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Theoretical Study of Physical and Chemical Properties of Pt Clusters Adsorbed on Single Wall Carbon Nanotubes

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Introduction

Platinum metal is widely used in environmentally and economically important processes, such as conversion of harmful gases into less harmful ones, hydrogenation and electrode-reactions of fuel cells, because of its high catalytic activity. It is, however, a precious metal, quite expensive, and limited supply. Therefore, reduction of the consumption of platinum metal is one of the most important issues. By using small clusters, the significant reduction of the consumption is expected. Recently, highly dispersed and size-controlled small Pt clusters (less than 1 nm) made from the dispersed single Pt atoms were achieved by using carbon nanotube (CNT) supports.

A miniaturization of materials usually results not only in the enhancement of relative surface area but also in the change of the property itself. Metal nanoclusters usually exhibit the unique catalytic properties that differ from those of extended flat surface or bulk materials. The properties of metal nanoclusters can be governed by several factors such as geometric factors (atomic arrangement, number of coordination), electronic factors (charge states, density of states), and support effects or dynamics structures. These parameters, however, are not always clearly distinct, since a change in the surroundings of a surface atom has a simultaneous influence on its electronic structure. Therefore, characterization and precise control of the properties of Pt clusters are among the outstanding challenges in the research fields of both physics and chemistry, and can lead to atom-by-atom design, tuning and control of chemical activity of Pt clusters.

The main purposes of our research are using first-principles density functional theory (DFT) simulation, which is presently established as a standard tool for large system with a reliable compromise between accuracy and efficiency:

- To explore the physical properties (geometry and electronic structure) of small Pt$_n$ ($n = 3, 4, 7, 10, 13$) nanoclusters adsorbed on single wall carbon nanotubes (SWNTs) support.
- To investigate the chemical reactivity of Pt$_n$ ($n = 3, 4, 7, 10, 13$) clusters adsorbed on single wall carbon nanotubes (SWNTs) support, for reactions in fuel cell particularly.

Computational Methodology

Physical and chemical properties of Pt$_n$ ($n = 3, 4, 7, 10, 13$) nanoclusters adsorbed on single wall carbon nanotubes (SWNTs) support have been investigated by density functional theory-based simulations in DMol3 code and OpenMX code. All calculations are carried out under the periodic boundary conditions. The PBE exchange-correlation functional is used to treatment the electron-electron interactions. Double numerical with polarization orbitals basis sets, and norm-conversing pseudo-potentials are used.

The initial structures of Pt$_n$ ($n = 3, 4, 7, 10, 13$) clusters are chosen from stable structures in gas phase. Then, adsorptions of single Pt atom and the Pt clusters on SWNTs support are performed, as shown in Fig. 1, to clarify their geometric structures. Electronic structures of Pt$_n$ ($n = 3, 4, 7, 10, 13$) clusters on SWNTs support are investigated based on the density of states and charge distribution analyses.

The chemical reactivity of Pt$_n$ ($n = 3, 4, 7, 10, 13$) clusters on SWNTs support are explored through the adsorption of O$_2$, CO, OH species, which are primary reactions in hydrogen fuel cell or direct methanol fuel cell (DMFC).
Moreover, the oxidation reaction of CO and OH, a rate-limiting step in DMFC is also investigated. The effects of geometry, electronic structure, and dynamic structure on the chemical reactivity of Pt clusters on SWNT are discussed.

**Results and Discussion**

1. **Morphology of Pt$_n$ (n = 3, 4, 7, 10, 13) clusters on SWNTs support**

   Figure 2 shows the optimized structures, electronic structure of single Pt atom on the (10, 0) SWNT support. The results show that the interaction between Pt and C is a weak covalent nature with hybridization between d-states of Pt and p-states of C. Pt adatoms can easily diffuse on the surface of SWNT and tend to form a cluster, than to disperse on the SWNT surface.

   Small Pt$_n$ (n = 3, 4, 7, 10, 13) clusters on single wall carbon nanotubes (SWNTs) support are preferred 3 dimensional structures, as shown in Fig. 3. The stability of Pt clusters strongly depends on the curvature and symmetry of the SWNTs surface. In addition, the Pt clusters adsorbed on SWNTs support exhibit several energetically accessible structural configurations with rather low energy barriers that Pt clusters fluctuate rapidly among the configurations, results a high degree of structural fluxionality.

2. **Electronic structures of Pt$_n$ (n = 3, 4, 7, 10, 13) clusters on SWNT support**

   Figure 4 shows the projected density of states of Pt$_3$ cluster in the Pt$_3$-(10, 0) SWNT. Due to a strong mixing between d-states of Pt clusters and p-states of carbon nanotube, electronic structures of Pt$_n$ clusters on SWNTs are

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**Figure 1.** Simulation models of Pt$_n$ (n = 3, 4, 7, 10, 13) nanoclusters on SWNTs support

**Figure 2.** Optimized structure, electronic structures of single Pt atom on the (10, 0) SWNT support
characterized by broadening in a wide range of $d$-states of Pt clusters, and become bulk-like density of states, at relatively small cluster size. Electron density is accumulated at Pt-C bonds, and a small amount of charge transfer from Pt clusters to SWNT support occurs, in an agreement with experimental measurements. Geometry and electronic structures of Pt clusters on SWNTs support are strongly interrelated.

3. Chemical reactivity of Pt$_n$ ($n = 3, 4, 7, 10, 13$) clusters on SWNT support

Mechanisms of the most important chemical reactions at anode and cathode electrodes of fuel cell are investigated. The results demonstrate that adsorption process of O$_2$ and CO is strongly depends on ‘local structure’ and cluster size of Pt clusters on SWNT. Oxygen molecule preferably adsorbs on the Pt-Pt bridge site that suggests the mechanism of oxygen reduction reaction is preferred the 4-electron direct pathway. The ‘Bronsted-Evans-Polanyi’ relation, a linear relation between adsorption energy and activation energy of the dissociation adsorption of O$_2$ and CO oxidation reactions on the Pt clusters adsorbed on SWNT support, is observed. These results suggest that adsorption energies of O$_2$ and CO can be used as descriptors for screening and designing better catalysts in O$_2$ activation process, and CO oxidation process.

Conclusions

We have studies the physical and chemical properties of small Pt$_n$ ($n = 3, 4, 7, 10, 13$) clusters on single wall carbon nanotubes (SWNTs) support. The results are summarized as follows:

- Interaction between Pt and C is a weak covalent nature with hybridizations between $d$-states of Pt and $p$-states of C.
- Small Pt clusters on SWNTs support are preferred 3 dimensional structures. The stability of Pt clusters strongly depends on the curvature of the SWNTs. Small Pt clusters adsorbed on SWNTs support exhibit a high degree of structural fluxionality.
- Small Pt clusters on SWNTs support exhibit high reactivity with O$_2$, CO, OH species. Adsorption energies of O$_2$ and CO can be used as descriptors for screening and designing better catalysts.

The results of this research provide insight into understanding behaviors of Pt clusters on SWNTs support at atomic-scale. These findings also give useful properties of hybrid materials: small transition metal clusters on carbon nanotube support. This research can potentially open a new aspect to atom-by atom design and control the properties of small Pt clusters.
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