# Hybrid Pd<sub>38</sub> nanocluster/Ni(OH)<sub>2</sub>-graphene catalyst for enhanced HCOOH **dehydrogenation: First principles approach**

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ment of high-efficiency materials and catalysts for storage and transportation of hydrogen energy must be achieved to realize hydrogen economy. Recently, catalyst systems such as Pd nanoclusters (Pd NCs) supported on nickel hydroxide (Ni(OH)2) have been reported to have advantages, including effective suppression of CO production and efficiency enhancement of HCOOH dehydrogenation. However, the reaction mechanism and multi-metallic interface system design of such systems have not been elucidated. Therefore, various Ni(OH)<sub>2</sub> surfaces supported on a graphene system were designed through density functional theory calculations, and the support material was combined with Pd<sub>38</sub>NC  $(Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>-G)$ . Subsequently, the adsorption behavior of HCOOH dehydrogenation intermediates was analyzed. We found a new adsorption configuration in which HCOOH<sup>\*</sup> (where \* and a single underline indicates the adsorbed species and adsorbed atom, respectively) was adsorbed in a more stable manner (adsorption energy,  $E_{ads}$ =  $-1.22$  eV) on the system than HCOOH\* (E<sub>ads</sub>=-1.10 eV) owing to the presence of Ni(OH)<sub>2</sub>-G. This affected the next step in HCOOH dehydrogenation, i.e., formation of HCOO\* species, and showed a positive effect on the HCOOH dehydrogenation. To fundamentally understand this phenomenon, electronic structure (d-band center and density of states) and stability (vacancy formation energy) analyses were performed.

Keywords: Formic Acid Dehydrogenation, Hydrogen Energy, Nickel Hydroxide, Density Functional Theory, Surface **Stability** 

## **INTRODUCTION**

The development of environmentally friendly energy sources is inevitable for solving the problem of global warming caused by the use of excessive fossil fuels. Among the various candidates, hydrogen  $(H<sub>2</sub>)$  is a promising energy source owing to its cleanness, sustainability, and high energy density [1]. Numerous studies have been conducted using fuel cell technology to practically apply hydrogen energy to our social infrastructure.

Formic acid (HCOOH) has been verified as an excellent liquid hydrogen storage material owing to its low toxicity and high hydrogen content  $(4.4 \text{ wt\%})$  [2,3]. Hydrogen can be controllably released through HCOOH dehydrogenation  $(\Delta G = -32.8 \text{ kJ mol}^{-1})$  at room gen content (4.4 wt%) [2,3]. Hydrogen can be controllably released through HCOOH dehydrogenation  $(\Delta G = -32.8 \text{ kJ mol}^{-1})$  at room temperature in the presence of a suitable catalyst [4,5]. Among various metal catalysts for HCOOH dehydrogenation, palladium (Pd) has been reported as the best catalyst, showing excellent hydrogen production [6,7]. However, HCOOH dehydrogenation on Pd catalysts is capable of generating CO, which can lower the catalytic efficiency. This phenomenon has been controlled by designing new Pd based catalysts (e.g., alloys, core-shall nanoparticles, and nanoclusters (NCs) supported on graphene), and even the catalysts were

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shown to be more efficient and economical than conventional Pd surfaces [8,9].

Recently, multi-metallic nanocatalysts have attracted considerable attention as catalysts for hydrogen production because of their significant synergistic effect from their multi-component interfacial active sites [10-15]. Precisely, the introduction of non-noble metal hydroxides to form multi-metallic nanostructures cannot only minimize the use of noble metals but also enhance the catalytic performance [10-13]. Previous studies reported that hybrid Pt/FeNi(OH) $_{x}$ nanoparticles exhibited high performance for CO oxidation [10],  $Pt/M(OH)_{2}$  (M=Ni, Co) nanoparticles were significantly able to enhance the activity of the hydrogen evolution reaction [12]. Additionally, subnanometric hybrid Pd-M(OH),  $(M=Ni, Co)$  clusters encapsulated within purely siliceous zeolites showed superior thermal stability at 600-700 °C and complete HCOOH dehydrogenation without using any additives at  $60^{\circ}$ C [15]. In addition, graphene supports, which can modify the electronic properties of a catalyst in positive ways, have been widely used to improve the activity and efficiency of metal catalysts [16-22]. For example, graphene-supported Cu nanoparticles showed the enhanced efficiency of conversion of  $CO<sub>2</sub>$  into hydrocarbon fuels, which was attributed to the electronic properties of graphene [23]. Shin et al. [24] reported that nitrogen-doped graphene-supports could improve the efficiency of formate (HCOO $\degree$ ) and bicarbonate (HCO $\degree$ ) reversible reactions by nitrogen-doped graphene-supports could improve the efficiency of ) and bicarbonate  $(HCO<sub>3</sub><sup>-</sup>)$  reversible reactions by modifying the electronic and geometrical properties of Pd nanoparticles. The graphene supports also had a positive influence on the formation of nickel hydroxide  $(Ni(OH<sub>2</sub>))$  nanoplates with excellent electrochemical properties and stability [25].

Innovative catalysts combining multi-metallic nanocatalysts and graphene support, with the aforementioned advantages, have tremendous potential for improving the efficiency of HCOOH dehydrogenation. However, analyzing the catalyst involves many difficulties due to the lack of related research and the complex structural properties of various elements. Therefore, it is necessary to clearly understand the structural properties of the new catalysts and the adsorption behavior of HCOOH dehydrogenation intermediates. In this study, we designed a large-scale metallic NC supported on a graphene system  $(Pd_{38}NC/Ni(OH), -G)$ , which can be compared with experiments, and we investigated the adsorption behavior of intermediates for enhancing the efficiency of HCOOH dehydrogenation, using density functional theory (DFT) calculations. This can provide important insight into future study on hydrogen storage and production related to HCOOH dehydrogenation as well as the development of complex structured multi-metallic catalysts applicable to various energy domains.

#### **COMPUTATIONAL METHODOLOGY**

All spin-polarized DFT+U calculations were carried out using the Vienna ab initio Simulation Package (VASP) [26,27] with the projector augmented wave (PAW) [27,28] method. The Perdew-Burke-Ernzerhof (PBE) functional [29] with the generalized gradient approximation (GGA) was used to describe the electron exchange correlation functions. A kinetic energy cutoff for the planewave basis sets was set at 300 eV in all calculations considering the large size of the system. Brillouin zone integration was performed on a 16×16×16 (bulk) and 4×2×1 (surface system) Monkhorst-Pack grid [30], respectively. The first-order Methfessel-Paxton smearing [31] with a value of 0.1 eV was applied. The Hubbard-U term of 3.8 eV was employed to correct the error of overdelocalization of electrons, of transition metal oxides [32]. The DFT-D2 method was selected to accurately simulate the van der Waals (vdW) forces [33].

The  $Pd_{38}NC-Ni(OH)_{2}(100)$ -graphene  $(Pd_{38}NC/Ni(OH)_{2}-G)$  system was designed based on a 13.06×25.76×25.00 Å orthorhombic supercell with periodic boundary conditions. The Pd<sub>38</sub>NC/  $Ni(OH)<sub>2</sub>$ -G system consisted of 277 atoms and the  $Pd<sub>38</sub>NC$  (38 Pd atoms) was adsorbed on a single-layer of  $Ni(OH)_{2}(100)$  (23  $Ni(OH)_{2}$ bulks) supported on graphene (124 carbon atoms). The octahedral Pd<sub>38</sub>NC was selected based on the magic numbers, considering the surface size of the designed  $Ni(OH)_{2}$ -G. Although Pd<sub>38</sub>NC may be included in an NC consisting of magic numbers (7, 13, 55, etc.) [34], it provides flat adsorption surfaces that can effectively prevent overestimation of adsorption behavior and energy. Furthermore, Pd<sub>38</sub>NC has a comparable stability to an NC consisting of atoms with magic numbers in terms of binding energy per atom  $( Pd_{13}, Pd_{38} \text{ and } Pd_{55} = 220, 274 \text{ and } 288 \text{ kJ/mol}, \text{ respectively})$  [35]. This consideration is supported by previous studies that have accurately analyzed the adsorption behavior by utilizing NCs providing flat adsorption sites [24,36].

The single-layer  $Ni(OH)_{2}(100)$  supported on graphene was de-

signed to not only minimize the lattice constant mismatch between Ni(OH)<sub>2</sub> and graphene but also to provide an appropriate adsorption site for adsorbing Pd<sub>38</sub>NCs. Tensile and compressive strain effects were observed between  $Ni(OH)$ <sub>2</sub> and graphene owing to the lattice mismatch.

All atoms of the  $Pd_{38}NC/Ni(OH)_{2}-G$  system were fully relaxed and optimized until the convergence criterion for electronic self-All atoms of the  $Pd_{38}NC/Ni(OH)_2$ -G system were fully relaxed<br>and optimized until the convergence criterion for electronic self-<br>consistent iteration was less than  $10^{-4}$  eV. However, when calculating the adsorption models, atoms were partially fixed except for interfacial sites between  $Pd_{38}NCs$  and  $Ni(OH)_{2}-G$  for the interaction of adsorbates, considering computing resources and time. The gas-phase molecules were optimized using Brillouin zone integration with only the  $\Gamma$ -point under the same conditions as those used of the surface system.

The adsorption energy  $(E_{ads})$  and vacancy formation energy  $(E_{vac})$ are defined as Eq. (1) and (2), respectively.<br>  $E_{ads} = E_{substrate-adsorbate} - E_{substrate} - E_{adsorbate}$  (1)<br>  $E_{vac} = E_{defective\;substate} + E_{atom} - E_{prisine\;substate}$  (2) the surface system.<br>The adsorption energy ( $E_{\text{zats}}$ ) and vacancy formation energy ( $E_{\text{vac}}$ )<br>cefined as Eq. (1) and (2), respectively.<br> $E_{\text{ads}} = E_{\text{substrate}} - E_{\text{substrate}} - E_{\text{substate}}$  (1)

$$
E_{ads} = E_{substrate+adsorbate} - E_{substrate} - E_{adsorbate}
$$
 (1)

$$
E_{vac} = E_{defective\; substrate} + E_{atom} - E_{pristine\; substrate}
$$
\n
$$
\tag{2}
$$

In Eq. (1),  $E_{substackedotate}$   $E_{substack}$  and  $E_{adsorbaite}$  are the total DFT energies of a substrate and adsorbate  $(Pd_{38}NC/Ni(OH), -G$  adsorbed HCOOH), substrate  $(Pd_{38}NC/Ni(OH),-G)$ , and gas-phase adsorbate (HCOOH), respectively. A negative adsorption energy value indicates that the adsorption is stable. In Eq. (2),  $E_{defective\ substates} E_{atom}$ and  $E_{pristine}$  substrate are the DFT energies of defective  $Pd_{38}NC/Ni(OH)_{2}$ -G, Pd atom divided by the number of atoms of bulk Pd, and pristine  $Pd_{38}NC/Ni(OH)_{2}-G$ , respectively. A positive vacancy formation energy indicates that the surface system is stable.

#### **RESULTS AND DISCUSSION**

In this section, we discuss the novel design of the  $Pd_{38}NC/$  $Ni(OH)_{2}$ -G systems and analyze the adsorption behavior of the intermediate species of HCOOH dehydrogenation on Pd<sub>38</sub>NC/  $Ni(OH)_{2}$ -G and Pd<sub>38</sub>NC systems, proposing a possible dehydrogenation pathway for HCOOH. In addition, we explain the results of electronic structure (density of states) and durability (Pd vacancy formation energy) analyses to understand the fundamental reason for the positive effect of  $Ni(OH)_{2}$ -G on HCOOH dehydrogenation. 1. Design of Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>-graphene System

To design  $Pd_{38}NC/Ni(OH)_{2}$ -graphene, we first optimized the bulk structures of Ni(OH)<sub>2</sub> and graphene. The optimized lattice constants of bulk Ni(OH), were  $3.16 \text{ Å}$  (a=b) and  $4.45 \text{ Å}$  (c), respectively (Fig. 1(a)), which were in agreement with the experimental values (3.12 Å and 4.66 Å, respectively) [37]. In the case of graphene, the C-C distance was calculated to be  $2.47 \text{ Å}$  (Fig. 1(c)), which agreed with the experimental value  $(2.46 \text{ Å})$  [38].

Graphene-supported  $Ni(OH)_2$  has been reported to show excellent cycling stability and electrochemical characteristics [25,39]. Because  $Ni(OH)_{2}$  has various surfaces, it is necessary to select a suitable surface to simulate it by using DFT calculations. As the most suitable surfaces for combining with graphene, we selected four  $Ni(OH)$ <sub>2</sub> surface models such as  $(001)$ ,  $(100)$ ,  $(101)$ , and  $(110)$  surfaces (Fig. S1 in Supporting Information (SI)) based on the previous X-ray diffraction (XRD) analysis [40] demonstrating the highest



Fig. 1. Optimized structures of (a) bulk  $Ni(OH)_{2}$  (side and top views), (b) clean Pd<sub>38</sub>NC (side view), and (c) bulk and extended graph**ene (top view). Light grey, dark grey, red, white, and brown spheres represent Ni, Pd, O, H, and C, respectively.**

XRD peak intensities of the selected facets. When designing  $Ni(OH)_{2}$ -G, the strain effect was applied to the Ni $(OH)_{2}$  surfaces which were placed on graphene because of the complex design properties such as the number of atoms, the size of the surface area, and the adsorption angle between the surfaces. A total of four candidates were designed for each surface, and the selection criteria of surfaces that could represent each surface were set to the minimum lattice constant mismatch and the number of atoms constituting the system. The designed  $Ni(OH)_{2}(001)$ , (100), (101), and (110)-G consisted of 214 atoms with 0.63% compressive strain, 239 atoms with 0.61-0.70% tensile strain, 213 atoms with 0.14% compressive strain, and 215 atoms with 0.53-0.73% tensile strain, respectively. Detailed structures of other candidates are shown in the SI section 1.

The  $Pd_{38}NC/Ni(OH)_{2}(100)$ -G system was selected as the surface model for this HCOOH dehydrogenation study (Fig. 2(a)). This was because other systems such as  $Pd_{38}NC/Ni(OH)_{2}(001, 101)$ and 110) showed the dissociation of hydrogen atoms from the  $Ni(OH)_{2}$ -G surface when  $Pd_{38}NC$  was adsorbed on the support, collapsing the Pd-Ni $(OH)$ <sub>2</sub> system due to the mismatch of the system components. Although the  $Ni(OH)_{2}(001)$  surface was reported to be the most stable surface [40], Qi et al. [41] reported that the (100) surface was also efficiently formed on graphene. In addition, the selected  $Ni(OH)_{2}(100)$  surface provided sufficient surface area for adsorbing Pd<sub>38</sub>NC.

#### **2. Adsorption Behavior of HCOOH Dehydrogenation Intermediates**

The adsorption configuration of HCOOH and the intermedi-



Fig. 2. Optimized structures of (a) the Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>(100)-graphene (side view) system, and (b) the Ni(OH)<sub>2</sub>(100)-graphene **(top view). Light grey, dark grey, red, white, and brown spheres represent Ni, Pd, O, H, and C, respectively.**

ate species in the HCOOH dehydrogenation reaction pathway are important factors for determining the dehydrogenation efficiency. Previous studies have reported that HCOOH dehydrogenation proceeds through two reaction pathways, HCOO (two O atoms are adsorbed) and COOH (O and H atoms are adsorbed) pathways, of which the HCOO pathway has shown a higher dehydrogenation efficiency than the COOH pathway [42]. This is because HCOOH dehydrogenation through the COOH pathway offers the possibility that CO can be produced, which can lower the efficiency of the catalysts [42]. Therefore, many researchers are developing catalysts suitable for HCOOH dehydrogenation through the HCOO pathway.

We focused on the analysis of HCOOH adsorption configurations, the most important adsorbate of HCOOH dehydrogenation, which determines further reaction pathways of the HCOOH dehydrogenation. Fig. 3 shows various adsorption models of HCOOH and their adsorption energy values. The notable points are that HCOOH was more strongly adsorbed on  $Pd_{38}NC/Ni(OH)_{2}-G$  than on Pd<sub>38</sub>NC. Additionally, the  $H$ COO $H_b^*$  (where  $*$  and a sign underline indicate the adsorbed species and adsorbed atom on the surface, respectively) in which two H atoms were adsorbed on  $Pd_{38}NC/$  $Ni(OH)<sub>2</sub>-G$  system was the most stable (E<sub>ads</sub>=-1.22 eV, Fig. 3(e)).

Another important finding is that in the  $Pd_{38}NC/Ni(OH)_{2}-G$ system, the adsorbates  $(HCOOH<sub>b</sub><sup>*</sup>, HCOO<sub>b</sub><sup>*</sup>, COOH<sup>*</sup>, and H<sup>*</sup>)$ that mainly interacted with the surface through their H atoms were strongly adsorbed, but those interacting through their O atoms



Fig. 3. Comparison of adsorption energy values of HCOOH on Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>(100)-G system (a)-(f) and clean Pd<sub>38</sub>NC (g)-(k).

**Table 1. Adsorption energy (E***ads***) of intermediates of HCOOH de**hydrogenation. HCOOH<sub>a</sub>, HCOOH<sub>b</sub>, HCOO<sub>a</sub>, and HCOO<sub>b</sub> **are adsorption configurations in which one H atom, two H atoms, two O atoms, and one O atom are adsorbed on a surface**



were weakly adsorbed (Table 1). These adsorption properties can facilitate their stable formation. However, in the case of the H\* adsorption, a stronger adsorption of H may negatively affect the desorption of H to produce  $H_2(g)$  as the final product.

## **3. Comparison of HCOOH Dehydrogenation in the HCOO Pathway**

The conventional HCOO pathway of HCOOH dehydrogenation has been reported as HCOOH,\* $\rightarrow$ HCOO,\* $+$ H\* $\rightarrow$ HCOO,\* $\rightarrow$  $CO_2^*+H_2^*$  (1 $\rightarrow$ 3 $\rightarrow$ 4 $\rightarrow$ 5 in Fig. 4). In this conventional HCOO pathway, HCOO<sub>3</sub><sup>\*</sup> rotates to form  $HCOO<sub>b</sub>$ <sup>\*</sup>, which is advantageous for proceeding to the  $CO_2^*+H_2^*$  step, thus improving the efficiency of  $H_2(g)$  production by decreasing the activation energy. Recently, Wang et al. [43] reported that the HCOOH dehydrogenation steps for  $H_2(g)$  production can be significantly influenced by the initial adsorption configuration of HCOOH. They proposed a direct HCOO pathway  $(2\rightarrow 4\rightarrow 5$  in Fig. 4) that did not involve the highest energy barrier step of the HCOO\* rotation, especially when a specific promoter was present. The direct HCOO pathway could significantly enhance the efficiency of HCOOH dehydrogenation by skipping the  $HCOO<sub>a</sub><sup>*</sup>+H<sup>*</sup>$  step, thereby shortening the



**Fig. 4. HCOOH dehydrogenation through the conventional HCOO**  $(1\rightarrow3\rightarrow4\rightarrow5)$ , direct HCOO  $(2\rightarrow4\rightarrow5)$ , and COOH  $(1\rightarrow$  $6 \rightarrow 5$  or 7) pathways on the Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>-G system. A sin**gle underline represents the adsorbed atom on the surface.**

reaction steps directly, such as  $HCOOH<sub>b</sub><sup>*</sup> \rightarrow HCOO<sub>b</sub><sup>*</sup> + H<sup>*</sup> \rightarrow$  $CO<sub>2</sub>*+H<sub>2</sub>*$  [43].

Fig. 4 indicates that the HCOOH dehydrogenation can occur through the HCOO or COOH pathway. According to the adsorption energy values of HCOO and COOH in both catalytic systems, the adsorption strength of COOH was significantly lower than that of HCOO, indicating that the HCOO pathway would be more preferred to the COOH pathway  $(1\rightarrow6\rightarrow5$  or 7 in Fig. 4). Examining the adsorbed configurations of HCOO\* in both catalytic

Adsorbates		$C-H(\AA)$	$O-H(\AA)$	$C=O(\AA)$	$C-O(A)$
HC <u>OOH</u> <sub>a</sub> (g)		1.0797	0.9917	1.2044	1.3599
$\underline{HCOOH}_b(g)$		1.1434	0.9647	1.2284	1.4276
$Pd_{38}NC/Ni(OH)_{2}-G$	$HCOOH*$	1.0896	1.0068	1.2347	1.2297
	$HCOOHb$ *	1.1581	1.0122	1.2158	1.3567
	$HCOOa$ *	1.1032	$\overline{\phantom{0}}$	1.2535	1.2024
	$HCOOb$ *	1.1194	$\overline{\phantom{0}}$	1.1973	1.3685
	$COOH*$	$\overline{\phantom{0}}$	1.0154	1.2276	1.2238
$Pd_{38}NC$	$HCOOH$ *	1.1048	1.0265	1.2606	1.3567
	$HCOOHb$ *	1.1631	1.0112	1.2225	1.3610
	$HCOOs$ *	1.0870	$\overline{\phantom{a}}$	1.2413	1.2853
	$HCOOb$ *	1.1258	$\overline{\phantom{0}}$	1.1956	1.3694
	$COOH*$	$\overline{\phantom{a}}$	1.0203	1.2267	1.2203

**Table 2. Comparison of bond lengths of adsorbed species**

systems, the  $Pd_{38}NC$  and  $Pd_{38}NC/Ni(OH)_{2}-G$  systems follow the conventional HCOO and direct HCOO pathways, respectively. In addition, the HCOO<sub>a</sub><sup>\*</sup>+H<sup>\*</sup> $\rightarrow$ HCOO<sub>b</sub><sup>\*</sup>+H<sup>\*</sup> step of Pd<sub>38</sub>NC was endothermic because the adsorption energy values of HCOO<sub>a</sub><sup>\*</sup> and  $\underline{HCOO}_h^*$  were  $-3.73$  and  $-3.10$  eV, respectively. However, in the case of  $Pd_{38}NC/Ni(OH)_{2}-G$ , the  $\underline{HCOO_{h}}^{*}$  adsorption strength was significantly stronger than that of  $HCOO<sub>a</sub><sup>*</sup>$ , indicating that  $HCOO<sub>b</sub><sup>*</sup>$ can be directly formed without forming the HCOO<sub>4</sub><sup>\*</sup> species, which implies that HCOOH dehydrogenation in the  $Pd_{38}NC/Ni(OH)_{2}-G$ system may follow the direct HCOO pathway. This finding suggests that the  $Pd_{38}NC/Ni(OH)_{2}$ -G system may improve the efficiency of HCOOH dehydrogenation by shortening the reaction step of the HCOO pathway without additional additives.

Despite the evidence of the adsorption behavior of HCOOH and intermediate species in both catalytic systems, a limitation may lie in the comparison of the HCOOH dehydrogenation efficiency. This may be due to the absence of activation energy calculations, which might be impractical for DFT calculations with such a large system of  $Pd_{38}NC/Ni(OH)_{2}-G$  (consisting of 277 atoms). However, the large adsorption energy difference in both catalytic systems may rationally support the current conclusion of this study.

Additionally, when the adsorbate is adsorbed on the surface, the change in the bond length between atoms having the adsorbate can be an indicator for predicting the next step [44]. In this study, the O-H bond of HCOO $H_*^*$  has been activated to form HCOO<sub>4</sub><sup>\*</sup>+  $H^*$  on Pd<sub>38</sub>NC, having a bond length higher (3.51%) than that of  $Pd_{38}NC/Ni(OH)_{2}-G$  (1.52%) system, whereas the O-H bond length of HCOOH $^*$  has been activated to form HCOO $^*$ +H $^*$  on the  $Pd_{38}NC/Ni(OH)_{2}-G$  (bond length: 4.92%) system (Table 2). These results imply that the first step of HCOOH dehydrogenation in Pd<sub>38</sub>NC proceeds to HC<u>OOH</u><sup>\*</sup>→HCOO<sub>a</sub><sup>\*</sup>+H<sup>\*</sup> and the first step of that in the Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>-G system proceeds to  $\underline{HCOOH}_{b}^{*} \rightarrow$  $HCOO<sub>b</sub>$ <sup>\*</sup>+H<sup>\*</sup>. Next, in Pd<sub>38</sub>NC, HCOO<sub>a</sub><sup>\*</sup>+H<sup>\*</sup> proceeds to  $HCOO<sub>b</sub>$ <sup>\*</sup>+H<sup>\*</sup>, but in the Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>-G system, it proceeds directly to  $CO_2^*+2H^*$  to produce  $H_2(g)$ . The C-H bond length of  $HCOO<sub>3</sub><sup>*</sup>$  on Pd<sub>38</sub>NC was decreased by 1.61%. As the final step, the C-H bond length of  $HCOO<sub>b</sub>$ <sup>\*</sup> was decreased by 3.34% in the  $Pd_{38}NC/Ni(OH)_{2}-G$  system and increased by 3.57% in  $Pd_{38}NC$ . In the final step, the C-H bond length of  $HCOO<sub>b</sub>$ <sup>\*</sup> in Pd<sub>38</sub>NC was activated more than in  $Pd_{38}NC/Ni(OH)_{2}$ -G system; it will not exceed the difference in activation energy for the shortened reaction step in the  $Pd_{38}NC/Ni(OH)<sub>2</sub>-G$  system.

When considering the C-H bond length of HCOOH\* to proceed to the COOH pathway, it increased by  $1.29\%$  in the  $Pd_{38}NC/$  $Ni(OH)_{2}$ -G system and by 2.32% in  $Pd_{38}NC$ . However, the increase was smaller than that of the previously described O-H bond length, which implies that both systems are favorable to the HCOO pathway. In addition, this is further supported by the stable adsorption of HCOO\* on both systems rather than that of COOH\* (Table 1).

Furthermore, in addition to the bond length comparison for the activity prediction, the activity may be affected by another factor like bending degree of molecules that can modify the polarity of the molecules [44]. Although there was no clear evidence indicating a strong relationship between the bending degree of molecules and the activation of adsorbates for the formic acid dehydrogenation in the current study, it would be necessary to take into account both bond length and bond bending degree for the activity prediction.

#### **4. Electronic Structure Analysis**

The d-band center and density of states (DOS) calculations were performed to accurately demonstrate the effect of  $Ni(OH)<sub>2</sub>-G$  on the adsorption behavior of the adsorbates. The d-band center position near the Fermi level indicates the relative adsorption strength

**Table 3. The d-band center values of Pd atoms for different locations of the catalytic surface (see Fig. 5)**

Locations	d-band centers <sup>a</sup>			
	$Pd_{38}NC/Ni(OH)_{2}-G$	$Pd_{38}NC$		
1st layer	$-2.03$	$-1.79$		
2nd layer	$-1.84$	$-1.83$		
3rd layer	$-1.66$	$-1.67$		
$H_{ads}$ site	$-1.76$	$-1.78$		
Interface	$-1.92$	$-1.82$		
Surface	$-1.84$	$-1.77$		

a The d-bands below the Fermi level were only considered in the center value calculation.



**Fig. 5. Projected density of states (PDOS) of Pd d-bands of the** Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>-G (red solid line) and Pd<sub>38</sub>NC (black dot**ted line) systems. The examined locations are marked in the** Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>-G structure (left bottom side); Red, blue, **and purple dotted lines represent the 1st, 2nd, and 3rd layers, respectively, and areas colored in orange, black, and green represent the H***ads***, interface and surface sites (Table 3).**

of adsorbates on the transition metals. A more shifted d-band center toward the Fermi level represents a stronger reactivity [45,46].

We confirmed that the presence of  $Ni(OH)<sub>2</sub>-G$  significantly influenced the activity of Pd<sub>38</sub>NC. As shown in Table 3, the d-band center values of the  $Pd_{38}NC/Ni(OH)_{2}$ -G system show a wider range of values than that of Pd<sub>38</sub>NC, which implies that the catalytic surface of  $Pd_{38}NC/Ni(OH)_{2}-G$  has various adsorption properties. It may be noted that at the interface site where the reaction mainly occurs, the d-band center of the  $Pd_{38}NC/Ni(OH)_{2}$ -G system is further (negatively) shifted from the Fermi level than that of  $Pd_{38}NC$ (Table 3). This can explain the weaker adsorption strength of  $HCOO<sub>a</sub><sup>*</sup>$ in the  $Pd_{38}NC/Ni(OH)_{2}-G$  system. Additionally, although the dband center values at the  $H_{ads}$  sites of both systems are almost the same, a slight proximity of  $Pd_{38}NC/Ni(OH)_{2}$ -G towards the Fermi level may explain the stronger adsorption of H.

Note that many electrons were filled at a stable energy level below  $-5$  eV in the Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>-G system compared to that in Pd<sub>38</sub>NC (Fig. 5). This implies that the electronic properties caused by  $Ni(OH)<sub>2</sub>-G$  may have a positive effect on the surface stability, but may be a negative factor in providing a site for adsorbates to be adsorbed on a surface [47]. As a simple method to confirm this phenomenon, we compared the atomic distances of Pd atoms depending on the adsorption strength of adsorbates in each system (e.g., weakly and strongly adsorbed  $HCOOa^*$  on  $Pd_{38}NC/$  $Ni(OH)<sub>2</sub>-G$  and Pd<sub>38</sub>NC, respectively). In the Pd<sub>38</sub>NC/Ni $(OH)<sub>2</sub>-G$ system, the distance between Pd atoms was larger than that of Pd<sub>38</sub>NC when H was adsorbed on the surface. The distance between Pd atoms was smaller than that of  $Pd_{38}NC$  when  $HCOO<sub>a</sub><sup>*</sup>$  was adsorbed on the surface, which was identically confirmed in the case of  $\underline{HCOOH}_b^*$ , which can shorten the reaction step. Detailed data are shown in the SI section 2. Thus, this implies that the electronic structure of  $Pd_{38}NC$  modified by  $Ni(OH)_{2}-G$  can have a positive effect on the change in the distance between the Pd atoms on the surface. This weakened the adsorption strength of adsorbates that interact with a surface by their O atoms, requiring a relatively large size of adsorption site as well as strengthening that of H atoms



Fig. 6. Vacancy formation sites on the  $Pd_{38}NC/Ni(OH)_{2}-G$  system. **(a), (b), and (c) indicate the vacancy sites where Pd atom was extracted.**

requiring a small surface area. Thus,  $Ni(OH)_{2}$ -G can provide favorable properties for the adsorption of adsorbates, thereby improving the efficiency of HCOOH dehydrogenation and positively affecting the stability of the Pd surface.

## 5. Surface Stability of Pd<sub>38</sub>NC/Ni(OH)<sub>2</sub>-G

The enhanced stability of  $Pd_{38}NC$  by  $Ni(OH)_{2}$ -G was also demonstrated by the results of Pd vacancy formation energy calculations. A Pd vacancy site in each system was selected at the site fairly similar to a flat surface (Fig. 6). The vacancy formation energy values of the  $Pd_{38}NC/Ni(OH)_{2}$ -G system were 0.76 (Fig. 6(a)), 0.46 (Fig. 6(b)), and 0.29eV (Fig. 6(c)), respectively, at each site designed by three vacancy models based on the distance between  $Pd_{38}NC$ and  $Ni(OH)<sub>2</sub>-G$ . The Pd<sub>38</sub>NC system showed a negative vacancy formation energy of  $-0.73$  eV (the vacancy site of the Pd<sub>38</sub>NC/  $Ni(OH)_{2}$ -G system is the same position in  $Pd_{38}NC$ ). The negative value of  $Pd_{38}NC$  would only be interpreted as a relatively weaker formation energy compared to those examined, because the formation energy value can be sufficiently varied depending on the definition of Pd atom energy (e.g., it can be obtained from the bulk Pd structure or from a single Pd atom). It may be noted that the farther the Pd location from  $Ni(OH)_{2}$ -G, the lower the vacancy formation energy. This may imply that  $Ni(OH)_{2}$ -G can play an important role in enhancing the Pd surface stability.

#### **CONCLUSION**

We have designed and optimized a  $Pd_{38}NC/Ni(OH)_{2}(100)$ -graphene catalyst for the dehydrogenation of formic acid (HCOOH). The most stable adsorption configuration,  $HCOOH$ <sup>\*</sup>, was found on the  $Pd_{38}NC/Ni(OH)_{2}-G$  system, which was attributed to the modification of the electronic structure by  $Ni(OH)_{2}$ -G. The initial adsorption behaviors of HCOOH\* and its intermediate species,  $HCOO<sub>s</sub>$ <sup>\*</sup> and  $HCOO<sub>b</sub>$ <sup>\*</sup>, which is the most important step for the entire HCOOH dehydrogenation reaction, was examined. It was confirmed that the HCOOH dehydrogenation on the  $Pd_{38}NC/$ Ni(OH)<sub>2</sub>(100)-graphene would proceed through the direct HCOO pathway, improving the efficiency of HCOOH dehydrogenation by shortening the reaction step of the HCOO pathway. These findings were supported by the density of states and d-band center calculations, and the results also demonstrated the possibility that  $Ni(OH)_{2}$ -G can enhance the surface stability of  $Pd_{38}NC$  according to the comparison of the vacancy formation energy. This study not only provides an insight for designing large-scale catalysts, but also is a milestone in solving the challenges of analyzing the reaction mechanisms of large catalysts.

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#### **NOTES**

The authors declare no competing financial interests.

## **SUPPORTING INFORMATION**

Additional information as noted in the text. This information is available via the Internet at http://www.springer.com/chemistry/ journal/11814.

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