Research Article
Preparation and Ordered Self-Assembly of Nano-Pd-Ga/PMMA by Ultrasonic

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Nano-Ga-Pd/poly methyl methacrylate (PMMA) composite materials were prepared with the palladium chloride solution containing metal gallium, MMA as monomer, and sodium dodecyl sulfate (SDS) as emulsifier without initiator or reducer. Pd, Ga, and Ga5Pd phase in PMMA matrix were identified by XRD. The characteristic absorption peak at 200 nm for nano-Ga/PMMA polymer solution, at 209 nm for nano-Pd/PMMA polymer solution were proved by UV-Vis; the binding energy changes of O1s, Ga2p3, Ga2d, and Pd3d were characterized by means of X-ray photoelectron spectroscopy. It is concluded that nano-Ga5Pd was produced based on segment electronics shifting from Gallium to Palladium, and coordination was formed on segment electronics from Gallium to oxygen of PMMA ester group. The anisotropism ordered assembly of PMMA around nano-Ga-Pd particles were illuminated by transmission electron microscopy; it is further interpreted that nano Ga-Pd particles had ordered-assembly induced effect.

1. Introduction
Nanocomposite materials have increasingly become a major part of new synthetic materials all over the world owing to their applications in mechanics, tribology, optics, magnetism, and electronics [1, 2]. Nanoalloy materials, which are different from the general alloy in both size and structure, have been one of the most promising and emerging research areas in advanced materials because of the unique properties [3, 4]. Lee and Shingu [5] in Tokyo University firstly prepared nanoalloy materials by ball milling. Subsequently, lots of new methods were used to prepare nanoalloy materials [6, 7], such as chemical reduction. However, there existed limitations in those methods: the reaction system was more complicated and the properties of the as-synthesized nanoalloy became bad because of initiator or metal reductant used in the preparation process. And in those methods, the as-synthesized nanosized alloy materials conglomerated easily, and so the nano-effect of nanomaterials weakened. To resolve these problems, the idea of the organic-based nanometal came into being and new nanocomposite alloys were reported to possess interesting properties [8, 9].

Ultrasonic radiation method was used widely to prepare nanomaterials because of its convenience, emulsification, dispersion, and cavitation. Works on nanometal particles synthesis in emulsion polymerization without initiator by means of ultrasonic radiation were reported [10], but literature on preparing Pd-Ga alloy nanocomposite particles by ultrasonic radiation cannot be found as far as our knowledge goes, let alone the ordered self-assembly of PMMA-based Pd-Ga nanocomposite. So-called self-assembly is the autonomous organization of components in system into patterns or structures without human intervention [11]. For polymer-based nanocomposites, the polymer chain would possess the ordered self-assembly phenomena due to nano-effect of nanometal particles. South Korean scholars [12] had reported that on nano-Fe2O3 particles into composite would induce polystyrene-block-polypropylene (PS-b-PI) as composite matrix to self-assembly; the results showed that size and content of nano-Fe2O3 into the nanocomposites had great impact on ordered self-assembly structure of PS-b-PI. Chinese scholars had studied that polyethylene (PE) crystallization would be induced by carbon nanotubes, of which the nanocomposites were composited [13]. XU group
palladium chloride (PdCl₂) was carried out as follow. A block of gallium (purity: 99.99%); high-purity palladium (II) chloride (PdCl₂) (Chen-zhou Xiangchen Hi-tech Industry Co.Ltd, China), and a little of sodium dodecyl sulfate (abbreviated as SDS, purity: 90.0%) were added into a beaker containing a proportional methyl methacrylate, then a proportional methyl methacrylate, which was supplied by Shanghai Chemical Reagent Company and washed with 5% aqueous sodium hydroxide and subsequently washed by distilled water to neutrality, was poured into the beaker. Then, N₂ was piped into the mixture for 15 min. The output power of the ultrasonic was adjusted to work efficiently before the ultrasonic generator (JY98-III ultrasonic generator, Ningbo Scientz Biotechnology Co., Ltd., China, frequency: 20 KHz, output power: 200~1200 W) was on. The reaction temperature was controlled between 40~45°C by circulating water. The system became emulsion and the color of the system became dusk after the ultrasound for several minutes. The ultrasonic generator was terminated when the emulsion polymerization of methyl methacrylate and reduction of the palladium ion (II) were disappeared in Pd-Ga/PMMA spectrum, which indicated the nanoalloy to prevent the nanoparticles from coagulating, so the Pd-Ga/PMMA nanoalloy composite particles with core-shell structure were prepared.

2. Experimental

Preparation of Pd-Ga/PMMA nanoalloy composite particles was carried out as follow. A block of gallium (purity: 99.99%); high-purity palladium (II) chloride (PdCl₂) (Chen-zhou Xiangchen Hi-tech Industry Co.Ltd, China), and a little of sodium dodecyl sulfate (abbreviated as SDS, purity: 90.0%) were added into a beaker containing a proportional distilled water, then a proportional methyl methacrylate, which was supplied by Shanghai Chemical Reagent Company and washed with 5% aqueous sodium hydroxide and subsequently washed by distilled water to neutrality, was poured into the beaker. Then, N₂ was piped into the mixture in the beaker to get rid of dissolved oxygen in the mixture for 15 min. The output power of the ultrasonic was adjusted to work efficiently before the ultrasonic generator (JY98-III ultrasonic generator, Ningbo Scientz Biotechnology Co., Ltd., China, frequency: 20 KHz, output power: 200~1200 W) was on. The reaction temperature was controlled between 40~45°C by circulating water. The system became emulsion and the color of the system became dusk after the ultrasonication for several minutes. The ultrasonic generator was terminated when the emulsion polymerization of methyl methacrylate and reduction of the palladium ion (II) were finished. The emulsion was refrigerated to demulsificate, then the precipitate was treated by filtering, washing, and drying in vacuum, and finally PMMA-based nano Pd-Ga composite particles were obtained.

The morphology of the nanoparticles was observed by means of high-resolution transmission electron microscope (TEM) (JEOL-2010, Japan Electronics Co., Ltd.) when a little of the powder samples was dispersed in acetone to form dispersion system, and the suspension was dropped to a copper grid. The crystalline phase of the powder was determined by using an X-ray diffractometer (XD-3, Beijing Pur-kinje General Instrument Co. Ltd, China) with DX-2000 (λCuKα = 0.15418 nm), using graphite-monochromatic Cu-Kα radiation (r = 0.154 nm) with an accelerating voltage of 40 KV, at scanning rate of 0.02°S⁻¹ in 2θ ranging from 10° to 85°. The UV spectrum of the samples were tested by TU-1901 dual-beam UV spectrophotometer (Beijing PUXI Co. Ltd in China). The chemical composition of Pd-Ga/PMMA nanocomposites was tested by Thermo ESCA-LAB 250 X-ray photoelectron spectroscopy (Thermo Corporation, American).

3. Results and Discussion

The infrared spectrum of the nanocomposites was collected after purification and pellet pressed together with potassium bromide, characteristic peak of C=O (1638 cm⁻¹) nearly disappeared in Pd-Ga/PMMA spectrum, which indicated the polymerization of MMA monomer occurred completely by ultrasonic radiation. Characteristic peak of C=O and C–H in the spectrum of PMMA was shifted to short wave region in that of the nanocomposites, which hinted that there existed interaction between nanoalloy particles with PMMA matrix to some degree.

The X-ray diffraction patterns of Pd-Ga/PMMA composite particles by ultrasonic irradiation were shown in Figure 1; the four strong characteristic peaks, shown as peaks 2, 5, 6, and 7 in Figure 1, of Pd-Ga/PMMA composite lied in 2θ = 40.31, 46.77, 68.31, and 82.23, which were corresponding to crystal face of [111], [200], [220], and [311] of Pd (PCPDFWIN#870639). The peaks at 2θ = 32.30, 41.37, and 44.48, respectively, belonged to [211], [213] and [310] of Ga₃Pd (PCPDFWIN#15-0577). But the peak of [114]
The peaks of Pd3d electrons were positioned at 335.43 eV, which were dropped down 2.07 eV than the standard energy spectrum peaks of Pd3d electrons that were positioned at 337.5 eV, that is, the energy spectrum peaks of Pd3d electrons within nanocomposites, moved to the low-binding energy region. This illustrated that chemical environment of palladium was changed due to getting some electronics from other atoms. For binding energy change of the gallium atom, on the one hand, in terms of interaction between gallium and palladium, the segment electron in 4d orbital of gallium flowed directly to 5s orbital of palladium in order to form a new alloy, namely, Ga$_5$Pd generated from Pd with Ga; on the other hand, coordination energy was based on segment electronics shifting from gallium to oxygen attached to ester group of PMMA into nanoalloy/PMMA composite. The curve of XPS spectrum of O1s pertain to the nano Pd-Ga/PMMA composite was fitted by means of Origin 6.1 shown in Figure 3(e); curve 1 peaked at 531.62 eV, in Figure 3(e) was originated from carboxyl (C=O), curve 2 peaked at 533.39 eV from ester group (C–O–C), and curve 3 from integration fitting peak of O1s in the two chemicals environment. Based on the standard dates of (532.62 eV) carboxyl (C=O) and (533.39 eV) ester group (C–O–C), the O1s binding energy peaks shift to lower region, which were reduced to 0.68 eV and 0.51 eV, respectively; it was probably due to carbonyl oxygen atom accepting electronics cloud to from gallium atom form nanoalloy of Pd-Ga into the nanocomposite. The XPS results proved that there existed interaction between carboxyl (C=O) oxygen and the nanoalloy.

The morphology of the nano Pd-Ga/PMMA composites was shown in Figure 4; it demonstrated that the nanocomposite was small in size about ~25 nm with nanometal particles as core and PMMA as shell. The interesting TEM images of the nanocomposites were formed when observed in high-resolution TEM; it hinted that there were ordered self-assembly of PMMA in addition to possessing in ordered distribution state around the nanoalloy in Figure 4; the ordered self-assembly of PMMA possibly had something to do with not only content of nanometal, but also distance and direction apart from surface of nanometal in distribution. The more the content of the nanometal is, the more the obvious is phenomenon of ordered self-assembly of PMMA, comparing Figure 4(a) based on metal content 0.15% with Figure 4(b) based on metal content 0.3%. For selectively of distance and direction apart from surface of nanometal in distribution, on the one hand, ordered direction of PMMA self-assembly was vertical near to the surface of nanometal particles; on the other hand, ordered direction of PMMA self-assembly was changed from vertical to slope even to horizontal direction apart from the surface of the nanometal particles, shown as the inset in Figure 4(b). The mechanism of the ordered self-assembly of PMMA was probably imagined as induced effect of nanometal Pd-Ga particles into nanocomposites due to interaction of nano Pd-Ga with PMMA illustrated above. Based on the interaction of palladium gallium nanocrystals as discontinuous phase with PMMA as continuous phase, why did nanometal contained gallium as one of components into polymer-based nanocomposites would occur the ordered self-assembly, the
Figure 3: XPS of nano-Ga-Pd/PMMA composites powder. (a) XPS survey spectrum of nanocomposites; (b) Ga2p; (c) Ga3d; (d) Pd3d; (e) fitting curve of O1s. (1) C=O; (2) C–O–C; (3) integrate fitting peak of O1s in the two chemicals environment.
possibly reason was suggested to come from stable atom-chain structure of gallium. Gallium itself possessed property to form stable atom-chain structure [15], and exhibit the most stable “Z” configuration. Gallium was mainly a component in the Ga₅Pd nanoalloy and was distributed possibly on the surface of the nanoalloy. On coordination of gallium atoms with oxygen of ester group attached to PMMA, oxygen atoms attached to PMMA were arranged along the direction of gallium atom-chain structure as a template; so, there were ordered oxygen atoms array and then ordered self-assembly of PMMA as polymer continuous phase into nanocomposites. The ordered self-assembly of PMMA as continuous phase into nano-Ag-Ga/PMMA composites possessed isotropic phenomenon in reported paper by XU group [14], which was closed ringlike around Ag-Ga particles as dispersing phase into nanocomposites; it was completely different to that of PMMA from the Pd-Ga nanocomposites. The possible reason was suggested that the nanosilver gallium alloy into polymer-based nanocomposites was presented to be isotropic based on hexagonal crystal, whereas the nano-palladium gallium alloy into polymer-based nanocomposites did not, it was found to be antitropic based on tetrahedron crystal.

4. Conclusions

Nano-Ga-Pd/poly methyl methacrylate (PMMA) composite materials were prepared with the palladium chloride solution containing metal gallium without initiator reducer, MMA as monomer, and sodium dodecyl sulfate (SDS) as emulsifier under ultrasonic generator, in which Pd²⁺ ion was reduced to nano-palladium particles, meanwhile produced Ga₅Pd with microdrop of liquid gallium, and the monomers were polymerized simultaneously by ultrasonic radiation. The structure of nanocomposite with nano metal was characterized by means of IR, XRD, UV-Vis, TEM, and XPS. Pd, and Ga₅Pd as dispersing phase in PMMA matrix was identified by XRD. The results of UV-Vis spectrum showed that there was a characteristic absorption peak at 200 nm for nano-Ga/PMMA polymer solution, at 209 nm for nano-Pd/PMMA polymer solution, and at 218 nm for nano Pd-Ga/PMMA; XPS proved that the O1s binding energy of C=O and C–O–C into nanocomposite was decreased compared to that of the standards, respectively, the meanwhile binding energy of Ga₂p₃, Ga₂d₃/₂, and Ga₂d₅/₂ into Pd-Ga nanoalloy of the nanocomposite were increased; however, Pd₃d binding energy was decreased; it is concluded that nano Ga₅Pd was formed on shifting of segment electronics of Gallium to Palladium, and coordination of gallium attached to nano Pd-Ga alloy and oxygen of ester group attached to PMMA was found on shifting of segment electronics of gallium to oxygen. So, there was interaction of nano metal particles with the polymer as matrix. Based on the interaction, TEM illuminated that ordered self-assembly of PMMA as continuous phase around nano-Ga-Pd particles into nanocomposites was presented and was found to be of anisotropism; it was further deduced that nano-Ga-Pd particles possessed induced effect on ordered self-assembly of polymer.

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References


