RESEARCH ARTICLE


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Abstract:

Background: In this paper, we report on the topics of one-dimensional (1D) and two-dimensional (2D) functional materials. Single-Walled Carbon Nanotubes (SWCNTs) are seamless hollow cylinders made of hexagonal lattice graphite sheets. The SWCNTs have attracted considerable attention due to the applicability of their enclosed nanospaces to engineering, and many types of guest materials are encapsulated inside their 1D space, expecting unusual properties. The poly Transition Metal (TM) phthalocyanine, in which phthalocyanine units are extended in two dimensions by sharing benzene rings, is one of the examples of the TM containing 2D carbon materials. Because of strong correlation between localized d-electrons in the TM atom and delocalized π-electrons on the poly phthalocyanine frame, it is expected that spin-polarized conduction, which is useful for the spintronic applications.

Objectives: The objective of the first work is to synthesize SWCNTs encapsulating oxygen molecules having spin one, whose O-O bond directions are aligned to the longitudinal direction of the SWCNTs. The objective of the second work is to synthesize Poly Cu Phthalocyanine (PCuPc) through a bottom-up method by using copper octacyanophthalocyanine as a building block and to elucidate its crystal structure and magnetic properties.

Methods: SWCNTs with inner diameter of ca 0.8 nm were prepared by the CoMoCAT method, and encapsulated together with oxygen molecules (~400 Torr) into a high-purity quartz tube. To subtract the background signals of the SWCNTs and the quartz tube, we prepared the same SWCNTs inducing He gas after evacuating oxygen molecules. Magnetization measurements of these SWCNTs samples were conducted by means of a SQUID magnetometer and a pulse magnet using an induction method. PCuPc were synthesized by a solid state reaction of octacyanophthalocyanine, tetracyanobenzene, and CuCl2·2H2O in glass ampoules sealed after evacuation. The as-synthesized samples were characterized using XRD analysis and TEM microscopy. Magnetization measurement of the samples were done by using a SQUID magnetometer.

Results: The intrinsic magnetization data from oxygen molecules inside the SWCNTs (temperature and magnetic field dependence) show magnetic properties typical of the spin-one Heisenberg antiferromagnet named a Haldane magnet. PCuPc and its half-filling counterpart were obtained by solid state reaction of octacyanophthalocyanine, tetracyanobenzene, and CuCl2·2H2O in glass ampoules sealed after evacuation. Both magnetic susceptibility and magnetization of PCuPc are larger than those of half-filling PCuPc, but the magnitudes of the former sample are about 1.5 times larger than those of the latter one, which is expected to be twice from the geometric superlattice structure.

Conclusion: We have studied magnetic properties (magnetic susceptibility and magnetization) of oxygen molecules encapsulated into Single Walled Carbon Nanotubes (SWCNTs) with diameters of about 0.8 nm, regarded as a 1D functional magnetic material, and Poly Copper Phthalocyanine (PCuPc) and poly half-filling copper phthalocyanine (half-filling PCuPc), regarded as 2D functional magnetic materials.

Keywords: Oxygen molecules, Single-walled carbon nanotubes, Magnetic susceptibility, High-field magnetization, Copper phthalocianine, Half-filling.
1. INTRODUCTION

The emergence of novel functionalities occasionally occurs by conventional materials or ions encapsulated in one-dimensional (1D) nanospaces and by those embedded in ideal two-dimensional (2D) sheets because of restricted spaces and low dimensionality. In the following, we will describe how carbon-based materials, which are usually diamagnetic, have novel magnetic features by introducing magnetic atoms into them.

1.1. Single-Walled Carbon Nanotubes (SWCNTs)

Single-Walled Carbon Nanotubes (SWCNTs) are formed by the $sp^2$ bonding carbon atoms and are seamless hollow cylinders made of hexagonal lattice graphitic sheets [1]. The typical size ranges from less than one nanometer (nm) to a few nanometers in diameter and a few micrometers in length. As mentioned above, the SWCNTs have attracted much attention due to the applicability of their enclosed nanospaces and surfaces to engineering [2 - 11]. Many types of guest materials are encapsulated inside their 1D space, and unusual phenomena have been observed. For example, water molecules inside SWCNTs named ice nanotubes [5] exhibit novel ferroelectric properties [8].

The most interesting magnetic molecule is the oxygen molecule which is a unique magnetic homonuclear diatomic molecule having spin one ($S=1$) and has been studied for more than one hundred years [12, 13]. Magnetic properties of oxygen molecules confined in nanospace have been studied considerably, e.g. in a graphitic slit-shaped nanospace [14] and in microporous metal-organic solids [15 - 17].

In previous numerical studies [11], five different structures are predicted depending on the diameter of the SWCNTs ranged from 0.6 to 2 nm. At the temperatures $T < 10$ K, the narrowest SWCNTs are filled with oxygen molecules having their O-O bond direction aligned to the longitudinal direction of the SWCNTs as shown in the inset of Fig. (1). This geometry of oxygen molecule alignment is named 1DL and the exchange interaction between the neighboring oxygen molecules is expected to be antiferromagnetic [18].

1.2. Phthalocyanine Cu Poly Sheets (PCuPc)

Graphene is a 2D carbon material formed by the $sp^2$ bonding carbon atoms [19], and is intrinsically diamagnetic due to the pairing of $\sigma$ and $\pi$ electrons, except in the states at the zigzag edges [20, 21]. Therefore, considerable efforts for further functionalization have been devoted to introduce Transition Metal (TM) atoms or ions with local magnetic moments into 2D carbon materials. Extensive studies have been made to explore TM-containing 2D carbon nanostuctures, and one of them is Poly TM Phthalocyanine (PTMpc) [22 - 24]. The strong correlation between localized $d$ electrons in TM atom or ion and delocalized $\pi$-electron on the poly phthalocyanine frame makes the spin-polarized conduction possible, which is useful for the spintronics application. Furthermore, the 2D porous structure of PTMpc ensures sufficient exposure of TM atoms to interact with reactants, and thus PTMpc may have functions applicable for the electrocatalysis and photo-catalysis. In addition, theoretical investigation suggests that the PTMpc have high-temperature ferromagnetically or antiferromagnetically long ranged ordered state [24], which is applicable for magnetic functionalized-materials. In the case of TM=Cu$^{2+}$ ions where the theory predicts antiferromagnetic coupling between Cu ions, we expect physical phenomena caused by quantum effects due to its small spin value ($S=1/2$) and two-dimensionality.

2. MATERIALS AND METHODS

2.1. Oxygen Molecules Encapsulated into SWCNTs

The SWCNTs sample with an inner diameter of $ca$ 0.8 nm was prepared by the CoMoCAT method [25] and encapsulated together with oxygen molecules (~400 Torr) into a high-purity quartz tube. After conducting all the magnetic measurements of this sample, oxygen molecules were evacuated and replaced with helium gas. Then, we performed the same magnetic measurements as for the oxygen loaded sample. Since we used the same SWCNTs and quartz tube, the difference between the two samples is nearly identical to the contribution of oxygen molecules [26]. We assumed that the increase of observed magnetic susceptibility at low temperatures would arise from the oxygen molecules outside the SWCNT. Then, we subtracted this contribution, which is fitted by the Curie-Weiss law for $S=1$ with a Weiss temperature of -3 K, from the total magnetic susceptibility to get the intrinsic magnetic susceptibility of oxygen molecules inside the SWCNT. For the magnetization data, we subtracted the extrinsic magnetization calculated by assuming the $S=1$ Brillouin function from the total magnetization given by the difference between the magnetization curve with and without oxygen molecules. Powder X-ray diffraction experiments using synchrotron radiation X-rays were conducted at beamline BL8B of the Photon Factory in Japan in order to check the encapsulation of the oxygen molecules inside the SWCNTs, and we found the 1DL alignment of oxygen molecules depicted in the inset of Fig. (1).

2.2. Poly Cu Phthalocyanine

Poly Cu phthalocyanine and its half-filling counterpart were synthesized according to the methods described in Ref [27]. 1,2,4,5-tetracyanobenzenes (tCB) and CuCl$_2\cdot$2H$_2$O, which were purchased from Aldrich Chemical Company, Inc., were...
Fig. (1). Temperature dependence of magnetic susceptibility of oxygen molecules encapsulated in SWCNTs. The solid line is the magnetic susceptibility calculated numerically for the $S=1$ 1D Heisenberg antiferromagnet. Inset: schematic of the alignment of oxygen molecules inside SWCNTs with inner diameter of ca 0.8 nm (chiral index (6,5)).

Fig. (2). (a) Geometric superlattice structure of PCuPc. Cu, C, and N atoms are denoted with red (large), black (small), and blue (middle) circles, respectively. Hydrogen atoms are omitted for clarity. (b) Geometric superlattice structure of half-filling PCuPc.

reacted at 240 °C for 4 hours in a minimal amount of sulfolane with the catalytic amounts of diaza-bicycloundecene [28] to produce Cu Octacyano-Phthalocyanine (oCCuPc). Then, Poly Cu phthalocyanine (PCuPc) was synthesized by a solid-state reaction of oCCuPc, tCB, and CuCl$_2$·2H$_2$O in glass ampoules sealed after evacuation (1 Pa). The half-filling PCuPc was synthesized without CuCl$_2$·2H$_2$O. The sealed ampoules were placed into a conventional box furnace and maintained at 425 °C for 48 hours, and then allowed to cool naturally to room temperature. The crystal structures of the PCuPc products were analyzed by the powder XRD method with a Bruker D8 ADVANCE ECO diffractometer, and their TEM images were collected with an FEI Technai G2 20 microscope operated at 120 KeV [27]. From these analyses, the expected structures of PCuPc and half-filling PCuPc are depicted in Figs. (2a) and (2b), respectively.

3. EXPERIMENTAL

Magnetic susceptibility of oxygen molecules encapsulated into SWCNTs were measured with a Superconducting Quantum Interference Device (SQUID) magnetometer (Quantum Design MPMS-7XL). To obtain the magnetic susceptibility only from the oxygen molecules (oxygen susceptibility), we also measured the same SWCNTs filled with He gas into the quartz tube after evacuating the oxygen molecules. The intrinsic magnetic susceptibility of oxygen molecules confined into the SWCNTs was obtained by the aforementioned procedure. High field magnetization was measured with a pulsed magnet by an induction method. As mentioned in the subsection 2.1, we subtracted the paramagnetic contribution assuming a Brillouin function from the measured magnetization to get the intrinsic magnetization coming from the oxygen molecules inside the SWCNTs [26].
Magnetic susceptibilities and magnetization curves in magnetic fields of up to ±7 T of PCuPc and half-filling PCuPc were measured with the same SQUID magnetometer. Diamagnetic corrections for these samples were not conducted.

4. RESULTS AND DISCUSSIONS

4.1. Oxygen Molecules Encapsulated into SWCNTs

The temperature dependence of intrinsic magnetic susceptibility (filled circles) is shown in Fig. (1). The intrinsic magnetic susceptibility is compared with the magnetic susceptibility calculated numerically for the S=1 1D Heisenberg antiferromagnet (solid line) [29]. The magnetic susceptibility shows a broad maximum at about 50 K, which is typical of the 1D antiferromagnet and falls down with decreasing temperature, reflecting a singlet ground state with an energy gap to the magnetic excited states. The difference in maximum values between the experimental and calculated magnetic susceptibilities may arise from the imprecise amount of oxygen molecules (6×10^6 mol) evaluated from the oxygen pressure (~400 Torr) and the inner volume of the quartz tube. In addition, the measured magnetic susceptibility decreases steeply at high temperatures and deviates largely from the calculated one. This must be caused by the escape of oxygen molecules absorbed inside the SWCNTs. Actually, oxygen density inside the SWCNTs with the diameter of 1.5 nm at 760 Torr decreases drastically upon heating as shown in Fig. (3). This observation solidifies the fact of the existence of oxygen molecules confined in the SWCNTs at low temperatures.

Next, we show in Fig. (4) the intrinsic magnetization curve at 1.4 K (solid line) which is compared with the magnetization curve calculated for the S=1 1D Heisenberg antiferromagnet using the same parameter values as in the magnetic susceptibility fitting (broken line). Nearly zero- magnetization appears up to about 10 T and increases almost linearly, which
is similar to the magnetization curve of Ni(C$_2$H$_7$N$_2$)$_2$NO$_3$(ClO$_4$)$_2$, abbreviated as NENP [30]. In contrast, oxygen molecules in micro-porous metal-organic solids exhibit a stepwise increase of magnetization [15 - 17]. A small hump observed below 4 T is caused by imperfect subtraction of extrinsic magnetization due to oxygen molecules outside the SWCNTs, which is made by assuming an $S=1$ Brillouin function. The agreement between experiment and calculation is considerably good. The zero-magnetization also indicates the existence of the energy gap between the singlet ground state and the excited magnetic state that is called the Haldane gap [31].

4.2. Poly Cu Phthalocyanine

Fig. (5) shows the temperature dependences of magnetic susceptibilities ($M/H$) of PCuPc (red filled circles) and half-filling PCuPc (blue filled squares) measured at 1 kOe. Both magnetic susceptibilities increase monotonically with decreasing temperature. Here, we used the following formula units: C$_{20}$H$_{4}$N$_{8}$Cu, MW=419.9 g/mol for PCuPc, and C$_{20}$H$_{5}$N$_{8}$Cu$_{0.5}$, MW=389.1 g/mol for half-filling PCuPc. As expected from the crystal structures of these materials, magnetic susceptibility of the PCuPc is larger than that of the half-filling PCuPc. We expect the magnetic susceptibility of PCuPc twice larger than that of half-filling PCuPc. But the half filling geometric superlattice structure was observed partially in PCuPc by the TEM [23], and thus our PCuPc sample is a mixture of full-filling and half-filling PCuPcs, resulting in a smaller magnitude of magnetic susceptibility. We merely observed paramagnetic Curie-like behavior probably because of large distance between neighboring Cu ions, resulting in weak exchange interactions between them.

The magnetization curves at 2 K in magnetic fields of up to ±7 T of the PCuPc and the half-filling PCuPc samples are shown in Fig. (6). As in the magnetic susceptibilities, we use the unit of emu/f.u. for the magnetization. Both samples show gradual upward changes near ±7 T. As mentioned above, the
PCuPc sample partly the half-filling area (Fig. 3b) as in the half-filling PCuPc sample. Thus, the magnitude of the magnetization at 7 T of the PCuPc sample is about 1.5 times larger than that of the half-filling PCuPc sample. From the g-value (2.06) determined by multi-frequency electron spin resonance measurements, we expected the saturation value of magnetization (5750 emu/f.u.) for full-filling PCuPc. But the observed value is less than half of this expected value. As described by Honda et al., breadth of the XRD peak suggests the partial amorphous nature of the reaction product, and therefore Cu atoms may not be contained in some areas of our PCuPc sample [27]. The same situation is observed in the half-filling PCuPc. We also observed paramagnetic magnetization curves in both samples due to the same reason as in the susceptibility measurements.

CONCLUSION

Novel magnetic properties would potentially emerge in conventional nonmagnetic materials when combining (encapsulating or embedding) with magnetic entities. As such trials, we have studied magnetic properties (magnetic susceptibility and magnetization) of oxygen molecules encapsulated into Single-Walled Carbon Nanotubes (SWCNTs) with diameters of about 0.8 nm, regarded as a one dimensional functional magnetic material, and Poly Copper Phthalocyanine (PCuPc) and poly half-filling copper phthalocyanine (half-filling PCuPc), regarded as two dimensional functional magnetic materials.

For the former, we observed magnetic susceptibility with a broad peak at about 50 K and a steep decrease towards zero upon further cooling below 50 K. The magnetization curve shows nearly zero magnetization up to 10 T and increases almost linearly. From these results, we have realized a Haldane magnet by arrayed oxygen molecules with spin one confined into the SWCNTs. This result paved the way to make noble magnetic material, and Poly Copper Phthalocyanine (PCuPc) using copper octa cyan phthalocyanine as a building block. Magnetic susceptibility and magnetization measurements were conducted for these samples. Both magnetic susceptibility and magnetization of PCuPc are larger than those of half-filling PCuPc, but the magnitudes of the former sample are about 1.5 times larger than those of the latter one, which is expected to be twice from the geometric superlattice structure. Furthermore, the saturation value of magnetization is also discussed, and the partial amorphous area suggested by the broad peak of XRD pattern may not contain Cu atoms in both samples.

CONSENT FOR PUBLICATION

Not applicable.

CONFLICT OF INTEREST

The authors declare no conflict of interest, financial or otherwise.

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