Preferential segregation of Pd atoms in the Ag-Pd bimetallic cluster: Density functional theory and molecular dynamics simulation

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Classical molecular dynamics simulations of the coalescence between an Ag cluster composed of 135 atoms and a Pd cluster of 16 atoms were performed at 500 K. All Pd atoms penetrated into the Ag cluster and preferentially segregated at the subsurface layer. The density functional theory calculations revealed that the center site is the most stable position for Pd atoms. However, the energy barrier for further penetration of Pd atoms located at the subsurface was too high to overcome, and thus a metastable structure with Pd atoms segregated at the subsurface did not evolve.

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It is well known that the bimetallic clusters composed of two different elements show unique optical and catalytic properties compared to single element clusters.1–4 Thus, investigation of the overall morphology and structural stability of bimetallic clusters has been one of the most important requirements for the practical application of bimetallic clusters.5–7 In general, the composite structure of bimetallic clusters such as core-shell or onionlike structures can be observed depending on the cluster system.6–10 It has been reported and believed that the significant difference in the atomic size and surface energy causes preferential surface segregation of the component having relatively large size and low surface energy, and gives rise to specific cluster morphology.9–11

Several computational methods such as genetic algorithms and Monte Carlo methods have been applied to investigate the globally stable structure of bimetallic clusters.6–8 It was confirmed that the core-shell structure is most stable in several bimetallic systems. In a recent report, Baletto et al. showed that a specific structure of a bimetallic cluster with subsurface segregated solute atoms can be found in Ag-based bimetallic cluster systems. They assumed that this is a metastable state, because all solute atoms penetrated into the cluster center at higher temperature.

In our previous molecular dynamics (MD) study of the coalescence between 135-Ag and 16-Pd clusters at 300 K, we observed penetration of Pd atoms into the Ag cluster and the formation of local fivefolded icosahedron (Ih) structures.5 However, some Pd atoms remained at the cluster surface, and thus did not segregate into the cluster core. As a result, the core-shell structure was not observed, which was explained by the strong mixing nature between Ag and Pd atoms and, more importantly, low kinetic energy. Thus, in the present work, we study the coalescence between 135-Ag and 16-Pd clusters at a somewhat high temperature of 500 K and compare the results with previous findings. We discuss the structural stability of the Ag-Pd cluster on the basis of density functional theory (DFT) and MD.

The MD simulation was carried out at 500 K under an NVT ensemble using a quantum Sutton-Chen (QSC) potential.12 The relevant potential parameters of Ag and Pd together with detailed MD simulation conditions were described in our previous study.5 An Ag cluster composed of 135 atoms and a Pd cluster of 16 atoms were individually prepared and fully relaxed at 500 K. They were later placed in a large simulation box (500 Å × 500 Å × 500 Å) and collided with each other. The MD simulation of the collision was carried out for 1000 ps (five million time steps).

The DFT calculations were performed using the DMol3 package in the Materials Studio.13 The Kohn-Sham equation was expanded in a local atomic orbital basis. The unrestricted Hartree-Fock (UHF) method with a double-numeric quality basis set with polarization functions (DNP) was applied. The effective core potential was applied to treat core electrons of heavy Ag and Pd atoms.14 The exchange-correlation functional in the generalized gradient approximation (GGA) was parametrized by the Perdew and Wang scheme.15 A Fermi smearing of 0.002 hartree (1 hartree = 27.2114 eV) was used.

The transition state calculations were performed using synchronous transit methods, the linear synchronous transit (LST), and quadratic synchronous transit (QST),16 in combination with the conjugate gradient (CG) minimization algorithm for subsequent refinement.17 In addition, the minimum energy path between the initial structure, transition structure, and final structure was investigated using the nudged elastic band (NEB) theory.18

Figure 1 summarizes the overall structure of the Ag-Pd cluster after 1 ns of MD simulation. Unlike at 300 K, all Pd atoms penetrated into the cluster, as seen in Fig. 1(a). This indicates that the average kinetic energy of Pd atoms was sufficient to overcome the energy barrier of penetration at 500 K. Almost (14 of 16) all of the penetrated Pd atoms preferentially segregated at the subsurface layer (one layer below the surface) of the cluster, as shown in Fig. 1(b), which is in partial agreement with the work of Baletto et al.5 They obtained a three-shell onionlike bimetallic nanocluster, and explained this abnormal cluster structure by referring to the most favorable site of impurities in an energetic manner. The penetrated Pd atoms induced surface disordering of the Ag cluster and the disordered Ag atoms at the surface were locally rearranged into Ih structures, Fig. 1(c).

Interestingly, in spite of relatively high simulation temperature, most Pd atoms (14 of 16) preferred to segregate at
bimetallic cluster study showed that the internal atomistic configuration of the cluster surface was expanded, developing a core-shell structure. In addition, penetrated Pd atoms were combined with the Ih structure at the surface, giving rise to Ih-Pd structures. One Ih-Pd is composed of six Ag atoms and one Pd atom embedded at the center, Fig. 1(d). The surface of the collided Ag-Pd cluster was covered with densely packed Ih-Pd and it persisted during the MD simulation. As a result, about two-thirds of the total surface area of the Ag-Pd bimetallic cluster was covered with Ih-Pd packages [Figs. 1(c) and 1(d)]. Once the Ih-Pd structure appeared, it maintained its morphology during the entire simulation period. Fourteen among a total of 16 Pd atoms were observed at the subsurface layer and 13 of them were involved in the Ih-Pd packages.

The local Ih-Pd structure was also observed at 300 K, though not as frequently. At 300 K, the Ih structure at the surface was expanded, developing a (111) plane. However, in this study at 500 K, all Pd atoms rapidly penetrated the cluster and many Ih structures were densely distributed on the cluster surface (61 surface atoms of the total 98 atoms that constitute the Ag-Pd cluster surface were involved in constituting Ih). This explains why an expanded (111) plane did not develop at 500 K.

Nam et al. reported that in studying the Au nanocluster, the liquid-to-Ih transition was initiated and affected by early penetration of Pd atoms. However, in the current study, this process occurred later. The fluctuation in average distance between Pd atoms and the cluster center with simulation time. It shows movement of Pd atoms along the subsurface layer of the Ag-Pd cluster. Although our MD simulation was performed at constant temperature, penetration of Pd atoms induced surface disordering and formation of Ih structures, followed by rearrangement of penetrated Pd atoms.

The Pd atoms eventually moved into specific positions below the Ih structures [Fig. 1(d)], giving rise to Ih-Pd packages. It is highly likely that the relative weak tendency of formation of a core-shell structure in the Ag-Pd system delayed further penetration of Pd atoms into the cluster center. This accounts for why the Pd atoms did not segregate at the cluster center in an early stage of the MD simulation.

It is notable that the penetrated Pd atoms mostly remained and moved on the near subsurface layer of the Ag-Pd cluster during the entire simulation period. According to Fig. 2 that shows this tendency, the average distance between the cluster center and Pd atoms did not change significantly after 200 ps and the Pd atoms moved along the subsurface layer of the cluster. Finally, Pd atoms moved into specific positions below the Ih structures.

In order to understand why the Ih-Pd structures are maintained stable, we modeled one Ih-Pd structure with a 55-Ih cluster composed of 54 Ag atoms and one Pd atom. There are three atomic positions along the cluster axis in the 55-Ih cluster from vertex to center: vertex, subsurface, and center sites. With respect to the Pd location, DFT calculations on the total energy of the cluster were performed. Initially, each cluster structure was geometrically optimized before total energy calculations. The total energy of the 55-Ih decreased by $-0.55$ and $-0.27$ eV, respectively, as the Pd atom moved from the vertex to subsurface site and from the subsurface to the center. This reflects a general tendency observed in the bimetallic system. The Gibbs free energy difference calculated between two 55-Ih structures, one with Pd at the subsurface and another with Pd at the center, showed that it is smaller than zero and decreases monotonically as a function of temperature. This also implies that the cluster center is the most stable site for Pd. However, the current study revealed that the Pd atoms preferentially segregated at the subsurface
energy in the Ag cluster was very similar between the sub-
atoms moved into specific positions, as shown in Fig. 1
MD simulations. However, it is not yet clear why the Pd
penetrated Pd atoms moved on the subsurface layer during
and another with Pd at the center. As mentioned earlier, the
between two 55-Ih structures, one with Pd at the subsurface
showed that there is reliable evidence of an energy difference
our DFT total energy calculations of the 55-Ih cluster
cannot be directly compared with their result.8 Nonetheless,
this study, we only employed a 55-Ih, and thus this result
sites and remained there. Thus, we assumed that once an
Ih-Pd package is formed, the energy barrier for further pen-
tration of Pd to the center is large enough to hinder move-
ment of Pd and to offset the driving force for minimization
of the total energy.
To test this assumption, the LST and/or QST and CG
methods were applied to determine the energy barrier for
penetration of Pd from the subsurface to the center and the
transient state structure. The minimum energy path was esti-
imated by the NEB method based on the LST and/or QST and
CG results. Figure 3 shows the structures at the transition
state and the energy barrier for Pd from the subsurface to the
center. The estimated minimum energy path is also shown.
The energy barrier was calculated as 4.655 eV. This is much
larger than the kinetic energy averaged for individual atoms
at 500 K, 0.065 eV. Thus, the large energy barrier prevents
further penetration of Pd atoms to the cluster center and, as a
result, the Ih structures with Pd at the subsurface were stabi-
lized, although at a metastable level.
Baletto et al. noted that in deposition of 200 Pd atoms
over a cluster of 201 or 147 Ag atoms the impurity (Pd)
energy in the Ag cluster was very similar between the sub-
surface and the center sites.8 In the case of the Ih-structured
Ag cluster, the center site was as favorable as the subsurface
site due to the small size mismatch between Ag and Pd. In
this study, we only employed a 55-Ih, and thus this result
cannot be directly compared with their result.8 Nonetheless,
our DFT total energy calculations of the 55-Ih cluster
showed that there is reliable evidence of an energy difference
between two 55-Ih structures, one with Pd at the subsurface
and another with Pd at the center. As mentioned earlier, the
penetrated Pd atoms moved on the subsurface layer during
MD simulations. However, it is not yet clear why the Pd
atoms moved into specific positions, as shown in Fig. 1(d).
This aspect is currently being investigated.
Recent studies on the Pd-Pt bimetallic cluster system re-
ported the appearance of an onionlike cluster and site pref-
ereence of the Pt solute atom.6,7 Lloyd et al. showed that Pt
solute atoms preferred the subsurface position in the Pd-Pt
55-Ih cluster.7 Referring to their report and considering our
result, it is likely that the nature of distribution of solute
atoms in the bimetallic cluster is different depending on the
kind of cluster systems.
Another point that must be considered is that the energy
barrier estimated by the synchronous transit method is an
upper limit to the barrier height and the NEB minimum en-
ergy path was also estimated based on the LST, QST, and CG
results. It is possible that there may be other pathways and
the corresponding energy barriers could be lowered. The
large gap between the calculated energy barrier and the av-
erage kinetic energy of individual atoms (4.59 eV) suggests
that even if there is a slight overestimation in our energy
barrier value, it does not seriously affect the overall results
obtained in this study. The estimated energy barrier was
based on the ideal 55-Ih cluster, and thus it may be lowered
by variation of local surface structures or the internal atomic
configuration of the real cluster. This can make the A-B-A to
A-B transition possible, as presented in the work of Baletto et
al.8
In general, the catalytic properties of metal surface and
clusters are very sensitive to their composition and atomic
distribution.20 It was asserted that the core-shell structure is
most stable in the Ag-Pd bimetallic cluster.10 However, our
results show that the metastable structure may be obtained
and maintains its stability under certain specific conditions.
The appearance of Ih and Ih-Pd at the surface must have
exerted a significant effect on atomic distribution of the
Ag-Pd cluster. Thus, it is highly likely that the catalytic prop-
erties of the Ag-Pd cluster may be different. As a result, our
study suggests that the kinetic aspects such as the energy
barrier between metastable and most stable structures or ev-
olution of the cluster surface and related solute distribution
should be carefully considered for the practical application
of bimetallic clusters.
In summary, we showed that Pd atoms preferentially seg-
regated at the subsurface layer of the Ag-Pd bimetallic clus-
ter. After collision between Ag and Pd clusters, two-thirds of
the surface Ag atoms were rapidly rearranged to Ih structures
and Pd atoms moved into specific positions below the Ih
thus, making Ih-Pd packages. It is found that the formation of
the Ih-Pd structure (equal to preferential segregation of Pd
atoms) and its stability were strongly affected by the surface
morphology of the cluster (existence of Ih structures at the
surface after collision). The local Ih-Pd structure is meta-
stable and the high-energy barrier hinders movement of Pd
atoms to more stable sites of the cluster center. This explains
why the bimetallic cluster structure with subsurface located
Pd atoms was preferred to the core-shell structure in this
work.
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