Synthesis of gold nanoparticles with different atomistic structural characteristics

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Abstract

A chemical reduction method was used to produce nanometric gold particles. Depending on the concentration of the main reactant compound different nanometric sizes and consequently different atomic structural configurations of the particles are obtained. Insights on the structural nature of the gold nanoparticles are obtained through a comparison between digitally-processed experimental high-resolution electron microscopy images and theoretically-simulated images obtained with a multislice approach of the dynamical theory of electron diffraction. Quantum molecular mechanical calculations, based on density functional theory, are carried out to explain the relationships between the stability of the gold nanoparticles, the atomic structural configurations and the size of nanoparticles.

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1. Introduction

Nanoparticles are defined as atomic arrangements with nanometric dimensions and usually with a small number of constituent atoms. The structural configurations of these small particles induce new types of physical properties. Therefore, the selection of a synthesis method which can give rise to a particular structural particle configuration is very important for new technological applications [1,2]. Depending on the nanometric particle size, the nanostructures can have different applications. These applications include fields such as: catalysts, photography, medicine, information storage in magnetic devices etc. In recent years, the high-resolution electron microscope (HREM) has been one of the main tools used to study the physical properties of nanometric particles. Metallic nanoparticles usually display simple geometric atomic arrangements, mainly with tetrahedral, octahedral and decahedral forms [3,4]. They even present atomic configurations which display five-fold symmetry [5]. In the reduction synthesis method used in this investigation, particles of different average sizes and different geometrical arrangements have been obtained.

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The morphological nature of the nanometric particles depends on the concentration of the main reagent compound used in the reduction process. In this study, the structural characterizations of gold nanoparticles with different structural morphologies have been carried out through a comparison between experimental and theoretical HREM images. The stability of the nanoparticles is also evaluated using density functional theory based on quantum mechanical calculations.

2. Experimental procedures

Ultra fine gold particles have been obtained using the chemical reduction method reported earlier [6,7]. Methanol solutions of gold ions were prepared by dissolving crystalline hydrogen tetrachloroaurate (HAuCl₄ · xH₂O) in methanol (0.033, 0.044, 0.099, 0.11 and 0.12 mmol in 25 ml of methanol). A methanol solution of PVP [poly (N-vinyl-2-pyrrolidone) (150 mg of PVP in 25 ml of methanol), was added to the metal ion mixture. To reduce the metal ions, 6 ml of an aqueous solution of NaBH₄ (0.066 M) was added to the mixture solution dropwise at room temperature. A homogeneous colloidal dispersion was formed after the addition of the NaBH₄ reductant in the solution containing the metal ions.

The structural and morphological characteristics of the dispersed metallic nanoparticles have been studied using a transmission electron microscope, Philips Tecnai F20, which has a field emission gun attachment, an operating voltage of 200 keV, a spherical aberration of 1 mm, and a direct maximum resolution dot to dot of 0.23 nm. TEM specimens were obtained from the homogeneous colloidal solution. Drops of this solution were deposited on a copper grid (3 mm in diameter) with an amorphous carbon film. The HREM images have been digitally processed. Theoretical simulations based on the multislice approach of the dynamical theory of electron diffraction [8] have been carried out to generate HREM images of gold nanoparticles.

Studies on the stability of the nanoparticles were carried out using quantum molecular calculations. These simulations were carried out using the Dmol³ software by Accelrys [9], which is based on the density functional
theory (DFT). In our case we used the generalized gradient approximation (GGA) with the Perdew–Wang (PW91) function [10] and a field consistent tolerance of $1 \times 10^{-6}$ eV. The configurations of 13, 19 and 38 atoms for cubo-octahedral and 13, 19 and 39 atoms for decahedral structures were studied; the selection of these clusters is focused because of the experimental results and previous reports that demonstrate these are the most commonly produced [11–13], even though from many reports suggest other more stable configurations [14–16]. The minimum energy for each of the structures was obtained by means of a geometry optimization task based on the DFT method. The geometry optimization was followed by a single point energy calculation to obtain the electrostatic potential, highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO), and Fukui functions.

3. Results and discussion

Fig. 1 shows low magnification electron microscope images of the nanometric gold particles obtained with different concentrations of the Au ions. These low magnification images just show the relative particle size increments as the concentration of gold ions in the reaction mixture is increased. The average particle size varied from a few nanometers at 0.033 mmol (in 50 ml of reaction mixture) to approximately 12 nm for the 0.12 mmol metal ion content. This is further illustrated in Fig. 2, where the average particle size values are displayed as a function of Au ion content in the solution. The nature of the atomistic structural configuration is also strongly dependent on the total ion content in the reaction mixture.

For example, for a metal ion content of 0.044 mmol, Fig. 3 shows some commonly found configurations. Fig. 3a illustrates a single twinned particle (stp) and Fig. 3b and c shows fcc-like particles. On the other hand, for a metal ion content of 0.11 mmol, Fig. 4 shows two fcc-like particles and one multi-twinned particle. A statistical summary of these structural characteristics is given in Fig. 5. At low concentrations of the metal ions
(0.033 mmol/50 ml), the more commonly found structures are decahedral and fcc-deformed; however, as the concentration is increased (0.044, 0.099 and 0.11 mmol/50 ml), the most commonly found structures change to single twinned particles and fcc-like particles. It is also important to mention that the size of the particles obtained for the lowest concentration of metal ions is smallest; only a few nanometers in diameter with less than 2 nm on average. Depending on the metal ion concentration in the reaction mixture, two main regions of nanoparticle growth with distinct characteristics in atomic ordering and hence of distinct structural features

![HREM images of gold nanoparticles with their corresponding power spectra](image1.png)

**Fig. 4.** HREM images of gold nanoparticles with their corresponding power spectra, prepared from a solution with 0.11 mmol/50 ml gold ion concentration; a) and b) fcc-like particles, c) multi-twinned particle.

![Graph showing statistical distribution of gold nanoparticles](image2.png)

**Fig. 5.** Statistical distribution of gold nanoparticles with different atomistic structural forms for different gold ion concentrations in the reaction mixture. These experimental results were obtained from the analysis of HREM images; fcc-like (fcc), fcc-deformed (fccd), decahedral (deca) and single twin particle (stp). The ion concentrations are expressed in mmol/50 ml.

![HREM images of a cubo-octahedron gold particle](image3.png)

**Fig. 6.** a) HREM image of a cubo-octahedron gold particle obtained from the reaction solution of lowest gold ion concentration (0.033 mmol/50 ml), b) same HREM image after digital processing, and c) theoretically-simulated image of a cubo-octahedron along the [103] crystalline orientation.
are found. Low concentrations (0.033 mmol/50 ml) induced the formation of decahedral and fcc-deformed gold particles. However, higher concentrations (0.044, 0.099 and 0.11 mmol/50 ml) induced the formation of single twinned particles and fcc-like particles.

A qualitative insight of the atomistic structural characteristics of the nanometric gold particles is obtained by comparing the experimental high-resolution images and theoretically-simulated images. Fig. 6a shows the HREM image of a nanometric gold particle produced under a low concentration condition of gold ions. Fig. 6b illustrates the same HREM image digitally processed. Fig. 6c shows the theoretically-simulated HREM image of a cubo-octahedron along the [103] crystalline orientation. A comparison between the experimental and simulated images suggests the possible nature of the atomistic structural configuration of the image shown in Fig. 6a. Therefore, Fig. 6a resembles the HREM image contrast produced by a cubo-octahedron.

Fig. 7a shows the HREM image of a typical gold nanoparticle obtained for a gold ion concentration of 0.044 mmol/50 ml and Fig. 7b displays the same image after it has been digitally processed. Fig. 7c shows the theoretical HREM simulation of an fcc-like particle oriented along the [011] crystalline axis. A comparison between the experimental image and the theoretical image suggests that the particle has a structure resembling fcc. Fig. 8 shows a single twinned gold particle obtained from the 0.044 mmol/50 ml gold ion concentration. The theoretical simulation was obtained assuming a model of two fcc gold crystals along the [011] orientation and joined planes of the (111) type. The image contrast obtained in the simulation resembles the contrast displayed in the experimental HREM images.

<table>
<thead>
<tr>
<th>Structure</th>
<th>Number of Au atoms</th>
<th>Binding energy per atom (eV)</th>
<th>HOMO (eV)</th>
<th>LUMO (eV)</th>
<th>Gap (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cubo-octahedral</td>
<td>13</td>
<td>-1.376</td>
<td>-3.715</td>
<td>-3.061</td>
<td>-0.654</td>
</tr>
<tr>
<td></td>
<td>19</td>
<td>-1.506</td>
<td>-3.650</td>
<td>-3.172</td>
<td>-0.479</td>
</tr>
<tr>
<td></td>
<td>38</td>
<td>-1.156</td>
<td>-3.791</td>
<td>-3.118</td>
<td>-0.673</td>
</tr>
<tr>
<td>Decahedral</td>
<td>13</td>
<td>-1.349</td>
<td>-3.460</td>
<td>-3.196</td>
<td>-0.264</td>
</tr>
<tr>
<td></td>
<td>19</td>
<td>-1.448</td>
<td>-3.414</td>
<td>-3.062</td>
<td>-0.352</td>
</tr>
<tr>
<td></td>
<td>39</td>
<td>-1.163</td>
<td>-3.252</td>
<td>-2.790</td>
<td>-0.461</td>
</tr>
</tbody>
</table>
When the concentration is increased, the structure changes. Structures are decahedral and fcc-deformed. However, gold ions in the reaction mixture increases. For a lower size of gold nanoparticles increases as the concentration of the material must have a highly dispersed activation energy suitable for wide range of catalytic applications.

4. Conclusions

Gold nanoparticles with narrow size distributions have been prepared using a chemical reduction method. The size of gold nanoparticles increases as the concentration of gold ions in the reaction mixture increases. For a lower concentration of the metal ions, the commonly found structures are decahedral and fcc-deformed. However, when the concentration is increased, the structure changes to single twinned and fcc. The theoretical analysis by a quantum mechanical approach reveals that the electronic structure of nanoparticles is significantly affected by their atomic configurations; inducing higher catalytic activity for decahedral configurations when five-fold faces are exposed. However, due to the coexistence of structures such as fcc-like and decahedral in the synthesized sample, the electronic configuration is dependent on the number of atoms, while the structure/configuration has no influence. From the energy gap values, we can observe that the decahedral structure Au13 have the lowest energy HOMO–LUMO gap, which is due to the higher density of electrons on its surface. Therefore, we can say that in the case of Au, the decahedral structures are favorable when the particle size is small, and when the particle size increases, the cubo-octahedral structures become more favorable. In Fig. 9, the electrophilic fields determined by the Fukui functions calculations 0.0025 eV iso-value distribution for the structure: a) cubo-octahedral Au13 and b) decahedral Au13.

Acknowledgments

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References