One-step synthesis of 3D dendritic gold/polypyrrole nanocomposites via a self-assembly method

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Abstract

Novel spherical three-dimensional (3D) dendritic gold–polypyrrole nanocomposites were successfully prepared in the presence of an amphiphilic p-toluene sulfonic acid (TSA) as dopant and surfactant via a self-assembly process which is based on the oxidation of pyrrole (Py) and the reduction of the chloroauroate ions, yielding PPy and Au(0) simultaneously. It was found that the probability of obtaining dendritic Au@PPy/TSA nanostructures depended on the concentration of TSA and the rate of addition of the oxidant (HAuCl4). It was also proposed that the supramolecular micelles formed by Py and TSA play the role of a ‘soft template’ to produce the dendritic Au@PPy/TSA nanocomposites.

1. Introduction

Nanocomposites of conducting polymers and metal nanoparticles have received considerable attention due to the potential possibilities to create suitable materials for electrocatalysis, chemical sensors, and microelectronic devices [1–3]. Conducting polymers are well known for their excellent electronic properties with conductivity covering the whole range from insulator to metal while retaining the light-weight, mechanical properties and processing advantages of polymers [4]. Compared with nanocomposites constructed by insulating polymers, conducting polymers can serve as a novel electroactive relay among these metal nanoparticles in the nanocomposites matrix [5, 6]. For example, it has been reported that conducting polymer-supported platinum and rhodium catalysts show potential applications to novel catalysts because the conducting polymers might shuttle the electronic charge to catalyst centres [7, 8]. On the other hand, metal nanoparticles incorporated into conducting polymers are also known to enhance the conductivity of the polymers [9, 10]. Therefore, the fabrication of novel nanocomposites based upon metal nanoparticles and conducting polymers would provide various interesting characteristics and new features in nano-technological applications.

Among these reported metal clusters, gold nanoparticles as well as nanorods have attracted much attention owing to their unique optical, electrical and photocatalytic properties depending on their sizes and shapes [11–13]. Recently, a variety of methods have been explored to achieve gold nanoparticle–conducting polymer composites. Overall, three different approaches have been used to date. The first one is the electrochemical method, which is based on the oxidation of the individual monomer and further incorporation of the metallic species deposited from an aqueous solution containing a suitable cation [14, 15]. The second technique consists of in situ preparation of the gold nanoparticles in the polymer matrix, which might open up much wider possibilities for the gold-conducting polymer applications. By this method, Hatchett and Chattopadhyay have reported the chemical synthesis of a polyaniline/gold composite using tetrachloroauroate [16, 17]. Liu has also demonstrated that the polypyrrole-stabilized gold nanostructures can be formed on roughened gold substrates through a similar reaction based on gold ions and pyrrole [18, 19]. In particular,
Selvan and co-workers have shown that the reduction of AuCl\textsuperscript{−}\textsubscript{4} ions sequestered in micelles of a diblock copolymer results in the formation of gold nanoparticles capped into polypyrrole bulk matrix [20]. In addition, they have also observed the Au–PPy dendritic nanostructures employing vapour phase polymerization of pyrrole onto a solution-cast film of block copolymer ionomers [21]. The third technique involves the synthesis of Au nanoparticles, and coating the Au nanoparticles on PPy templates [22] or polymerization of Py using the Au nanoparticles as a template [23]. In the above-mentioned works, much emphasis was focused on the structure and properties of gold-conducting polymer composite; however, there are very few reports addressed to the fabrication of unusual dendritic gold nanostructures in a matrix of polypyrrole [21] by a simple liquid phase reaction.

Herein, a novel method is reported to synthesize 3D dendritic gold/polypyrrole (Au@PPy/TSA) nanocomposites via a self-assembly process in which pyrrole monomer is oxidized by chlorauric acid (HAuCl\textsubscript{4}) in the presence of TSA.

2. Experimental section

16.7 µl of distilled pyrrole monomer was mixed with 25 ml TSA aqueous solution (0.1 M) under stirring for 0.5 h to obtain a uniform emulsion. 5 ml of aqueous solution of HAuCl\textsubscript{4} (5 mM) was added to the emulsion. The mixture was allowed to react at 0–5 °C (in an ice bath) for 10 h under stirring. Then, the product was washed with de-ionized water, methanol, and ethyl ether several times, dried under vacuum for 24 h, and finally a dark powder of Au@PPy/TSA nanocomposites was obtained. The synthetic route of Au@PPy/TSA composites is illustrated as follows:

\[ \text{CH}_2\text{CH}_2\overset{\text{SO}_3\text{H}}{\longrightarrow} + 7/4 \text{HAuCl}_4 \rightarrow \text{Au@PPy/TSA} \]

The morphology of Au@PPy/TSA nanocomposites was proved by scanning electron microscope (SEM, XL30 ESEM FEG) and transmission electron microscope (TEM, JEOL2000). Ultraviolet–visible (UV–vis) absorption spectra of Au@PPy/TSA nanocomposites dispersed in aqueous solution were recorded on a UV-3100 spectrometer. FTIR (Perkin-Elmer system) and x-ray diffraction (MAC Science, Japan M-18AHF) measurements were performed to characterize the structure of Au@PPy/TSA nanocomposites.

3. Results and discussion

Figure 1 shows typical SEM images of such 3D dendritic Au@PPy/TSA nanocomposites. The Au@PPy/TSA nanocomposites show evident spherical morphology (yield >90%), as seen in figure 1(a), and these spherical Au@PPy/TSA nanocomposites are well dispersed in amorphous colloidal polypyrrole matrix. Electron micrographs of an individual spherical Au@PPy/TSA nanocomposite at higher magnification were shown in figure 1(b). Interestingly, on the surface of nanoscale spherical Au@PPy/TSA composites, many dendritic Au@PPy/TSA nanorods can be observed. In particular, the surface of spherical Au@PPy/TSA nanocomposites shows a bionic nano-nanoscale binary structure, similar to the natural lotus leaf, where every microscale papilla on the leaf surface is covered by nanoscale papillae (figure 1(c)) [24].

The 3D dendritic structure of spherical Au@PPy/TSA nanocomposites is more clearly seen in figures 2(a) and (b). The average diameter of the spherical Au@PPy/TSA nanocomposites was estimated to be ca 80–150 nm, and they were rather polydisperse. The detailed structure of the 3D dendritic Au@PPy/TSA nanocomposites is further revealed by high-resolution TEM images and an electron diffractogram. Figure 2(b) shows the sphere structure at higher magnification.
aggregates could be observed, as shown in figures 3(a)
the TSA concentration was high (1 M), the spherical dendritic
concentration on the morphology measured by TEM. When
Figure 3 shows a typical example of the effect of the TSA
morphology of the obtained Au@PPy/TSA nanocomposites.
However, irregular dendritic-like grain morphology was
observed when the TSA concentration further decreased to
0.01 M (figures 3(c) and (f)). Furthermore, in the absence
of TSA dopant, the Au agglomeration phenomenon occurred
(figures 3(g) and (h)). These results indicate that a change
in morphology from dendritic aggregates, to 3D dendrites,
to dendritic-like grains, and to agglomerations, took place
when the concentration of TSA changed from 1 to 0.1 M, to
0.01 M, and to 0 M, respectively. As described in previous
reports [25, 26], the self-assembled micelles play a ‘soft-
template’ role in the formation of the nanostructured PPy.
Here, TSA can act as dopant and surfactant at the same time due
to the ionized –SO3H group and the amphiphilicity of TSA.
In addition, pyrrole monomer in TSA aqueous solution easily
forms PPy/TSA micelles by hydrophobic interaction, while
addition of an oxidant (H[AuCl4]), the nucleation and growth
of Au, and polymerization of pyrrole take place simultaneously
at the water/micelle interface due to the hydrophilicity of
this AuCl4−. As the reaction continues, a 3D dendritic
Au@PPy/TSA nanocomposite with high aspect ratio is
obtained. It can be proposed that such Py/TSA supramolecular
micelles might serve as a ‘soft template’ and kinetically
control the growth rates of various faces of Au nanoparticles
by selectively adsorbing onto the crystallographic planes,
thus resulting in the dendritic morphology. The results
also suggest that such Py/TSA ‘soft-template’ supramolecular
micelles might be destroyed when the concentration of TSA
increases or decreases, and thus the morphology of the obtained
Au@PPy/TSA composite changes, which is in agreement with
our results (as shown in figures 2(a) and (d)).
In this synthesis, the reduction of pyrrole monomer
and the oxidation of the chloroaurate ions (AuCl4−) occur
simultaneously, yielding PPy and zero-valent Au. The
polymerization rate of the pyrrole monomer was affected by
the polymerization temperature, rate of addition of the oxidant,
concentration of the oxidant, and so forth. Here, only the rate
of addition of the oxidant is changed to further investigate
the influence of the polymerization rate on the morphology
of Au@PPy/TSA composites. Figure 4 represents the influence
of the polymerization rate on the resulting composite morphology.
Spherical granular Au@PPy/TSA nanocomposite was obtained when the addition was very
quick, ca 1 ml s−1 (figures 4(a) and (b)). On the other hand, 3D
dendritic Au@PPy/TSA nanocomposites were prepared when the
addition was slow, ca 0.1 ml s−1. However, the aspect ratio
of the obtained Au@PPy/TSA nanorod arms is low (ca 2–4)
(figures 4(c) and (d)). Only when the addition was very slow
(0.01 ml s−1) could the regular 3D dendritic Au@PPy/TSA
nanocomposites with the high aspect ratio of the fractal-
shaped nanorod arms (5–10) be obtained (figures 2(a) and (b)). This indicates that when the polymerization rate was quick,
too much PPy–TSA and Au was produced in a very short
time, and PPy–TSA and Au did not have enough time to
form the regular dendritic morphology via self-assembly; in
contrast, when the rate was slow, they easily formed the regular
3D dendritic Au@PPy/TSA nanocomposites through a self-
assembly process. The results confirm that the formation of
3D dendritic Au@PPy/TSA nanocomposites described in this
study is a slow self-assembly process.

The dark dendritic Au@PPy/TSA nanocomposites could be
readily dispersed into water to form a clear and dark
suspension, which was quite stable even for several more
Figure 3. Influence of the concentration of TSA on the morphology of Au@PPy/TSA nanocomposites: ((a), (b)) 1 M; ((c), (d)) 0.1 M; ((e), (f)) 0.01 M; ((g), (h)) 0 M (other reaction conditions: [Py] = 0.1 M, [HAuCl₄] = 1.67 M, reaction time = 10 h, temp. = 0–5°C).

The presence of Au in the Au@PPy/TSA nanocomposites was further confirmed by powder XRD measurements, as shown in figure 6. In the XRD curve of the Au@PPy/TSA nanocomposites, a broad peak centred at 2θ = 23.3° related to the amorphous PPy was observed, which is similar to the

Figure 4. Influence of polymerization rate on the morphology of Au@PPy/TSA nanocomposites synthesized by the self-assembly method: ((a), (b)) very quick and ((c), (d)) slow (other reaction conditions: [TSA] = 0.1 M [Py] = 0.1 M, [HAuCl₄] = 1.67 M, reaction time = 10 h, temp. = 0–5°C).

The molecular structure of Au@PPy/TSA nanocomposites was characterized by FTIR spectroscopy (data not shown here) and XRD (in figure 6). In FTIR results, the characteristic bands of polypyrrole, the pyrrole ring fundamental vibrations centred at 1541 and 1448 cm⁻¹, the C–H in-plane vibrations at 1303, 1161, and 1037 cm⁻¹, and N–H stretching vibration at 3432 cm⁻¹, can be observed well in the Au@PPy/TSA nanocomposites. These results indicate that the structure of the polypyrrole backbone in Au@PPy/TSA nanocomposites is similar to that of conventional PPy synthesized by the well established method [28].

The UV–vis absorption spectrum of the Au@PPy/TSA nanocomposites dispersed in aqueous medium is shown in figure 5. Two typical absorption peaks can be observed at ca 440 and 800 nm, which corresponds to the π–π* transition of the PPy chain and to a polaron transition similar to that of an ordinary PPy [27]. The absence of the typical surface plasmon resonance of Au(0) around 520 nm for Au/PPy nanocomposites reveals that Au is dispersed as ultra-small clusters of elementary gold within the Au/PPy nanocomposites. This type of Au is characterized by a weak and broad absorption maximum at about 440 nm. But it should be noted that the π–π* transition of the PPy chain at 400–500 nm overlaps this region. This is consistent with the idea that elementary Au is formed concurrently with PPy. A similar observation was also reported by Liu et al [19] and Selvan et al [20, 21].

The presence of Au in the Au@PPy/TSA nanocomposites was further confirmed by powder XRD measurements, as shown in figure 6. In the XRD curve of the Au@PPy/TSA nanocomposites, a broad peak centred at 2θ = 23.3° related to the amorphous PPy was observed, which is similar to the
The unit cell constant of Au can be obtained from (111), (200), (220) and (311) planes of Au. Furthermore, the method described in this study provides a nano-nanoscale binary structure similar to the natural lotus leaf. Moreover, four strong bands appeared with maximum intensity at 38.1°, 44.2°, 64.6° and 77.7° representing Bragg’s reflections from (111), (200), (220) and (311) planes of Au. The unit cell constant a of such an fcc lattice can be obtained as 4.086 Å, which means that the Au nanodendrites had grown well along all planes.

In conclusion, a simple and novel method has been introduced to prepare 3D dendritic Au@PPy/TSA nanocomposites using HAuCl₄ as an oxidant, and TSA as dopant and surfactant. A model of a supramolecular micelle ‘soft template’ based on pyrrole and TSA is proposed to interpret the formation mechanism of 3D dendritic Au@PPy/TSA nanocomposites. In particular, the synthesized Au@PPy/TSA nanocomposites show a bionic nano-nanoscale binary structure similar to the natural lotus leaf. Furthermore, the method described in this study provides a simple and inexpensive route to prepare inorganic–organic supermolecular nanocomposites.

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