26} structures (Tables 3–8), let us first look at in detail geometrical features of Pt\textsubscript{1}-C\textsubscript{96–n}H\textsubscript{26}. These key geometrical parameters in the Pt\textsubscript{1}-C\textsubscript{96–n}H\textsubscript{26} configurations (lengths of newly formed Pt-C bonds and of lengthening CC bonds) are listed in Tables 3–5.

In these tables, we can distinguish two types of the formed Pt-C bond, by whether a Pt atom binds into orange or red atoms. When the single Pt atom binds into the mono-vacancy defect, it inserts between the orange atoms in the five-membered ring, and then two P-C (orange) bonds are formed newly. As a result of the Pt addition, the separation between the orange atoms lengthens from 1.754 to 2.764 Å. At the same time, the Pt atom also coordinates to the unsaturated red atom. The binding Pt atom lifts from the sp\textsuperscript{2} surface, because the hole is not large enough to accommodate the Pt atom. Similar Pt lifting can be seen in the Pt\textsubscript{1}-C\textsubscript{94}H\textsubscript{26} configuration, where the Pt atom inserts between orange atoms in both five-membered rings, and it breaks the connections. The degree of Pt lifting in Pt\textsubscript{1}-C\textsubscript{94}H\textsubscript{26} is less significant than that in Pt\textsubscript{1}-C\textsubscript{95}H\textsubscript{26}, due to relatively larger hole in C\textsubscript{94}H\textsubscript{26}.

In contrast, the hole of C\textsubscript{93}H\textsubscript{26}, surrounded by ten carbon atoms, can house the Pt atom, and therefore the binding Pt atom is on the sp\textsuperscript{2} surface. Then, four Pt-C bonds are formed, accompanying the cleavage of the bonds between orange atoms in the five-membered rings. Moreover, the Pt binding into the unsaturated C atom was also seen. When a Pt\textsubscript{k} cluster binds into a vacancy-type defect, slightly longer separations of the nearest Pt atom from reactive (orange and red) atoms were found. Despite the stabilization operated between a Pt\textsubscript{k} cluster and C\textsubscript{96–n}H\textsubscript{26}, slightly longer Pt-C bonds imply weakening interactions of the nearest Pt atom from the reactive carbon atoms compared with Pt\textsubscript{1}-C\textsubscript{96–n}H\textsubscript{26} case.

Compensating the weakening of the interactions, remaining Pt atoms of a clusters are additionally bound to carbon atoms of a defective site to maximize the Pt-C interactions. Then their clusters are more or less deformed from the most stable configuration in the gas-phase [56–58]. The degree of cluster deformation was estimated by using \(\Delta E(\text{Pt}_k)\), defined as \([E(\text{Pt}_k \text{ on surface}) – E(\text{Pt}_k)]\), where
\( E(\text{Pt}_k \text{ on surface}) \) is the total energy of \( \text{Pt}_k \) cluster taken from an optimized \( \text{Pt}_k-\text{C}_{96-n}\text{H}_{26} \) structure and \( E(\text{Pt}_k) \) is that of the optimized geometry for the bare \( \text{Pt}_k \) cluster. Positive \( \Delta E(\text{Pt}_k) \) values in Tables 3–8 suggest destabilization from cluster deformation upon the interactions with a vacancy-type defect. Although we cannot find a clear correlation between \( E_{\text{bind}} \) and \( \Delta E(\text{Pt}_k) \) values, the balance between the stabilization from the Pt-C bond formation and the destabilization from the cluster deformation is a key in determining the stability. From Figures 5–7 and 11, we found clear differences between \( \text{Pt}_k-\text{C}_{93}\text{H}_{26} \) and \( \text{Pt}_k-\text{C}_{95}\text{H}_{26} \) (\( \text{Pt}_k-\text{C}_{94}\text{H}_{26} \)) in terms of the number of Pt atoms binding directly into orange atoms in the defective site to cleave connections between adjacent orange atoms.

**Figure 11.** Optimized geometries for the singlet Pt atom on the mono-, di-, and tri-vacancy-type defects in the triplet state (\( \text{Pt}_1-\text{C}_{95}\text{H}_{26}, \text{Pt}_1-\text{C}_{94}\text{H}_{26}, \text{and Pt}_1-\text{C}_{93}\text{H}_{26}, \) respectively). Optimized bond lengths are given in Tables 3–5.

In \( \text{Pt}_k-\text{C}_{95}\text{H}_{26} \) (\( \text{Pt}_k-\text{C}_{94}\text{H}_{26} \)), one Pt atom participates in cleaving the orange connection(s), irrespective of the cluster size. In the tri-vacancy cases containing larger ten-membered ring, two Pt atoms in a cluster bind to the defective site to split two orange connections. The accommodation of two Pt atoms cannot be seen in the \( \text{Pt}_1-\text{C}_{94}\text{H}_{26} \) structure, and thus the significant enhanced stabilization in \( \text{Pt}_k-\text{C}_{93}\text{H}_{26} \) is understandable. Moreover the acceptability of the ten-membered-ring to trap Pt atoms differentiates \( \text{C}_{93}\text{H}_{26} \) from \( \text{C}_{95}\text{H}_{26} \) and \( \text{C}_{94}\text{H}_{26} \) in terms of their reactivity toward \( \text{Pt}_k \) clusters. Due to the strong interactions between a \( \text{Pt}_k \) cluster and a vacancy-type defect, we can see unique orbital features, which cannot be seen in \( \text{C}_{96-n}\text{H}_{26} \) (Figure 12).

In fact, 5d(Pt)-based orbitals, given by blue bars in Figure 12, appear in the frontier orbital regions of the \( \text{Pt}_6-\text{C}_{94}\text{H}_{26} \) and \( \text{Pt}_6-\text{C}_{93}\text{H}_{26} \) configurations. As the most striking case, we can see in Figure 13 that the \( \text{Pt}_6(\text{i})-\text{C}_{94}\text{H}_{26}(\text{B}) \) configuration has the HOMO and LUMO consisting of 5d(Pt) orbitals. On the other hand, levels of 5d(Pt)-based orbitals in the \( \text{Pt}_6-\text{C}_{95}\text{H}_{26} \) strongly depend on their cluster-shape. In the \( \text{Pt}_6(\text{i})-\text{C}_{95}\text{H}_{26}(\text{A}) \) and \( \text{Pt}_6(\text{ii})-\text{C}_{95}\text{H}_{26}(\text{C}) \) configurations, such 5d(Pt)-based orbital lies larger than 1.3 eV above the LUMO, whereas the LUMO+1 consists of 5d(Pt)-based orbitals in the other \( \text{Pt}_6-\text{C}_{95}\text{H}_{26} \) configurations.

### 2.3.2. Triplet State

As shown in Figure 12, the all optimized geometries in the single state have relatively small HOMO-LUMO gaps (0.29 ~ 0.40 eV). Thus, higher spin states can be energetically stable relative to the singlet states. Along the assumption, we obtained their triplet states, and estimated the energy difference between the two spin states (\( \Delta E_{\text{state}}(\text{Pt}_k-\text{C}_{96-n}\text{H}_{26}) \)), as tabulated in Table 9.
Figure 12. Orbital energies (eV) in the frontier orbital region of the optimized Pt₆-C₉₅H₂₆, Pt₆-C₉₄H₂₆, and Pt₆-C₉₃H₂₆ configurations whose structures are given in Figures 5–7. The HOMO-LUMO gaps are given. Orbitals originated from 5d(Pt) orbitals are denoted by blue bars, and those with no or less 5d(Pt) orbital contribution are denoted by black bars.

Figure 13. Frontier orbitals (the HOMO and LUMO) in the Pt₆(i)-C₉₄H₂₆(B) configuration (Figure 6) are given as a representative Pt₆-C₉₆–ₙH₂₆ configuration.

Table 9. Energy difference between the singlet and triplet states in the Pt₆-C₉₆–ₙH₂₆ configurations (ΔE_{state} in kcal/mol).\(^a\)

|                 | Pt₆(i)-(A) | Pt₆(i)-(B) | Pt₆(ii) | Pt₆(iii) |
|-----------------|------------|------------|---------|----------|
| Clusters on C₉₅H₂₆ | −14.3      | −17.8      | −14.0   | \(\_\)   |
| Clusters on C₉₄H₂₆ | −23.3      | −16.2      | −23.3   | −38.5    |
| Clusters on C₉₃H₂₆ | −15.0      | −23.7      | −13.7   | \(\_\)   |

\(^a\) ΔE_{state}(Pt₆-C₉₆–ₙH₂₆) = E_{total}(triplet state) − E_{total}(singlet state). Negative ΔE_{state} values indicate that the triplet state of a Pt₆-C₉₆–ₙH₂₆ configuration is energetically stable relative to the singlet state. ΔE_{state} in kcal/mol; \(^b\) we could not obtain the optimized geometry in the triplet state.
Table 9 shows that their triplet spin states are energetically stable relative to the singlet states, as expected. According to DFT calculations, most Pt₆-C₉₆–ₙH₂₆ configurations have radical Pt₆ clusters on defective graphene patches. As representative cases, spin density distributions on the Pt₆(i) cluster binding into C₉₅H₂₆ or C₉₄H₂₆ are shown in Figure 14. Figure 14 shows that substantial spin densities appear on the adsorbed clusters in the three Pt₆-C₉₆–ₙH₂₆ configurations (Pt₆(i)-C₉₅H₂₆(B) and two Pt₆(i)-C₉₄H₂₆ configurations), whereas the Pt₆(i)-C₉₅H₂₆(A) configuration does not have such radical cluster due to the absence of 5d(Pt)-based frontier orbitals (Figure 12). In the three configurations with radical clusters, a variety of the spin density distributions was found. In the Pt₆(i)-C₉₅H₂₆(B) and Pt₆(i)-C₉₄H₂₆(B), the spin density distributions are strongly deviated from those in the bare Pt₆ cluster, whereas the Pt₆(i)-C₉₄H₂₆(B) has similar distributions. Furthermore, we found a relationship between spin densities of Pt clusters and those on the defective sp² surface. When spin density distributions of the Pt cluster of a Pt₆-C₉₆–ₙH₂₆ configuration are (not) similar to the bare case, spin densities are (not) delocalized over the carbon surface. Similar tendencies were found in spin density distributions of the stable triplet Pt₁₃-C₉₆–ₙH₂₆ configurations [59], because radical Pt clusters exist on the defective graphene patches as shown in Figure 15.

**Figure 14.** Spin density distributions of representative Pt₆-C₉₆–ₙH₂₆ configurations (Pt₆(i) cluster on C₉₅H₂₆ or C₉₄H₂₆ in two binding fashions, displayed in Figures 5 and 6). Isosurface α- and β-spins are given by pink and blue, respectively.

Finally, let us compare the spin density distributions on Pt₆ clusters on defective sp² surfaces (Figure 14) with the pristine case (Figure 2). As mentioned above, spin density maps on Pt₆ cluster on pristine C₉₆H₂₆ are similar to those in the bare cluster. However, such similarity cannot be always found in the defective graphene cases. The different tendencies come from whether underlying carbon atoms have radical characters or not in Figures 2 and 4. In the pristine patch, underlying carbon atoms do not have radical characters, and they cannot perturb spin density distributions of the absorbed cluster. In contrast, unpaired electrons exist on underlying carbon atoms in the defective sites, and thus spin density distributions of the adsorbed clusters are modulated by the Pt-C interactions. Therefore, perturbation of the radical sp² surface by introduction of vacancy-type defects can change characters of adsorbed Pt clusters.
Figure 15. Spin density distributions of representative Pt_{13}-C_{96-n}H_{26} configurations. Isosurface \( \alpha \)- and \( \beta \)-spins are given by pink and blue, respectively.

The DFT findings are important in the catalytic activity of Pt clusters on sp\(^2\) surface, because the supported clusters can serve as active site for catalytic reactions. For example, if radical Pt clusters exist on carbon surface, it can cleave the H-H bond of hydrogen molecules in a homolytic manner. Otherwise, the H-H bond is activated by the clusters via a non-radical mechanism in [22]. Thus, the DFT calculations propose that one can change chemical reactivity of Pt clusters on graphene patches by introducing of vacancy-type defects on the surface.

3. Experimental

To investigate interactions of Pt clusters with C\(_{96}\)H\(_{26}\), we carried out DFT calculations implemented in the Gaussian 03 and 09 program packages [60,61]. Adsorbed clusters that we considered consist of one, six, and thirteen Pt atoms. To perform the calculations, a hybrid Hartree–Fock/density functional theory method, B3LYP [62–66] was used. The B3LYP method consists of the Slater exchange, the Hartree–Fock exchange, the exchange functional of Becke [62–64], the correlation functional of Lee, Yang, and Parr (LYP) [65], and the correlation functional of Vosko, Wilk, and Nusair (VWN) [66]. In general the hybrid B3LYP method has been reported to provide excellent descriptions of various properties. The Gaussian-type basis set we used for the C and H atoms is 6-31G* [67], and that for the Pt atom is the quasi-relativistic effective core potential RECP and valence basis sets recommended by Stuttgart group (SDD) [68]. The SDD RECP is adjusted to total valence energies of a multitude of atomic references states, which are quantum mechanical observables [68]. As indicated in the previous papers [69–76], the B3LYP/6-31G* calculations correctly reproduce experimental data for C\(_{60}\), especially its IR and Raman vibrational frequencies. According to the theoretical report by Nova et al. [77], the method of our choice (B3LYP/SDD + 6-31G*) is appropriate to reproduce experimental values in terms of Pt-C bonds in Pt complexes [77]. The computational method is also suitable to study transition metals adsorbed on graphene. In fact, Pt-C bond lengths obtained from the B3LYP/SDD + 6-31G* calculations fall in the range reported from other theoretical reports [24,30].
4. Conclusions

Density functional theory (DFT) B3LYP calculations were employed to investigate the adsorption of Pt\(_k\) cluster (\(k\) is 1–6, and 13) into a nanometer-size graphene patch (C\(_{96}\)H\(_{26}\)) with or without vacancy-type defects. According to the DFT calculations, removing a few carbon atoms (\(n\)) from C\(_{96}\)H\(_{26}\) results in the formation of five-membered rings as well as coordinatively unsaturated carbon atoms. Introduction of a vacancy-type defect on C\(_{96}\)H\(_{26}\) strongly affects spin density distributions in its stable triplet state. Although spin densities appear only on edge carbon atoms of the triplet C\(_{96}\)H\(_{26}\) structure, defective graphene patches have radical carbon atoms at the center where the reactive carbon atoms exist. These spin density distributions differentiate characters of Pt clusters adsorbed on defective graphene patches from those on the pristine. According to the DFT calculations, spin density maps of Pt clusters on C\(_{96}\)H\(_{26}\) are similar to those of the corresponding bare clusters. In contrast, Pt clusters interact strongly with radical carbon atoms in defective graphene patches, and thus spin density distributions of the adsorbed Pt clusters are usually deviated from the bare cases. Consequently, DFT calculations propose that characters of Pt clusters adsorbed on the sp\(^2\) carbon surface can be modulated by introducing vacancy-type defects.

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