Magnetic properties of one-dimensional iron-sulfur compounds formed inside single-walled carbon nanotubes

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Abstract  
In this work, we perform the filling of single-walled carbon nanotubes (SWCNTs) with sulfur and study the magnetic properties of the formed nanomaterials. Encapsulation of sulfur species results in the appearance of a specific magnetic ordering in the system due to the formation of nanoscopic grains composed of sulfur and residual catalytic Fe nanoparticles contained in the SWCNTs. We study the magnetic character of the obtained 1D nanostructures using SQUID magnetometer and reveal a sequential ferromagnetic-antiferromagnetic ordering in the material. Magnetic and optical properties are strongly dependent on the synthesis protocols. We obtain a significant Raman intensity increase related to the encapsulated nanostructures when filling is performed at high-pressure high-temperature conditions. Simultaneously, the magnetic susceptibility gets strongly reduced for high-pressure filling which is related to the escape of iron particles from the nanotube interior, and the magnetic properties of the material are governed by a weak ferromagnetic ordering of Fe-S structures remained inside SWCNTs. Sulfur encapsulation provides the new route for controlling the magnetic properties in 1D nanomaterials that pave the way for advanced magnetooptical applications.

Text  
Carbon-encapsulated magnetic nanoparticles are very prospective in the context of biomedical applications [1], memory devices [2,], and magnetic sensors [3]. Single-walled carbon nanotubes (SWCNTs) stand out among other different protective materials. They not only serve as a template for nano-dimensional growth [4] and promote interactions between the encapsulated species, but also can influence the properties of the formed host-guest structures [5,6,7]. Indeed, the SWCNT ensures a fine dispersion of magnetic species along the longitudinal axis thus restricting the magnetic coupling between them and thereby enhancing the inherent properties of nanoparticles [8–11]. A reduction of intermolecular dipole-dipole interactions between the neighboring encapsulated molecules leading to enhanced magnetic properties was recently demonstrated for single-molecule magnets [12]. 1D arrangement of single-molecule magnets provides the suppression
of quantum tunneling of the magnetization \cite{12,13} that is crucial for practical applications. Initially, filling of nanotubes with magnetic species was studied for cobalt \cite{8,14,15}, iron \cite{9,10,16,17,18} and nickel \cite{19} nanoparticles. These metals are frequently used as catalysts for the growth of nanotubes \cite{20}, and often detected as side products even after the purification process \cite{21}. Briones-Leon \textit{et al.} \cite{22} specifically studied the effects coming from iron filling of SWCNTs differed by the type of conductivity. It was revealed that magnetic properties are affected by both the metallicity and the diameter of nanotubes, encapsulating nanoclusters of different sizes. Namely, the iron-filled metallic SWCNTs demonstrated the pronounced ferromagnetic behavior, while the semiconducting ones were paramagnetic. It is worth noting that magnetic moments associated with structural defects in pristine, non-filled SWCNTs can also order ferromagnetically with the transition temperature being higher in metallic tubes than in semiconducting ones due to the spin-spin interaction via conduction electrons \cite{23,24}.

Recently, sulfur confined inside SWCNTs has attracted a lot of much attention \cite{25,26}. Researchers focus, in particular, on the study of the chemical properties of the encapsulated material and the identification of chemical reactions with lithium ions in order to improve the performance of batteries \cite{26}. Interaction of iron and sulfur can lead to formation of various functional materials, whereas the confinement provides improvement of properties required for solar energy conversion based on iron pyrite \cite{27}, electrode materials for lithium-ion batteries \cite{28,29} or applications based on hydrogen evolution reactions \cite{30}.

Iron-sulfur compounds include a rich diversity of phases with various magnetic properties that make them very attractive for filling of SWCNTs. Conventional synthesis of Fe – S compounds in bulk results in various products including mackinawite (tetragonal FeS), cubic FeS, troilite (hexagonal FeS), pyrrhotite (Fe1-xS), greigite (cubic Fe3S4), pyrite (cubic FeS2), and marcasite (orthorhombic FeS2). Both forms of bulk FeS2 are non-magnetic. In pyrite, Fe2+ ions are in a low spin state leading to a diamagnetic response \cite{31}. At the same time, Fe1+xS materials (x=0-0.2) exhibit magnetic ordering. Iron-rich structures could be ferromagnetic (hexagonal FeS \cite{32}), antiferromagnetic (expanded tetragonal FeS \cite{33}) and monoclinic Fe1-xS \cite{34}) or ferrimagnetic (Fe3S4 and Fe1-xS \cite{35}). The magnetic ordering of iron sulfides depends on crystallinity, the presence of sulfur vacancies \cite{36,37}, impurity phases (i.e., Co doping), and particle size in the nanometer regions \cite{38}. All these factors can lead to a modified magnetic response. Implementation of SWCNTs with known diameters can provide a model structure for studying the properties of ordered magnetic nanomaterial. One of the routes for construction of iron sulfide structures inside the nanotubes is the use of metallic nanograins that are located inside the nanotube channels due to the growth protocols and encapsulation of sulfur through the vapor phase. The properties of resulting iron-sulfur structures may exhibit dependence not only on their particular arrangement and composition, but also on the charge transfer between the encapsulated species and nanotube walls.

In this work, we investigate the magnetic properties of nanoscopic iron-sulfur grains synthesized inside of SWCNTs from encapsulated sulfur and residual iron that was initially used to catalyze the growth of these SWCNTs. We compare the morphology, composition and magnetic properties of products obtained by heating a mixture of SWCNTs with sulfur under high-temperature (HT) and high-temperature high-pressure (HTHP) conditions. In the former case, it is shown that along with sulfur, nanotubes contain encapsulated pyrite nanoparticles, which ensure the antiferromagnetic response of the host-guest nanostructures. The material obtained under HTHP conditions consists of SWCNTs tightly filled with sulfur as indicated by the giant Raman response from the encapsulated species. However, high pressure applied to a mixture of SWCNTs and sulfur along with the heating significantly deforms nanotubes and results in the reconfiguration of the inner structure and escape of the majority of large magnetic particles from the nanotubes thereby quenching the magnetic response of the material.

Experiments were performed on commercially available SWCNTs (TUBALL™, OCSiAl, Novosibirsk) with an average diameter distribution of 1.7-1.9 nm. In order to remove amorphous carbon and residual external catalyst nanoparticles composed mainly from Fe (~7 wt%), SWCNTs were purified using the procedure described elsewhere \cite{31}. The remaining Fe content in the purified

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SWCNTs was estimated by atomic absorption spectroscopy (AAS) to be about 0.3 wt%. These nanotubes were continuously heated with sulfur in a sealed ampoule (HT method) followed by the toluene treatment to remove sulfur from the outside.
of the nanotubes, while the sulfur located inside the nanotubes remained intact \[^{[39]}\]. Energy dispersive X-ray (EDX) spectroscopy revealed ca. 12.7 wt% of sulfur, the content of iron estimated by AAS was about 0.3 wt%. Due to a quite low iron concentration, its visualization using microscopy methods is a rather challenging task. However, sulfur chains placed inside the SWCNTs are discernible in the high-resolution transmission electron microscopy (HR TEM) images (indicated by red arrows in Figure 1a). The homogeneous filling of nanotubes with sulfur by the ampoule method was also previously demonstrated \[^{[39]}\]. Both TEM and Raman spectra have shown that SWCNTs filled with sulfur keep their integrity (Figure 1a,b); no metallic particles can be seen. However, when sulfur is filling the SWCNTs that contain encapsulated residual iron catalyst, iron sulfides are also expected to be formed \[^{[39]}\] as confirmed by X-ray diffraction (XRD) analysis (Figure 1c). One can see the pattern of pyrite FeS\(_2\), which is one of the most probable compounds to be formed according to the iron-sulfur phase diagram \[^{[40]}\]. Note that the fingerprint from graphite is due to graphite foil used to prepare the sample for the XRD study. However, Raman spectroscopy identifies mostly the spectral peaks related to sulfur chains \[^{[42]}\] while no considerable response from pyrite or Fe related vibrations \[^{[42]}\] is seen (vibrations of FeS \[^{[43]}\] and FeS\(_2\) \[^{[44]}\] are shown by bars), making the evaluation of the specific structure of the encapsulated material challenging. One of the ways to detect the Raman signal should be in using different excitation wavelengths in order to resonantly excite the inner structure, similarly to the case of graphene nanoribbons inside SWCNTs \[^{[45]}\].

Figure 1d,e shows the SQUID magnetization data for sulfur-filled SWCNTs in comparison with those for the sulfur-free purified SWCNTs. For the sulfur-free SWCNTs, we detect a weak ferromagnetic response at temperatures at least up to 330 K – a contribution originating from iron encapsulated in nanotubes, which accompanies the intrinsic diamagnetic response of SWCNTs. This behavior is similar to the results reported for iron-filled SWCNTs \[^{[46]}\]. According to the magnetization value, the iron content is estimated as ~0.5 wt% which reasonably agrees with the value obtained by AAS method. Filling the nanotubes with sulfur results in a drastic change of magnetic properties. The magnetization significantly decreases for the filled samples, by about an order of a magnitude in the temperature range 175 – 330 K. Above 175 K the scaled temperature dependence of magnetization well match that of sulfur-free nanotubes, while an additional pronounced feature appears at temperatures below 175 K. It can be seen explicitly after the subtraction of the background related to pristine SWCNTs and encapsulated Fe nanoparticles (Figure 1f). Note, that the magnetization curve of sintered sulfur is featureless in this temperature region (Supporting Figure 1). When temperature decreases, the magnetization of sulfur-filled SWCNTs grows significantly reaching maximum values around 75 K with further continuous decrease at temperatures below 50 K. We identify two characteristic temperatures marked with blue arrows in Figure 1e. The higher one being about 175 K marks the inception of the magnetization increase that we attribute to the formation of ferromagnetic response of encapsulated sulfur-containing nanostructures. The second one (~75 K) corresponds roughly to the point where the magnetization reaches its local maximum, thus indicating where an additional magnetic ordering mechanism starts to be pronounced. At 75 K the impact of external magnetic field on encapsulated ferromagnetic nanoclusters is apparently exceeded by the magneto-dipole or antiferromagnetic-type exchange interaction between the nanoclusters. This results in the appearance of antiferromagnetic
ordering, which prevails at temperature below ~75 K. The magnetization peak position shifts towards lower temperatures with increasing magnetic field (Figure 1f), since stronger antiferromagnetic interactions are needed to override the external field influence. Rather broad features in the M(T) dependences can be explained by the distribution in size and also geometry of the encapsulated nanostructures and therefore different Curie temperatures due to finite-size effects \[47\]. The observed peculiar magnetic ordering is apparently produced by the nanostructures formed due to the reaction of residual Fe nanoparticles with sulfur inside SWCNTs.

An examination of the M(H) curve at 10 K (Figure 1g) reveals that ferromagnetic fragments partially survive down to low temperatures providing the magnetic irreversibility and considerable remnant magnetization. It is worth noting also that the sample’s magnetization saturates at rather low field of about 5 kOe giving an additional evidence for fairly large correlated nanoparticles. Given the possible size of encapsulated particles and the observed coercive field, one can infer that ferromagnetic nanoparticles should possess a monodomain state \[48, 49\], while the relative arrangements of their moments is governed by an external field and interactions within the host-guest structure.

As we mentioned above, iron-sulfur compounds exhibit very different magnetic responses including antiferromagnetic ordering for FeS and Fe\(_{1-x}\)S. However, our XRD data have revealed the patterns only from non-magnetic pyrite. In order to understand the morphology of iron sulfide synthesized inside nanotubes, we performed a control experiment of sintering iron with sulfur excess (1:2) in the same conditions as when filling the SWCNTs. The Raman spectrum for this sulfide exhibits the vibrations characteristic of pyrite FeS\(_2\) (Supporting Figure 2). However, the magnetization curves measured upon heating after zero-field cooling and field cooling both demonstrate a complicated magnetic response and give an additional understanding of processes that take place upon sulfur interaction with Fe nanoparticles (Supporting Figure 3). In the sintered sample the Curie temperature is higher than 350 K and there is no specific inception of the magnetization increase at lower temperatures. However, some features are similar, including the magnetization maxima with a rather small magnetization value of less than 0.08 \(\mu_B/\text{Fe}\), and further decrease of magnetization upon reducing the temperature, which supports the assumption that the dominant response is related to iron sulfide nanostructures in the studied material. Importantly, we detect that magnetization is reduced when the sample is cooled in the presence of magnetic field (Supporting Figure 3b), which is a clear indication of the antiferromagnetic response in the temperature range 150 – 40 K. It seems that the sintered sulfide is composed of pyrite domains with additional surface Fe\(_{S}\) arrangement.

The general similarity between the magnetization curves of the sintered sample and filled SWCNTs suggests that the encapsulated pyrite nanoparticles contains thin surface layers composed of magnetic low-coordinated iron. The exact configuration should be governed by the diameter of the nanotube, similarly to the case of organosulfur compounds \[50,51\] or coronene-based molecules \[52\] inside the nanotubes, where the narrow diameter tubes provide the ordering of molecules stacks \[53\], while larger diameter nanotubes render the formation of graphene nanoribbons \[5,50,54\]. As a driving force for the antiferromagnetic ordering one can consider the long-range magneto-dipole interaction between encapsulated pyrite nanoparticles that can be established even between individual filled nanotubes. However, we note that in the systems with the distances between the separated magnetic nanoclusters around 1 nm the exchange interaction becomes strong and should also be taken into account \[16,55\]. Troilite and recently reported expanded FeS with short range antiferromagnetic order \[53\] have magnetic moments pointing along the c-axis and in the case of formation inside the nanotube moments would be pointing along the tube axis. Another interesting arrangement that was demonstrated inside the nanotubes is the sulfur containing linear polymerized chains \[53\]. The diameter dependent configurations of these chains, including one, two or three aligned 1D structures, also suggest the possibility for antiferromagnetic ordering. More studies need to be performed in order to corroborate the type of encapsulated structure with the detected magnetic response. Moreover, the presence of nanotube structural defects and remained non-reacted
Fe \[^{38,56,57}\], both may influence significantly the properties of the material including its magnetic response.

Finally, we demonstrate how the high pressure applied during the annealing of SWCNTs with sulfur affects the magnetic response of the obtained material. Note that HTHP conditions promote liquid-phase filling of SWCNTs, while the HT protocol results in vapor-phase diffusion of sulfur inside nanotubes. According to EDX analysis, the content of sulfur in the HTHP sample is about 37.5 wt\%. AAS spectroscopy detected about 0.2 wt\% of iron. HR TEM also showed filling of SWCNTs with polymeric chains (indicated by red arrows in Figure 2a), which can be assigned to encapsulated sulfur species. The applied pressure partially breaks the nanotubes shortening them (see inset in Figure 2a) and creating additional structural defects in their walls that correlates with an increase in the ratio of the integrated intensities of D to G peaks in the Raman spectrum (Figure 2b). This is in contrast with the unaffected morphology of SWCNTs filled with sulfur using HT synthesis (Figure 1a and 1b). The Raman spectrum of the pressed sample showed a giant response from the encapsulated sulfur (peaks at \textit{ca.} 319, 394, 645, 714, 788, 1033 and 1100 cm\(^{-1}\)), while the ratio of RBM peaks to G peak intensity is close to the values for initial SWCNTs and nanotubes filled without application of pressure. A similar spectrum previously reported for small diameter sulfur-filled HiPCO-SWCNTs was attributed to a strong van der Waals interaction of the confined species with the walls of the nanotubes \[^{41}\]. We used nanotubes of larger diameter, therefore, the significantly enhanced intensity of the vibrations related to the encapsulated sulfur species suggests a highly dense filling of the SWCNTs cavities under pressure. Unfortunately, these Raman spectral lines hinder the possible response from Fe or iron sulfide particles if they are present. The XRD pattern of the sample does not show any clear fingerprints from phases other than SWCNTs and graphite (Figure 2c).

Turning to the magnetic data, the susceptibility of HTHP-obtained sulfur-filled SWCNTs is significantly (by about an order of magnitude) reduced with respect to that of SWCNTs filled without additional pressure (Figure 2d and 2e). In the magnetization hysteresis measurements (Figure 2f), a saturation around 8 kOe is still visible. However, no clear signature of an antiferromagnetic response is presented. A weak ferromagnetic ordering below 110 K shows up in data taken at H=1 kOe, yet becomes hardly seen at H=10 kOe. We suggest that the ferromagnetic response and the reduced Curie temperature can be explained by the difference in amounts and sizes of the encapsulated magnetic nanostructures. Taking into account the finite-size scaling theory \[^{47}\] we suppose that larger Fe nanoparticles that participated in the Fe-S structure formation were removed from the nanotube interior under elevated pressure. According to the previous report \[^{58}\], in the case of encapsulated FeC\(_2\) molecules the diffusion of iron outside the nanotube and its aggregation into nanoparticles can be completed already at 873 K. Liquid sulfur acts as a transport medium promoting extraction of Fe from the cavity through open ends of SWCNTs and defects in the walls created under high pressure. At the same time, sulfur is much less soluble in this carbon rich environment in comparison with iron, explaining the high filling density of sulfur inside the SWCNTs. When iron gets out, it dissolves into high-pressure graphitic cell escaping from the sample. That agrees with the lowering of the iron content and the magnetization value of the HTHP sample compared with the values for initial and HT SWCNTs.

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Figure 2. SWCNTs filled with sulfur under HTHP conditions: (a) HR TEM image (cyan and red arrows indicate SWCNTs' walls and sulfur chains), insert shows TEM image, (b) Raman spectra, and (c) XRD patterns for pristine (grey) and sulfur-filled SWCNTs (red). (d) ZFC magnetization data taken at H=1 kOe (e) at H=10 kOe for pristine (grey) and sulfur-filled SWCNTs (red). (f) M (H) measured at 2 K.
To conclude, we examined the magnetic properties of SWCNTs with encapsulated iron sulfide nanoparticles obtained by the reaction of residual encapsulated catalytic species with sulfur at elevated temperatures with and without applied high pressure. The material obtained under merely high temperatures contains predominantly pyrite nanoparticles endowing the host-guest structure with a ferromagnetic-type behavior below 175 K and an antiferromagnetic ordering below ~75 K. Such an unusual magnetic behavior for encapsulated sulfur-based nanomaterial material can be assigned to the presence of Fe,S nanolayers and to peculiar ordering due to confinement. We suggest that the antiferromagnetic ordering arises from the long-range magneto-dipole interaction between encapsulated nanoparticles through nanotubes walls and from exchange interaction of closely located nanoparticles. HPHT filling brings about a significant increase of the Raman response from encapsulated sulfur that we attribute to the improvement of the filling of SWCNTs cavities, although simultaneously the iron partially diffuses outside the nanotubes. As a result, the filled SWCNTs obtained by HPHT method exhibit a significantly reduced ferromagnetic response related to trace concentrations of Fe that avoided extraction or conversion into non-magnetic state in pyrite nanoparticles. 1D ordered sulfur compounds inside SWCNTs can be used as a new platform for nanoscale magnetophotonic applications. Properties of the encapsulated structures can be governed by the synthesis parameters and diameters of the hosting nanotubes.

Methods

SWCNTs (OCSiAl) with a diameter range of about 1.7–1.9 nm purified from iron\[^{21}\]\ and pre-opened were used in this work, which we refer to as pristine. Non-purified ones are called raw in this work. The Fe content in the pristine SWCNTs was estimated to be below 0.3 wt% by AAS\[^{21}\]. Filling of nanotubes with sulfur was carried out using two approaches: (1) ampoule method described elsewhere\[^{59}\]\ and (2) hot pressing\[^{59-61,62}\]. Briefly, in the first approach mixture of SWCNTs and sulfur with a ratio of 1:4 was heated in a sealed quartz ampoule at 750°C for 1 h and 600°C for 14 h. Hot pressing process was carried out using a split-sphere multi-anvil apparatus\[^{59}\]. A mixture of SWCNTs and sulfur taken with ratio 1:2 was embedded into a graphite capsule, which was loaded into a high-pressure cell. The experiments were performed at 5 GPa and temperature of 800°C for 2 h. The products synthesized by the ampoule and pressing method were cleaned in toluene for 10 min using an ultrasound bath to remove the excessive sulfur. For the control experiment, a mixture of iron and sulfur taken with ratio 1:2 was heated in a sealed quartz ampoule at 600°C for 15 h.

The structure of materials was characterized by TEM using a JEOL 2010 microscope and scanning electron microscopy using a JEOL JSM 6700F microscope equipped with an EDX detector (TM-3000). XRD patterns were obtained at room temperature using a Shimadzu XRD-7000 diffractometer (Cu Kα radiation, Ni filter on the reflected beam). In order to decrease the background contribution, the powders were deposited on a zero-diffraction plate of a SiO2 single-crystal. For XRD measurements, powders of SWCNTs were pressed between graphite foils under 200 atm for 15 min (residual foil gives the intensive XRD peak located at 26.6°). The concentration of iron in the initial and purified samples was quantified by atomic absorption spectroscopy (AAS) on iCAP-6500 (Thermo Scientific, USA) spectrometer with the addition of sample through a pneumatic nebulizer and a cyclone-type spray chamber. Raman analysis was performed at room temperature with a LabRAM HR Evolution spectrometer using a 514 nm excitation. Magnetization data were taken in the temperature range 1.7 – 330 K, at magnetic fields up to 10 kOe using a MPMS-XL SQUID magnetometer. Temperature dependences of the magnetization were measured on heating after cooling the sample in zero magnetic field (ZFC) or after cooling in magnetic field (FC) as well as upon cooling the sample.

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Supporting information

Supporting Figure 1. Magnetic properties of pristine sulfur measured at 10 kOe.

Supporting Figure 2. Raman spectrum of iron sulfide sintered in an ampoule at 600°C for 15 h.

Supporting Figure 3. Magnetic properties after sintering iron with sulfur without nanotubes (a) ZFC and FC magnetization data at H=1 kOe. (b) Magnetization data taken upon cooling in magnetic field H=10 kOe. (c) M(H) data measured at 2 K revealing apparent remnant magnetization.

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Filling of SWCNTs containing encapsulated iron nanoparticles with sulfur results in the formation of one-dimensional iron sulfide nanoparticles, which demonstrate sequential ferromagnetic-antiferromagnetic ordering. The addition of high pressure treatment results in the partial extraction of iron from SWCNTs’ cavities and reduction of the magnetic susceptibility of the material.