Synthesis of chalcogenide ternary and quaternary nanotubes through directed compositional alterations of bacterial As–S nanotubes†

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Received 9th February 2011, Accepted 23rd May 2011
DOI: 10.1039/c1jm10601e

Semiconductor nanostructures have become intensively investigated by both experimentalists and theoreticians because of their unique size dependent electronic and optical properties.1 One group of the most investigated semiconductors is chalcogenide compounds (MX, M = As, Cd, Zn; X = S, Se, Te) because their band gap can be easily fine-tuned from zero (like the semi-metal HgTe) to a large band gap (e.g. ZnS (Eg = 3.8 eV)).2 Recently, diverse one-dimensional (1D) chalcogenide nanostructures including the semiconducting nanowires and nanotubes have been synthesized and used as important building blocks for many potential applications.3,4 However, majority of the nanostructures were synthesized through chemical or physical methods which typically require harsh reaction conditions such as high operating temperature, extremely high or ultra-low pressure, catalyst and toxic precursors.5,6 In contrast, bio-inspired or biomimic methods allow synthesizing nanoengineered materials with “greener” precursors under mild ambient conditions.7,8 It is well-known that microorganisms play essential roles in the biogeochemical cycling of elements and in the formation of unique minerals/materials1,9–13 through altering the valence/oxidation state of heavy metals and metalloids for anaerobic respiration.14–16 Recent research has shown new insight on the reducing capabilities of certain anaerobic bacteria which offer significant utility in both heavy metal remediation and nano-manufacturing.17–19 Among the bacteria, Shewanella strains have been shown to contribute to the formation of diverse nano-scaled minerals by virtue of their respiratory mechanisms.20,21 For example, Shewanella sp. HN-41 showed the biological synthesis of one-dimensional As–S nanotubes which exhibited photoactive and semiconducting properties via the reduction of As(V) and thioulate under ambient anaerobic culture conditions.3 It has been reported that diverse semiconductor inorganic hybrid nanotubes were synthesized via ion exchange reaction to enhance the functionality and applicability.22–24 It is also known that electrical conduction is closely associated with the structures such as the grain size, defects and impurities. Especially, the conduction of semiconductors is mainly governed by the grain boundary scattering where amorphous/nanocrystalline materials have much lower carrier concentration and mobility than single or polycrystalline materials with larger grains.25 As the grain size increased, the contribution of grain resistance would be reduced, resulting in smaller thermal activation energy, Ea.26 This suggested that the biologically photactive As–S nanotubes can be transformed into tunable structures with varying compositions and ideal electrical property via kinetically controlled solution-phase ion exchange reaction and crystallization. Thus, in this study, various biological activities of dissimilatory metal-reducing bacteria, including the reduction of Se(v) and the formation of the photoactive As–S nanotubes, were applied for the synthesis of the versatile ternary and quaternary chalcogenide (i.e. As–S–Se, As–Cd–S and As–Cd–S–Se) nanotubes with the aid of biological and/or abiological activities. Se and/or Cd were incorporated either by biogenic deposition or ion exchange onto As–S nanotubes to control their electrical properties, which may open-up the possibility to integrate these nanotubes in nanoelectronics, optoelectronics, and solar cells. The mineralogical, crystal structure, morphology and electrical properties of the nanotubes were characterized, thereby understanding the influence of the ratio and different elemental compositions.

Shewanella sp. HN-41 produced the As–S nanotubes via concomitant reduction of As(v) to As(III) and S4O62− to S2− when both 5 mM As(v) and 5 mM thioulate were present in the anaerobic medium as previously described.3 The purified As–S nanotubes were resuspended in the same medium supplemented with 10 mM lactate and 2 mM sodium selenite as the electron donor and acceptor, respectively. After 24 h incubation with the bacterial inoculum, the concentration of dissolved Se in the culture decreased from 2 to
0.9 mM (Fig. 1a) and a filamentous product with smooth surface morphology was formed (Fig. 1b and d). Cross-sectional TEM images showed tube features and EDX line profile analysis showed the presence of As, S and Se in a ratio of 2 : 1 : 2 (Fig. 1c) whereas the ratio of As : S in the As–S nanotubes was 2 : 3. These results implied that the sulfur in the As–S nanotubes was replaced by Se and the composition of the As–S–Se nanotubes can be tuned as As$_2$S$_x$Se$_{3-x}$, depending on the different reaction rates. Since it has been reported that some bacteria including *Shewanella oneidensis* can reduce Se(IV), through Se(0), down to Se(–II),$^{27-29}$ sulfur in the As–S nanotubes appeared to be replaced by Se(–II) which was produced by *Shewanella* sp. HN-41. In order to observe an oxidation state of selenium in the synthesized nanotubes, the XANES peak feature of the Se-ion exchanged nanotube is compared to those of standards Se(IV) and Se(–II) (ESI, Fig. S5a†). As shown in the ESI, Fig. S5a†, the main peak of the Se K-edge data from our samples (As–S–Se nanotubes) and the commercially available reference sample (As$_2$Se$_3$ powder) appeared at approximately 12658 eV which is identical with previously reported As$_2$Se$_3$ from chalcogenide films,$^{30}$ and clearly showed a different absorption peak position as compared to that of Na$_2$SeO$_3$ (Se(IV)). This result suggested that sulfur in the biogenic As–S nanotubes can be replaced by Se(–II) through the reduction of Se(IV) by bacterial culture under anaerobic conditions.

In contrast to As–S–Se, the As–Cd–S nanotubes were synthesized through an abiotic process. The purified As–S nanotubes were resuspended in 2 mM CdCl$_2$ solution. A filamentous product with rough surface morphology (Fig. 2b and d) was formed and the concentration of Cd in the liquid phase decreased from 2 to 0.4 mM coupled to increase of As from 0 mM to 1.1 mM (Fig. 2a) after 2 h incubation. EDX line profile analysis of the tubular structure showed that the ratio of As, Cd and S was approximately 1 : 4 : 5 (Fig. 2c). The significantly lower ratio of As/S indicated Cd incorporation into the As–S nanotubes via cation exchange reaction. XRD spectral analysis showed several diffraction peaks of CdS with the preferred crystal orientation in the (444) and (107) direction (ESI, Fig. S3b†). Similar to Se anionic exchange process, Cd K-edge XANES characterization for the As–Cd–S nanotube directly presents successful ion-exchange reaction between host arsenic and guest cadmium ions (ESI, Fig. S5b†).

Furthermore, consecutive Cd and Se dual-ion exchange can be successfully performed for the As$_2$S$_3$ nanotube to synthesize quaternary As–Cd–S–Se nanotubes. The As–Cd–S synthesized via cation As/Cd exchange reaction was purified and resuspended in the medium containing bacteria, lactate and Se(IV). After 24 h incubation, the concentration of soluble Se decreased from 2 to 1.2 mM (Fig. 3a) and the product with similar rough surface shape was formed (Fig. 3b and d). The EDX line profile analysis of the tubular structure showed that the ratio of As, Cd, S and Se was about 1 : 4 : 4 : 1 (Fig. 3c). The Cd and Se K-edge XANES after dual-ion exchange only showed a Se–As chemical interaction without any Se–Cd chemical bonding, whereas those for the Cd ion directly present distinct Cd–S chemical interaction and an absence of Cd–Se bonding (ESI, Fig. S5c†). This fact indicated that after the first Cd cationic exchange, the second Se anionic exchange occurred not in the Cd site but only the As site. As a result, there coexist arsenic sulfide, cadmium sulfide and arsenic selenide phases in the As–Cd–S–Se nanotube.

In determining the electric properties of the synthesized nanostructures, the temperature coefficient of the resistance (TCR) and...
develop composition and structure dependent nanomaterials and selective, and tunable synthesis can provide new opportunities to genide ternary and quaternary nanotubes by the incorporation of Cd under environmentally benign conditions, producing the chalco-As--S nanotubes can be tuned by biological and abiological processes electrical properties can be tuned by controlling the composition of S7, and S8†, respectively. The comparison of their electrical prop-

In summary the, chemical composition of the biogenic photoactive As–Cd–S network showed almost linear $I$–$V$ characteristics (Fig. 4a), indicating that the As–Cd–S nanotubes formed an ohmic contact. However, as the temperature decreased, the $I$–$V$ curves became non-linear which might be caused by the decrease of the carrier concentration resulting from lower tunneling probability. Fig. 4b shows the temperature dependent resistance. The small activation energy of 13.4 meV was obtained from 270 to 210 K from the As–Cd–S nanotubes which implied a low density of deep charge traps and subsequent high channel conductivity. To further investigate the electrical properties, FET transfer characteristics were measured. As shown in Fig. 4c, the source–drain current ($I_{DS}$) was strongly dependent on the gate bias where a clear off-state was observed at positive bias voltage. From these results it can be inferred that the As–Cd–S nanotubes are p-type semiconductors with the carrier concentration and field effect mobility of 1.1 ± 0.4 × 10^{10} cm^{-1} and 0.08 ± 0.01 cm² V^{-1}s^{-1}, respectively (ESI, Fig. S9b†). Similar analysis was performed on the As–S, As–S–Se, and As–Cd–S–Se nanotubes which are shown in the ESI, Fig. S6, S7, and S8†, respectively. The comparison of their electrical properties is summarized in the ESI, Fig. S9† which clearly indicated the electrical properties can be tuned by controlling the composition of nanotubes.

In summary the, chemical composition of the biogenic photoactive As–S nanotubes can be tuned by biological and abiological processes under environmentally benign conditions, producing the chalco-

**Notes and references**