New Magic Au$_{24}$ Cluster Stabilized by PVP: Selective Formation, Atomic Structure, and Oxidation Catalysis

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ABSTRACT: An unprecedented magic number cluster, Au$_{24}$Cl$_x$ ($x = 0$–3), was selectively synthesized by the kinetically controlled reduction of the Au precursor ions in a microfluidic mixer in the presence of a large excess of poly(N-vinyl-2-pyrrolidone) (PVP). The atomic structure of the PVP-stabilized Au$_{24}$Cl$_x$ was investigated by means of aberration-corrected transmission electron microscopy (ACTEM) and density functional theory (DFT) calculations. ACTEM video imaging revealed that the Au$_{24}$Cl$_x$ clusters were stable against dissociation but fluctuated during the observation period. Some of the high-resolution ACTEM snapshots were explained by DFT-optimized isomeric structures in which all the constituent atoms were located on the surface. This observation suggests that the featureless optical spectrum of Au$_{24}$Cl$_x$ is associated with the coexistence of distinctive isomers. X-ray photoelectron spectroscopy and Fourier-transform infrared spectroscopy of CO adsorbates revealed the electron-rich nature of Au$_{24}$Cl$_x$ clusters due to the interaction with PVP. The Au$_{24}$Cl$_x$:PVP clusters catalyzed the aerobic oxidation of benzyl alcohol derivatives without degradation. Hammett analysis and the kinetic isotope effect indicated that the hydride elimination by Au$_{24}$Cl$_x$ was the rate-limiting step with an apparent activation energy of 56 ± 3 kJ/mol, whereas the oxygen pressure dependence of the reaction kinetics suggested the involvement of hydrogen abstraction by coadsorbed O$_2$ as a faster process.

KEYWORDS: magic number gold cluster, poly(N-vinyl-2-pyrrolidone), isomerization, aerobic alcohol oxidation, aberration-corrected transmission electron microscopy, density functional theory calculation

INTRODUCTION

Metal clusters with diameters smaller than the critical dimension of ∼2 nm are known to show unique chemical reactivities that cannot be predicted by a simple scaling law deduced from those of the extended surfaces and nanoparticles (NPs) of the corresponding metal.1−10 The emergence of the novel catalytic properties in metal clusters is ascribed to the quantized electronic structures, specific surface structures, and structural fluxionality. To understand the origin of novel catalysis and to optimize the catalytic performances of metal clusters, it is crucial to control their sizes with atomic precision. Atomically precise synthesis of Au clusters has been achieved using protecting ligands such as thiolates, alkynyls, halides, phosphines, and N-heterocyclic carbenes.2,3,11 Successful synthesis relies on size-focusing by chemical etching and subsequent isolation by chromatography and fractional precipitation. The main advantage of using atomically precise Au clusters for establishing the correlation between structures and catalytic performances is that their geometrical structures are determined by single-crystal X-ray diffraction (SCXRD).2,3 Nevertheless, these ligand-protected Au clusters are in general not suitable for catalytic applications because all the surface atoms of the metal core are masked by the ligands, although ligand-induced promotion of catalysis has been reported for some reactions.12

For the catalytic application of metal clusters, stabilization by linear polymers is suitable, since the cluster surface is inevitably exposed owing to the steric repulsion between the polymers.13−21 For example, it has been demonstrated that small (<3 nm) Au clusters stabilized by poly(N-vinyl-2-pyrrolidone) (PVP) show size-specific and high catalytic activities for oxidation and coupling reactions under aerobic conditions.22−26 The average diameter of the Au NPs can be tuned using conventional wet-chemical methods such as seed-mediated growth.27 It was reported that the catalytic activity of Au:PVP was monotonically enhanced with the decrease in the average diameter.22,24 However, postsynthetic size-focusing methods,2 size-selective fractionation,28 and SCXRD are not applicable because of the weak Au–polymer interactions and polydisperse...
structures of the polymer layers. Thus, major challenges for the catalytic studies of polymer-stabilized Au clusters are (1) atomically precise control of the cluster size and (2) determination of the atomic structure. The first challenge is tackled by controlling the formation kinetics of Au clusters in the presence of an excess amount of PVP using a microfluidic mixer, followed by matrix-assisted laser desorption/ionization (MALDI) mass spectrometry. Previous studies have shown that Au:PVP prepared by microfluidic mixing were a mixture of magic number Au clusters such as Au$_{35}$Cl$_x$, Au$_{43}$Cl$_x$, and Au$_{55}$Cl$_x$. Therefore, there is a possibility of producing atomically precise Au:PVP through further control of the formation kinetics. The second issue can be addressed by using aberration-corrected transmission electron microscopy (ACTEM). A three-dimensional (3D) atomic structure can be determined from the 2D projection image with the help of density functional theory (DFT) calculations. ACTEM video imaging with a high frame rate allows us to monitor the dynamic change in the structures at the atomic resolution.

In this study, we achieved atomically precise synthesis of an unprecedented magic number cluster, Au$_{24}$Cl$_n$ stabilized by PVP through the kinetically controlled reduction of Au precursor ions in the presence of a large amount of PVP. ACTEM video imaging with the help of DFT calculations revealed the coexistence of structural isomers in which all the Au atoms constituted the cluster surface. Au 4f X-ray photoelectron spectroscopy (XPS) and Fourier-transform infrared (FT-IR) spectroscopy of adsorbed CO indicated that Au$_{24}$Cl$_n$ was negatively charged. Kinetic studies were conducted for the aerobic oxidation of benzyl alcohol derivatives catalyzed by Au$_{24}$Cl$_n$. The clear kinetic isotope effect (KIE) and negative reaction constant in the Hammett plot indicated that the hydride elimination by Au$_{24}$Cl$_n$ was rate-limiting: the apparent activation energy was estimated to be 56 ± 3 kJ/mol by the Arrhenius equation. The reaction kinetics under pure oxygen (1 atm) suggested the involvement of direct hydrogen abstraction from adsorbed alkoxide by coadsorbed O$_2$. We believe that this work will open up a new research paradigm for atomically precise nanocatalysts.

## RESULTS AND DISCUSSION

### Atomically Precise Synthesis of Au$_{24}$Cl$_n$

Three samples of Au:PVP (a–c) were obtained at the [Au]:[PVP] mixing ratios of 1:40, 1:100, and 1:200, respectively. The size distributions of a–c were evaluated by MALDI mass spectrometry, based on our previous observations that Au clusters could be desorbed in the naked anionic form from the PVP matrix as a result of the MALDI processes. Figure 1 shows typical MALDI mass spectra of a–c in the negative ion mode recorded under minimal laser fluence. Sample a contained Au$_{24}$Cl$_{5}$ (x = 0–4) and Au$_{24}$Cl$_{1}$ (x = 0–4), whereas sample b was a mixture of Au$_{24}$Cl$_{4}$ (x = 0–3), Au$_{33}$Cl$_{1}$ (x = 0–2), and Au$_{34}$Cl$_{2}$ (x = 0–3). Most notably, the mass spectrum of sample c is dominated by the progression of peaks assigned to Au$_{34}$Cl$_{2}$ (x = 0–3). According to classical nucleation theory, formation of atomically monodisperse Au clusters in sample c is attributed to a kinetically controlled nucleation of Au atoms and a suppression of the subsequent growth. Such a condition was achieved by instantaneous and homogeneous reduction of all the AuCl$_{n}$ precursors by mixing with the strong reducing agent (BH$_4^-$) in the micromixer. The trend in Figure 1 indicates that

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**Figure 1.** Negative-ion MALDI mass spectra of Au:PVP (a–c).

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PVP also plays an important role in atomically precise synthesis by retarding the nucleation of Au atoms. The ligation by Cl originating from AuCl$_{4}^-$ has been frequently observed for small Au clusters such as Au$_{34}$Cl$_{16}$, Au$_{43}$Cl$_{16}$ [Au$_{35}$((PPh$_3$)$_2$Cl)], and Au$_{43}$PPh$_3$Cl$_{6}$. Elemental analysis of sample c determined the average number of the Cl ligands on Au$_{24}$ to be 9.4, suggesting that the MALDI process is accompanied by dissociation of the Cl ligands. Such selective dissociation suggests that the Cl ligands are weakly bound to Au$_{24}$ as electron withdrawing ligands as implied by weak magic behavior at x = 1 and 3 (Figure 1c). Noncontamination of sample c with Au NPs larger than 2 nm was confirmed by the following results: (1) absence of the localized surface plasmon resonance (LSMR) band at ~520 nm in the optical absorption spectrum (Figure 2a); (2) observation of a single broad peak in the powder X-ray diffraction (XRD) pattern (Figure 2b); (3) small coordination number (CN) of the Au–Au bond (4.2 ± 0.5) obtained by the curve-fitting analysis of the Au L$_3$-edge extended X-ray absorption fine structure (EXAFS) (Figure 2c, Table S1); and (4) mean diameter of the Au clusters estimated to be 1.0 ± 0.2 nm by ACTEM (Figure 2d). We concluded from these results
that single-sized Au_{24}Cl_{4} was selectively formed in sample c. In the following, we focus on sample c, which is hereafter referred to as Au_{24}Cl_{4}:PVP.

**Geometrical Structure of Au_{24}PVP**

It is well-known that the magic stability of naked Au clusters is governed by electronic shell closure based on a jellium model. The formation of Au_{24} and Au_{24}Cl_{4} in PVP reported in our previous studies is explained by the same model. In contrast, Au_{24}Cl_{4} in Au_{24}:PVP corresponds to an unprecedented magic cluster. This result implies that the Au_{24}Cl_{4} cluster does not have a spherical shape and does not provide a spherical potential to confine the valence electrons. To gain insight into the atomic structure of Au_{24}Cl_{4}, we first conducted DFT calculations on Cl-free Au_{24} clusters. Figure 3 lists the DFT-optimized structures of Au_{24} (1–7) in the order of the relative stability. The frontier Kohn–Sham orbitals of 1–7 are depicted in Figure S1. The relative energies (ΔE) and energy gaps between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) are summarized in Table 1. Interestingly, all the constituent Au atoms of structures 1–6 are located on the surface: 1 and 3 have flat cage structures; 4 and 6 have empty tubular shapes; and 2 is constructed by attaching an Au_{4} unit to the well-known pyramidal Au_{20} magic cluster. In contrast, the least stable structure 7 contains two Au atoms inside the Au_{24} cage. The dimensions of 1–7 fall within the diameter distribution shown in Figure 2d. The projection of 4 or 6 along the longitudinal axis gave the smallest diameter (~0.7 nm), whereas that of 1 or 3 from the flattened surface normal gave the largest diameter (~1.3 nm) (Figure S3). This semiquantitative agreement supports the validity of the structural models in Figure 3. These results suggest that various disordered structures such as 1–6, in which all the constituent Au atoms constitute the cluster surface, are energetically accessible. Then, the effect of Cl ligation on the geometrical structures of Au_{24} was studied by using Au_{24}Cl_{4} with 20 valence electrons as a model. Figure 4 lists the isomeric structures 1Cl–6Cl obtained by optimizing the initial structures in which four Cl ligands are bonded to low coordination sites of low energy isomers 1–6 in Figure 3, respectively. The frontier Kohn–Sham orbitals of 1Cl–6Cl are depicted in Figure S2. Cl ligation induced the slight distortion of the Au frameworks and as a result the relative stability of the isomer was altered. Nevertheless, it is safe to conclude that Au_{24}Cl_{4} can take isomeric structures in which all the Au atoms are located on the surface, similarly to the naked Au_{24} clusters. The CNs and average lengths of the Au–Au bonds in 1–7 and 1Cl–6Cl are summarized in Table S2. The CN values of the Au–Au bonds in 1–7 (5.0–5.4) and 1Cl–6Cl (4.2–4.8) qualitatively agreed with that (4.2 ± 0.5) determined by Au L_{3}-edge EXAFS analysis (Table S1), while the calculated Au–Au bond lengths of 1–7 (2.87–2.90 Å) and 1Cl–6Cl (2.85–2.87 Å) are longer than that experimentally determined (2.71 ± 0.01 Å). In the following discussion of the geometrical structures, we ignore the Cl ligands.

**Theoretical prediction of energetically comparable isomers**

In this paper, we explore the structural diversity and dynamic behavior of Au clusters. Theoretical prediction of energetically comparable isomers suggests that Au_{24} clusters in PVP are composed of distinct structural isomers and/or interconvert among isomers under ambient temperature. To address this interesting issue, atomic structures were probed by ACTEM video imaging at 0.04 s per frame. The ACTEM images of Au_{24}:PVP did not show any signs of atom ejection during continuous observation for several minutes. Figure 5 shows representative ACTEM snapshots of different particles (#1–#4) of Au_{24}:PVP. These images could not be reproduced by considering a single isomeric form in Figure 3 but could be by using different isomers 1, 2, 2, and 5, respectively. Other snapshots of different particles (#5–#7) in Figure S4 reasonably agree with the images simulated for isomers 4, 5, and 7, respectively. No images could be assigned to the icosahedral nor face-centered cubic structures observed in the selenolate (RSe)- or thiolate (RS)-protected gold clusters Au_{24}(SeR)_{14} and Au_{144}(SR)_{60}. Furthermore, the images of a single particle (#8) of Au_{24}:PVP fluctuated every 0.08–0.4 s during the observation period, as exemplified in Figure S5 and Movies S1–S3. Figure 6 shows snapshots of particle #8 taken from the images in Figure S5 at different times (t_{1}–t_{4}). Comparison with a set of images simulated by systematically changing the projection directions revealed that the fluctuation of the images was associated with structural isomerization rather than simple rotation of a single isomer: the images of Figure 6a–6c are consistent with a six-fold rotation and those of Figure 6d–6f correspond to a two-fold rotation. These results suggest that the dynamics of Au clusters in PVP are driven by interconversion among structural isomers.
of Au24:PVP interconverted among structural isomers possibly h), respectively. Thus, we concluded that the individual particles exhibited a broad peak of 12CO adsorbed on Au24:PVP at 2106 cm−1, in addition to that of free 12CO at 2136 cm−1 (Figure 7b).

The redshift is ascribed to the π back-donation from the electron-rich Au sites of Au24Cl4.63 To gain deeper insight into the geometrical and electronic structures of Au24:PVP, CO-adsorbed structures were theoretically studied for two isomers 1 and 4 with charge states of 0 and −1. DFT-optimized structures are illustrated in Figure S6: structural isomers 1COa=1COa and 1COa=1COa with different CO adsorption sites were obtained for neutral and anionic 1, respectively, whereas 4COc=4COc and 4COa=4COa were for neutral and anionic 4, respectively. The adsorption energy and stretching frequency of CO adsorbed on different sites of 1 and 4 are summarized in Tables S3 and S4, respectively. The binding energies are in the range of 0.1–0.4 eV regardless of the structures, charge states, and adsorption sites of Au24 clusters. The CO stretching frequencies on neutral 1 and 4 (green plots in Figure 7b) differ only slightly (<20 cm−1) depending on the adsorption sites and can explain the experimental result. On the other hand, the CO stretching frequencies on anionic 1 and 4 (blue plots in Figure 7b) are significantly red-shifted compared to those for neutral 1 and 4. Although these results could not identify the structure of the Au24:Cl4 core and adsorption site of CO, they imply that the negative charge on the Au24:Cl4 core is smaller than 1e as a result of a balance between the electron-withdrawing nature of Cl and the electron-donating nature of PVP.

Aerobic Oxidation Catalysis of Au24:PVP

Since Haruta’s discovery of CO oxidation catalyzed by Au NPs on metal oxide supports,49 the catalytic properties of nanosized Au have attracted considerable interest in nanoscience.57–60 In particular, alcohol oxidation is catalyzed efficiently by Au-based catalysts5,51,52 and has been studied as a model reaction to investigate the effects of structural parameters on the catalysis. For example, we have demonstrated that Au:PVP shows size-specific and high catalytic activities for aerobic alcohol oxidation.23,24 However, these studies were conducted using monodisperse Au clusters but with distributions in the number of constituent atoms. Au24:PVP synthesized here provides a novel opportunity to study atomistic details of the catalytic mechanism.

Kinetic measurements were conducted for the aerobic oxidation of p-substituted benzyl alcohol X-C6H4CH2OH (X = OCH3, CH3, H, CF3) catalyzed by Au24:PVP (Figure 8a). The selectivity of products between the aldehydes and carboxylic acids is as a function of the substituent X. The aerobic oxidation of p-substituted benzyl alcohol catalyzed by Au24:PVP was monitored using GC analysis and the selectivity of products was determined by comparing the peak areas of the aldehydes and carboxylic acids.
It was also reported that the Cl ligands of PVP-stabilized Au34Cl2 di出售 to O2. In this mechanism, the resulting superoxo-like species (Figure 8a) was explained by the hydride elimination mechanism by Au24 (Scheme 1b). To gain further insight into the mechanism by Au24:PVP, the effect of O2 pressure was investigated. The time course of the alcohol oxidation under 1 atm of O2 (Figure 9a) showed two characteristic behaviors: (1) for t > 0.05 h, the rate constant under pure O2 (1.18 h⁻¹) was larger than that under air (O2: 0.21 atm, 0.87 h⁻¹), although the rate constants were not proportional to the oxygen pressure; (2) the rate constant at the initial stage (t < 0.05 h) was apparently larger than that for t > 0.05 h. The former can be explained by the hydride elimination mechanism (Scheme 1b) with the assumption that the catalyst recovery by O2 is not much faster than the hydride elimination. However, it was difficult to rationalize the latter only by hydride elimination mechanism. We propose that hydrogen abstraction by O2 and (c) hydride elimination by Au24.

Two alcohol oxidation pathways on Au24:PVP: (b) hydrogen abstraction by O2 and (c) hydride elimination by Au24, adsorbed hydrogen is removed by O2. This hydride elimination mechanism has been accepted for alcohol oxidation by Au NPs supported on metal oxides such as TiO2, CeO2, and Al2O3. A similar reaction mechanism has been proposed by Kaneda for alcohol oxidation catalyzed by hydroxyapatite-supported Pd clusters.

The reaction constant for Au24:PVP determined by the Hammett plot under air (Figure 8b, −1.2 ± 0.2) is comparable to those reported for Au NPs supported on metal oxides (−0.2 to −1.4). This fact also lends support for the hydride elimination mechanism by Au24 (Scheme 1b). Two alcohol oxidation pathways on Au24:PVP: (b) hydrogen abstraction by O2 and (c) hydride elimination by Au24.

The negative reaction constant (ρ = −1.2 ± 0.2) indicates that the benzyl carbon has cationic characteristics at the transition state of the rate-limiting step. Furthermore, a large KIE (kH/kD = 4.1) was observed for α-deuterated benzyl alcohol (C6H5CH2CD3OH). Thus, it was concluded that the C–H bond cleavage at the benzylic position was the rate-limiting step and that the H atom was abstracted as a hydride. The apparent activation energy for X = H was estimated to be 56 ± 3 kJ/mol from the Arrhenius plot (Figure 8c).

Importantly, MALDI-MS, optical spectroscopy, and Au L3-edge EXAFS confirmed that the cluster size of Au24:PVP was retained during the catalytic reactions under the present conditions (Figures S9–S19). In contrast, MALDI-MS and Au L3-edge EXAFS suggested the loss of Cl ligands during the reaction (Figures S9–S19). These results indicate that the catalysis is governed by Au24 and not a different number of Cl ligands showed a similar activity (Figures S9 and S15). It was also reported that the Cl ligands of PVP-stabilized Au43Cln and Au46Cln were easily released during the reactions and did not prevent the catalytic alcohol oxidation.

Two different mechanisms have been proposed for aerobic alcohol oxidation in which O2 plays different roles. We have proposed that O2 is reductively activated upon adsorption onto the negatively charged Au:PVP based on the correlation between the electron affinities of naked Au and the reactivity to O2. In this mechanism, the resulting superoxo-like species abstracts hydrogen from the α-carbon of coadsorbed alkoxyde (Scheme 1a). A similar mechanism has been proposed by Ebitani in the alcohol oxidation by hydrotalcite-supported metal (AuPd, AuPt or Pt) NPs costabilized by polymers such as PVP and starch. The other mechanism, proposed by Kobayashi, Chechik, and co-workers is based on the observation of an Au–H intermediate during the alcohol oxidation on Au NPs stabilized by styrene-based copolymers (Au:PS): hydride is directly transferred to the Au NP surface (Scheme 1b) and

**Figure 8.** (a) Aerobic oxidation of p-substituted benzyl alcohol X-CsH11CH2OH (X = OCH3, CH3, H, CF3) catalyzed by Au24:PVP. (b) Hammett plot for the aerobic oxidation of p-substituted benzyl alcohol catalyzed by Au24:PVP dispersed in H2O. (c) Arrhenius plot for the benzyl alcohol oxidation (X = H). Reaction conditions: substrate 80 μmol; Au24:PVP 5 atom %; K2CO3 300 mol %; H2O 20 mL; air; 303 K (unless specified).

**Figure 9.** (a) Natural logarithm of benzyl alcohol concentration (normalized by initial concentration) during the catalytic oxidation reaction under air (blue, O2: 0.21 atm) or pure O2 (red, O2: 1 atm). Conditions other than O2 pressure were the same as those for Figure 8. Two alcohol oxidation pathways on Au24:PVP: (b) hydrogen abstraction by O2 and (c) hydride elimination by Au24.

**Scheme 1.** Previously Proposed Reaction Mechanisms of Alcohol Oxidation Catalyzed by (a) Au:PVP and (b) Au:PS.
concentrator (Vivaspin20, Vivascience) having a membrane.

benzyl alcohol (C₆H₅CD₂OH) was purchased from Sigma-Aldrich.

described by assemblies of PVP in which Au clusters can be

unprecedented magic number is associated with the cavity size

calculations. We speculate that the observation of an

constituent Au atoms on the surface are suggested by DFT

imaging revealed the polydispersity and

step by the presence of a large amount of PVP. ACTEM video

hydration of benzyl alcohol derivatives while retaining the size. Kinetics studies

revealed that the hydride elimination from benzyllic carbon by

Au₄₃ corresponds to the rate-limiting step with an apparent

activation energy of 56 ± 3 kJ/mol. It was also suggested from the

O₂ pressure dependence on the reaction kinetics that the

hydrogen abstraction by the adsorbed O₂ is involved as well. We

propose that the oxidation is initiated by the hydrogen

abstraction by O₂ adsorbed on Au₄₃, followed by the rate-

limiting hydride elimination by Au₄₃.

CONCLUSION

In summary, a novel magic number cluster, Au₄₃:PVP, was

serendipitously and reproducibly obtained by homogenizing the formation process using a micro mixer and limiting the growth step by the presence of a large amount of PVP. ACTEM video imaging revealed the polydispersity and fluxionality of the atomic structure of Au₄₃:PVP. Noncrystalline structures with all constituent Au atoms on the surface are suggested by DFT calculations. We speculate that the observation of an

unprecedented magic number is associated with the cavity size

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METHODS

Chemicals

All chemicals were commercially available and were used without further purification. Hydrogen tetrachloroaurate(III) trihydrate, sodium borohydride, PVP (K30, average molecular weight: ∼40 000), benzyl alcohol, p-methylbenzyl alcohol, p-methoxybenzyl alcohol, and potassium carbonate were purchased from Wako Pure Chemical Industries. trans-2-[3-(4-tert-Butylphenyl)-2-methyl-2-

propenylidene]malononitrile (DCTB) and p-(trifluoromethyl)benzyl alcohol were purchased from Tokyo Chemical Industry. Deuterated benzyl alcohol (C₆H₅CD₂OH) was purchased from Sigma-Aldrich.

Deionized water used was Milli-Q grade (>18 MΩ cm).

Synthesis

Au clusters were synthesized by microfluidic mixing of two aqueous solutions of Au precursors and a reducing agent as reported previously.⁷⁻⁻¹⁻⁴⁻⁵⁻⁻¹ The ice-cooled aqueous solution of HAuCl₄ and PVP was mixed with that of NaBH₄ and PVP at a flow rate of 200 mL/h in a micro mixer (SIMM-V2, IMM GmbH) placed in an ice bath. Three samples of Au:PVP (a–c) were obtained by tuning the [Au₃⁺]:[PVP] mixing ratios as listed in Table 2. The resulting brown solution was collected in an Erlenmeyer flask, stirred for 1 h at 273 K, and then deionized four times at 273 K by using a centrifugal ultrafiltration concentrator (Vivaspin20, Vivascience) having a membrane filter with a
cutoff molecular weight of 10 kDa (for a) or 100 kDa (for b and c). The deionized solution was lyophilized to obtain Au:PVP in a powder form. The yields of a–c were 79, 73, and 79%, respectively, based on the amount of Au. Optical absorption spectroscopy and thermogravimetry showed that the amount of PVP was reduced to ∼40 equiv (monomer unit to Au atom) by ultrafiltration. PVP was added to adjust the equivalence to 50 in the catalytic tests.

Characterization

MALDI mass spectra were recorded using a time-of-flight mass spectrometer (AXIMA-CFR, Shimadzu): DCTB was used as a matrix. The specimens for MALDI-MS were prepared by drop-casting a methanol dispersion of Au:PVP and DCTB ([Au₃⁺]:[DCTB] = 1:18) onto a sample plate. Optical absorption spectra were recorded by using a spectrophotometer (V-670, JASCO). The PXRD pattern was obtained using a diffractometer with CuKα radiation (SmartLab 3, Rigaku). ACTEM was carried out on a JEM-ARM200F instrument (JEOL) at an acceleration voltage of 80 kV, under 1 × 10⁻⁵ Pa at 298 K in the specimen column. A series of ACTEM images were continuously recorded at 25 frames per second (fps) on a CMOS camera (Gatan OneView with in situ option, output image size: 2,048 × 2,048 pixels operated in the binning 2 mode, pixel resolution: 0.01 nm at x2,000,000 magnification). We adjusted the spherical aberration value, electron dose rate, and defocus value to 1–3 μm, 2.7–5.6 × 10⁻⁶ e⁻·nm⁻²·s⁻¹, approximately −4 nm (underfocus), respectively. Acquired images were filtered by a bandpass filter using ImageJ software,⁶⁶ structures larger than 40 pixels or smaller than 3 pixels were filtered with the tolerance of direction of 5%. The specimens for ACTEM were prepared by drop-casting a methanol dispersion of Au:PVP onto a thin carbon-coated copper grid (SHR-C075, Rigaku). The yields of Au₄₃:PVP catalyzed the aerobic oxidation of benzyl alcohol by using a spectrophotometer (FTIR-4200, Jasco) in the transmission mode at a resolution of 2 cm⁻¹. For FT-IR measurements, Au:PVP (Au 15 μmol) was dispersed in dichloromethane (3 mL) under CO (1.0 atm) and the colloidal dispersion was introduced into the IR cell (CaF₂ window, 0.5 mm path length) after the stirring for 50 min. Au L₃-edge X-ray absorption spectroscopy (XAS) was conducted in the transmission mode using the BL01B1 beamline at Spring-8 of the Japan Synchrotron Radiation Research Institute (JASRI). The incident X-ray beam was monochromatized by a Si (111) double crystal monochromator. The spectral data were analyzed by using the EXCURVEX program (Rigaku). The k¹-weighted χ spectra in the k range of 3–16.5 Å⁻¹ were Fourier-transformed into the r space. The Au–Au and Au–Cl bonds were considered in the curve-fitting analysis over the r range of 1.7–3.0 Å.

Calculations and Simulations

DFT calculations of the model structures of Au₄₃, Au₄₃Cl₄, and Au₄₃(CO) were performed using the Gaussian 09 program.⁷² B3LYP was used as a functional.⁷³ A double-ζ basis set with scalar relativistic effective core potential (ECP), LANL2DZ, was applied for Au and the 6–31G(d) basis set was used for C, O, and Cl. Structures were obtained by reoptimization of those reported previously.⁶⁷⁻⁻¹⁻⁷⁻⁻¹ PVP was not included in the calculations since the impact on the atomic structure of Au clusters is negligibly small.⁷⁻⁻¹ Neutral clusters (Au₄₃, Au₄₃(CO), and Au₄₃Cl₄) were optimized with the singlet spin state, whereas the doublet spin state was applied for the anionic cluster, Au₄₃(CO)⁻. The vibrational frequencies were computed for the optimized structures to ensure that they corresponded to the local minima of the potential energy surface. The relative energy of the clusters presented in this study includes the zero-point energy. For the stretching frequencies of CO adsorbed on Au₄₃, a scaling factor of 0.970 was applied so that the calculated frequency of isolated CO (2203 cm⁻¹) reproduced that of free CO in dichloromethane (2136 cm⁻¹). TEM simulation images of DFT-optimized structures were generated by using a multislice procedure implemented in a BioNet

Table 2. Conditions of Synthesizing Au:PVP

| Sample | [HAuCl₄] [mM] | [PVP] [mM] | [NaBH₄] [mM] | [PVP] [mM] | [Au₃⁺] | [NaBH₄] | [PVP] |
|--------|---------------|------------|--------------|------------|--------|--------|-------|
| a      | 10            | 200        | 50           | 200        | 1:5:40 |        |       |
| b      | 10            | 500        | 50           | 500        | 1:5:100|        |       |
| c      | 10            | 1000       | 50           | 1000       | 1:5:200|        |       |

*Concentrations of monomer unit.*
elbis software. DFT-optimized structures were converted to a series of simulated TEM images with the observation direction varying every 1° around the x and y axes, producing 18 × 36 = 648 images. A defocus value of ~4 nm and a spherical aberration coefficient (Cs) of ~3 µm were applied for the simulation. To quantify the similarity of the experimental and simulated images, a cross-correlation analysis was conducted. The cross-correlation function of two images (a and b), \( \gamma(a, b) \), is defined as follows:

\[
\gamma(a, b) = \frac{\sum_{i,j} [I_a(r_{ij}) - \bar{I}_a] [I_b(r_{ij}) - \bar{I}_b]}{\sqrt{\sum_{i,j} [I_a(r_{ij}) - \bar{I}_a]^2 \cdot \sum_{i,j} [I_b(r_{ij}) - \bar{I}_b]^2}}
\]

where \( I_a, I_b, \) and \( \bar{I} \) represent the intensity of the image, pixel position, and mean intensity, respectively. The value of this function is an index of the similarity of images; \( \gamma \) of 1 or 0 corresponds to a perfect match or no match, respectively. Therefore, the cross-correlation function of the experimental and simulated images was calculated for the efficient screening of the large number of simulated images. Candidate images obtained by the screening were then carefully compared to the experimental images.

Catalytic Test

Catalytic properties of Au:PVP for aerobic oxidation of benzyl alcohol derivatives in water were studied by using a temperature-controlled personal organic synthesizer, PPS-2510 (Eyela). The catalytic reaction was initiated by adding the aqueous dispersion (8 mL) of Au:PVP (4 µmol) to the aqueous solution (12 mL) of alcohol (80 µmol) and \( \text{K}_2\text{CO}_3 \) (240 µmol) magnetically stirred at 800 rpm. Aliquots (5 mL) were sampled at the given time and hydrochloric acid (HCl 1 mol) was added to quench the reaction. Reaction products were extracted by ethyl acetate (1 mL) three times. The extract thus collected was dried over Na2SO4 and analyzed by a gas chromatograph (GC).

**Notes**

The authors declare no competing financial interest.

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