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Importance of Solution Equilibria in the Directed Assembly of Metal Chalcogenide Mesostructures

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Abstract: We describe the new nanostructured Pt/Ge/Se materials prepared from the molecular units $[Ge_2Se_6]^{4-}$ and $[GeSe_4]^{4-}$ and linking Pt²⁺ ions in the presence of surfactant micelles. X-ray diffraction coupled with transmission electron microscopy images reveals hexagonal pore symmetry. The solvent dependence and solution speciation of these building blocks were investigated by means of multinuclear NMR spectroscopy and by fast atom bombardment (FAB) mass spectroscopy and it is shown that rapid exchange equilibrium is reached between species like $[Ge_4Se_{10}]^{4-}$, $[Ge_2Se_6]^{4-}$, and $[GeSe_4]^{4-}$ in both water and formamide. This results in multiple Ge/Se anions being incorporated in the mesostructured materials which is supported by Raman and IR spectroscopic data. It is likely that the presence of multiple building units both in water and formamide solutions favors the assembly of mesostructured metal chalcogenides with good pore order. Systematic variation of both surfactant headgroup and chain length modulates the optoelectronic properties of the mesostructures. The Pt/Ge/Se materials show sharp band gap transitions in the range of 1.24–1.97 eV. Finally, the materials exhibit reversible ion-exchange properties and a marked inorganic framework flexibility that enables a contraction—expansion process in response to the exchange. The Pt/Ge/Se framework possesses a very high surface area as estimated by small-angle X-ray scattering techniques.

Introduction

Mesoporous solids are open-framework materials with pores in the 2–50 nm size range. They are now some of the most actively studied materials because of their promise for applications in a wide variety of fields ranging from separations, ^{1–5} catalysis, ⁶ and host guest chemistry ^{7,8} to low-*k* dielectric coatings. ⁹ Though oxide-based materials have attracted most of the attention, ^{10,11} porous nonoxidic counterparts are emerging as a unique subclass ¹² that offers the possibility to combine porosity and optoelectronic functionality in a single system promising new capabilities in sensing, photo- and electrocatalytic applications. Prominent members of this subclass are the chalcogenides and a number of elements. ^{13–17} To date,

mesoporous metal chalcogenides with accessible pores are scarce, $^{1.18,19}$ but materials with occupied pores are known and have band-gaps starting from $\sim\!\!3$ to less than 0.7 eV. $^{20-35}$ The synthesis of metal chalcogenide mesostructured materials involves supramolecular assembly of suitable anionic clusters around liquid crystal templates in the presence of appropriate linking metal ions. The solvent and temperature can strongly affect the overall pore order of the mesostructured materials.

Materials with mesopores have been reported using the chalcogenide building blocks $[SnSe_4]^{4-}$, $[Sn_2Se_6]^{4-}$, $[Sn_2Se_6]^{4-}$, $[Ge_4S_{10}]^{4-}$, $[Ge_4Se_{10}]^{4-}$, $[Sn_4Se_{10}]^{4-}$, $[SnTe_4]^{4-}$, $[MoS_4]^{2-}$, and $[SbSe_4]^{3-}$ and different linking metal ions $(Mn^{2+}$, Fe^{2+} , Co^{2+} , Ni^{2+} , Zn^{2+} , Cd^{2+} , Hg^{2+} , Ga^{3+} , In^{3+} , Sb^{3+} , Sn^{4+} , Pt^{2+} , Pt^{4+} ,

- Bag, S.; Trikalitis, P. N.; Chupas, P. J.; Armatas, G. S.; Kanatzidis, M. G. Science 2007, 317, 490–493.
- (2) Kisler, J. M.; Dahler, A.; Stevens, G. W.; O'Connor, A. J. Microporous Mesoporous Mater. 2001, 44, 769–774.
- (3) Feng, X.; Fryxell, G. E.; Wang, L. Q.; Kim, A. Y.; Liu, J.; Kemner, K. M. Science 1997, 276, 923–926.
- (4) Han, Y. J.; Stucky, G. D.; Butler, A. J. Am. Chem. Soc. 1999, 121, 9897–9898.
- (5) Brown, J.; Mercier, L.; Pinnavaia, T. J. Chem. Commun. 1999, 1, 69–
- (6) Climent, M. J.; Corma, A.; Iborra, S.; Miquel, S.; Primo, J.; Rey, F. J. Catal. 1999, 183, 76–82.
- (7) Moller, K.; Bein, T. Chem. Mater. 1998, 10, 2950–2963.
- (8) Yiu, H. H. P.; Wright, P. A.; Botting, N. P. Microporous Mesoporous Mater. 2001, 44, 763–768.
- (9) Wang, Z. B.; Wang, H. T.; Mitra, A.; Huang, L. M.; Yan, Y. S. Adv. Mater. 2001, 13, 746–749.
- (10) Kresge, C. T.; Leonowicz, M. E.; Roth, W. J.; Vartuli, J. C.; Beck, J. S. *Nature* **1992**, *359*, 710–712.
- (11) Davis, M. E. Nature 2002, 417, 813-821.
- (12) Kanatzidis, M. G. Adv. Mater. 2007, 19, 1165-1181.

- (13) (a) Armatas, G. S.; Kanatzidis, M. G. *Science* **2006**, *313*, 817–820. (b) Armatas, G. S.; Kanatzidis, M. G. *Nature* **2006**, *441*, 1122–1125.
- (14) Sun, D.; Riley, A. E.; Cadby, A. J.; Richman, E. K.; Korlann, S. D.; Tolbert, S. H. *Nature* **2006**, *441*, 1126–1130.
- (15) Attard, G. S.; Bartlett, P. N.; Coleman, N. R. B.; Elliott, J. M.; Owen, J. R.; Wang, J. H. Science 1997, 278, 838–840.
- (16) Attard, G. S.; Goltner, C. G.; Corker, J. M.; Henke, S.; Templer, R. H. Angew. Chem., Int. Ed. Engl. 1997, 36, 1315–1317.
- (17) Johnson, S. A.; Ollivier, P. J.; Mallouk, T. E. Science 1999, 283, 963–965.
- (18) Vanchura, B. A.; He, P. G.; Antochshuk, V.; Jaroniec, M.; Ferryman, A.; Barbash, D.; Fulghum, J. E.; Huang, S. D. J. Am. Chem. Soc. 2002, 124, 12090–12091.
- (19) Mohanan, J. L.; Arachchige, I. U.; Brock, S. L. Science 2005, 307, 397–400.
- (20) (a) Trikalitis, P. N.; Rangan, K. K.; Bakas, T.; Kanatzidis, M. G. *Nature* 2001, 410, 671–675. (b) Rangan, K. K.; Billinge, S. J. L.; Petkov, V.; Heising, J.; Kanatzidis, M. G. *Chem. Mater.* 1999, 11, 2629–2632.
- (21) Trikalitis, P. N.; Bakas, T.; Papaefthymiou, V.; Kanatzidis, M. G. *Angew. Chem., Int. Ed.* **2000**, *39*, 4558–4562.
- (22) Rangan, K. K.; Trikalitis, P. N.; Kanatzidis, M. G. J. Am. Chem. Soc. 2000, 122, 10230–10231.

 Pd^{2+}). $^{20-36}$ Among these metals, Pt^{2+} has yielded materials of high pore order presumably because of its slower ligand substitution kinetics during the linking reaction that allows the structure more time to organize. Although the adamantane $[Ge_4Se_{10}]^{4-}$ cluster and various linking metals resulted in high quality mesostructured materials, 24 the corresponding dimeric $[Ge_2Se_6]^{4-}$ and monomeric $[GeSe_4]^{4-}$ species have not been studied. We will show here that unlike the adamantane these two species undergo transformation equilibria that complicate the assembly process.

We describe the synthesis of platinum-based hexagonal mesostructured systems from $[Ge_2Se_6]^{4-}$ and the $[GeSe_4]^4$ clusters and explore the role of solution equilibria in controlling mesophase composition and pore organization. We find that systematic variation of surfactants changes the pore size and pore-pore spacing, the overall symmetry and even the energy band gaps of the mesostructured materials. Surprisingly, compared to the materials with cubic pore $Ia\bar{3}d$ symmetry obtained with the corresponding [Sn₂Se₆]⁴⁻/Pt²⁺ combination, ²⁶ the analogous [Ge₂Se₆]⁴⁻ did not produce cubic pore symmetry. ⁷⁷Se NMR spectroscopy of [Ge₂Se₆]⁴⁻, in water and formamide (FM) as well as fast atom bombardment mass spectroscopy (FABS) show multiple species present in equilibrium all of which can be incorporated into the Pt-Ge-Se framework. This is in contrast to the [Sn₂Se₆]⁴⁻ system which shows only a single species. The implications of multiple building unit participation to assemble hexagonally ordered materials is discussed. The new materials described here exhibit reversible ion-exchange properties and flexible inorganic networks. Small angle X-ray scattering (SAXS) experiments on selected systems suggest a very high mesoporous surface for the inorganic framework.

Experimental Section

Starting Materials. $K_4Ge_2Se_6$ and K_4GeSe_4 were prepared according to a literature procedure. 37,38 K_2PtCl_4 was purchased from Strem Chemical Inc. Cetylpyridinium bromide monohydrate $(C_{16}PyBr.H_2O)$ and formamide (FM) were purchased from Aldrich. The surfactants $C_nH_{2n+1}N(C_5H_5)Br$ (denoted as C_nPyBr ; n=14, 18, and 20), $C_nH_{2n+1}N(CH_3)_2(CH_2)_3N(CH_3)_2Br$ (denoted as $C_nT-MPDABr$; n=20, 22), and $C_nH_{2n+1}N(CH_3)_2(CH_2)_2N(CH_3)_2Br$

- (23) Rangan, K. K.; Trikalitis, P. N.; Bakas, T.; Kanatzidis, M. G. Chem. Commun. 2001, 9, 809–810.
- (24) Trikalitis, P. N.; Rangan, K. K.; Kanatzidis, M. G. *J. Am. Chem. Soc.* **2002**, *124*, 2604–2613.
 (25) Rangan, K. K.; Trikalitis, P. N.; Canlas, C.; Bakas, T.; Weliky, D. P.;
- Kanatzidis, M. G. *Nano Lett.* **2002**, 2, 513–517. (26) Trikalitis, P. N.: Rangan, K. K.: Bakas, T.: Kanatzidis, M. G. *J. Am*
- (26) Trikalitis, P. N.; Rangan, K. K.; Bakas, T.; Kanatzidis, M. G. J. Am. Chem. Soc. 2002, 124, 12255–12260.
- (27) Trikalitis, P. N.; Ding, N.; Malliakas, C.; Billinge, S. J. L.; Kanatzidis, M. G. J. Am. Chem. Soc. 2004, 126, 15326–15327.
- (28) Trikalitis, P. N.; Bakas, T.; Kanatzidis, M. G. J. Am. Chem. Soc. 2005, 127, 3910–3920.
- (29) Trikalitis, P. N.; Kerr, T. A.; Kanatzidis, M. G. Microporous Mesoporous Mater. 2006, 88, 187–190.
- (30) Ding, N.; Takabayashi, Y.; Solari, P. L.; Prassides, K.; Pcionek, R. J.; Kanatzidis, M. G. Chem. Mater. 2006, 18, 4690–4699.
- (31) MacLachlan, M. J.; Coombs, N.; Ozin, G. A. *Nature* **1999**, *397*, 681–684
- (32) MacLachlan, M. J.; Coombs, N.; Bedard, R. L.; White, S.; Thompson, L. K.; Ozin, G. A. J. Am. Chem. Soc. 1999, 121, 12005–12017.
- (33) Riley, A. E.; Tolbert, S. H. J. Am. Chem. Soc. 2003, 125, 4551-4559.
- (34) Korlann, S. D.; Riley, A. E.; Kirsch, B. L.; Mun, B. S.; Tolbert, S. H. J. Am. Chem. Soc. 2005, 127, 12516–12527.
- (35) Riley, A. E.; Korlann, S. D.; Richman, E. K.; Tolbert, S. H. Angew. Chem., Int. Ed. 2006, 45, 235–241.
- (36) Trikalitis, P.; Rangan, K. K.; Kanatzidis, M. G. Mater. Res. Soc. Symp. Proc. 2002, 703, 1–6.
- (37) Eisenmann, B.; Hansa, J. Z. Kristallogr. 1993, 203, 301-302.
- (38) Klepp, K. O. Z. Naturforsch., B: Chem. Sci. 1985, 40, 878–882.

(denoted as C_n TMEDABr; n=20, 22) were synthesized by refluxing the corresponding alkyl-bromide with an excess of pyridine, N,N,N',N'-tetramethyl-1,3-propanediamine (denoted as TMPDA) and N,N,N',N'-tetramethyl-1,2-ethanediamine (denoted as TMEDA), respectively, in either ethanol or acetone. The solvent was removed using a rotary evaporator and pure compounds were obtained in high yield (>90%) after a single recrystallization from CHCl₃-ethyl acetate. Surfactant purity was checked by ¹H NMR after dissolving the solid sample in CDCl₃. The materials we describe here are generically formulated as $Surf_Pt_Ge_2Se_6$ and $Surf_Pt_Ge_2Se_6$ or $C_nTMPDAPt_Ge_2Se_6$ or $C_nTMPDAPt_Ge_2Se_6$ or $C_nTMEDAPt_Ge_2Se_6$ or $C_nTMEDAPt_Ge_2Se_6$

Mesostructured Platinum Germanium Selenides. All reactions were carried out inside a glovebox under nitrogen. In a typical preparation, 0.4 g of surfactant (for pyridinium based) was dissolved in 8 mL of FM under stirring. In another vial, 0.078 g (0.1 mmol) of K₄Ge₂Se₆ was dissolved in 1 mL of hot FM forming a clear deep yellow solution. An amount of 0.042 g (0.1 mmol) of K₂PtCl₄ was dissolved in another 1 mL of FM to make a deep orange-red solution. These two solutions were added dropwise at the same time to the surfactant solution over a period of 10 min using pipets. A dark brown-red solid formed within a few minutes and the mixture was stirred overnight at 75 °C. The dark pinkish brown product was isolated by filtration, washed with copious amount of warm (75 °C) FM and H₂O, and dried under vacuum (overall yield $\sim 90\%$ based on K₄Ge₂Se₆). For the other surfactants, 0.25 g was dissolved in 4 mL of hot FM and 0.15 mmol platinum salt was added instead of 0.1 mmol.

Synthesis of mesostructured C_{16} PyPt_Ge₂Se₆ in water was carried out using the same procedure as above, except 0.4 g of surfactant C_{16} PyBr.H₂O was dissolved in 4 mL of water (overall yield \sim 90% based on K_4 Ge₂Se₆). However, slight variation of surfactant concentration did not affect the overall pore order of the final mesostructures.

In the case of $[GeSe_4]^{4-}$, 0.05 g of K_4GeSe_4 (0.1 mmol) and 0.4 g of surfactant was dissolved in 1 and 4 mL of hot FM respectively. An amount of 0.04 g (0.1 mmol) of K_2PtCl_4 dissolved in 1 mL of FM was added at the same time with K_4GeSe_4 solution to the surfactant solution. A dark brown precipitate was collected after aging overnight at 75 °C followed by washing with 30 mL each of hot FM and water with overall yield of ~75% based on K_4GeSe_4 . The $C_nPyPt_Ge_2Se_6$ materials are stable in air for a couple of days, while those prepared from $[GeSe_4]^{4-}$ are stable only for couple of hours after which the pore structure collapsed.

Ion-Exchange Experiments. Ion exchange reactions were carried out in nitrogen atmosphere at 75 °C in FM. Typically, 1.2 g of C_n PyBr (e.g., n=12) surfactant was dissolved in 6 mL of hot FM and then 0.15 g of mesostructured C_{18} PyPt_Ge₂Se₆ was added to it. The mixture was stirred for 22 h, filtered hot, washed subsequently with hot FM and water and then dried under vacuum. Ion exchange with smaller cationic species such as NH_4^+ was performed in a similar fashion as above using 0.12 g of CH_3COONH_4 dissolved in 12 mL of ethanol and adding 0.208 g of mesostructured C_{18} PyPt_Ge₂Se₆ material at room temperature. The NH_4^+ exchanged material is denoted as (NH_4) Pt_Ge₂Se₆.

Synthesis of Hexagonal Mesostructured Pt/Ge/Se Materials Using a Mixture of Selenogermanate Anions. An amount of 0.4 g of surfactant (C_n PyBr; n = 16, 18) was dissolved in 8 mL of FM under stirring. In another vial, 0.025 g (0.05 mmol) of K_4GeSe_4 and 0.07 g (0.05 mmol) of $(TMA)_4Ge_4Se_{10}$ were dissolved together in 1 mL of hot formamide. An amount of 0.042 g (0.1 mmol) of K₂PtCl₄ was dissolved in another 1 mL of FM to make a deep orange-red solution. These two solutions were added dropwise at the same time to the surfactant solution over a period of 10 min using pipets. A dark brown-red solid formed within a few minutes and the mixture was stirred overnight at 75 °C. The dark pinkish brown product was isolated by filtration, washed with copious amount of warm (75 °C) FM and H2O, and dried under

vacuum. The powder X-ray diffraction of the product indicate a hexagonal order similar to the other hexagonal materials described here

Physical Measurements. Powder X-ray Diffraction (XRD), Elemental Analysis, Thermogravimetric Analysis (TGA) and Mass Spectroscopy. Powder X-ray diffraction patterns were acquired on a Rigaku Rotaflex diffractometer (45 kV, 100 mA) using a rotating anode Cu Kα rotation source. The scan rates were 0.15 deg/min and a step size of 0.02 deg. Quantitative microprobe analyses were performed on a JEOL JSM-6400 V scanning electron microscope equipped with a Noran energy dispersive spectroscopy (EDS) detector. Data acquisition was performed with an accelerating voltage of 25 kV and 60 s accumulation time. Elemental C, H, N analyses were obtained on a Perkin-Elmer Series II CHNS/O Analyzer 2400. Thermogravimetric analyses were performed using a computer controlled Shimadzu TGA-50 analyzer. Typically, 40 mg of sample was placed in a silica bucket and heated in a nitrogen flow of 50 mL/min with a rate of 10 °C/min. Pyrolysis mass spectra were obtained with a TRIO-1 mass spectrometer. Samples were heated at 20 °C/min and the volatile products were ionized by electron ionization. A fast atom bombardment (FAB) mass spectrum was acquired using JEOL JMS HX 110 double focusing magnetic sector mass spectrometer and p-nitrobenzyl alcohol (NBA) was used as matrix.

Band Gap Measurements. UV/vis/near-IR diffuse reflectance spectra were obtained at room temperature on a Shimadzu UV-3010 PC double beam, double monochromator spectrophotometer in the wavelength range of 200–2500 nm. BaSO₄ powder was used as a reference (100% reflectance) and base material on which the powder sample was coated. The reflectance data were converted to absorption using the Kubelka-Munck function, and the band edge for each sample was estimated from the intercept of the line extrapolated from the high-energy end of the absorption to the baseline.³⁹

Infrared and Raman Spectroscopic Measurements. Spectra in the far-infrared (Far-IR) region (600–100 cm⁻¹) were recorded with a computer-controlled Nicolet 750 Magna-IR series II spectrometer equipped with a TGS/PE detector and silicon beam splitter in 2 cm⁻¹ resolution. Samples were diluted by making CsI pellets and then spectra were recorded in transmittance mode under inert nitrogen atmosphere. Raman spectra (3500–100 cm⁻¹) were recorded on a computer-controlled BIO-RAD Fourier transform FT-Raman spectrometer with a Spectra-Physics Topaz T10–106c 1064-nm YAG laser. The instrument was configured in 180° backscattering mode and samples were loaded into melting point capillary tubes.

Transmission Electron Microscopy (TEM). Transmission electron micrographs were obtained with a JEOL 2200FS instrument (field emission) operating at 200 kV using the facilities of the Center of Advanced Microscopy at Michigan State University. TEM samples were prepared by suspending the precipitate in diethyl ether, then casting on a holey carbon grid after 10 min of sonication.

Nuclear Magnetic Resonance (NMR) Spectroscopy. All ⁷⁷Se NMR spectra were recorded on a Varian UnityPlus-500 NMR spectrometer equipped with a 5-mm broadband probe over the frequency range 50-220 MHz and were obtained without locking. The spectrometer frequency was 95.367 MHz for ⁷⁷Se. Typical measurement conditions were a spectral width of 39024.4 Hz, a 90° pulse angle, an acquisition time of 0.5 s with data point resolution of 1.0 Hz/pt and 0.2 s relaxation delay between scans. The pulse width setting was $6.0 \, \mu s$. An exponential multiplication of free induction decay of 5 Hz was used.

The ⁷⁷Se two-dimensional exchange spectroscopy (2D-EXSY) homonuclear exchange spectra were obtained at 65 °C. The pulse sequence for the 2D spectra was standard nuclear Overhauser

enhancement spectroscopy (NOESY) parameters. $^{40-42}$ The spectral width in both dimensions was 39024.4 Hz. The acquisition time was 0.21 s, and a relaxation delay of 0.5 s was used between transients. A total of 92 transients were collected for each of the 400 longitudinal relaxation time (t_1) increments. The mixing time (t_m) employed was 0.15 s. The F_1 dimension was zero-filled to 16384 points, a Gaussian function was applied to both dimension prior to Fourier transformation. A 3-fold linear prediction in the F_1 dimension was employed. The 77 Se nucleus was externally referenced to a sample of neat (CH_3)₂Se at 25 °C.

Solution ¹¹⁹Sn NMR spectra were obtained without locking on Varian Inova 400 NMR spectrometer equipped with a 5-mm broadband probe. The spectrometer frequency was 149.341 MHz for ¹¹⁹Sn. Typical measurement conditions were a spectral width of 69991.3 Hz, an acquisition time of 0.3 and 0.1 s relaxation delay between scans. The pulse width setting was 8.0 μ s. The ¹¹⁹Sn nucleus was externally referenced to (CH₃)₄Sn at room temperature. All NMR samples were prepared by dissolving 0.2 mmol of the sample in 2 mL of solvent (FM or H₂O) at about 60 °C.

Small Angle X-Ray Scattering (SAXS) Analysis. Small angle X-ray scattering (SAXS) was carried out with a Bruker Hi-star twodimensional area detector (512 × 512 pixels) coupled with an Elliott GX3 microfocusing Cu rotating anode operated at 40 kV and 13 mA $(0.01 \le q \le 0.6 \text{ Å}^{-1})$. Measurements were performed by transmission in gently grounded samples and held in a custom built cell. The sample cell was a Kapton-windowed (~4 mm width) cell in \sim 40 μ m thick zirconium metal foil (1 cm \times 2 cm). The sample to detector distance and the center of the beam were precisely determined by calibration with Ag-behenate diffraction standard $(d_{001} = 5.838 \text{ nm})$. The diffraction intensities of two-dimensional (2-D) collected images were integrated to yield one-dimensional (1D) diffraction patterns as a function of the wave vector q with the Fit2D program.⁴³ The wave vector q is defined as $q = 4\pi \sin \theta$ θ/λ , where 2θ is the scattering angle. Scattering data were corrected for dark current and empty cell scattering.

Preliminary attempts to determine the Porod constant (K) in the pristine C_{18} PyPt_Ge₂Se₆ sample were unsuccessful with the Bruker Hi-star system. The appearance of the first (100) Bragg diffraction peak in the relatively low q region masked the Porod region. To determine well the Porod constant (K) in the pristine C_{18} PyPt_Ge₂Se₆ sample, measurements were performed with a Rigaku X-Max 3000 High Brilliance SAXS system equipped with Cu rotating anode operated at 40kV and 86mA and equipped with a multilayer mirror and 2D wire detector. The sample to detector distance was set to ~150 cm to increase resolution of the scattering curve in the low q region (0.01 < q < 0.15 Å $^{-1}$).

Pair Distribution Function (PDF) Analysis. Diffraction experiments for pair distribution function analysis (PDF) were performed at the Advanced Photon Source (APS) located at Argonne National Laboratory, Argonne, Illinois using the high energy X-rays with the powder samples packed in 1 mm glass capillary. For data collection, an X-ray energy of 77.528 keV ($\lambda=0.15992$ Å) was used to record diffraction patterns to high values of momentum transfer while eliminating fluorescence from the sample. The two-dimensional images were integrated within Fit2D to obtain the one-dimensional powder diffraction pattern, masking areas obscured by the beam stop arm. The PDFs, $G(r)=4\pi r[\rho(r)-\rho_{\rm o}]$ where $\rho(r)$ and $\rho_{\rm o}$ are the instantaneous and average densities, were

⁽³⁹⁾ Mccarthy, T. J.; Ngeyi, S. P.; Liao, J. H.; Degroot, D. C.; Hogan, T.; Kannewurf, C. R.; Kanatzidis, M. G. *Chem. Mater.* **1993**, *5*, 331–340

⁽⁴⁰⁾ Perrin, C. L.; Dwyer, T. J. Chem. Rev. **1990**, 90, 935–967.

⁽⁴¹⁾ Milne, J.; Williams, A. J. *Inorg. Chem.* **1992**, *31*, 4534–4538.

⁽⁴²⁾ Wynants, C.; Vanbinst, G.; Mugge, C.; Jurkschat, K.; Tzschach, A.; Pepermans, H.; Gielen, M.; Willem, R. Organometallics 1985, 4, 1906–1909.

⁽⁴³⁾ Hammersley, A. P.; Svensson, S. O.; Hanfland, M.; Fitch, A. N.; Hausermann, D. *High Press. Res.* **1996**, *14*, 235–248.

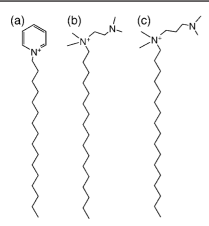


Figure 1. Surfactants used in this study: (a) C_nPyBr (n = 14 to 20), (b) $C_nTMEDABr$ (n = 20, 22), and (c) $C_nTMPDABr$ (n = 20, 22).

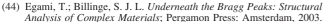
extracted using PDFgetX2, 44,45 subtracting the contributions from the sample environment and background to the measured diffraction intensities. Corrections for multiple scattering, X-ray polarization, sample absorption, and Compton scattering were then applied to obtain the structure function S(Q). Direct Fourier transform of the reduced structure function F(Q) = Q[S(Q) - 1] yielded G(r), the pair distribution function.

Density Measurements. The density of the materials was measured by a Micromeritics AccuPyc 1340 gas pycnometer (1 cm³ model) using ultra high purity (UHP) helium gas. About 200 mg sample was used for the measurements. Analysis run cycles were continued until a standard deviation of \pm 0.0002 cm³ was obtained for the sample volume.

Results and Discussion

New mesostructured materials starting from [Ge₂Se₆]⁴⁻/Pt²⁺ and [GeSe₄]⁴⁻/Pt²⁺ combinations were prepared having hexagonal pore order, good thermal stability, reversible ionexchange properties and strong absorption in the visible spectrum. Previous work with the monomeric $[SnQ_4]^{4-}$ (Q = Se, Te) clusters indicated the presence of solution equilibria and multiple building units, some or all of which could be incorporating in the final structure thereby playing a role in influencing the pore order.²⁸ Many questions however still remain unanswered. The temperature dependence of equilibrium is one of the most important issues to be understood since all templated assembly reactions are performed within a certain temperature range (between 60 and 100 °C). To study the precursor anions in solution and gain additional insights regarding speciation in water and FM solutions, we used ⁷⁷Se NMR and FAB MS.

Long Range Periodicity: Role of Surfactant and Inorganic Cluster. In this work, we employed surfactants with three different head-groups and variable chain lengths to observe their effect on long-range pore periodicity and other properties of the templated materials. In addition to the most commonly used pyridinium surfactant,²⁸ we also employed surfactants with tetramethyl-propanediamine (TMPDA) and tetramethyl-ethanediamine (TMEDA) as the head groups, Figure 1. The effective headgroup areas⁴⁶ of these two surfactants are significantly larger than that of the pyridinium-based ones. The $C_nPyBr/$



⁽⁴⁵⁾ Qiu, X.; Thompson, J. W.; Billinge, S. J. L. J. Appl. Crystallogr. **2004**, 37, 678

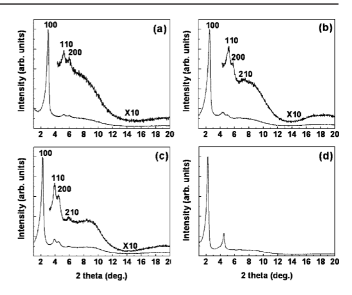


Figure 2. Low angle X-ray powder diffraction patterns of C_n PyPt_Ge₂Se₆, where: (a) n = 14, (b) n = 16, (c) n = 18, and (d) n = 20 (lamellar).

 $K_4Ge_2Se_6/Pt^{2+}$ system gave hexagonal pore symmetry from n=14 to n=18 and lamellar structure at n=20. In the case of $C_nTMPDABr$ and $C_nTMEDABr$ surfactants, hexagonal materials were formed even at n=22. This can be rationalized in terms of surface curvature of the cylindrical micelle formed. The larger head groups of $C_nTMPDABr$ and $C_nTMEDABr$ create cylindrical micelles with larger surface and greater curvature and can thus accommodate the inorganic clusters around it without forming lamellar phases up to n=22. The decrease in cylindrical surface area for C_nPyBr allows the $[Ge_xSe_y]^{z-}$ anions to pack more tightly resulting in lamellar phases for n>18.

In X-ray diffraction, all samples (except n=20) showed three or four well-defined Bragg reflections in the $2^{\circ} < 2\theta < 7^{\circ}$ region as shown in Figure 2, characteristic of a regular hexagonal pore arrangement similar to those of mesostructured silica MCM-41. In the K₄Ge₂Se₆/Pt²⁺/C_nPyBr system, the highest hexagonal pore order was found when n=18, as evidenced by the more intense and sharper higher order reflections (110) and (200), as well as the observation of the fourth (210) reflection, Figure 2c. The materials do not show Bragg reflections above 10 deg in 2θ indicating the inorganic frameworks are nonperiodic.

The extended hexagonal pore structure is revealed by TEM, Figure 3. The wall thickness of the $C_{18}PyPt_Ge_2Se_6$ compound is \sim 2.2 (\pm 0.1) nm and pore diameter \sim 2.4(1) nm. Pore-pore spacing of 4.3(2) nm obtained from the TEM image is in good agreement with the X-ray diffraction data. In the case of $[Ge_4Se_{10}]^{4-}/Pt^{2+}/C_nPyBr$ combination described previously, however, hexagonal pore symmetry was observed even when n=22 due to the larger cluster size of $[Ge_4Se_{10}]^{4-}$ compared to $[Ge_2Se_6]^{4-}$. 24

The C_n TMPDABr family of surfactants also gave hexagonal structures with the highest degree of order observed for n = 20. This is indicated by the four sharp well resolved Bragg peaks in the X-ray diffraction pattern of Figure 4, and also the clear TEM image shown in Figure 5. The pore-pore distance, here increased slightly to 4.54 nm compared to the pyridinium analogue. For the C_n TMEDABr class of surfactants, the sharpest powder X-ray diffraction pattern was obtained when n = 22, but still in this case the (110) and (200) peaks were not as well resolved as with the other family of surfactants, Figure 6.

⁽⁴⁶⁾ Huo, Q. S.; Margolese, D. I.; Stucky, G. D. Chem. Mater. 1996, 8, 1147–1160.

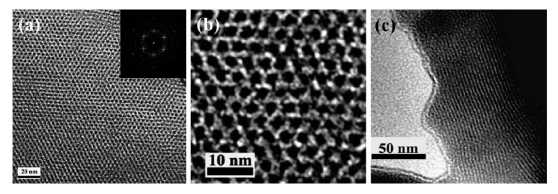


Figure 3. TEM images (a and b) of C_{18} PyPt_Ge₂Se₆ prepared in FM down the pore channel axis and (c) perpendicular to pore channel axis. Inset of (a) shows corresponding Fourier transform.

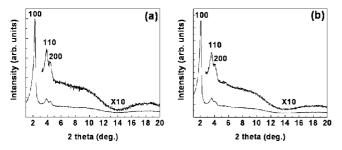


Figure 4. Low angle X-ray powder diffraction patterns of C_n TMPDAPt_Ge₂Se₆, where: (a) n = 20 and (b) n = 22.

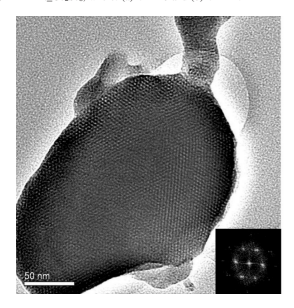


Figure 5. Typical TEM image of C₂₂TMPDAPt_Ge₂Se₆ along the (100) direction and the corresponding Fourier transform; larger particle showing hexagonal arrangement extending over its entire body.

Nevertheless, clear hexagonal pore organization was observed from TEM images in Figure 7.

Reactions with the [Ge₂Se₆]⁴⁻/Pt²⁺/C₁₆PyBr combination in water also led to similar ordered hexagonal pore symmetry as evidenced by four sharp low angle Bragg peaks and the definitive hexagonal TEM image in Figure 8. The unit cell constant of this material was found to be 0.9 Å longer than that of the corresponding material made in FM under similar conditions.

In the K₄GeSe₄/Pt²⁺/ C_n PyBr system good hexagonal order was observed when n = 14. Four well resolved, intense, sharp peaks in the range of $2^{\circ} < 2\theta < 8^{\circ}$ were obtained in the powder

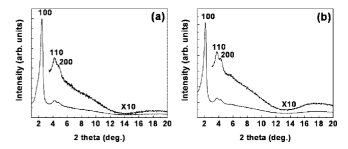


Figure 6. X-ray powder diffraction patterns of $C_nTMEDAPt_Ge_2Se_6$ (a) n=20 and (b) n=22.

XRD as shown in Figure 9. Moving to higher surfactant chain length did not improve the degree of pore organization, while a lamellar phase was obtained for n = 20. This underscores the necessity of higher surface curvature to obtain good long-range periodicity for this smaller cluster.

Table 1 shows trends of hexagonal unit cell constants with surfactant chain length and starting building blocks. The cell a constant gradually increases from 33.7(1) to 44.6(1) Å along the $C_nPyPt_Ge_2Se_6$ series ranging from C_{14} to C_{18} hydrocarbon chain length.

Optical Properties. The optical absorption properties of the title materials show clear energy gaps are in the range of 1.24–1.97 eV, see Figure 10 and Table 2. There is a significant variation in the energy gap considering the inorganic framework is always "Pt/ Ge/Se". As we will argue below, the exact Pt/Ge/Se composition changes with surfactant and with carbon tail length (n). In the C_n PyPt_Ge₂Se₆ system, starting from n = 14 to n = 20 the energy gap gradually increases from 1.24 to 1.54 eV. This suggests that the inorganic walls change in thickness and or composition and it is consistent with the elemental analysis results shown in Table 2. For hexagonal pore packing, the inorganic wall is made up with two different structural regions: (a) wall lying between two adjacent pores (2-WR) and (b) wall lying between three such pores (3-WR).²⁸ The amount of Pt/Ge/Se material found in the 2-WR and 3-WR regions gradually changes with surfactant length and this creates changes in band gap energy. In other words changing the value of n does not produce rigorous analogs as the formulas C_nPyPt_Ge₂Se₆ would imply but homologues with varying composition. A similar trend was previously observed in the c-C_nPyPtSnSe (n = 18 and 20) system, ²⁶ where moving to higher surfactant chain length band gap also increased. Figure 10b further shows increase in band gap energies of C₂₂TMPDAPt_Ge₂Se₆/C₂₂TMEDAPt_Ge₂Se₆ compared to those of C₂₀TMPDAPt_Ge₂Se₆/ C₂₀TMEDAPt_Ge₂Se₆.

In addition to changing the surfactant chain length, the effective tuning of the band gap can also be achieved by

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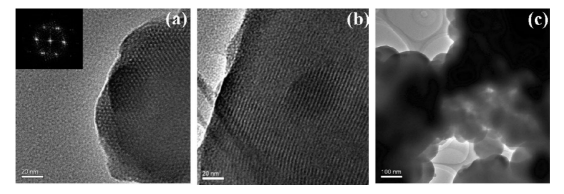


Figure 7. TEM images of C_{22} TMEDAPt_ Ge_2Se_6 (a) down the pore tunnel axis and corresponding Fourier transform (inset), (b) perpendicular to the pore tunnels, and (c) showing spherical particles of \sim 100 nm size.

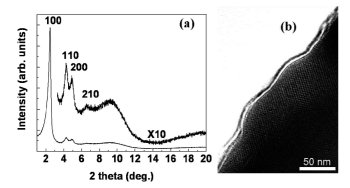


Figure 8. (a) Low angle X-ray powder diffraction patterns of C_{16} PyPt_Ge₂Se₆ prepared in water and (b) its representative TEM image.

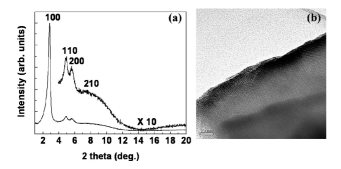


Figure 9. (a) Low angle X-ray powder diffraction patterns of C_{14} PyPt_GeSe₄ and (b) its representative TEM image.

changing the building unit in the precursor solution. Starting with [GeSe₄]⁴⁻ unit, material with lower band gap and well ordered structure was observed in C₁₄PyPt_GeSe₄. Differences in band gap energies (1.31 vs 1.78 eV) for C₁₆PyPt_Ge₂Se₆ prepared in FM and water were also observed and they might also arise from higher fraction of [Ge₄Se₁₀]⁴⁻ incorporation in water (consistent with the NMR spectra discussed in the following section). The band gap energies obtained in this study lie in the same range as several important semiconductors (e.g., InP, GaAs, CdSe, CdTe etc.) suggesting possible interest in optoelectronic investigations.

Solution Behavior of [Ge₂Se₆]⁴⁻. A study of the solution behavior of the precursor chalcogenido ions can answer several important questions: does the precursor building block remain intact in solution? What happens with the species during mesophase synthesis at 75 °C? What is the fate of the dissolved species with the addition of a Lewis acid (i.e., linking metal)?

Table 1. Observed Bragg Reflections and Indexing for Mesostructured Platinum Germanium Selenides

mesophase	d ₁₀₀ , Å	d ₁₁₀ , Å	<i>d</i> ₂₀₀ , Å	<i>d</i> ₂₁₀ , Å	hexagonal unit cell constant, Å ^a
C ₁₄ PyPt_Ge ₂ Se ₆	29.1	16.8	14.6		33.7
C ₁₆ PyPt_Ge ₂ Se ₆	35.0	20.2	17.5	13.2	40.4
$C_{16}PyPt_Ge_2Se_6^b$	35.7	20.7	17.8	13.5	41.3
C ₁₈ PyPt_Ge ₂ Se ₆	38.7	22.3	19.4	14.6	44.6
C ₂₀ PyPt_Ge ₂ Se ₆	39.4		19.6		lamellar
C ₂₀ TMPDAPt_Ge ₂ Se ₆	39.1	22.9	19.7		45.4
C ₂₂ TMPDAPt_Ge ₂ Se ₆	42.9	24.6	21.5		49.5
C ₂₀ TMEDAPt_Ge ₂ Se ₆	36.2	21.1	18.4		42.1
C ₂₂ TMEDAPt_Ge ₂ Se ₆	41.6	23.8	20.7		47.8
C ₁₄ PyPt_GeSe ₄	30.9	17.8	15.4	11.6	35.6

 $[^]a$ Estimated standard deviation \pm 0.1 Å. b Sample prepared in water. All others prepared in FM.

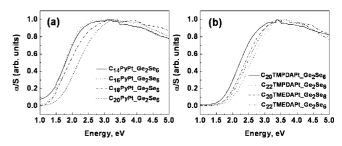


Figure 10. Solid-state UV—vis absorption spectra of (a) $C_nPyPt_Ge_2Se_6$ and (b) $C_nTMPDAPt_Ge_2Se_6$ and $C_nTMEDAPt_Ge_2Se_6$ materials (all prepared in FM).

Answers to these issues may help in understanding the composition and origin of pore order in these materials.

Although, some multinuclear NMR spectroscopy studies have been reported with $[SnSe_4]^{4-}$, $[SnTe_4]^{4-}$, $[Sn_2S_6]^{4-}$, $[SnS_4]^{4-}$, similar studies with the germanium analogs are few. 25,28,47 Earlier literature on the Ge/S system in water by Krebs et al. pointed out the presence of multiple species. 48 To our best knowledge the present work includes the first solution studies ever reported on the $[Ge_xSe_y]^{z-}$ system.

The 77 Se NMR spectrum of K₄Ge₂Se₆ in FM at 25 °C contained several signals among which those at \sim 620 and \sim 265 ppm are assigned to respective bridging and terminal selenium atoms of the [Ge₂Se₆]⁴ anion, see Figure 11a.⁴⁹ The remaining peaks however belong to different species. The tiny peaks at

⁽⁴⁷⁾ Melullis, M.; Dehnen, S. Z. Anorg. Allg. Chem. 2007, 633, 2159– 2167.

⁽⁴⁸⁾ Krebs, B. Angew. Chem., Int. Ed. Engl. 1983, 22, 113-134.

⁽⁴⁹⁾ The peak positions are temperature and concentration dependent. Varying the temperature and or concentration causes spectral shift.

Table 2. Elemental Analysis, Colors, and Energy Band Gaps for Mesostructured Platinum Germanium Selenides

% C, H, N	Pt/Ge/Se Ratio ^a	energy gap ^b (eV)
24.40:3.52:1.54	1.4:2:6.1	1.24
30.51:4.81:1.82	1.4:2:6.2	1.31
29.26:4.62:1.76	1.3:2:6.3	1.78
36.31:5.75:1.89	1.3:2:6.2	1.41
34.67:5.37:1.70	1.4:2:6.1	1.54
29.34:5.48:2.46	1.7:2:6.2	1.63
31.03:5.76:2.78	1.6:2:6.1	1.97
24.57:4.44:2.29	1.5:2:6.1	1.79
31.89:5.82:2.65	1.6:2:6.2	1.85
28.38:4.44:1.82	1.4:1:3.9	1.27
	24.40:3.52:1.54 30.51:4.81:1.82 29.26:4.62:1.76 36.31:5.75:1.89 34.67:5.37:1.70 29.34:5.48:2.46 31.03:5.76:2.78 24.57:4.44:2.29 31.89:5.82:2.65	24.40:3.52:1.54

^a Based on EDS results. Listed values are an average of five measurements on different areas of powder samples. ^b Determined by optical spectroscopy. ^c Sample prepared in water. All others prepared in FM. Color of all samples: dark brown red.

 ${\sim}157$ and ${\sim}212$ ppm are fingerprints of the respective terminal and bridging selenium atoms of the adamantane $[Ge_4Se_{10}]^{4-}$ anion in FM. 50 In addition, several weaker peaks at ${\sim}129, {\sim}307,$ and ${\sim}340$ ppm are consequently ascribed to respective terminal, Se_t, bridging $\mu_2\text{-Se}_{mb},$ and bridging diselenide Se_{db} atoms of the $[Ge_2Se_7]^{4-}$ anion which is present as a minor impurity. The very weak resonance at ${\sim}27$ ppm is that of the tetrahedral $[GeSe_4]^{4-}$ anion. The coexistence of $[Ge_2Se_6]^{4-}$ and $[Ge_4Se_{10}]^{4-}$ implies that they are in equilibrium (eq 1)

$$2[Ge_{2}Se_{6}]^{4-} \rightleftharpoons [Ge_{4}Se_{10}]^{4-} + 2Se^{2-}$$
 (1)

The $[Ge_2Se_7]^{4-}$ species likely was formed in solution by the reaction of Se^0 (traces of which could come from the starting materials) with the $[Ge_2Se_6]^{4-}$ anion (eq 2) or by the presence of some $K_4Ge_2Se_7$ in the starting material. This is consistent with earlier observations that $[Ge_2S_6]^{4-}$ is always found in equilibrium with $[Ge_2S_7]^{4-}$ and $[Ge_4S_{10}]^{4-}$.

$$[Ge_{2}Se_{6}]^{4-} + Se^{0} \rightleftharpoons [Ge_{2}Se_{7}]^{4-}$$
 (2)

The 77 Se NMR spectrum of $K_4Ge_2Se_6$ at 25 °C in aqueous solution shows a higher proportion of $[Ge_4Se_{10}]^{4-}$ compared to that in FM with no evidence of [GeSe₄]⁴⁻, Figure 11b. This is consistent with the greater Bronsted acidity of water which favors the conversion of the more basic [GeSe₄]⁴⁻ species to [Ge₂Se₆]⁴⁻. This hypothesis is supported by the observed change in relative ratio of [GeSe₄]⁴⁻ to [Ge₄Se₁₀]⁴⁻ in FM after the addition of Ba²⁺ ions as a weak Lewis acid. Upon addition of Ba(NO₃)₂, the resulting NMR spectrum showed the disappearance of [GeSe₄]⁴⁻, and the appearance of a significant higher fraction of [Ge₄Se₁₀]⁴⁻. Conversely, the equilibrium can be reversed with the addition of base in the solution. Indeed, by increasing the pH to $\sim 10-12$ through the addition of KOH we observed no peaks of $[Ge_4Se_{10}]^{4-}$ in the NMR spectrum. At the same time the concentration of [GeSe₄]⁴⁻ increased rapidly as shown in Figure 11c. In FM solution, the equilibrium could also be shifted quantitatively to the [GeSe₄]⁴⁻ anion by adding K_2Se . The impurity $[Ge_2Se_7]^{4-}$, can be converted to $[Ge_2Se_6]^{4-}$ by adding PPh₃ which is a well-known chalcogen scavenger. The phosphine abstracts Se^0 from the Se_2^{2-} group forming Ph₃PSe, Figure 11d.

To confirm the above equilibria, we investigated the system further using two-dimensional NMR techniques. Magnetization transfer experiments can distinguish between species exchanging in solution. ⁴⁰ 2D EXSY is a powerful experiment capable of observing exchange between species. ^{41,42} At 60 °C, when we saturated the peaks of the anion $[Ge_2Se_6]^{4-}$, we observed saturation of the peaks of $[Ge_2Se_7]^{4-}$, confirming rapid exchange between the two. However, we did not observe any saturation of the peaks of $[Ge_4Se_{10}]^{4-}$ at this temperature. It is possible that the exchange is too fast to be observed in this time scale. In 2D EXSY experiments at 60 °C; however, we clearly observed off-diagonal cross peaks between $[Ge_2Se_6]^{4-}$ and $[Ge_2Se_7]^{4-}$ and also between $[Ge_2Se_6]^{4-}$ and $[Ge_4Se_{10}]^{4-}$, see Figure 12. We did not observe off-diagonal cross-peaks between $[Ge_2Se_7]^{4-}$ and $[Ge_4Se_{10}]^{4-}$ indicating no equilibrium between these species.

Variable temperature experiments showed that at 60 °C in FM the major species are $[Ge_2Se_6]^{4-}$ and $[Ge_4Se_{10}]^{4-}$ in almost equal proportion; whereas at 100 °C, the system gains $[Ge_4Se_{10}]^{4-}$ at the expense of $[Ge_2Se_6]^{4-}$. Cooling the solution to 25 °C, caused the characteristic peaks of the species present initially (i.e., $[Ge_2Se_6]^{4-}$, $[Ge_2Se_7]^{4-}$, $[Ge_4Se_{10}]^{4-}$, and $[GeSe_4]^{4-}$) to reappear, indicating complete reversibility and thus equilibria. Similar effects were observed in water as well. The spectroscopic observations described above are consistent with the solution equilibria shown in Scheme 1.

The spectroscopic results provide certain insights regarding mesostructured materials made in FM starting from [Ge₂Se₆]⁴ precursor. The presence of multiple species in solution during the preparation of mesophases and their rapid interconversion with temperature and presence of Lewis acid centers suggest it is unlikely that only a single species is incorporated in a given mesostructure. More probably, multiple species are incorporated simultaneously in the framework structure. Each of those species likely plays a vital role in building and stabilizing the overall mesostructure. A control experiment in which a mixture of selenogermante anions [GeSe₄]⁴⁻ and [Ge₄Se₁₀]⁴⁻ was used in a synthesis reaction (see Experimental Section) showed that the product obtained with both C₁₆Py and C₁₈Py surfactants possessed a hexagonal pore order identical to that observed in the materials described here prepared from single selenogermante anions.

Comparison of $^{119} Sn\ NMR\ spectrum of\ [Sn_2Se_6]^{4-}, [SnSe_4]^{4-}, and\ [SnTe_4]^{4-}$ in FM with the presence of K_2PtCl_4 at least indicates that the cubic mesostructure is possible only when $[M_2Q_6]^{4-}$ is the single building unit present in FM (Supporting Information). 28 Even though $[SnSe_4]^{4-}$ almost completely converts to $[Sn_2Se_6]^{4-}$ in FM, traces of $[SnSe_4]^{4-}$ and additional unassigned species prevent the system from yielding cubic structures; hexagonal mesophases were formed instead. $[SnTe_4]^{4-}$ also behaved similarly and again no cubic structures were observed. The fact that $[Ge_2Se_6]^{4-}/Pt^{2+}$ combination does not afford any cubic structure analogous to the $[Sn_2Se_6]^{4-}/Pt^{2+}$ system is attributed to the participation of the $[Ge_2Se_6]^{4-}$ unit in equilibria producing additional species inappropriate for the cubic motif. For the hexagonal structures, however, the equilibria of $[Ge_2Se_6]^{4-}$ appears to be an advantage helping to achieve good pore order. 51

FAB Mass Spectroscopic Support. FAB mass-spectroscopy of FM solutions of the precursor $K_4Ge_2Se_6$ in positive ionization mode showed peaks at m/z 776 and 814.5 due to $[Ge_2Se_6]^{4-}$ anion, Figure 13. On the basis of isotopic distribution patterns,

^{(50) &}lt;sup>77</sup>Se NMR spectra of (TMA)₄Ge₄Se₁₀ in FM consists of two distinct peaks at 156.6 and 212.7 ppm.

⁽⁵¹⁾ Attempts to prepare mesophases after controlled suppression of [Ge₄Se₁₀]⁴⁻ from solution with the addition of base (KOH) resulted disordered pore arrangement.

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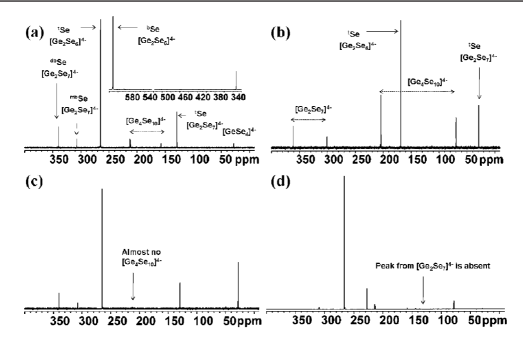


Figure 11. ⁷⁷Se NMR spectra of K₄Ge₂Se₆ dissolved in (a) FM, (b) H₂O, (c) FM and KOH, and (d) FM and PPh₃ (all spectra were taken at 25 °C).

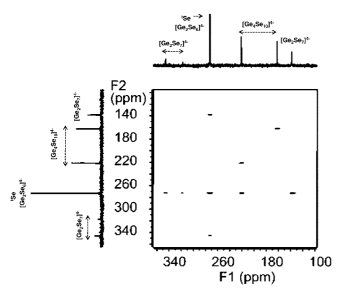
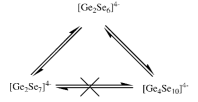


Figure 12. 2D-EXSY 77 Se NMR spectra (at 65 °C) of K_4 Ge $_2$ Se $_6$ dissolved in FM.

Scheme 1. Solution Equilibrium of [Ge₂Se₆]⁴⁻ in FM



these peaks could be assigned to $[H(K_4Ge_2Se_6)]^+$ and $[K(K_4Ge_2Se_6)]^+$, respectively. Other peaks at m/z 855.5, 894, and 1275.5 could be matched with the isotopic distribution patterns of $[H(K_4Ge_2Se_7)]^+$, $[K(K_4Ge_2Se_7)]^+$, and $[K(K_4Ge_4Se_{10})]^+$, respectively. This supports the presence of those multiple species in FM solution.

Raman and Far-IR Spectroscopy of Surf_Pt_Ge₂Se₆. To probe the fate of $[Ge_2Se_6]^{4-}$ in the final mesostructured

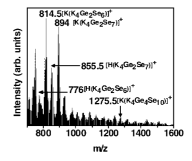
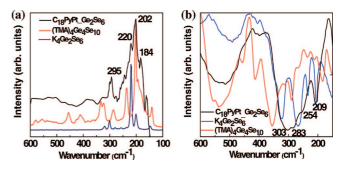


Figure 13. FAB mass spectra of the precursor K₄Ge₂Se₆ in FM. Data was taken in the positive ionization mode.



materials, we compared the Raman spectra with those of $K_4Ge_2Se_6$ and $(TMA)_4Ge_4Se_{10}$, Figure 14a. For $C_{18}PyPt_Ge_2Se_6$, at least four peaks were observed in the $100-600~cm^{-1}$ region. The sharp peak at $202~cm^{-1}$ is due to the combination of totally symmetric v_2 (A_1) breathing mode of the Ge $_4Se_6$ cage from the $[Ge_4Se_{10}]^{4-}$ unit and terminal v_{15} (B_3) Se-Ge-Se bending mode (might be overlapped with the v_6 (B_1) asymmetric Ge-Se stretching mode) from the $[Ge_2Se_6]^{4-}$ unit. $^{52-54}$ The peak at 220

⁽⁵²⁾ Krebs, V. B.; Muller, H. Z. Anorg. Allg. Chem. 1983, 496, 47–57.
(53) Campbell, J.; Devereux, L. A.; Gerken, M.; Mercier, H. P. A.; Pirani, A. M.; Schrobilgen, G. J. Inorg. Chem. 1996, 35, 2945–2962.

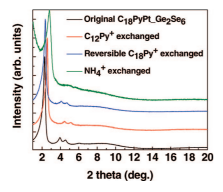


Figure 15. Low angle XRD of ion-exchanged materials. Reversible ion-exchange property of mesostructured $C_{18}PyPt_Ge_2Se_6$ is demonstrated by the shift of Bragg diffraction peaks.

cm⁻¹ resembles the symmetric v_2 (A₁) ring mode of the Ge₂Se₂ core in [Ge₂Se₆]⁴⁻. The less intense peak centered around 295 cm⁻¹ could be from the symmetric v_I (A₁) and asymmetric v_{II} (B₂) terminal Ge-Se stretching modes. These were originally at 301 and 320 cm⁻¹ respectively in K₄Ge₂Se₆; however, they shift slightly to lower frequency due to binding with the linking Pt²⁺ ion.

Comparison of vibrational frequencies of C₁₈PyPt_Ge₂Se₆ with those of the K₄Ge₂Se₆ precursor and (TMA)₄Ge₄Se₁₀ in the far-infrared (IR) region clearly indicates the mesostructured material contains both [Ge₂Se₆]⁴⁻ and [Ge₄Se₁₀]⁴⁻ units, Figure 14b. Because of the presence of different bonding environments in the nonperiodic inorganic framework, the spectra are much broader compared to the free clusters, however, characteristic peaks from both $[Ge_2Se_6]^{4-}$ and $[Ge_4Se_{10}]^{4-}$ are evident. The slight shifts of the frequencies to lower energy are attributed to binding with the metal ions. The sharp peak at 209 cm⁻¹ likely arises from the totally symmetric v_2 (A₁) ring mode from a Ge_2Se_2 fragment since it highly resembles that in $[Ge_2Se_6]^{4-.52-54}$ It could be also possible that this comes from the spectral shift of the v_6 (B₁) vibration of [Ge₂Se₆]⁴⁻. The v_{11} (T₂) and v_{12} (T₂) vibrational modes at 354 and 290 cm⁻¹ wave numbers of $[Ge_4Se_{10}]^{4-}$ unit are shifted to 324 and 283 cm⁻¹ respectively. Similarly, peaks from [Ge₂Se₆]⁴⁻ at 322 and 267 cm⁻¹ due to respective v_{11} (B₂) and v_1 (A₁) vibrations give respective signals at 303 and 254 cm⁻¹ in the mesostructured material. Because the mesophase lacks long-range symmetry, clustering of these vibration modes causes a broad spectral hump.

Ion-Exchange Properties and Porosity. The pore accessibility of the chalcogenide materials was investigated by studying their ion-exchange properties with other cationic species. The powder diffraction data showed that the main Bragg diffraction peak of C₁₈PyPt_Ge₂Se₆ is clearly shifted from 38.7 to 34.2 Å after exchange with C₁₂PyBr, Figure 15. The corresponding hexagonal unit cell parameter was reduced from 44.6(1) to 39.6(1) Å. Pyrolysis mass spectroscopy of the C₁₂Py⁺ exchanged material showed a large peak at m/z = 248.3 (C₁₂H₂₅Se) and only a tiny one at m/z = 332, confirming the presence of mainly $C_{12}Py^+$ in the pores. Interestingly, the ion-exchanged material can expand its unit cell again after reverse exchange with C₁₈Py⁺. The reverse exchanged material caused the main Bragg diffraction peak to shift from 34.2(1) Å to 37.4(1) Å corresponding to a unit cell expansion to 43.3(1) Å. The long-range pore order was maintained in C_nPyPt_Ge₂Se₆ after reverse ion-exchange

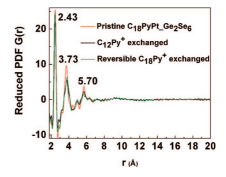


Figure 16. Reduced atomic pair distribution function G(r) of mesostructured C_{18} PyPt_ Ge_2 Se $_6$ and its ion-exchanged materials as a function of interatomic distance r.

as evidenced by the three sharp Bragg peaks. This reversible cell contraction and expansion, known as "breathing" behavior, is a rare property in framework materials and has been previously observed in mesostructured *c*-C_nPyPtSnSe and *c*-/*h*-C_nPyInSbSe systems.^{27,30} The "breathing" behavior of these mesostructured materials is remarkable and appears to be a general property of chalcogenide mesostructured materials. Silica and other oxide mesostructured materials are rigid in this regard.

Attempts to exchange the surfactant inside the pore with very small cationic species such as $\mathrm{NH_4}^+$ resulted in even larger unit cell contraction to 36.3(1) Å and the long-range pore ordering deteriorated. Elemental analysis indicated that almost all surfactant molecules were replaced with $\mathrm{NH_4}^+$. Again $(\mathrm{NH_4})\mathrm{Pt_Ge_2Se_6}$ did not show open porosity because of the contraction of the flexible chalcogenide framework.

Pair Distribution Function (PDF) Analysis. To probe the local structures of the nonperiodic mesostructured framework, we used the PDF technique which analyses both diffuse and Bragg scattering.⁵⁵ This allows to determine the interatomic distances between various neighbors in the structure and thus gain local structure information about the building units present in the structure. Figure 16 shows PDF plots as a function of interatomic distance for pristine and ion-exchanged C₁₈PyPt_Ge₂Se₆ material. The PDF of pristine C₁₈PyPt_Ge₂Se₆ material showed up to three sharp correlation vectors. The first correlation vector at \sim 2.43 Å arises from the first neighbor Ge-Se and Pt-Se bond distances. The peak at \sim 3.73 Å could be from the second neighbor Ge-Ge or Se-Se or Pt-Ge distances. The lack of well defined features in the PDF beyond ~6 Å suggests the local order is limited in a volume of material with \sim 6 Å diameter. Beyond this distance the randomness of the inorganic network predominates. The material exchanged with $C_{12}Py^+$ and then reverse ion-exchanged with C₁₈Py⁺ displays similar PDF correlations at the same distances as observed in the pristine C₁₈PyPt_Ge₂Se₆ material indicating that the local inorganic framework remains intact during the ion-exchange process. This suggests that the observed flexibility of the Pt/Ge/Se network that allows it to expand and contract in response to ion-exchange is due to changes in bond angles and dihedral angles and not to changes in bonding.

Stability. Investigations of the thermal stability of C₁₆PyPt_Ge₂Se₆, C₂₀TMPDAPt_Ge₂Se₆, C₂₀TMEDAPt_Ge₂Se₆ and C₁₄PyPt_GeSe₄ were carried out with TGA and pyrolysis MS-experiments. The TGA curves in Figure 17 indicate that

⁽⁵⁴⁾ Campbell, J.; Diciommo, D. P.; Mercier, H. P. A.; Pirani, A. M.; Schrobilgen, G. J.; Willuhn, M. *Inorg. Chem.* 1995, 34, 6265–6272.

⁽⁵⁵⁾ Billinge, S. J. L.; Kanatzidis, M. G. Chem. Commun. 2004, 7, 749–760.

Importance of Solution Equilibria ARTICLES

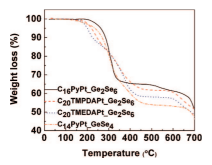


Figure 17. Typical TGA curves of C_{16} PyPt_Ge₂Se₆, C_{20} TMPDAPt_Ge₂Se₆, C_{20} TMEDAPt_Ge₂Se₆ and C_{14} PyPt_GeSe₄ at 20 °C per minute heating rate under nitrogen flow.

all materials undergo a two-step (at least) weight loss between 160 and 470 °C and it is due to surfactant decomposition. Common trends were observed for each surfactant class. The pyridyl headgroup and $C_nH_{2n+1}Se^+$ are typical decomposition species for all C_nPy mesostructured chalcogenides. The $C_nTMPDABr$ and $C_nTMEDABr$ templated compounds show different decomposition products during thermal treatment.

For C_nTMPDAPt_Ge₂Se₆ between 160 and 470 °C, the major decomposition products evolved were N(CH₃)₃, N(CH₃)₂-CH₂CH₂CH₃, N(CH₃)₂CH₂CH₃ and C_nH_{2n+1}N(CH₃)₂. In the case of C_nTMEDAPt_Ge₂Se₆, the inorganic framework also started decomposing in this temperature region eliminating fragments of R-Se⁺ along with N(CH₃)₃, N(CH₃)₂CH₂CH₃ and C_nH_{2n+1}N(CH₃)₂. The difference in thermal behavior between the two surfactant types can be rationalized in terms of the different degree of interaction with the inorganic framework. Above 570 °C rapid collapse of the inorganic structures occurs with the fragments of (CH₃)₂(CH₂CH₃)N-Se⁺ and N(CH₃)₂+ being the main detectable species. At 750 °C, a mixture of amorphous and partly crystalline PtGeSe phase was detected.⁵⁶

Small Angle X-ray Scattering. Because of the presence of surfactants in the pore of C_nPyPt_Ge₂Se₆ direct gas absorptions measurements could not be made to determine the surface area. In such cases small-angle X-ray scattering (SAXS) is a powerful technique that can be used to obtain the underlying surface area of the inorganic framework. It is important to note that SAXS is sensitive to differences in electron density at the wall-pore interface and for this reason it can be used to determine the area of the surfactant/Pt—Ge—Se interface by analyzing the Porod region. The latter can give information about both closed and open porosity, thereby giving a good estimation of the potential free space and surface area in the system. ^{57–62}

The SAXS curve usually displays fractal behavior as expressed by the intensity (I(q)) vs scattering vector, q plot according to the equation, $I(q) = k.q^{-\alpha}$, where $q = 4\pi \sin \theta/\lambda$, θ is the scattering angle, λ is the wavelength, k is a constant, α

(56) Abrahams, S.; Bernstein, J. L. Acta Cryst. B 1977, 33, 301.

is related to the fractal dimension. It follows Porod's law at a large value of the q vector and a limiting behavior is observed: $\lim_{q\to\infty} I(q) = K/q^3$ (K is the Porod constant). The specific surface area ($S_{\rm SAXS}$) is then determined using the method of the invariant (where b is the background intensity)

$$Q = \int_{q_{\min}}^{q_{\max}} (I(q) - b)q^2 dq + \frac{K}{q_{\max}}$$
 (3)

$$S = \pi \varphi (1 - \varphi) \left(\frac{K}{\rho Q} \right) = \pi \varphi \left(\frac{K}{\rho_s Q} \right) \tag{4}$$

where the Porod constant $K = \lim_{q \to \infty} \{I(q)q^3\}$, φ is the volume fraction of framework voids, ρ , ρ_s are the bulk and skeletal mass density respectively (where $\rho/\rho_s = 1 - \varphi$), and Q is the invariant parameter defined as the q^2 weighted integrated intensity I(q) of the scattering curve according to eq

Because almost all surfactant molecules were replaced by NH₄⁺ ions in (NH₄)Pt_Ge₂Se₆, SAXS surface area (S_{SAXS}) analysis was performed on this sample and compared with that of the pristine C₁₈PyPt_Ge₂Se₆. To determine the specific surface area of the pristine C₁₈PyPt_Ge₂Se₆, $I(q)q^3$ vs q^3 was plotted in the Porod region (q of 0.032 to 0.066 Å⁻¹) and a K value of 0.09 was obtained from the intercept (Supporting Information).⁵⁸ The Q value was determined from eq 3 where the upper limit, q_{max} was evaluated by arbitrarily choosing a q value and fitting with $\int_{q_{\text{min}}}^{\infty} I(q) - b)q^2 dq \approx K/q_{\text{max}}$.^{61,62}

Figure 18a shows the intensity I(q) falls gradually for pristine $C_{18}PyPt_Ge_2Se_6$ until q=0.08Å $^{-1}$ followed by the Bragg peak at $q \sim 0.15$ Å $^{-1}$. Based on the measured skeletal density 63 of ~ 5.1 g/cm 3 and bulk density of 1.67 g/cm 3 for the pristine $C_{18}PyPt_Ge_2Se_6$, we obtained 331 m 2 /g surface area (S_{SAXS}) for the inorganic framework. Similar measurement with the ion-exchanged material (NH₄)Pt_Ge₂Se₆, 64 shows 250 m 2 /g surface area (S_{SAXS}). These potential surface areas are very large and correspond to ~ 1688 and ~ 1275 m 2 /g silica equivalent areas, respectively. 65

From the $\ln[I(q)]$ vs $\ln[q]$ plot in Figure 18b in the Porod region, two distinct slopes of 2.38 and 1.27 were obtained, indicating two different mass fractal behaviors. ⁶⁶ Thus though particles are connected in a three-dimensional like pattern, the low mass fractality (<3) reflects the hollow nature of the framework and thereby the enormous surface area of the inorganic framework available in (NH₄)Pt_Ge₂Se₆. Pristine C_{18} PyPt_Ge₂Se₆ showed a mass fractality of 2.90.

Average chord length $(l_{\rm p})^{59}$ of the pores of the ion-exchanged sample as determined by $l_{\rm p}=l/\varphi_{\rm s}$ (where $l=4Q/\pi K$ and $\varphi_{\rm s}$ is the volume fraction of solid part) turns out to be ~ 2.5 nm which is consistent with the pore diameter measured by TEM for the pristine C_{18} PyPt_Ge₂Se₆. The average pore diameter can also

⁽⁵⁷⁾ Ne, F.; Zemb, T. J. Appl. Crystallogr. 2003, 36, 1013-1018.

⁽⁵⁸⁾ Li, Z. H.; Gong, Y. J.; Wu, D.; Sun, Y. H.; Wang, J.; Liu, Y.; Dong, B. Z. Surf. Interface Anal. **2001**, 31, 897–900.

⁽⁵⁹⁾ Dieudonne, P.; Delord, P.; Phalippou, J. J. Non-Cryst. Solids 1998, 225, 220–225.

⁽⁶⁰⁾ Donatti, D. A.; Ruiz, A. I.; Vollet, D. R. J. Non-Cryst. Solids 2001, 292, 44–49.

⁽⁶¹⁾ Ehrburger-Dolle, F.; Fairen-Jimenez, D.; Berthon-Fabry, S.; Achard, P.; Bley, F.; Carrasco-Marin, F.; Djurado, D.; Moreno-Castilla, C.; Morfin, I. Carbon 2005, 43, 3009–3012.

⁽⁶²⁾ Fairen-Jimenez, D.; Carrasco-Marin, F.; Djurado, D.; Francoise, B.; Ehrburger-Dolle, F.; Moreno-Castilla, C. J. Phys. Chem. B 2006, 110, 8681–8688.

⁽⁶³⁾ The skeletal density was measured using helium pycnometry. Here it is assumed that helium penetrates throughout the pores and accounts only the volume of the inorganic wall. However, since helium cannot penetrate through the surfactant occupied pores of pristine C₁₈PyPt_Ge₂Se₆, the bulk density of this material was assumed to be its helium density.

⁽⁶⁴⁾ The bulk and skeletal densities of (NH₄)Pt_Ge₂Se₆ are 2.16 and 5.10 g/cm³ respectively.

⁽⁶⁵⁾ The empirical formulas of the inorganic framework were normalized to two selenium atoms to compare well with SiO₂ for equivalent surface area calculation.

⁽⁶⁶⁾ Vollet, D. R.; Donatti, D. A.; Ruiz, A. I. J. Non-Cryst. Solids 2001, 288, 81–87.

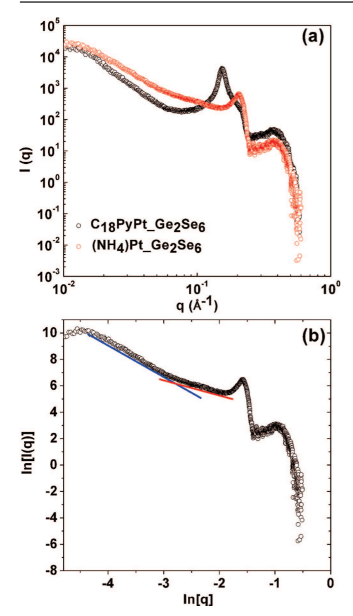


Figure 18. Small angle X-ray scattering (SAXS) data of pristine C_{18} PyPt_Ge₂Se₆ and ion-exchanged material (NH₄)Pt_Ge₂Se₆. (a) Intensity I(q) versus q plot (data presented in the log scale). (b) Porod's plot of ln[q] vs ln[I(q)] for (NH₄)Pt_Ge₂Se₆.

be calculated by plotting $I(q)^{-1/2}$ vs θ^2 (in radians)⁶⁷ and was found to be ~2.6 nm for (NH₄)Pt_Ge₂Se₆ in good agreement with the TEM observations described above. The SAXS results clearly imply that despite the collapse of pore order, the inorganic framework in the ion exchanged material preserves

the characteristic features of the pristine parent sample and retains its enormous surface area.

Concluding Remarks

In the synthesis of mesophases, starting with the anionic $\left[\text{Ge}_2\text{Se}_6\right]^{4-}$ chalcogenide cluster, a single species is not involved; rather multiple species are present and likely participate in the assembly. Even though temperature can favor one species over another, varying the temperature or adding the linking metal ion in the system alters the relative concentration of those species. Surfactant variation influences the composition of the mesophase materials thereby tuning their band gap transitions. This could be an attractive route to changing systematically the electronic functionality of these materials. Depending upon the specific conditions, the mesostructured materials can possess various anionic building blocks and the optoelectronic properties are dictated mainly by relative population of those building units. Thus, mesostructured chalcogenides with variable energy gap and pore symmetry are now available for further manipulation.

Our current and previous work on $Pt^{2+}/[SnSe_4]^{4-}$, $Pt^{2+}/[SnTe_4]^{4-}$, $Pt^{2+}/[Sn_2Se_6]^{4-}$, $Pt^{2+}/[Sn_4Se_{10}]^{4-}$ systems suggests that the adoption of a particular symmetry, for example, wormhole vs cubic vs hexagonal, in mesostructured chalcogenides is largely inherent in the selection of building units. The fact that surfactant size, shape, concentration, solvent, coordination preference of the linking metal ion and kinetics of the metathesis reaction also contribute to the final pore symmetry underscore the great complexity of these directed self-assembly reactions.

Even though surfactant-containing crystalline mesoporous chalcogenides have not been reported to date, the more appreciated role of solution equilibria during directed assembly of these structures could provide useful insights on selecting building units and solvents for controlling pore order and ultimately crystallinity. The demonstrated reversible ion-exchange property of these systems illustrates that optically and electronically active guest species could be incorporated into the pores to create novel organic—inorganic junctions.

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Supporting Information Available: ¹¹⁹Sn NMR spectra of neat (TMA)₄Sn₄Se₁₀, neat K₄Sn₂Se₆ and K₄Sn₂Se₆ in the presence of K₂PtCl₄ in FM. Determination of Porod's constants for C₁₈PyPt_Ge₂Se₆ and (NH₄)Pt_Ge₂Se₆. This material is available free of charge via the Internet at http://pubs.acs.org.

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⁽⁶⁷⁾ Longman, G. W.; Wignall, G. D.; Hemming, M.; Dawkins, J. V. Colloid Polym. Sci. 1974, 252, 298–305.