Wet chemical synthesis and magnetic properties of single crystal Co nanochains with surface amorphous passivation Co layers

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Abstract: In this study, for the first time, high-yield chain-like one-dimensional (1D) Co nanostructures without any impurity have been produced by means of a solution dispersion approach under permanent-magnet. Size, morphology, component, and structure of the as-made samples have been confirmed by several techniques, and nanochains (NCs) with diameter of approximately 60 nm consisting of single-crystalline Co and amorphous Co-capped layer (about 3 nm) have been materialized. The as-synthesized Co samples do not include any other adulterants. The high-quality NC growth mechanism is proposed to be driven by magnetostatic interaction because NC can be reorganized under a weak magnetic field. Room-temperature-enhanced coercivity of NCs was observed, which is considered to have potential applications in spin filtering, high density magnetic recording, and nanosensors.

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Introduction

In the last decade, diverse technological applications of magnetic nanostructures in magnetofluid, recording tape, flexible disk recording media, permanent magnets, microwave oscillators as well as biomedical materials, and catalysts have provided an impetus for extensive research in nanometer scale magnets [1-17]. Most of these applications rely on the stability of ferromagnetic ordering with temperature. In nanometer scale magnets, the thermal fluctuations randomize the magnetic moment by overcoming the anisotropy energy leading to unstable state of paramagnetism (non-magnetic materials) or superparamagnetism (corresponding coercivity and hysteresis fall to zero). Cobalt (Co) is superior to other ferromagnetic materials because of its highest Curie temperature ($T_c$) of about 1394 K, which is crucial for thermal stability in high-temperature nanodevice applications [1,5-16]. For realizing high $T_c$, magnetization, and coercivity, the aim must be directed toward increasing the amount of the ferromagnetic Co phase [7-10]. Owing to its basic metallic characteristic, pure cobalt, especially for nanosized Co, is very reactive and must be unstabilized in ambient air [11], and therefore, its use has been limited to prepare Co nanostructures in the absence of the shell [11-16]. A very simple bottom-up method is to produce stable film-assisted synthesis for a surface with slightly controlled Co to passivate the surface of the host materials, including organic and inorganic templates, and alloyed technique [1,2,5-10,12-16]. For example, recently, control of magnetism in cobalt nanoparticles by oxygen passivation was reported by Srikala’s research group [14,15], and cobalt nanowires with controlled diameters have been synthesized using electrochemical deposition in etched-ion-track polycarbonate membranes [16]. In this latter case, however, corresponding magnetic properties are inevitably decreased by the addition of non-magnetic materials or natural oxide layers [1,5-16]. As far as we know, amorphous phases lack long-range crystalline order and have unique electronic, magnetic and corrosion-resistant properties. In this article, based on our earlier study [17], we report for the first time a wet chemical synthesis of high-pure Co nanochains (NCs) without any oxide shells and templates. The amorphous Co covering layer would be able to protect the active Co core from oxygen in atmosphere. In particular, the room-temperature coercivity (up to 355.8 Oe) of the NCs is larger than that (93.6 Oe) of pure single-crystal
Co (PSC) metal, which will make them as promising candidates for advanced magnetic media and investiga-
tive studies on novel great magnetoresistive properties.

**Experimental section**

In a typical experiment, 10.0 mL glycerine was heated to the boiling point (approx 560 K) and refluxed for 3 min. Then, 50 mL of hydrazine monohydrate was added drop-
wise to the boiling solution. After 1 min, 10.0 mL of Co
(NO$_3$)$_2$ 6H$_2$O solution (0.5 mol/L, in glycerine) and
10.0 mL of hydrazine hydrate solution (0.5 mol/L, in gly-
cerine) were added rapidly to the boiling solution under
vigorous magnetic stirring. After refluxing for approx 80
min, as proposed in our previous article [17], the final
products in the form of loose powders (large quantities
of light-gray wool-like products) were obtained by centri-
fugation under permanent-magnet (approx. 0.5 T). The
powders were rinsed repeatedly with absolute ethanol for
several times, followed by the removal of the residual sol-
vent through evaporation in vacuum at 500 K. The yield
of the as-prepared Co specimens is about 60% according
to our calculation. We noticed that no sign of oxidation
was observed on the as-synthesized metal Co NCs even
after aging for over 1 month under ambient conditions.
This indicates that the Co NCs are very stable after sur-
face modification with amorphous Co, which is very
important for future applications. The samples were
characterized extensively for morphology, phase, and
chemical composition using scanning/transmission elec-
tron microscopy (SEM/TEM), energy dispersive X-ray
spectroscopy (EDS), X-ray powder diffraction (XRD),
selective area electron diffraction (SAED), high resolution
TEM (HRTEM), and X-ray photoelectron spectroscopy
(XPS). The temperature dependence of magnetization
and room temperature (RT) hysteresis curves were car-
ried out by means of vibrating sample magnetometer
(VSM, Model 4 HF) and physical properties measure-
ment system (PPMS, Quantum Design PPMS-7).

**Results and discussion**

Figure 1a shows a typical SEM image, in which high
dense (high-yield) chain-like nanostructures are observed.
One can observe that the diameter and length of single
NCs are approx. 60 nm and several micrometers, respec-
tively. A high-resolution SEM image is demonstrated in
the inset of Figure 1a where a nanoparticle array, taken
from a single NC, is clearly seen. Based on SEM statis-
tic analysis, the yield of the NCs is for approx. 60%. The
XRD patterns of bulk Co NCs (Figure 1b) reveal that the
two sharp diffraction peaks can be assigned to the Co
face-centered cubic (FCC/Fm3m) structure ($a = 0.354$
nm) (see Card No. 15-0806, JCPDS-ICDD, June 2002).
Two very broad peaks are noticed in 2 theta angles (20-
40 and 45-60), which may result from the amorphous
passivation Co layers and substrate. In addition, no
impurity phases such as cobalt oxides or precursor com-
ounds have been confirmed within instrumental error.
More accurately, XPS was used to determine the compo-
sition of the bulk Co-NC samples. As shown Figure 1c, a
range of XPS spectrum is indicated, in which the inten-
sive peaks located at 778.3 eV (Co2p3/2) and 793.3 eV
(Co2p1/2) correspond to the respective electronic states
of metallic Co. As indicated with the arrows, these wide
peaks (from 831 to 838 and from 777 to 698) originate from auger lines of monochromated Al for XPS characterization. It is clear that the NCs are high-pure Co$^0$ with other elements absent, which is in very good agreement with the results of XRD. As shown in Figure 2a, a typical slight enlargement TEM image of the as-synthesized large-scale NCs is exhibited. From the TEM image, one can, by closer observation, conclude that perfectly aligned nanoparticles (chain-like nanostructures) were produced and the diameter of individual Co NCs is approx. 60 nm. For microanalysis, HRTEM, SAED, and EDS were employed for phase, and composition of single NCs. A HRTEM image taken from a single nanoparticle is shown in Figure 2b, in which the presence of the gray edge without any stripes and the center with perfect continuous lattice fringes reveals the amorphous passivation shell (marked with the large white arrow) and high-quality crystal core growth. The lattice spacing of 8.86 Å ($5 \times 1.772$ Å) is consistent with that of the [200] planes of single-crystalline face-centered Co, which is compatible with the data of XRD. The SAED patterns (see the inset in Figure 2b) are composed of the regular, clear diffraction dots, which reveal the single crystalline nature and can be indexed to the FCC Co. Diffraction patterns taken from different parts along the NC axis show the same features, indicating the same periodical orientation along the single-crystalline Co NC. Based on micro-composition-analysis, Figure 3a gives a typical EDS pattern, in which only the Co peaks can be indexed within experimental error, and the peaks of elements Cu and C are attributable to copper grid with carbon film.

The synthesized NCs can be reorganized in a weak magnetic field. Typically, the purified Co NCs were dispersed into de-ionized water by ultrasonic agitation. A drop of the Co NCs solution was dripped on a copper grid with holes and carbon film to characterize the TEM in the absence or the presence of the weak external magnetic field (about 0.5 T) and dried naturally. The result reveals that the NCs in the weak magnetic field have aligned according to the direction of the magnetic field as shown in Figure 3b, whereas without the external magnetic field, nonaligned NCs appear as shown in Figure 3c. Regarding the mechanism for the growth of the highly branched Co nanoparticle chains, we believe that magnetostatic interaction plays an important role. The magnetic dipole-dipole interaction displayed behavior similar to that of soft templates. Initially, very small Co nanoparticles were formed. With the increase of the growth time, presumably, the small Co nanoparticles diffused and aggregated to form larger nanoparticles. The Co nanoparticles were then assembled into neck-like chains with multiple branches because of the stronger anisotropic magnetic forces, and these findings are in agreement with those of our earlier article [17].

Despite the presence of the amorphous buffer layer, the NCs display a strong ferromagnetic behavior. Figure 4a shows the magnetization curve as a function of temperature for the NCs and PSC metal. As can be seen, for the NCs, a sharp magnetic transition is observed for $T = \sim 600$ K as determined from the inflection point of the magnetization versus temperature curve; the inflection point may result from the enhanced single-crystal Co mass from amorphous Co because the corresponding magnetic transition disappears in PSC under the same conditions. Figure 4b shows the magnetic hysteresis loop, measured at RT with the applied magnetic field perpendicular to the substrate surface. Based on both these curves, coercive fields ($H_c$) of 355.8 and 93.6 Oe, respectively, for NCs and PSC, saturation magnetizations ($M_s$) of 125.7 and 162.5 emu/g, respectively, for NCs and PSC, and remnant magnetization ($M_r$) up to 46.6, and 4.5 emu/g, respectively, for NCs and PSC were determined. A detailed analysis of the magnetic properties as a function of the NC with various amorphous shell sizes will be published separately.
Conclusion
In this study, this proposed method provides a simple and inexpensive method for the preparation of stable, magnetic Co NCs with the complete absence of impurities. The as-synthesized NCs are produced with increasing $H_c$ and $M_r$ at room temperature. In addition, the large $H_c$, coupled with the controllable coercive field, makes these NC arrays the preferable candidates for probe-based data-storage systems. Introducing long-range, 1D translational order over macroscopic distances among the NCs will undoubtedly be a key driver in this respect, and is an area that clearly warrants future exploration.

Abbreviations
EDS: energy dispersive X-ray spectroscopy; HRTEM: high-resolution TEM; NCs: nanochains; PSC: pure single-crystal Co; SAED: selective area electron diffraction; SEM: scanning electron microscopy; transmission electron microscopy (TEM); XPS: X-ray photoelectron spectroscopy; XRD: X-ray powder diffraction; RT: room-temperature; VSM: vibrating sample magnetometer; PPMS: physical properties measurement system.

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Authors’ contributions
SM: carried out the experimental and numerical calculations, as well as drafted the manuscript. VSM and PPMS: performed an analysis and interpretation of results, and gave final approval of the version to be
published. SY and YQ: partly carried out the TEM and HRTEM experiments, as well as drafted the manuscript. XL, LS, and HL: conceived of the study, and participated in its design and coordination. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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