Engineering New Layered Solids from Exfoliated Inorganics: a Periodically Alternating Hydrotalcite – Montmorillonite Layered Hybrid

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Two-dimensional (2D) nanosheets obtained by exfoliating inorganic layered crystals have emerged as a new class of materials with unique attributes. One of the critical challenges is to develop robust and versatile methods for creating new nanostructures from these 2D-nanosheets. Here we report the delamination of layered materials that belong to two different classes - the cationic clay, montmorillonite, and the anionic clay, hydrotalcite - by intercalation of appropriate ionic surfactants followed by dispersion in a non-polar solvent. The solids are delaminated to single layers of atomic thickness with the ionic surfactants remaining tethered to the inorganic and consequently the nanosheets are electrically neutral. We then show that when dispersions of the two solids are mixed the exfoliated sheets self-assemble as a new layered solid with periodically alternating hydrotalcite and montmorillonite layers. The procedure outlined here is easily extended to other layered solids for creating new superstructures from 2D-nanosheets by self-assembly.

One of the main challenges in the design of materials is the self-directed assembly of ordered superstructures from basic structural motifs. In the organic solid state and, to a lesser extent, in framework solids the concept of crystal engineering using synthetic building blocks has emerged as a successful concept for rational design. In most other crystalline solids the ability to de-construct a solid into basic structural blocks and re-construct new solids from them is yet to be realized. These ideas are most likely to succeed in the case of layered solids where deconstruction, in this case exfoliation or delamination, procedures are reasonably well established. Here we report the delamination of layered materials that belong to two different classes – the cationic clay, montmorillonite, and the anionic clay, hydrotalcite - to their ultimate constituent, intact, nanometer thick, single layers. We then show that these exfoliated sheets can be made to self-assemble as a new solid with periodically alternating hydrotalcite and montmorillonite layers.

Delamination of inorganic layered solids by exfoliation resulting in the formation of highly dispersed monolayer colloidal phases has attracted considerable interest, particularly as building blocks for the construction of functional nano-composites and nanostructures. The spontaneous swelling and exfoliation of smectite clays in water to give stable colloidal dispersions is well documented. Delamination has been artificially realized for a variety of different layered materials, layered oxides, di-chalcogenides, layered double hydroxides, metal-organic framework solids and more recently carbides and carbonitrides, by controlling interlayer interactions. The term 'nanosheets' has been widely used to classify these nanometer thick inorganic sheets whose lateral dimensions can extend to microns. Most reported methods have achieved exfoliation by use of highly polar solvents. The solvent molecules replace the solvation shell of the interlamellar counter ions resulting in the dilation of the layers and subsequent modification of interlayer interactions. These methods result in the formation of either positively or negatively charged nanosheets with solvent polarity playing a crucial role in ensuring that the inorganic macro-ions remain as stable dispersions. On solvent evaporation the original solid can be recovered. There have been attempts to create new layered hybrids from a combination of dispersions of two different layered materials. Most approaches have used the “layer-by-layer” method, a thin-film fabrication technique that uses sequential adsorption of oppositely charged species on a suitable substrate. New functional nanoarchitectures formed by the electrostatic layer-by-layer assembly either of negatively charged montmorillonite layers and polycations or positively charged layered double hydroxide nanosheets in combination with polyanionic counterparts (polymer, DNA, metal oxide, metal dichalcogenide...
The spontaneous flocculation of positively charged layered double hydroxide nanosheets with the complimentary, negatively charged oxide or metal dichalcogenide layers have been reported. By judicious choice of the individual components, charge balanced ordered nanostructures were obtained. What is clearly desirable is a procedure that allows for the spontaneous self-assembly of new ordered layered structures from combinations of dispersions of exfoliated inorganic nanosheets.

The layered materials chosen for this study are two ionic lamellar solids, the cationic clay montmorillonite and the anionic clay hydrotalcite. The smectite clay, montmorillonite, is a 2:1 layered silicate; two tetrahedral sheets containing Si\textsuperscript{4+} sandwiching a sheet of octahedrally coordinated Al\textsuperscript{3+} ions. Part of the Al\textsuperscript{3+} ions in the octahedra are substituted by Mg\textsuperscript{2+} and the charge deficit of the negatively charged layers compensated by interlamellar cations that are usually hydrated. The anionic clays or layered double hydroxides (LDH) are derived from the brucite (Mg(OH)\textsubscript{2}) structure by the isomorphous substitution of M\textsuperscript{2+} ions by M\textsuperscript{3+} ions. The LDH layers are positively charged and charge neutrality is realized by the presence of interlamellar anions. When M\textsuperscript{3+} is Al\textsuperscript{3+} the mineral hydrotalcite is obtained. It may be argued, that if the charges on the cationic montmorillonite and anionic hydrotalcite are tuned, by suitable substitutions, to be identical then their exfoliated sheets could, in principle, self-assemble in an alternating fashion to give an electrically neutral crystalline hybrid. Such a scenario is, however, most unlikely for although the total charge on these electrically insulating sheets can be made identical, there is no way to control the distribution of the positive charges on the montmorillonite sheets or the negative charges on the hydrotalcite sheets as the substitutions that create these charge on the layers are random. Here we adopt a different approach to circumvent this problem. We first prepare neutral exfoliated nanosheets of montmorillonite and hydrotalcite separately and show that they self-assemble as a periodically alternating layered structure.

Recently a simple approach to exfoliate layered double hydroxide solids to their ultimate constituent, intact single sheets of nanometer thickness and micron size in non-polar solvents like toluene had been outlined. The method involved intercalation of an anionic surfactant to form a hydrophobic anchored surfactant bilayer in the interlamellar space of the LDH. Delamination was effected by simply stirring the surfactant intercalated layered solid in the solvent. The method was rapid but at the same time gentle enough to produce exfoliated nanosheets of regular morphology. A distinctive feature of the procedure was that the exfoliated nanosheets are electrically neutral as the ionic surfactant chains remain tethered to the inorganic sheets. It was shown that delamination occurs because of the inclusion of nonpolar solvent molecules in the interlayer space. These solvent molecules weaken the attractive dispersive interactions between the chains tethered to opposing inorganic layers thereby facilitating separation of the layers. The method is general, and should, in principle, be applicable to other layered solids that have been intercalated with appropriate long chain surfactants that render their interlamellar space hydrophobic.

**Results**

The anionic clay Mg\textsubscript{0.66}Al\textsubscript{0.33}(OH)\textsubscript{2} (NO\textsubscript{3})\textsubscript{0.33} (Mg-Al LDH) with NO\textsubscript{3} as the interlamellar anion was prepared by reported procedures. Na-montmorillonite (Na-MMT) was obtained from Southern Clay Product Inc., USA (Na-Cloisite) and has the formula Al\textsubscript{1.49}Mg\textsubscript{0.24}Fe\textsubscript{0.27}Si\textsubscript{10}O\textsubscript{31} (OH\textsubscript{1}) (Na\textsubscript{0.31}). The Co-Al LDH (Co\textsubscript{0.62}Al\textsubscript{0.38}(OH)\textsubscript{2}(NO\textsubscript{3})\textsubscript{0.33}) was also prepared to help in establishing the stoichiometry of the hybrid and to demonstrate that the procedures, described subsequently, are universal and can be applied to other anionic clays. Procedures for the intercalation of ionic surfactants in these materials as well as their characterization by powder X-ray Diffraction (XRD) are well established.

Intercalation occurs with an increase in the interlayer separation that depends on the chain length of the surfactant. The cationic surfactant intercalated in the smectite clay, montmorillonite, was the di-octadecyl-dimethyl ammonium (DODMA) ion while in the LDH it was the dodecyl sulfate (DDS) anion. The X-ray determined interlayer spacings of the intercalated MMT-DODMA and Mg-Al LDH-DDS are 3.91 nm and 2.81 nm respectively. This inter-layer spacing is characteristic of a bilayer arrangement of the intercalated surfactant chains with the ionic head group anchored to the inorganic sheets and with no interdigitation of the surfactant alkyl chains.

Delamination of the surfactant layered solids was effected by dispersing 10.0 to 50.0 mg of the solid in 100 ml of chloroform or any other non-polar organic solvent followed by either stirring or sonication for a maximum of 10 minutes. The observed Tyndall light scattering indicating the presence of the exfoliated inorganic nanosheets dispersed in the organic solvent (Fig. 1).

Dispersions of the nanosheets with the surfactant chains anchored to the inorganic sheets were characterized by proton NMR spectroscopy. The NMR spectra of dilute solutions of the nanosheets dispersed in deuterated chloroform show well-resolved resonances of the anchored surfactant chains (Fig. 1). The spectrum of the surfactant along with the assignment of the resonances is also shown. For the dodecyl sulfate ions tethered to the Mg-Al LDH sheets the \( \alpha \)-CH\textsubscript{2}, \( \beta \)-CH\textsubscript{2}, the CH\textsubscript{2}-chain as well as the terminal \( \omega \)-CH\textsubscript{3} resonances appear at the same position as that of the free Na-DDS chains in chloroform solution. The same is true for the cationic DODMA chains that are anchored to the montmorillonite sheets; most resonances appear at the same position as that of free DODMA surfactant chain (Fig. 1). A significant difference, however, between the anchored and free DODMA chains is in the position of the \( \alpha \)-CH\textsubscript{3} and the methyl head-group resonances. For the anchored DODMA the \( \alpha \)-CH\textsubscript{3} resonance is up-field shifted and the head-group CH\textsubscript{3} resonance absent. This is a consequence of the proximal charge of the layers and the restricted rotational freedom of the head-group CH\textsubscript{3} units as consequence of the anchoring of the DODMA chains to the inorganic sheets. The NMR spectra provides evidence that the schematic cartoon of the surfactant tethered delaminated inorganic sheets as depicted in Figure 1 is essentially correct. Further support that the nanosheets are neutral was provided by zeta potential measurements. The zeta potential values of Mg-Al LDH-DDS and MMT-DODMA dispersions are ~ -2.3 mV and 5.6 mV, respectively. The results highlight one of the main differences of the present delamination strategy as compared to most reported methods that yield nanosheets of either positive or negatively polarity, depending on the nature of the solid being delaminated. In the procedure outlined here, however, the dispersed nanosheets are electrically neutral as the ionic surfactant chains remain tethered to the layers even after exfoliation.

The morphology and size of the exfoliated nanosheets were examined by TEM, SAED and tapping mode AFM. Representative TEM images of the delaminated nanosheets are shown in Figure 2. The TEM images show two-dimensional ultra-thin sheets of micron size. A distinctive feature of the images is that the sheets are characterized by a well-defined morphology. The Mg-Al LDH, for example, shows hexagonal morphology. The selected area electron diffraction pattern of individual sheets exhibits the characteristic diffraction patterns with six-fold symmetry establishing their single crystal nature. The conservation of lateral morphology of the nanosheets is the distinctive feature of the delamination process described here. In contrast, most reported exfoliation procedures result in fractured sheets with poorly defined morphologies. The tapping mode AFM images shows two- dimensional ultra-thin sheets with lateral dimension up to 200 nm. The height profile reveals that the sheets have a fairly flat, smooth terrace with an average thickness that corresponds to that of a single inorganic layer. For the layered double hydroxides, Mg-Al LDH, the thickness of the sheets in the tapping mode AFM images is typically 0.5 nm. The thickness of the hydrotalcite sheet is
0.48 nm. The softer surfactant layer is not seen in the tapping-mode AFM image of the Mg-Al LDH-DDS because of the stiff cantilever used for recording the images. The typical thickness of the MMT-DODMA sheets in the AFM images is 1.02 nm. The thickness of a montmorillonite sheet is 0.98 nm. It may be pointed out that the observation of single sheets in the AFM images is a consequence of the fact that the delaminated sheets are electrically neutral.

We found that when the chloroform dispersions of the surfactant hydrotalcite and montmorillonite were mixed and the solvent allowed to evaporate, the X-ray diffraction pattern of the resulting solid showed additional reflections different from that of either the surfactant intercalated hydrotalcite or montmorillonite. Figure 3a shows the X-ray diffraction patterns of the resultant solid obtained on mixing different molar ratios of chloroform dispersions of Mg-Al LDH-DDS and MMT-DODMA and allowing the solvent to evaporate. As mentioned earlier, on solvent evaporation from the single component dispersion the original solid is recovered. The pure Mg-Al LDH-DDS dispersion gives a set of seven 00l Bragg reflection on solvent evaporation corresponding to an interlayer spacing of 2.81 nm (Fig. 3c(i)). This interlayer spacing is identical to that of the starting Mg-Al LDH-DDS (Supplementary Fig. S1) that had been shown to correspond to an anchored bilayer arrangement with the intercalated surfactant chains tilted away from the interlayer normal and adopting an all-trans conformation41. For the pure MMT-DODMA dispersion a set of 12 00l Bragg reflection are observed (Fig. 3c(ii)), on solvent evaporation, corresponding to an interlayer spacing of 3.91 nm, identical to the starting layered solid (Supplementary Fig. S3). It may be seen, however, that on addition of montmorillonite to the Mg-Al LDH dispersion a new set of reflections (shaded peaks in Fig. 3a) are observed that do not correspond to either that of MMT-DODMA or Mg-Al LDH-DDS. At a molar ratio of 1:1, peaks corresponding to neither the surfactant intercalated Mg-Al LDH nor montmorillonite are observed and only the new set of reflections are observed. The stoichiometry of this phase, Mg$_{0.39}$Al$_{0.54}$Fe$_{0.07}$Si$_{0.02}$O$_{2.50}$($\text{OH})_{1.50}$(DDS)$_{0.17}$(DODMA)$_{0.12}$.0.75H$_2$O, corresponds to a 1:1 molar composition of Mg-Al LDH-DDS and MMT-DODMA. For this composition the Bragg reflections may be indexed as 00l reflections corresponding to an interlayer spacing of 5.85 nm. A plot of the d-spacing versus 1/l is linear confirming a unique interlayer spacing (Fig. 3b). The unique interlayer spacing rules out the occurrence of a randomly interstratified layered structure and indicates a structure that is periodically ordered along the interlayer normal.

The results suggest that the 1:1 hybrid has an alternating Mg-Al LDH-DDS – MMT-DODMA layered structure. We found that the X-ray diffraction patterns generated for a structure in which hydrotalcite and montmorillonite sheets alternate in a periodic fashion with an interlayer repeat of 5.85 nm are similar to the experimentally recorded diffraction pattern of the 1:1 hybrid (Fig. 3c (iii)) (Supplementary Fig. S8). It may be noted that the intercalated surfactant chains in both Mg-Al LDH-DDS and MMT–DODMA are randomly anchored and conformationally disordered and therefore do not contribute to Bragg reflections in the experimental diffraction pattern. For the calculated diffraction patterns surfactant chains are not included in the structure and also lateral registry between alternating sheets is absent. If the anchored surfactant bilayers were to adopt a similar structure as in their parent Mg-Al LDH-DDS and MMT-DODMA compounds an interlayer spacing of 6.72 nm may be anticipated for the 1:1 alternating hybrid. The fact that the spacing is 5.85 nm suggests that the surfactant bilayers adopt a different structure in which the DDS and DODMA chains in opposing bilayers are interdigitated. The X-ray diffraction patterns of the solid recovered after solvent evaporation from mixtures of dispersions of Co-Al LDH -DDS and MMT-DODMA show a similar behaviour (Supplementary Figure S5). At a 1:1 molar ratio of Co-Al LDH-DDS and MMT-DODMA the X-ray diffraction shows a set Bragg reflections with an unique interlayer spacing of 5.81 nm; the interlayer spacing of Co-Al LDH- DDS is 2.62 nm (Supplementary Fig. S2) and that of MMT-DODMA is 3.91 nm (Supplementary Fig. S3).

The TEM images of the hybrid provide additional evidence of a layered structure that is periodically ordered along the interlayer normal. The cross-sectional TEM of the starting layered solids and the 1:1 hybrid are shown in Figure 4 along with an expanded Fourier filtered image. The HRTEM image of the anionic clay Mg-Al LDH-DDS is shown in Figure 4a. The FFT generated diffraction pattern is shown in the inset. The absence of diffraction spots with hexagonal symmetry indicating that the cross-section of the sample is indeed in the electron beam direction. The spacing measured between the two lattice fringes, 2.6 nm, is in good agreement with the spacing determined from XRD measurements. The cross-sectional HRTEM of the
cationic clay, MMT-DODMA, is shown in Figure 4b. The spacing measured between the fringes matches with the \( d \)-spacing determined from the XRD pattern, 3.91 nm. The electron diffraction pattern of the 1:1 hybrid shows diffraction rings of both Mg-Al LDH-DDS and MMT–DODMA (Supplementary Fig. S7). What is interesting, however, is the cross-sectional TEM of the 1:1 hybrid (Fig. 4c) that shows a structure that is periodically ordered along the interlayer normal. The interlayer spacing measured from the TEM image of Figure 4 5.7 nm is comparable with the interlayer spacing 5.85 nm, determined from the X-ray diffraction pattern. The distance between the spots of the FFT generated diffraction pattern (inset of Fig. 2c) is 2.9 nm corresponding to 002 reflection of the hybrid structure. Contrast does not permit us to distinguish between the individual hydrotalcite and montmorillonite sheets, but an Energy-dispersive X-ray spectroscopy (EDS) elemental analysis of individual layers of the corresponding 1:1 Co-Al LDH-DDS-MMT-DODMA hybrid reveals alternation in composition (Supplementary Fig. S6) indicating that the two types of sheets alternate.

We have also attempted to address the question as to why the periodically ordered 1:1 alternating hybrid is obtained on solvent evaporation from a mixture of the Mg-Al LDH-DDS and MMT–DODMA
dispersions, rather than a mixture of the individual solids. We believe that hybrid formation is a consequence of favourable electrostatic interactions between the ‘tails’ of the DDS anionic chains anchored to the positively charged hydrotalcite sheets and the cationic DODMA surfactant chains tethered to the negatively charged montmorillonite sheets. It may be noted that although both the Mg-Al LDH-DDS and MMT–DODMA nanosheets are electrically neutral, because the surfactant chains remain anchored even on delamination, the disposition of charges in the two nanosheets are opposite.

We believe that it is the residual charges, of opposite sign, located on the ‘tails’ of the DDS and DODMA alkyl chains that is responsible for hybrid formation. To test this hypothesis we took an equimolar dispersion of Mg-Al LDH-DDS and MMT–DODMA in chloroform and increased the polarity of the solvent by incremental addition of methanol. On addition of methanol the material precipitates out of the dispersion. The X-ray diffraction pattern of the precipitate shows Bragg reflections corresponding to both Mg-Al LDH-DDS and MMT–DODMA indicating that the product of precipitation is indeed a mixture of the parent compounds (Fig. 5). These observations support the role of electrostatic forces in favouring the self-assembly of the periodically ordered alternating 1:1 hydrotalcite-montmorillonite hybrid.

Discussion

In summary we have shown that the surfactant intercalated hydrotalcite and montmorillonite layered solids are easily delaminated in non-polar solvents to give stable dispersions. The solids are delaminated to their ultimate constituent, single layers of atomic thickness with the ionic surfactants remaining tethered to the inorganic sheets and consequently the nanosheets are electrically neutral. We show
here that when dispersions of the two solids are mixed and the solvent allowed to evaporate a new solid is obtained in which the hydrotalcite and montmorillonite sheets alternate in a periodic fashion with a repeat length different from that of the starting surfactant intercalated hydrotalcite and montmorillonite (Fig. 6). The procedure outlined here has the advantage over thin-film techniques like the layer-by-layer method for the design of new layered hybrids as these methods require precise control and manipulation to achieve identical atomic-level thickness of the individually deposited layers, if truly periodic superstructures are to be realized. In our method attractive forces between the neutral exfoliated nanosheets of montmorillonite and hydrotalcite ensures self-assembly of a perfectly periodically alternating layered structure. The procedure outlined here may easily be extended to other layered solids for creating new superstructures from 2D exfoliated inorganic nanosheets by self-assembly. Additionally, we believe that the significance of the present work is that it illustrates the application of the concept of crystal engineering using structural building blocks to the inorganic solid state, in much the same way as what has been achieved in the organic solid state.

Methods

Materials. Mg(NO$_3$)$_2$·6H$_2$O, Al(NO$_3$)$_3$·9H$_2$O, Co(NO$_3$)$_2$·6H$_2$O, ammonia solution (25%), chloroform, methanol (SD Fine Chemicals), sodium dodecyl sulfate (Na-DDS), and dioctadecyldimethylammonium bromide (DODMA-Br) (Sigma-Aldrich) were used as received without further purification.

Preparation of Mg-Al LDH-DDS. Nitrate ion intercalated Mg-Al LDH [Mg-Al LDH-NO$_3$], was used as the starting material for the preparation of the surfactant

Figure 4 | Cross-sectional HRTEM of (a) Mg-Al LDH-DDS (b) MMT-DODMA and (c) the 1:1 Mg-Al LDH-DDS MMT-DODMA hybrid. The FFT generated diffraction pattern and an enlarged view of the Fourier filtered image obtained by an inverse FFT of the diffraction pattern is shown in the panels on the right along with a structural model for each of the solids.

Figure 5 | X-ray diffraction patterns of the solid on precipitation from dispersions of (a) Mg-Al LDH-DDS, (b) MMT-DODMA and (c) 1:1 Mg-Al LDH-DDS: MMT-DODMA in chloroform on increasing the polarity of the medium by addition of methanol.
negatively charged inorganic is montmorillonite and the positively charged layers hydrotalcite. Negative charged inorganic sheets are mixed and on solvent evaporation a periodically alternating layered hybrid is obtained. In the present study the compliments the charge of the inorganic. Exfoliation in a non-polar solvent gives stable dispersions of neutral nanosheets that consist of the charged inorganic sheet with the inorganic surfactant chains remaining anchored to the layers. Dispersions of the neutral nanosheets of the positive charged and negatively charged inorganic sheets are mixed and on solvent evaporation a periodically alternating layered hybrid is obtained. In the present study the negatively charged inorganic is montmorillonite and the positively charged layers hydrotalcite.

Preparation of Montmorillonite-DODMA. Na-montmorillonite (Na-MMT) was obtained from Southern Clay Product Inc., USA (Na-Cloisite) and has the formula Al\(_{1.5}\)Mg\(_{0.5}\)Fe\(_{0.07}\)Si\(_{1.02}\)O\(_{2.55}\)(OH)\(_{1.53}\)(DDS)\(_{0.17}\)(DODMA)\(_{0.12}\). The carbonate ions were removed and the \('\) parameter dilated to 8.43 nm. The montmorillonite structure was generated from that of hydrotalcite that has a unit cell composition Mg\(_{36}\)Al\(_{5.5}\)Si\(_{2.5}\)O\(_{6}\) (OH)\(_{2}\) \(\cdot\) 5.85 nm. In the unit cell of hybrid, the number of metal (Mg \+ Al \+ Fe) ions in the octahedral layer of 2 \(\cdot\) 1 montmorillonite and the number of metal (Mg \+ Al) ions in the octahedral layer of the LDH was taken to be identical. This would correspond to a 1 \(\cdot\) 1 molar ratio of montmorillonite and hydrotalcite layers.

TEM Bright-field images and SAED patterns were recorded on a JEOL JEM-2100F, fitted with a bottom mounted Olympus View G2 CCD camera and an Oxford Physical characterization. Powder X-ray diffraction patterns were recorded on a Bruker D8 Advance machine. X-ray diffractometer using Cu K\(_\alpha\) radiation of wavelength 1.54 Å. In the region 1 \(\rightarrow\) 5, a scintillation detector was used while at higher angles, a Lynxeye solid state detector was used. X-ray diffraction patterns were calculated for the inorganic layered lattice by using the Reflex module of Materials Studio (Version 5.5.0.0, Accelrys Software Inc. (2010)). Patterns were calculated for the Bragg-Brentano geometry with a Lorentz function for peak profiles and peak broadening accounted by varying crystallite size. The Mg-Al LDH structure was generated from that of hydrotalcite that has a unit cell composition Mg\(_{36}\)Al\(_{5.5}\)Si\(_{2.5}\)O\(_{6}\) (OH)\(_{2}\) \(\cdot\) 5.85 nm with lattice parameters \(a = b = 0.304\) nm, \(c = 2.281\) nm, \(\beta = 90^\circ\), \(\gamma = 120^\circ\), and space group \(\text{P6}_3\) (ICSD No. 81963)\(^*\). The carbonate ions were removed and the \('\) parameter dilated to 8.43 nm. The montmorillonite structure was generated from a unit cell composition Al\(_{1.5}\)Mg\(_{0.5}\)Fe\(_{0.07}\)Si\(_{1.02}\)O\(_{2.55}\)(OH)\(_{1.53}\)(DDS)\(_{0.17}\)(DODMA)\(_{0.12}\). This indicates that the hybrid is formed when the mole ratio of hydrotalcite to montmorillonite is 1 \(\rightarrow\) 1. The infrared spectra of the 1 \(\rightarrow\) 1 hybrid is included as part of the Supplementary Information (Supplementary Fig. S4). The spectra show bands due to the anchored DDS and DODMA surfactant chains.

Preparation of hydrotalcite-montmorillonite hybrid. Mg-Al LDH-DDS and MMT-DODMA were sonicated in chloroform separately for 15 minutes. The concentration of hydrotalcite or montmorillonite in dispersion is 2 mg/ml. The dispersions of Mg-Al LDH-DDS and MMT-DODMA were mixed and sonicated for 20 minutes. This dispersion was drop coated on a glass substrate and chloroform allowed to evaporate slowly. The composition of the resultant solid was determined by ICP-OES, TGA and elemental analysis. The composition of the hybrid that gave a unique XRD pattern is Mg\(_{36}\)Al\(_{5.5}\)Si\(_{2.5}\)O\(_{6}\) (OH)\(_{2}\) \(\cdot\) 5.85 nm. This indicates that the hybrid is formed when the mole ratio of hydrotalcite to montmorillonite is 1 \(\rightarrow\) 1. The infrared spectra of the 1 \(\rightarrow\) 1 hybrid is included as part of the Supplementary Information (Supplementary Fig. S4). The spectra show bands due to the anchored DDS and DODMA surfactant chains.

The Co-Al LDH-DDS and MMT-DODMA hybrid was prepared and characterized by procedure outlined in Supplementary Method S2 and Supplementary Figure S5 & Figure S6. The stoichiometry and compositions of the layered solids and hybrids are summarized in Supplementary Table S1.
INCA Energy EDAX attachment. Analysis of the TEM images was done using the Olympus-SIS item software package. For TEM analysis, dispersions of surfactant intercalated solids in toluene were cast on a holey carbon grid. Tapping mode AFM experiments were performed using a (Digital Instruments, Santa Barbara, CA) nanoscope IVA multimode AFM equipped with a A-scanner. 1H-NMR spectra of the experiments were performed using a (Digital Instruments, Santa Barbara, CA) intercalated solids in toluene were cast on a holey carbon grid. Tapping mode AFM INCA Energy EDAX attachment. Analysis of the TEM images was done using the Brookhaven-Zetapals instrument. Zeta potential measurements were done on Brookhaven-Zetapals instrument.

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