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Flexible Ribbon-shaped Coaxial Electrical Conductive Nanocables Array Endowed with Magnetism and Photoluminescence

Qianli Ma, Jinxian Wang, Xiangting Dong,* Wensheng Yu and Guixia Liu

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A new type of flexible [FeOx/polyaniline (PANI)/PMMA][@Tb(BA)3 phen/PMMA] ribbon-shaped coaxial conductive nanocables array possessing conductivity, magnetism and photoluminescence has been successfully fabricated by electrospinning technology using a specially designed coaxial spinneret. Every single ribbon-shaped coaxial nanocable in the array is assembled with a FeOx/PANI/PMMA electric conductive-magnetic bifunctional core with tunable width and a Tb(BA)3 phen/PMMA insulating photoluminescent sheath. All the ribbon-shaped coaxial nanocables are insulated with each other and arranged in array alignment. The conductivity of the core of each ribbon-shaped coaxial nanocable reaches up to the order of $10^4$ S·cm$^{-1}$, and the electric conductance of each single ribbon-shaped coaxial conductive nanocable can be calculated by formula and available experimental data. The ribbon-shaped coaxial nanocables array is simultaneously endowed with magnetic and photoluminescent characteristics for improving controllability and visibility. The electrical conductivity, magnetic and photoluminescent properties of ribbon-shaped coaxial nanocables array can be tunable by adjusting the contents of PANI and FeOx nanoparticles and rare earth complexes, or by varying the width of the cores of the every ribbon-shaped coaxial nanocable in the array. This nanostructure is ideally suited to be applied in high density electrical connection in narrow space, such as in mobile phones, subminiature integrated circuits, microchips and nano-machines.

Introduction

Ribbon-shaped one-dimensional (1D) materials have attracted increasing interest from scientists owing to their anisotropy, large width-thickness ratio, and unique optical, electrical, and magnetic properties. More and more manufacturers use ribbon-shaped cables as conductive lines because the ribbon-shaped cables have better anti-entangling performance and smaller winding volume than the traditional cables in round section (See Figure 1a and b). Multi-wire planar cable is a cable with many conducting wires running parallel to each other on the same flat plane. Multi-wire planar cables are usually seen for internal peripherals in computers, such as hard drives, CD drives and floppy drives (See Figure 1c and d).

With the rapid development of nanotechnology in recent years, fabrication and application of electroconductive nanowires and nanocables are becoming more and more crucial. Up to now, some types of nanocables have been synthesized. The researchers have adopted carbon fibers or tubes, metal materials, conductive polymers and other conductive materials as the inner conductor of nanocables.[1-6] The sheath must have good insulation property and, sometimes, have other functions. Many methods are used for the fabrication of nanocables, including hydrothermal, template, deposition and electrospinning method, etc. Guo et al. synthesized well-defined silica-coated carbon nanotube coaxial nanocable via the combination of sonication and sol-gel techniques.[7] Zhou et al. investigated the properties of BN nanotubes filled with close-packed Cu nanowires by gradient-corrected density functional theory computations.[8] Song et al. fabricated core-sheath silver nanowire/polyvinylpyrrolidone nanocables via an efficient single-spinneret electrospinning method.[9] Jang et al. fabricated polymeric/poly(methyl methacrylate) coaxial nanocable through the sequential polymerization of methyl methacrylate and pyrrole monomers inside the channels of mesoporous SBA-15 silica, followed by the removal of the silica template.[10] Based on the characterizations of the products described in references, it is found that the most of the reported conductive nanocables have some deficiencies. Firstly, some kinds of nanocables are quite brittle, resulting in the poor deformation, bending and stretch ability; secondly, the lengths of most types of the products are very small (even less than 1 cm) due to the limitation of preparation methods; thirdly, in order to obtain the nanocables with core-sheath structure, two steps are usually needed by most of the preparation methods, leading to the long production period and high cost. Fourthly, the nanocables are hard to install to the instruments, because it is hard to observe them due to their too small diameters. All of these deficiencies restrict the practical applications of conductive nanocables.

Electrospinning technique has been widely adopted to
effectively fabricate 1D nanomaterials, and has been applied in many areas such as filtration,[11] optical and chemical sensors,[12] biological scaffolds,[13] electrode materials,[14] luminescent materials,[15] photocatalysts.[16,7] The electrospun products have some unique advantages, such as long length (up to several meters), good flexibility (owing to the polymer template) and low-power consumption. The order of the electrospun products can also be arranged by controlling the collector.[18-20] In recent years, our research group has made some progress in the electrospun flexible 1D magnetic-photoluminescent bifunctional nanomaterials.[21-23] The obtained 1D magnetic-photoluminescent bifunctional nanomaterials simultaneously possess controllability (by magnetic field) and visibility (by emitting visible light).

Inspired by the unique properties of the flexible 1D magnetic-photoluminescent bifunctional nanomaterials, the controllability, visibility, flexibility and length of the nanocables can be improved once endowing the nanocables magnetic and photoluminescent properties and fabricating via electrospinning technique. Thus, in this work, novel ribbon-shaped flexible coaxial conductive nanocables with magnetic and photoluminescent properties were fabricated via electrospinning technique, and a new kind of coaxial spinneret was designed and manufactured to fabricate the ribbon-shaped coaxial nanocables. The tunable core of the ribbon-shaped coaxial nanocables is composed of conductive polyaniline (PANI), magnetic Fe₃O₄ nanoparticles (NPs) and PMMA template; the sheath consists of photoluminescent Tb(BA)₃phen complexes and PMMA template. In order to enhance the conductivity of PANI in the core, the aniline (ANI) was polymerized in the mixture solution of PMMA, DMF and CH₂Cl₂. Furthermore, the ribbon-shaped coaxial conductive nanocables were arranged in array alignment by using a rolling aluminum rotary drum as the collector. The ribbon-shaped coaxial nanocables, as well as their arrays, were firstly fabricated in this study. Furthermore, it is the first time that the electric conductive nanocables were endowed with magnetic and photoluminescent properties. The structure, conductivity, fluorescence, and magnetism of the ribbon-shaped coaxial conductive nanocables arrays were studied, and some new results were obtained. The obtained ribbon-shaped coaxial conductive nanocables arrays take over many deficiencies of the existing conductive nanocables as mentioned above. The synthesized multifunctional ribbon-shaped coaxial conductive nanocables arrays will have important potential application perspective in high-density interconnection. Furthermore, the design concept and preparation method of the ribbon-shaped coaxial nanocables arrays are of universal significance for the fabrication of other types of ribbon-shaped coaxial nanocables arrays.

**Experimental Sections**

**Chemicals**

Methylmethacrylate (MMA), benzoylperoxide (BPO), Tb₃O₄, benzoic acid (BA), 1,10-phenanthroline (phen), FeCl₃, 6H₂O, FeSO₄·7H₂O, NH₄NO₃, polyethylene glycol (PEG, Mr=20000), ammonia, oleic acid (OA), aniline (ANI), (IS)-(+)-camphor-10-sulfonic acid (CSA), ammonium persulfate (APS), anhydrous ethanol, CHCl₃, N,N-dimethylformamide (DMF), and deionized water were used. All the reagents were of analytical grade. The purity of Tb₃O₄ was 99.99 %. The deionized water was homemade.

**Preparation of OA modified Fe₃O₄ NPs**

Fe₃O₄ NPs with the saturation magnetization of 62.02 emu g⁻¹ were obtained by using a facile coprecipitation synthetic method.[25] To improve the monodispersity, stability, and solubility of Fe₃O₄ NPs in the spinning solution, the as-prepared Fe₃O₄ NPs were coated with OA as below: 1.5 g of the as-prepared Fe₃O₄ NPs were ultrasonically dispersed in 50 mL of deionized water for 20 min. The suspension was heated to 80 °C under an argon atmosphere with vigorous mechanical stirring for 30 min and then 0.5 mL of OA was slowly added. The reaction was stopped after heating and stirring the mixture for 40 min. The precipitates were collected from the solution by magnetic separation, washed with ethyl alcohol for three times, and then dried in an electric vacuum oven at 60 °C for 6 h. Then, all of the products were used to prepare the core spinning solution.

**Synthesis of Tb(BA)₃phen complexes**

Tb(BA)₃phen powders were synthesized according to the traditional method as described in the literature.[26]

**Preparation of PMMA**

PMMA used in this study was prepared by oxidative polymerization of MMA. The detailed synthetic process was provided in the reference.[23]

**Preparation of spinning solutions for fabricating magnetic-photoluminescent ribbon-shaped coaxial conductive nanocables array**

Two different kinds of spinning solutions were prepared to fabricate the coaxial nanocables array. The spinning solution for the core of coaxial nanocables array was composed of CSA doped PANI, OA modified Fe₃O₄ NPs, PMMA, CHCl₃, and DMF (core spinning solution). In the preparation of core spinning solutions, 0.5000 g of PMMA was dissolved into the mixed solution of 9.3750 g of CHCl₃ and 0.6250 g of DMF under magnetic stirring for 48 h. The mixture was then cooled down to 0 °C in an ice-bath. In order to find the optimum adding dosage of ANI, various amounts of ANI and CSA were added into the above mixture and kept stirring for 2 h, followed by introducing of APS and stirring for 30 min. The final mixture was allowed to react at 0 °C for 24 h and then the as-prepared OA modified Fe₃O₄ NPs were added into the mixture under mechanical stirring for 2 h at room temperature. Thus, the core spinning solution was prepared. For performing the electrical conductivity tests, 2 mL of the core spinning solution was dripped onto a piece of glass and the solvent was allowed to completely volatilize at room temperature for 48 h. The dosages of ANI, CSA, APS and the corresponding conductivities were shown in Table 1. One can see that the conductivity reaches up to the order of 10² S cm⁻¹, and it increases not much with adding more ANI than 0.1500 g. Thus the core spinning solution containing 0.1500g ANI was adopted for the following study.

**Table 1** Dosages of ANI, CSA, APS and corresponding conductivities

<table>
<thead>
<tr>
<th>ANI (g)</th>
<th>CSA (g)</th>
<th>APS (g)</th>
<th>Conductivity (S cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.0500</td>
<td>0.0625</td>
<td>0.1225</td>
<td>1.974×10⁻⁹</td>
</tr>
<tr>
<td>0.1000</td>
<td>0.1249</td>
<td>0.2450</td>
<td>3.931×10⁻³</td>
</tr>
<tr>
<td>0.1500</td>
<td>0.1873</td>
<td>0.3676</td>
<td>1.092×10⁻²</td>
</tr>
<tr>
<td>0.2000</td>
<td>0.2497</td>
<td>0.4900</td>
<td>1.475×10⁻²</td>
</tr>
<tr>
<td>0.2500</td>
<td>0.3125</td>
<td>0.6125</td>
<td>1.874×10⁻²</td>
</tr>
</tbody>
</table>

Another spinning solution for the sheath of the coaxial nanocables array consisted of Tb(BA)₃phen, PMMA, CHCl₃, and DMF (sheath spinning solution). It has been known through our previous work that the optimum ratio of Tb(BA)₃phen to PMMA is 1:10. [27] Thus for the preparation of the sheath spinning solutions, 0.5000 g of PMMA and 0.0500 g of Tb(BA)₃phen were added into the mixed solution of 9.3750 g of CHCl₃ and 0.6250 g of DMF.
Coaxial electrospinning technology has recently matured for fabricating coaxial nanofibers and hollow nanofibers. Nearly all types of spinnerets for coaxial electrospinning are composed of two stainless-steel needles of different diameters using one inside the other. We found the traditional coaxial spinnerets have some disadvantages when we attempted to use them to fabricate coaxial nanoribbons in our previous work, and a modified coaxial spinneret was designed and manufactured to fabricate coaxial nanoribbons. The assembly diagram of the modified coaxial spinneret is shown in Figure S1 in Supporting Information. As illustrated in Figure S1, two kinds of spinning solution had formed a coaxial structure before they flowed to the tip of the plastic nozzle, and the Taylor cone remained in this coaxial structure. This illustrates how the structure of the final products reflected the structure of the spinning solutions approaching the tip of the spinneret and the Taylor cone. We have successfully fabricated coaxial nanoribbons via electrospinning using the modified coaxial spinneret.

Another advantage of the modified coaxial spinneret was that the width of the core of the fabricated ribbon-shaped coaxial nanocables could be tunable easily only by changing the diameter of the inner stainless-steel needle without readjusting other spinning parameters. In this work, the ribbon-shaped coaxial nanocables with three different widths of their cores were fabricated using inner stainless-steel needles with three different diameters, including 7 # (inner/outer diameter: 0.41/0.71 mm), 12 # (0.90/1.26 mm) and 16 # (1.25/1.61 mm). Unless otherwise indicated, the products were fabricate using 16 # inner stainless-steel needle in the following text. To our knowledge, this is the first report that the width of the core of an electrospun 1D coaxial nanomaterial can be tuned just by changing the inner stainless-steel needle.

The equipment for the electrospinning process is presented in Figure 2. The core spinning solution was injected into the inner plastic syringe while the sheath spinning solution was loaded into the outer one. In order to obtain nanocables array, an aluminum rotary drum was used as a nanocables collector, which was put about 18 cm away from the tip of the plastic nozzle. The aluminum rotary drum was 20 cm in length, 7 cm in diameter, fixed in horizontal, and the rotation speed was 1500 r·min⁻¹. A positive direct current (DC) voltage of 6 kV was applied between the spinneret and the collector to generate stable, continuous PMMA-based nanocables array at 20-22 °C and the relative humidity of 20-30 %.

Electrospinning equipment for fabricating magnetic-photoluminescent ribbon-shaped coaxial conductive nanocables array

In the electrospinning process, the two spinning solutions at the tip of the plastic nozzle have formed a coaxial structure, and they are then stretched by electric field force to form a coaxial Taylor cone and a coaxial jet. The coaxial jet is stretched to ribbon shape by electrostatic repulsion and solidified with the volatilization of solvents, thus ribbon-shaped coaxial nanocable is obtained. A more detailed formation mechanism of electrospun ribbon-shaped coaxial nanocable is proposed in Supporting Information. Furthermore, because the coaxial nanocable swings in the air due to the instability of electrosprowing process, an array with a certain width can be collected on the surface of the aluminum rotary drum.

Characterization

The phase compositions were identified by an X-ray powder diffractometer (Bruker, D8 FOCUS) with CuKα radiation. The operation voltage and current were kept at 40 kV and 20 mA, respectively. The morphologies and internal structures were observed by a field-emission scanning electron microscope (FESEM, XL-30) and a biological microscope (CVM500E), respectively. The elemental analysis was performed by an energy-dispersive spectrometer (Oxford Instruments) attached to the FESEM. The conductivity properties were measured by a Hall effect measurement system (ECOPIA HMS-3000) using Van der Pauw method. The fluorescent properties were investigated by Hitachi fluorescence spectrophotometer F-7000. The UV-vis absorption spectra were tested by a UV-Vis spectrophotometer (SHIMADZU UV mini 1240). Then, the magnetic performances were measured by a vibrating sample magnetometer (VSM, MPMS SQUID XL).

Results and Discussion

Phase analyses

The XRD patterns of the as-prepared Fe₃O₄ NPs and [Fe₃O₄/PANI/PMMA]@[Tb(BA)₃phen/PMMA] ribbon-shaped coaxial conductive nanocables array were characterized by means of X-ray diffractometry (XRD) analysis, as shown in Figure 3. The XRD patterns of the as-prepared Fe₃O₄ NPs conform to the cubic structure of Fe₃O₄ (PDF 74-0748), and no characteristic peaks are observed for other impurities such as Fe₂O₃ and Fe(OH). The XRD analysis results of the nanocables array demonstrate that the nanocables array contain Fe₃O₄ NPs, and the broad diffraction peak at 15 ° can be attributed to the amorphous PMMA, PANI and Tb(BA)₃phen.

Morphology and internal structure

Fig. 3 XRD patterns of the Fe₃O₄ NPs (a) and [Fe₃O₄/PANI/PMMA]@[Tb(BA)₃phen/PMMA] ribbon-shaped coaxial conductive nanocables array (b)
The morphological observation of the as-prepared Fe$_3$O$_4$ NPs was performed by means of SEM, as presented in Figure 4a. The size distribution of the spherical Fe$_3$O$_4$ NPs is almost uniform, and the particle size of the NPs is 9.06±0.79 nm (Figure 4b). The macro-appearance of the [Fe$_3$O$_4$/PANI/PMMA]$[@$Tb(BA)phen/PMMA] ribbon-shaped coaxial conductive nanocables array is shown in Figure 1e. The ribbon-shaped coaxial conductive nanocables array is flexible, and its length is greater than 10 cm. The SEM image of the [Fe$_3$O$_4$/PANI/PMMA]$[@$Tb(BA)phen/PMMA] ribbon-shaped coaxial conductive nanocables array was shown in Figure 4c. The width of every single ribbon-shaped coaxial conductive nanocable is ca. 10 µm and the thickness is about 879 nm. All the ribbon-shaped coaxial conductive nanocables are arranged in the same direction. Depending on the transmission light of the BM, the inner structures of the ribbon-shaped coaxial conductive nanocables can be observed. As revealed in Figure 4d, a clear coaxial structure can be seen in the ribbon-shaped coaxial conductive nanocables. The cores of them contain large quantities of dark-colored PANI and Fe$_3$O$_4$ NPs, and the sheaths are transparent.

In order to further demonstrate the coaxial structure of the ribbon-shaped coaxial conductive nanocable array, EDS line-scan analysis was performed, in which S, Fe and Tb elements represent PANI, Fe$_3$O$_4$ and Tb(BA),phen, respectively, as shown in Figure 5. Elemental S and Fe only exist in the middle domain of the ribbon-shaped coaxial conductive nanocable. The amount of elemental Tb in the middle domain of the ribbon-shaped coaxial conductive nanocable is lower than that in both sides of the ribbon-shaped coaxial conductive nanocable because Tb only exists in the top and bottom surfaces of the middle domain of the ribbon-shaped coaxial conductive nanocable. It is further found that only elemental Tb without elemental S and Fe is dispersed in both sides of the ribbon-shaped coaxial conductive nanocable. These results are consistent with the core-sheath structure.

The electric conductance of the core of each single ribbon-shaped coaxial conductive nanocable was measured by the formula $G=\kappa(A/l)$, in which $G$, $\kappa$, $A$ and $l$ respectively represent electric conductance, electric conductivity, cross-sectional area and length. The value of $\kappa$ can be obtained by testing the dried core spinning solution, as indicated in Experimental Section. The value of $A$ can be calculated by the formula $A=T_sW_s^2/W_w$, in which $T_s$, $W_s$ and $W_w$ respectively represent the thickness of the single ribbon-shaped coaxial nanocable, the width of the single ribbon-shaped coaxial nanocable and width of the core of the ribbon-shaped coaxial conductive nanocable in array.

The BM images of the ribbon-shaped coaxial conductive nanocables respectively fabricated using 7#, 12# and 16# inner stainless-steel needles are revealed in Figure 6. It can be seen that the width of the core is increased with the increase of the diameter of the inner stainless-steel needle, demonstrating that the width of the core of the fabricated ribbon-shaped coaxial nanocables can be tunable easily only by changing inner stainless-steel needles without readjusting other spinning parameters.

From the SEM and BM observations, and the EDS line-scan analysis, we have clearly proven that the [Fe$_3$O$_4$/PANI/PMMA]$[@$Tb(BA)phen/PMMA] ribbon-shaped coaxial conductive nanocables array with a new and interesting nanostructure has been successfully fabricated.

**Electrical conductivity analyses**

Figure S4 shows the schematic diagram of electrical property of ribbon-shaped coaxial conductive nanocables array. Generally, a conductor and an insulator respectively refer to the materials with the conductivity range of $>10^3$ S cm$^{-1}$ and $<10^2$ S cm$^{-1}$, and a semiconductor is a material whose electrical conductivity is intermediate between that of a conductor and an insulator. As described in the previous section, the conductivity of the cores we used in the [Fe$_3$O$_4$/PANI/PMMA]$[@$Tb(BA)phen/PMMA] ribbon-shaped coaxial conductive nanocables in the array is $1.092\times10^2$ S cm$^{-1}$. In order to investigate the insulativity of each ribbon-shaped coaxial conductive nanocable with each other in the array, the surface conductivities of the arrays, including fabricated using 7#, 12# and 16# inner stainless-steel needles, were also measured by Van der Pauw method. The results (Table 2) reveal that the surface conductivity of all the arrays are lower than the order of $10^{-9}$ S cm$^{-1}$, indicating that every ribbon-shaped coaxial conductive nanocable in array is well insulated with each other.

**Table 2** The surface conductivities of ribbon-shaped coaxial conductive nanocables arrays fabricated using 7# (a), 12# (b) and 16# (c) inner stainless-steel needles

<table>
<thead>
<tr>
<th>Sample</th>
<th>Conductivity (S cm$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>$4.151\times10^{-11}$</td>
</tr>
<tr>
<td>b</td>
<td>$6.274\times10^{-11}$</td>
</tr>
<tr>
<td>c</td>
<td>$6.581\times10^{-11}$</td>
</tr>
</tbody>
</table>

The electric conductance of the core of each single ribbon-shaped coaxial conductive nanocable can be calculated by the formula $G=\kappa(A/l)$, in which $G$, $\kappa$, $A$ and $l$ respectively represent electric conductance, electric conductivity, cross-sectional area and length. The value of $\kappa$ can be obtained by testing the dried core spinning solution, as indicated in Experimental Section. The value of $A$ can be calculated by the formula $A=T_sW_s^2/W_w$, in which $T_s$, $W_s$ and $W_w$ respectively represent the thickness of the single ribbon-shaped coaxial nanocable, the width of the single ribbon-shaped coaxial nanocable and width of the core of the
ribbon-shaped coaxial nanocable (The deriving process of this formula is provided in Supporting Information), and these parameters can be obtained through SEM and BM observation. After determination of these parameters, the value of electric conductance of each single ribbon-shaped coaxial conductive nanocable with a given length can be obtained by calculation. Furthermore, the κ can be tuned by adding different amount of ANI, CSA and APS into the core spinning solution, as well as the A can be tuned through changing the width of the core by using different inner stainless-steel needles, so that the electric conductance of each single ribbon-shaped coaxial conductive nanocable with a given length is tunable.

Photoluminescence properties

The sheath of a single coaxial conductive nanocable not only can keep insulting from other nanocables, but also provides photoluminescence property in order to improve the visibility of the ribbon-shaped coaxial conductive nanocables array. As seen in the Figure 7, the ribbon-shaped coaxial conductive nanocables array can emit green light under the excitation of 333-nm ultraviolet light.

![Digital photos of [FeO/PANI/PMMA][Tb(phen)/PMMA] ribbon-shaped coaxial conductive nanocables array taken in natural light (a) and under the excitation of 333-nm ultraviolet light in the darkness (b)](image)

It has been proven in our previous work that the optimum exciting ultraviolet light for Tb(phen)/PMMA composite materials is at 333 nm, thus the 333-nm ultraviolet light was used to excite the [FeO/PANI/PMMA][Tb(phen)/PMMA] ribbon-shaped coaxial conductive nanocables arrays in this work. The left figure in Figure 8 shows the photoluminescence emission spectra of the ribbon-shaped coaxial conductive nanocables arrays fabricated using different inner stainless-steel needles. Characteristic emission peaks of Tb$^{3+}$ are observed in these products under the excitation of 333-nm ultraviolet light and ascribed to the energy levels transitions of $^5D_4\rightarrow^7F_5$ (495 nm), $^5D_4\rightarrow^7F_3$ (544 nm), $^5D_4\rightarrow^7F_4$ (588 nm), and $^5D_4\rightarrow^7F_2$ (624 nm), and the $^5D_4\rightarrow^7F_1$ hypersensitive transition at 544 nm is the predominant emission peak. One can see that with the increase of the width of the core, the photoluminescence intensity is decreased. This phenomenon can be explained by two reasons: (1) Because the sheaths of the ribbon-shaped coaxial conductive nanocables are transparent, the exciting and emitting light are absorbed by the dark-colored PANI and FeO NPs in the cores. From the UV-vis absorption spectra of PANI and FeO NPs illustrated in the right figure in Figure 8, it can be seen that both the PANI and FeO NPs can absorb visible light (400-700 nm) and much more easily absorb the ultraviolet light (< 400 nm). Thus, the exciting light and emitting light are absorbed by the FeO/PANI/PMMA core and the intensities are decreased. Furthermore, the light absorption would become stronger with increasing the width of the core. (2) As seen in the Figure 6, the ribbon-shaped nanocables fabricated using different inner stainless-steel needles almost have the same width, so that the content of the photoluminescent Tb(phen) complex in single ribbon-shaped coaxial conductive nanocable almost have the same width, so that the content of the photoluminescent Tb(phen) complex in single ribbon-shaped coaxial conductive nanocables with different width of cores. The exciting light and emitting light which are not affected by the cores contribute most to the photoluminescence intensity ("strong" section), whereas the exciting light and emitting light which are absorbed by the cores contribute only a little ("weak" section). The overall result is that the photoluminescence intensity is reduced with the reduction of "strong" section.

![Photoluminescence emission spectra (excited by 333-nm ultraviolet light) of the ribbon-shaped coaxial conductive nanocables arrays fabricated using 7 # (a), 12 # (b) and 16 # (c) inner stainless-steel needles (left figure) and UV-vis absorption spectra of PANI/PMMA and FeO NPs (right figure)](image)

Magnetic properties

The typical hysteresis loops for ribbon-shaped coaxial conductive nanocables arrays fabricated using different inner stainless-steel needles are shown in Figure 10. It is well known that the saturation magnetization of a magnetic composite material depends on the mass percentage of the magnetic substance in the magnetic composite material. The saturation magnetization of the ribbon-shaped coaxial conductive nanocables array is increased with the increase of width of the cores of the ribbon-shaped coaxial conductive nanocables in the array, indicating that the magnetic performance of the nanocables array can be tuned by adding various amount of FeO NPs. The magnetic properties
of the ribbon-shaped coaxial conductive nanocables arrays indicate that the ribbon-shaped coaxial conductive nanocables arrays can be easily manipulated by controlling the applied magnetic field.

In summary, novel flexible \([\text{Fe}_{3}\text{O}_{4}/\text{PANI/PMMA}@\text{[Tb(BA):phen]/PMMA}}\) ribbon-shaped coaxial conductive nanocables arrays with electric conductivity, magnetism and photoluminescence have been successfully fabricated by electrospinning using specially designed coaxial spinneret. The arrays are composed of ribbon-shaped coaxial conductive nanocables arranged in the same direction. Each ribbon-shaped coaxial conductive nanocable in the array consists of a \(\text{Fe}_{3}\text{O}_{4}/\text{PANI/PMMA}\) conductive-magnetic bifunctional core and a \(\text{Tb(BA):phen}/\text{PMMA}\) insulating photoluminescent sheath, so that each ribbon-shaped coaxial conductive nanocable is insulated with each other in the array. The width of every single ribbon-shaped coaxial conductive nanocable is ca. 10 µm and the thickness is ca. 879 nm. The width of the core of a ribbon-shaped coaxial conductive nanocable can be easily tuned just by changing the inner stainless-steel needle in the coaxial spinneret. The electric conductance of each single ribbon-shaped coaxial conductive nanocable can be calculated by given formula and available experimental data. The ribbon-shaped coaxial conductive nanocables arrays can emit green light under ultraviolet light and have response in the magnetic field. The conductivity, magnetism and photoluminescence of the ribbon-shaped coaxial conductive nanocables arrays can be tunable by adjusting the contents of PANI and \(\text{Fe}_{3}\text{O}_{4}\) NPs and rare earth complexes, or by varying the width of the cores of the every ribbon-shaped coaxial nanocable in the array. The ribbon-shaped coaxial conductive nanocables arrays resolve many deficiencies of the existing nanocables, such as high brittleness, short length, poor visibility and controllability, to accord with practical applications.

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**Notes and references**

Key Laboratory of Applied Chemistry and Nanotechnology at Universities of Jilin Province, Changchun University of Science and Technology, Changchun 130022. Fax: 86 0431 85383815; Tel: 86 0431 85582574; E-mail: dongxiangting888@163.com


Graphical Abstract

A new type of flexible ribbon-shaped coaxial conductive nanocables array possessing conductivity, magnetism and photoluminescence can be achieved by one-pot electrospinning using a specially designed coaxial spinneret. All the ribbon-shaped coaxial nanocables are insulated with each other and arranged in array alignment. The ribbon-shaped coaxial nanocables array is simultaneously endowed with magnetic and photoluminescent characteristics for improving controllability and visibility.