INTRODUCTION

Research on nanomaterials has demonstrated that nanocapsules can have special characteristic compared to the bulk materials, largely due to their ultrafine size and high surface to volume ratio (Gleiter, 1989; Uyeda, 1991; Siegel, 1993). It is recognized that nanocapsules, which area bridge between the bulk and the atomic state, have novel characteristics for electrical, magnetic, and chemical properties. Besides the interest in understanding the nature and mechanisms of such new phenomena, there is a technologically driven force due to the immediate applications of these systems, such as catalysts, battery materials, color imaging, drug delivery systems, pigments in paints, magnetic tapes, ferrofluids, magnetic refrigerants, giant magnetoresistance (Hainfeld et al., 2006; McMahon et al., 2008; Guidelli et al., 2013), etc.

Carbon-coated Ni, Cu and Sn nanocapsules were investigated by means of X-ray diffraction (XRD), transmission electron microscopy (TEM) and a four-point probe device. All of these nanocapsules were prepared by an arc-discharge method, in which the bulk metals were evaporated under methane (CH₄) atmosphere. Three pure metals (Ni, Cu, Sn) were typically diverse in formation of the carbon encapsulated nanoparticles and their different mechanisms were investigated. It was indicated that a thick carbon layers formed on the surface of Ni(C) nanocapsules, whereas a thin shell of carbon with 1~2 layers covered on Cu(C) nanocapsules, and the Sn(C) nanocapsules was, in fact, a longer multi-walled carbon nanotubes partially-filled with metal Sn. As one typical magnetic/dielectric nanocomposite particles, Ni(C) nanocapsules and its counterpart of oxide-coated Ni(O) nanocapsules were compared in the electrically conductive behaviors for further applications as the electromagnetic materials.

Key Words: Nanocapsules, Carbon, Metal, Core/shell, Electric conductivity
as nanowires and suspensions of magnetic particles. Lots of studies (Seraphin et al., 1993; Guerret-Piecourt et al., 1994; Saito et al., 1994; Seraphin, 1995) revealed that not all materials introduced into the arc plasma from the composite target can form the encapsulated structures, which have been summarized in four categories. First, elements that can be encapsulated in the form of their carbides. Second, elements that are not encapsulated but to form carbon cages. Third, elements that form molecular carbides, competing with the carbon supply for the graphitic cage formation and suppressing them. Fourth, the ferromagnetic metals that stimulate catalytically growth of single-walled tubes and strings of spherical beads under a modified conditions of preparation (Dravid et al., 1995; Jiao et al., 1996).

In the present work, several typical carbon-coated metal nanocapsules, i.e., Ni(C), Cu(C) and Sn(C), were prepared by arc discharge method in the mixture atmosphere of methane and argon gases. The morphologies and formation mechanisms of these nanocapsulation structures were characterized and discussed. The electric conductivities of comparable Ni(C) nanocapsules and its counterpart of Ni(O) nanocapsules were measured. The effects of quite different shells of carbon and nickel oxide on electric conductivity were also emphasized.

MATERIALS AND METHODS

Using the same synthesis method of arc-discharge as in our previous work (Dong et al., 1998a), pure Ni, Cu, Sn bulk metals to be evaporated served as the anode, while a carbon rod served as the cathode. After the chamber was evacuated, methane gas was introduced and acted as a reactant gas during evaporation of Ni, Cu and Sn bulks. The evaporating atmospheres for Ni(C), Cu(C) and Sn(C) nanocapsules are listed in Table 1. The distance between the two electrodes can be adjusted from outside of the chamber, so that the arc can be started and controlled for a continuous operation. X-ray diffraction (XRD) was performed to identify the phases existing in the as-prepared nanopowders. The particles size and morphology was determined by analyzing transmission electron microscopy (TEM) micrographs. The as-prepared nanopowders of Ni(C) and its counterpart of Ni(O) were compacted into round plates with 10 mm in diameter and about 0.3 to 0.5 mm in thickness. The electronic conductivities of both plates were measured under the stress pressures by a four-point probe device, then the relationship between conductivity of nanoparticles assembly and exoteric pressure was revealed.

RESULTS AND DISCUSSION

XRD patterns of the Ni(C), Cu(C) and Sn(C) nanocapsules are shown in Fig. 1 (Zhang et al., 2007; Liu et al., 2014). It is seen from Fig. 1 ‘a’ and ‘c’ that no diffraction peaks of carbon or carbides occurred in Ni(C) and Sn(C) nanopowders indicating that no carbon assemblies with larger grain sizes existed in powders, nonetheless, a distinct peak of graphite is found in Cu(C) nanopowders (Fig. 1 ‘b’) implying a bigger carbon phase may exist inside. The diffraction peak of graphite emerged at 2θ=26.7° corresponds to (002) planes of its crystal lattice. All lattice constants of metal (Ni, Cu, Sn) phases are the same with those of bulks. The diffraction peaks of Ni(C) nanopowders (Fig. 1 ‘a’) at 2θ=44.37°, 51.75°, and 76.35° are in accordance with the (111), (200) and (220) crystallographic planes of Ni (JCPDS No. 00-001-1258; Hull, 1921); the peaks at 2θ=43.48°, 50.44°, and 74.13° are of (111), (200) and (220) planes of Cu (JCPDS No. 00-001-124; Hanawalt et al., 1938); likewise, peaks at 2θ=30.78°, 32.19°, and 45.12° reflect the (200), (101) and (211) planes of Sn (JCPDS No. 01-086-2265; Lee & Raynor, 1954).

In order to observe the morphologies and microstructures of all kinds of nanoparticles, the TEM with low and high magnifications were used to get the images of nanocapsules of Ni(C) (Fig. 2A and D), Cu(C) (Fig. 2B and E) and Sn(C) (Fig. 2C and F). It is found that the Ni(C) nanocapsule is of an integrated carbon-coated particle with ~15 layers of graphite-like shell encapsulating the Ni core. The lattice planes of

Table 1. The preparation atmospheres for Ni(C), Cu(C), and Sn(C) nanocapsules

| Sample | CH4 (MPa) | Ar (MPa) |
|--------|-----------|----------|
| Ni(C)  | 0.0133    | -        |
| Cu(C)  | 0.0133    | -        |
| Sn(C)  | 0.01      | 0.02     |

Fig. 1. X-ray diffraction patterns of the carbon-coated nanocapsules: (a) Ni(C); (b) Cu(C); and (c) Sn(C).
graphite is quite visible and the contiguous fringes around the Ni nanocrystal are good evidence for complete encapsulation by carbon layers. These nanocapsules are about 25 to 40 nm in diameter and the carbon protective shells are about 5 to 6 nm in thickness, corresponding to the (002) planes of crystal graphite phase. As shown in Fig. 2A, Ni(C) nanocapsules are spherical in shape and do not show any well-developed facets. One significant morphological characteristic to be noted from the image of Fig. 2D, is that neither gaps nor intermediate phases are observed between the outer carbon layer and the Ni core, which imply that the carbon atoms were released from the interior Ni, i.e., a solid solution of Ni-C.

A dissimilar core/shell structure with respect to Ni(C) is found in Cu(C) nanocapsules as shown in Fig. 2B and E. A great amount of empty carbon cages exist in Cu(C) nanocapsules, indicating some of Cu cores have escaped from the carbon shells in formation of the nanocapsules which may be an evidence for Cu-C solution with less carbon content than Ni-C solution. Moreover, from HRTEM image of Fig. 2E, it is clearly noted that the carbon shell on Cu(C) nanocapsules is so thin to be just 2 to 3 graphite layers with around 1 to 2 nm in thickness. It is different with the case of Ni(C) nanocapsules, e.g., the empty carbon cages and thin carbon layers, all come from the diversity between Cu and Ni metals in solubility of the carbon atoms. As can be seen, Cu(C) nanocapsules are spherical in shape, with the mean diameter ranging in 20 to 50 nm. The lattice spacing of graphite shell is approximately close to that of the bulk graphite (002) planes, which is accordance with the previous results (Dong et al., 1998a, 1998b; Dong et al., 1999). No Cu carbide phase existed (detected by XRD profile) indicates the inner core is of metallic copper. The interface of core/shell is clearly distinguished and an inverse fast fourier transform image of the giant onion-like fullerenes is presented in the inset of Fig. 2B, where the lattice spacing corresponds to the graphite phase. Moreover, many lattice defects occur, such as collapses, unevenness and dislocations among the carbon shells.

Distinctive images of Sn(C) nanocapsules are presented in Fig. 2C and F. It is well shown, in fact, the Sn(C) nanocapsules are of Sn-filled carbon nanotubes (CNTs) with the average diameter of about 40 nm and 200 to 300 nm in length. The CNTs have multi walls of 5 to 7 graphene layers and are partially-occupied by Sn metal. The electron diffraction pattern of the as-synthesized Sn/CNTs is shown in the inset of Fig. 2F which exhibits a pattern of poly-crystalline rings.
assigning to the (200), (220), (301) planes of Sn. Such a structure of Sn-filled CNTs implies a weak combination between Sn and carbon atoms, in other word, the carbon is difficult to be dissolved in Sn substrate.

Gas-phase nucleation is the basis for a gas-to-solid conversion process. In this work, atoms of Ni, Cu, Sn were evaporated from the raw bulk with the assistance of high temperatures under arc plasma conditions. At the area of arc-discharge, the high temperature results in the decomposition of methane into hydrogen and carbon elements. Hydrogen atoms serve as a source of hydrogen plasma and carbon atoms act as reactant and condensation gas. Large thermal gradients were created between the hot arc and water-cooled walls of the chamber. The driving forces and nucleation energy were provided by supercooling and lead to a series of transitions. Generally, the atoms of metals and carbon can form a solid solution by random collisions, and subsequently continue until a supersaturated state is reached. Then, the nucleus forms and grows by the addition of monomer units. With the decline of temperature, the carbon is precipitated from solid solution in a supersaturated state and then form a coated layer on the solid, finally the core/shell structure is formed.

In this work, the metals of (Ni, Cu, Sn) are typical to form the diverse carbon-coated core/shell nanocapsules, and their formation mechanisms can be understood in the binary alloy phase diagrams of Ni-C, Cu-C and Sn-C, as shown in Fig. 3. It is shown that the solubility limit of C in Ni-C solid solutions reaches 0.8 wt%, in Cu-C solid solution become 0.007 wt%, nevertheless, the carbon absolutely can not be dissolved into Sn and no Sn-C solid solution existed. It is obvious that the thickness of carbon layers on nanocapsules would be controlled by the solubility of C in a solid solution. More carbon atoms dissolved, thicker the carbon shells. The thickness of carbon layer on Ni(C) nanocapsule is about 5 to 6 nm, typically reflecting a growth history of this nanocapsules (Saito et al., 1993), while only 1 to 2 nm thickness of carbon layer is formed on the surface of Cu(C) nanocapsules and empty carbon cages are also created. It can be deduced that less carbon content in Cu-C results in thin graphite shell and quite a hollow graphite ball. It is known from Cu-C phase diagram that no copper carbide can be formed (Seraphin et al., 1996), that is, the metallic Cu cores are able to migrate out of their original cages in a solidification process and leave a mass of carbon cages behind. Because the carbon is greatly higher in melting point than Sn (Sn, 305 K; C, 3,773 K), so molten Sn will stay in a short time during the rapid solidification, meanwhile the catalysis of Sn (Zou & Wang, 2011; Xu et al., 2013) can promote the free carbon phase isolated from each other, to grow carbon atoms along sp^2 sites and form CNTs with the existance of liquid Sn (Lu et al., 2010). During the process, CNTs can adsorb the liquid Sn by the capillary force. As the temperature fall down to the melting point of Sn, the liquid Sn inside tube wall will be shrunk and solidified, finally the structure of Sn-partially filled CNTs was shaped.

In order to study the effects of outer layers (shells) on electrical properties, the carbon-coated Ni(C) nanocapsules and nickle oxide-coated Ni(O) nanoparticles are chosen to measure their electrical resistivities. The resistivities of the compacted Ni(C) and Ni(O) nanopowders slices were detected using a four-point probe method. From Fig. 4, it is shown that both nanopowders samples exhibit a gradually decreased resistivity with increasing of stress pressure. The resistivities of Ni(C) nanocapsules are higher than those of Ni(O) nanoparticles, they are 4.7×10^{-3} \ \Omega \cdot cm and 5.5×10^{-5} \ \Omega \cdot cm at 30 MPa, respectively. The intrinsic resistivity of bulk metal Ni is 7.0×10^{-6} \ \Omega \cdot cm, which means the resistivity of Ni(C) nanocapsules slices is 1,000 times greater than Ni bulk and 100 times higher than Ni(O) nanoparticles. In our previous work, it had been proved that Ni(O) nanoparticles was coated by Ni_2O_3 shells with 2 to 3 nm in thickness (Dong

![Fig. 3. Binary alloy phase diagram of metals-carbon.](image)

![Fig. 4. Electrical resistivities of the compacted Ni(C) nanocapsules and Ni(O) nanoparticles, detected at different stress pressures.](image)
et al., 1998a). The dielectric characters of Ni(C) nanocapsules had also been confirmed and investigated. It can be found from the TEM images of Ni(C), a mass of amorphous carbon with defections surround the Ni cores and would be main factors to affect the electrical conductivities.

CONCLUSIONS

The carbon-coated Ni(C), Cu(C) and Sn(C) nanocapsules were prepared by an arc-discharge method under methane (CH₄) atmosphere. Ni(C) nanocapsule is of an integrated carbon-coated particle with ~15 layers of graphite-like shell encapsulating the Ni core. Cu(C) nanocapsules have just 2 to 3 graphite layers on its surfaces and a great deal of empty carbon cages. Sn(C) nanocapsules are of Sn-filled CNTs with the average diameter of about 40 nm and 200 to 300 nm in length. The CNTs have multi walls of 5 to 7 graphene layers. The electrical resistivity of compacted Ni(C) nanocapsules is about 1,000 times and 100 times higher than Ni bulk and compacted Ni(O) nanoparticles, respectively. The defections in surrounded carbon shells would be main factors to affect the electrical conductivities of Ni(C) nanocapsules.

CONFLICT OF INTEREST

No potential conflict of interest relevant to this article was reported.

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