





The creation, analysis, and use of organic guest-containing layered double hydroxides

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Abstract:

Interest in creating inorganic-organic host-guest supramolecular structures using layered inorganic solids as host materials is on the rise. This article provides an overview of layered double hydroxides (LDHs) including organic guests, including their synthesis, characterisation, and applications. Stacks of mixed metal hydroxide layers, each with a positive charge, make up an LDH, and the presence of anions between the layers is necessary to maintain the charge neutrality of the structure as a whole. Porphyrin and phthalocyanine derivatives, sulfonates, phosphonates, aromatic and aliphatic carboxylates, and a host of other organic guests are all within the realm of possibility. We highlight potential uses of organo-LDHs in photochemistry, electrochemistry, sorption, and catalysis. The ability to systematically construct new inorganic-organic supramolecular structures is shown to be achievable using LDHs.

1 Introduction

Layered materials that are able to intercalate neutral guest molecules or to exchange inorganic and organic ions for inter-layer ions have attracted considerable attention.1h3 Through the incorporation of a guest species into a layered host, novel solids may be engineered with desirable physical and chemical properties. Various layered materials such as clay minerals, graphite, transition metal dichalcogenides, and metal phos-phates and phosphonates have the ability to act as host materials. These materials have been studied as potential cata-lysts, selective sorbents and hosts for nanoscale reactions.4h8 A particularly attractive feature of such layered hosts is that they serve as a template for the creation of intercalated supra-molecular arrays. Layered double hydroxides (LDHs), also known as anionic clays, are host-guest materials that, owingto their potential application in these and other areas, have recently gained much attention.9h11 LDHs are available as naturally occurring minerals and as

synthetic materials. They were first prepared in the laboratory in 1942 when Feitknecht12,13 reacted dilute aqueous metal salt solutions with base, although the first detailed structural analyses of LDHs were not carried out until the late 1960s by Allmann14,15 and Taylor16,17 and their co-workers. Briefly, LDHs consist of stacks of positively charged mixed metal hydroxide layers that require the presence of interlayer anionsto maintain overall charge neutrality. One subset of LDHs is that in which the charge balancing anion is organic. The incorporation of organic molecules into inorganic layered hosts in general has been extensively investigated18,19 and presents a method of obtaining novel nanocomposite materials. This review discusses the preparation, properties and applications of organo-LDHs. In particular, we indicate the range of organic guests that may be incorporated intoLDHs and describe how it is possible to create, in a system- atic way, inorganic Eorganic supramolecular structures.

1.1 Structural properties of LDHs

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The most important group of LDHs may be represented by the formula $[M2^{}_{1\sim x}M3^{}_{x}(OH)_{2}]An \sim_{\chi \parallel n} \ell mH_{2}O$, where M2 and M3 are divalent and trivalent cations, respectively; χ is equal to the ratio M3/(M2 M3) and A is an anion of valence *n*. The structure of LDHs is most clearly described by considering the structure of brucite, Mg(OH), which consists of Mg2 ions co-ordinated octahedrally by hydroxyl groups. The octahedral units share edges to form infinite, charge neutral layers. In an LDH, isomorphous replacement of a fraction of the Mg2 ions with a trivalent cation, such as Al3, occurs and generates a positive charge on the layers that necessitates the presence of interlayer, charge balancing, anions. The remaining free space of the interlayer is occupied by water of crystallization (Fig. 1).

Di†erent stacking arrangements of the hydroxide layers are observed, giving rise to polytypism. The mineral of composi- tion Mg Fe (OH) (CO) \acute{E} 4H O, for example, may have either a three-layer repeat (Pyroaurite, the 3R polytype) or a two-layer repeat (Sjo[°] grenite, the 2H polytype).17 An investiga- tion of the polytype diversity of LDHs has recently been con- ducted.20,21

In general, M2[×] and M3[×] ions that have an ionic radiusnot too di[†]erent from that of Mg2[×] (lonic ²/₄adius of Mg2[×] 0.65 Å) can be accommodated in the holes of the close-packed OH groups in the brucite-like layers to form LDHs. Observed M2[×] and M3[×] species include Mg2[×], Fe2[×], Co2[×], Cu2[×], Ni2[×], Cd2[×] or Zn2[×] and Al3[×], Cr3[×], Ga3[×] or Fe3[×], respectively. Furthermore, LDHs containing more than two species of the second cation have been prepared.22,23 The charge density on the hydroxide layers of the LDH will clearly depend upon the M2[×]/M3[×] ratio. A particular advantage of LDHs, therefore, is that the anion-exchange capacity of theLDH, and hence the number and arrangement of the charge



Fig. 1 Schematic representation of the interlayer structure of an LDH

balancing anions in the LDH, may be controlled by varyingthe M2'/M3'ratio.

A second class of LDHs, which have not been as extensively studied, contains monovalent and trivalent matrix cations. An example is [LiAl (OH)] $A \sim E mH$ O, which consists of sheets of aluminium octahedra with vacancies filled by lithiumatoms.24h27

In naturally occurring LDHs, the most common anion found is carbonate. In practice, however, there is no signifi- cant restriction to the nature of the charge balancing anion that can occupy the interlayer region; for example, halides,24,25,28,29 oxo-anions,24,25,28,29 silicates,30v32 polyoxo- metalate anions,24,25,33h40 complex anions,41h43 as well as organic anions, may all be incorporated.

In the remainder of this review the composition of LDHs with divalent and trivalent cations will be represented with the convenient notation M2`M3`(A), where A is the interlayer anion. Similarly, LDHs containing Li and Al matrix cations will be represented as LiAl(A).

1.2 Characterization of LDHs

A variety of techniques are used to characterize LDHs. Powder X-ray di[†]raction (PXRD) and infrared spectroscopy (IR), for example, are routinely used whilst others such as ESR,44 XAS,39,45,46 and computer modelling,47,48 although reported, are less extensively employed.

The most frequently employed technique is PXRD. Di[†]rac- tograms are normally obtained using randomly oriented pow-dered samples although in certain cases oriented samples or samples under solution have been studied (it will be seen in Section 3 that the di[†]ractogram obtained for an organo-LDH is critically dependent upon the degree of hydration of the sample). A typical di[†]ractogram obtained for an LDH con- sists of sharp basal (00*l*) reflections at low values of 2h corre- sponding to successive orders of the interlayer spacing. In addition, relatively weak non-basal reflections at higher values of 2h are present. In certain cases, the non-basal reflections arequite broad, which may be attributed to turbostratic dis- ordering of the hydroxide layers.

Generally, the patterns are indexed on the basis of a hex- agonal unit cell. The interlayer spacing of the LDH is equiva- lent to 1/n of the *c* parameter, where *n* is the layer repeat of the unit cell, which depends upon the stacking sequence of the layers. Subtracting the hydroxide layer thickness of approx- imately 4.8 Å from the interlayer spacing yields the gallery height (Fig. 1). The gallery height will depend on the size and orientation of the charge balancing anion.

The M2[/]/M3[°] ratio of the LDH may be inferred from the *a* parameter of the unit cell, which is determined using the relationship $a \setminus 2d$, where *d* is the *d* spacing of the 110 reflection. For a MgAl LDH, for example, and as a result of the smaller ionic radius of Al3[°] (0.50 Å) compared to Mg2[°] (0.65 Å), the *a* parameter decreases as the aluminium content of the LDH increases. The measurement of *d* for determining the M2[°]/M3[°] ratio of the LDH along with elemental analysis techniques, enables an approximate chemical formula for the LDH to be deduced. The possibility of precipitating M(OH) and/or M(OH) phases during synthesis, however, may mean that bulk chemical analysis would not be appropri- ate. Such phases may be amorphous and therefore notdetected by PXRD.

IR is a useful technique for confirming the presence of the charge balancing anion in an LDH. The incorporation of a carboxylic acid anion into an LDH, for example, would be identified bM0trong antisymmetric and symmetric carboxylatestretching bands at approximately 1560 and 1400 cm~1, respectively. Alternatively, intercalation of the undissociated acid form would be identified by the strong absorption of the carbonyl stretch of the acid at approximately 1700 cm~1. In

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addition, IR allows the presence of impurity inorganic charge balancing anions, such as carbonate and/or nitrate, to be con- firmed. The thermal properties of organo-LDHs are commonly studied using thermogravimetry (TG), di†erential scanning calorimetry (DSC) and di†erential thermal analysis (DTA). In certain cases these techniques are used in combination with a mass spectral analyser, to study the nature of the evolvedgases during the thermal treatment. The thermal properties of organo LDHs are discussed in Section 3.6.

2 Synthesis procedures

There are four general approaches to the preparation of organo-LDHs: (I) anion-exchange of a precursor LDH; (II) direct synthesis by coprecipitation; (III) rehydration of a cal- cined LDH precursor and (IV) thermal reaction. A common problem with all the methods is that in preparations of LDHs with anions other than carbonate it is important to avoid con- tamination from CO, since the carbonate anion is readily incorporated and tenaciously held in the interlayer. Conse-quently, decarbonated and deionized water is often used and exposure of the reacting material to the atmosphere is kept to a minimum.

The synthesis of LDHs containing carboxylic acid deriv- atives has recently been reviewed by Carlino.49

2.1 Anion exchange

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The anion-exchange method is the most common method for the preparation of organo-LDHs. The anion-exchange proper- ties of LDHs containing simple inorganic anions are well documented.28,29,50 The ease of exchange of monovalent anions is in the order OH~ \Box F~ \Box Cl~ \Box Br~ \Box NO ~. Divalent anions such as SO 2~ and CO 2~ have higher selec- tivity than monovalent anions. LDHs containing nitrate anions are, therefore, the most suitable precursors for anion- exchange syntheses due to the relative ease with which the nitrate anions can be displaced from the interlayer. In general, the anion-exchange reaction is carried out by simply disper- sing the precursor LDH in aqueous solution containing an excess of the organic anion that is to be incorporated. The organic anion of interest must be stable at the pH of exchange and, if the M2[×]/M3[×] ratio of the precursor LDH is to be retained in the anion-exchanged LDH, the hydroxide layers must also be stable.

Exchange of interlayer anions by an organic anion (acetate) was first reported by Miyata and Kumura41 in 1973, although the publication gives no details of the characterization of the exchanged LDH. Later, Boehm *et al.*51 reported the anion exchange of a ZnCr LDH containing nitrate or chloride anions with short- and long-chain sodium alkyl sulfate anions. A variety of organo-LDHs, with di†erent combinations of matrix cations, as well as di†erent interlayer organic anions, have been prepared by Meyn *et al.*52 using an anion-exchangeroute.

A large variety of other organic molecules have been incorporated into LDHs using anion exchange. These include porphyrins functionalized with sulfonate or carbonategroups,53h55 MII phthalocyanines (M \ Cu or Ni) functional- ized with sulfonate groups,42,56 malonate,57 aromatic diphosphonate37 and 5-benzoyl-4-hydroxy-2-methoxy-benzensulfonate anions.58 In addition, the incorporation of ananionic b-cyclodextrin derivative into a MgAl LDH *via* anion exchange has recently been reported.59

2.2 Direct synthesis by coprecipitation

The coprecipitation method for the direct synthesis of LDHs containing simple inorganic anions has been developed over a number of years12,13,60h63 and a similar procedure is used for

organo-LDHs: the M2[°] and M3[°] hydroxide layers are nucle- ated and grown from an aqueous solution containing the anion that is to be incorporated into the LDH. The anion that is to be introduced must have a high affinity for the hydroxide layers, otherwise the counter anions of the metal salts may be incorporated, thus contaminating the LDH. For this reason, metal nitrate or chloride salts are commonly utilized because of the low selectivity of LDHs towards these anions.28

In general, the coprecipitation is performed by adding an

aqueous solution of the metal nitrate or chloride salts to an aqueous solution of the organic anion. Usually, the coprecipi-tation is performed at constant pH by simultaneously addingNaOH . A particular advantage of the coprecipitation method is that it allows for accurate control of the chargedensity (M2[/]M3 ratio) on the hydroxide layers of theproduct LDH. In order to ensure simultaneous precipitation of the matrix cations it is necessary to perform the synthesis at a pH at which both the M2 and M3 hydroxides precipitate. To ensure that the M2[/]M3 ratio in the starting solution is retained in the product LDH it is important to control the pH. Without pH control, the formation of impurity M(OH) and/or M(OH) phases, and consequently a product LDH with an undesired M2[/]M3 ratio, is probable.64,65 It has been reported that the optimum coprecipitation pH is depen-dent upon the combination of matrix cations utilized and the organic anion present.11,66 To obtain phase-pure LDHs gen-erally only M2[/]M3 ratios between 2 and 4 are possible (*i.e.*, values of x in the range 0.33È0.2).10 Ratios lower than 2 leadto a high density of M3 octahedra in the hydroxide layer, which act as nuclei for the formation of M(OH) ; similarly, atratios higher than 4 the formation of M(OH) is probable. The usual direct synthesis procedure involves crystallization of the coprecipitation product for 18 h at a temperature in the region of 65È75 ;C. In certain cases, however, this does notlead to a wellcrystallized organo-LDH phase, in which casehydrothermal treatment may be used to improve the crystallinity of these materials.67,68

2.3 Rehydration of calcined LDH precursor

Miyata69 reported that a hydrotalcite-like LDH [MgAl(CO)] transforms into a mixed magnesium aluminium oxide after heat treatment at $500\dot{E}800$ _jC, which rehydrates and combines with anions to re-form a LDH in the presence of water and anion. This so-called *memory e†ect* property of LDHs provides an e†ective synthetic route for inserting inor- ganic and organic anions into LDHs. An advantage of this method is that the incorporation of competing inorganic counter anions into the LDH is prevented, although the inclu- sion of carbonate from atmospheric CO remains a problem. The ease and extent of reconstruction of the calcined LDH are controlled by the properties of the matrix cations as well as the temperature of calcination.70,71

The reconstruction of an LDH from a mixed metal oxide

precursor is believed to occur *via* a topotactic reaction. To improve the crystallinity of the rehydrated product, therefore, Carlino *et al.*72 recommend heating the precursor LDH using a slow ramping technique (D1 $_{\rm i}$ C min~1). This technique pre- vents the rapid expulsion of carbon dioxide and water from the LDH, which occurs upon direct and sudden heating, dis- rupting the layered structure of the calcined product.

Chibwe and Jones43 used the rehydration method to prepare a series of MgAl LDHs containing various organic

guests *via* the rehydration route. In this procedure, a 1:2 (v/v) water E glycerol solution was used for the reconstruction of a calcined MgAl(CO) precursor in a nitrogen atmosphere. The reaction products obtained were single phase and crystallo- graphically well-ordered. In the absence of glycerol mixed phases were observed.

A variety of organo-LDHs have been prepared *via* the rehydration route, including LDHs containing naphthalenecarboxylate75 or cobaltII phthalocyaninetetrasul- fonate anions.44,76,77

2.4 Thermal reaction

Thermal reaction is a relatively new approach to the prep- aration of organo-LDHs. The procedure was first reported in 1994 by Carlino and Hudson78 who reacted molten sebacic acid with a MgAl(CO) LDH. In this procedure an intimate mixture of MgAl(CO) and sebacic acid was heated at a tem- perature $20\dot{E}30$ _iC above the melting point of the acid.

A similar procedure for incorporating the caprate or phenylphosphonate anions into a MgAl LDH was also used.72,79 In addition to the corresponding organo-LDH phase, the thermal reaction product was found to contain in each case an unreacted MgAl(CO) phase. Although phase- pure organo-LDH products have not been obtained using thermal reaction, this approach nevertheless provides an interesting alternative to the conventional *wet* methods.

3 Organization of anions ithin the LDH

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3.1 Aromatic carboxylate anions

Organo-LDHs in which the charge balancing anionis an aromatic carboxylate anion are among the most widely studied systems. LDHs containing terephthalate33,34,39,40,47,52,64h68,73,80h86 or ben- zoate42,52,64h68,73,83,84,87 anions in particular have received much attention. For terephthalate anions an inter- layer spacing of between 14.0 and 14.4 Å, corresponding to a gallery height of between 9.2 and 9.6 Å, is generally reported for a variety of matrix cations and synthesis methods. The physical length of the terephthalate anion, including the van der Waals radii of the terminal oxygen atoms, is approx- imately 9.9 Å. Allowing for the fact that the terminal oxygen atoms of the terephthalate anion are most likely hydrogen bonded to the hydroxide layers, dimensional analysis suggests that for such a spacing the terephthalate anion adopts an orientation with its long axis perpendicular to the hydroxide layers.

For LDHs containing the benzoate anion, a gallery height of between 10.4 and 10.7 Å is generally observed. The length of the benzoate anion, including van der Waals radii of the terminal oxygen and hydrogen atoms is approximately 8.8 Å and two interlayer arrangements of the benzoate anion have been proposed to account for the observed gallery height. First is a bilayer-like structure with the carboxylate groups of the anions attached to adjacent inwardly facing hydroxide layers and the aromatic groups interleaved within the mid- plane of the gallery.64,65,68 Moggridge *et al.*87 have suggested, from NEXAFS measurements, that in this arrangement the benzoate anion is inclined at an angle of $35 \land 10$; to the hydroxide surface of the layers. Alternatively, a vertical arrangement in which a water molecule is retained between

anions. In certain cases impurity MgAl(CO₂) phases were the benzene ring and the hydroxide surface has been sug-

detected in addition to the corresponding organo-LDH phase. If the rehydration is performed in a nitrogen atmosphere car- bonation of the product LDH may still remain a problem due to the high affinity of the mixed oxide for carbonate anions.73

Dimotakis and Pinnavaia74 showed that the use of glycerol

as a swelling agent assists the incorporation of the organic gested.52

Kooli *et al.*64,65 have investigated the properties of a series of MgAl LDHs containing terephthalate or benzoate anions and possessing varying layer charge (*i.e.*, varying Mg/Al ratio). Two extreme orientations were observed for the organic anions between the hydroxide layers: either vertical (*i.e.*, gallery heights of approximately 9.2 and 10.4 Å for terephtha-late or benzoate, respectively) or horizontal. For the tereph- thalate or benzoate anions in a horizontal orientation a gallery height of approximately 3.5 Å was observed, approx- imately the thickness of a benzene ring.

At room temperature, it was found that the orientation of the terephthalate anion is dependent upon the charge density on the layers and the degree of hydration of the LDH. For samples prepared with initial Mg/Al ratios of 1 or 2, PXRD indicated that the terephthalate anion adopts a vertical orien- tation (interlayer spacing $\14.2$ Å) before and after drying. For the sample prepared with an initial Mg/Al ratio of 3, the same vertical orientation was observed before drying. After drying in air overnight at 60 iC, however, it was found that the terephthalate anion adopts a horizontal orientation, as determined by a reduction of the interlayer spacing from 14.2 to 8.3 Å. This process was found to be completely reversible upon repeated cycles of drying and rehydration.

The apparent change in orientation of the terephthalate anion with increasing Mg/Al ratio demonstrates that a balance of forces determines the adopted interlayer arrange- ment of the anion. Coulombic repulsion and anion packing density, which are dependent upon the M2[×]/M3[×] ratio, are important factors. For a rigid, linear dicarboxylate anion such as terephthalate a vertical monolayer arrangement allows for maximum separation of the positively charged layers, whilst also maintaining a strong interaction between the anion and the layers. This arrangement also ensures maximum lateral separation between the anions in the interlayer. The charge density on the hydroxide layers of the LDH decreases with increasing Mg/Al ratio and hence there is a lower packing density of the anions at high Mg/Al ratios (*e.g.*, Mg/Al ratio3), which allows the terephthalate anions to adopt a horizon- tal orientation in the interlayer. In addition, coulombic repul- sion between the hydroxide layers will be reduced as the Mg/Al ratio of the LDH increases. The ability of excess waterto maintain the vertical orientation of the terephthalate anion at Mg/Al ratio 3 indicates that hydrophobic/hydrophilic inter-actions are also important.

The existence of a 3a hexagonal superlattice in the *ab* plane for a MgAl LDH containing benzoate anions and pre- pared with an initial Mg/Al ratio of 2 has recently been reported.67,68 The formation of a superlattice could result from an ordered distribution (*i.e.*, no Al3` neighbouring each other) of the cations within the hydroxide layer, and has also been reported for a MgAl LDH containing sulfate anions.21 Furthermore, the observation of a superlattice suggests anordered distribution of the interlayer anions, since Al and Mg have similar scattering powers. Cation ordering in synthetic LDHs, including a MgFe LDH containing benzoate has recently been investigated by Vucelic *et al*.88

3.2 Aliphatic carboxylate anions

A series of ZnAl LDHs containing terminal linear aliphatic dicarboxylates, $C_n H_n(CO \sim)$, was prepared by direct syn-

thesis by Miyata and Kumura.41 The interlayer spacing of the organo-LDH was found to increase linearly from 9.4 Å for oxalate $(n \ 0)$ to approximately 18È19 Å for sebacate $(n \ 8)$ (Fig. 2). The gallery height of the ZnAl(sebacate) LDH, obtained by subtracting the brucite layer thickness of 4.8 Å from the interlayer separation, is close to the

physical length of the sebacate anion. Dimensional analysis suggests, there- fore, that the sebacate anion is oriented with its long axis approximately perpendicular to the hydroxide layers in a monolayer arrangement.

Chibwe and Jones43 reported interlayer spacings of 16.3 and 18.8 Å for a MgAl LDH containing the sebacate anion prepared *via* rehydration and coprecipitation routes, respec- tively. Similarly, Carlino and Hudson78 observed interlayer



Fig. 2 Relationship between carbon number of dicarboxylic acid anions and the d spacing of the (006) plane of the organo-LDH (after Miyata and Kumura41)

spacings of 19.3 and 17.6 Å for the same material prepared *via* thermal reaction and coprecipitation, respectively. These inter-layer spacings are also consistent with a monolayer arrange- ment of the sebacate anion between the hydroxide layers. The range of values observed, however, show that the interlayer spacing of the organo-LDH depends on the history (*e.g.*, the method of synthesis and drying conditions *etc.*) of the product. The di†erences between the values observed may be due to additional inclusion of water or to small di†erences in the orientation and packing of the alkyl chains (possibly as a result of di†erent layer charge densities on the product LDHs). Carlino and Hudson78 attributed the di†erent separa- tions that they observed to a change in the angle of orienta- tion (from 90; to approximately 60_i) of the sebacate dianion with respect to the surface of the hydroxide layers.

A linear relationship between the observed interlayer spacing and the chain length of linear aliphatic dicarboxylates was also reported by Meyn *et al.*52 for series of ZnCr and ZnAl LDHs. The interlayer spacing of ZnCr(NO) increased from 8.9 to 9.9 Å when oxalate anions were exchanged for the interlayer nitrate anions. The interlayer spacing was then found to increase linearly with the chain length of the exchanged anion up to 18.0 Å for sebacate. A similar linear relationship was observed by Raki *et al.*89 for a MgFe LDH containing a series of linear dicarboxylate anions (from $n \ 0$ to 12), prepared *via* coprecipitation.

An analogous trend is reported by Meyn *et al.*52 for a ZnCr LDH containing a series of linear aliphatic monocarboxylates, $C_n H_n$, CO

 \sim prepared *via* an anion-exchange route. The interlayer spacing was found to increase linearly from 11.2 Å

for formate $(n \setminus 0)$ to 31.6 Å for nonadecanoate $(n \setminus 18)$. It is important to note, Aowever, that the interlayer spacings were determined for the LDH in equilibrium with the exchange solution. Upon drying in air (30 $\grave{E}60$ iC), it was found that the layer structure becomes poorly ordered and the intensity of the basal reflections is significantly reduced. The disordering was attributed to decarboxylation of the anions.

For a linear monocarboxylate anion the formation of a ver- tical bilayer arrangement is probable because the interlayer of a LDH has two ionic surfaces, top and bottom, to which the carboxylate group is equally likely to attach. The bilayer may be arranged with the alkyl chains of the anions end-to-end or interleaved. In addition, the anion may also be inclined at an angle from the normal to the layers. Carlino and Hudson79 prepared a MgAl LDH containing the caprate anion $(n \setminus 9)$ via a coprecipitation route. The interlayer separation, afterair-drying at room temperature, was found to be 19.6 Å, which was attributed to the formation of an end-to-end bilayer-like arrangement with the caprate anion at a slant angle of 38; to the surface of the hydroxide layers.

3.3 Sulfates and sulfonates

A large number of di†erent organic anions containing sulfate(SO ~) or sulfonate (SO ~) groups have been incorporated into LDHs. A series of alkyl sulfate anions, $C_n H_2$, SO ~, forexample, was incorporated into a ZnCr LDH *via* anion exchange by Boehm *et al.51* The products of the exchangewere found to have basal spacings in the range 21.1 ($n \setminus 8$) to 32.6 Å ($n \setminus 18$) and underwent substantial additional swelling

in the presence of *n*-alkyl alcohols or *n*-alkylamines. A detailed study of alkyl sulfate anions incorporated *via* anion exchange into ZnCr LDHs has been carried out by Kopka *et al.*90 It was proposed that the alkyl sulfate anions adopt monolayer arrangements between the hydroxide layers. In equilibrium with the exchange solution, a near perpendicular orientation of the alkyl chains was proposed. Following washing and drying at 60 $_{i}$ C, however, a reduction in the interlayer spacing revealed that the chains adopt a slant angle of 56; to the surface of the hydroxide layers.

The incorporation of dodecyl sulfate anions, CH (CH) SO ~, into series of NiAl, MgAl and ZnCr LDHs with varying layer charge has been studied by Clearfield *et al.*91 For comparison, the dodecyl sulfate anion was incorpor- ated *via* two methods: anion-exchange and correcipitation. Depending upon the method of preparation, the products yielded interlayer spacings distributed about three mean values: 26, 36 and 47 Å. The diterent spacings revealed the existence of three diterent interlayer arrangements of the dodecyl sulfate anion, with the shortest interlayer spacing (26 Å) corresponding to a perpendicular monolayer arrangement of the anion. Similar values have been obtained by other researchers for LDHs containing the dodecyl sulfate anion.33,51 The interlayer

spacing of approximately 36 Å was attributed to a bilayer arrangement in which the anion is tilted at an angle of approximately 40; to the surface of the hydroxide layers. An alternative suggestion, that an inter- leaving vertical bilayer arrangement is adopted by the dodecylsulfate anion, was also proposed. The 47 Å spacing was considered consistent with a vertical end-to-end bilayer arrangement.

A particularly interesting result has been reported for an LDH containing secondary alkanesulfonates.52 The anions are a mixture of di†erent isomers, with the sulfonate group positioned at di†erent carbon atoms along the alkyl chain. The PXRD pattern obtained for this organo-LDH showsseveral orders of sharp basal reflections, indicating that the layer structure is extremely well-ordered, *i.e.*, the interlayer



Fig. 3 Formation of bilayers consisting of isomers of secondary alk- anesulfonates between the hydroxide layers of an LDH (after Meyn et al.52)

The incorporation of 9,10-anthraquinone-2,6-disulfonate anions into a MgAl LDH led to two distinct conformations of the organic anion.92 The LDHs were prepared *via* coprecipi- tation (using a controlled pH of $10\dot{E}11$) with various initial Mg/Al/organic anion ratios. The products were single phase with, depending on the orientation of the anion, interlayerspacings of approximately 19 or 12 Å. The 19 Å phase was selectively obtained under Al-rich preparation conditions, although the initial Mg/Al ratio was not retained in the product LDH. It was found that the Mg/Al ratio of the product LDH, as determined by chelatometric titration, depends neither on the phase to which the sample belongs northe initial ratio in the sample preparation. (It should be noted, however, that an *a* parameter of 3.02 Å was reported for the 19 Å phase, whereas an *a* parameter of 3.06 Å was reported for the 12 Å phase. This would seem to suggest that the 19 Å phase is, in fact, Al-rich compared with the 12 Å phase. The resulting di†erent layer charge densities of the two phases would then account for the di†erent interlayer arrangements observed.)

Franklin et al.58 reported the presence of three distinct and

interconvertible phases when the 5-benzoyl-4-hydroxy-2-methoxybenzenesulfonate (BHMBS) anion was incorporated into a ZnAl LDH. The selected anion is dibasic, having both a strong sulfonate group and a much weaker phenolate group. It was found that anion exchange of the divalent anion (in which both the sulfonic and phenolic groups are ionized) formed an LDH with an interlayer spacing of 13.8 Å. The monovalent form (in which only the sulfonic group is ionized)

spacing is highly regular. This is a surprising result consider-and

ing that the charge balancing anions have diterent structures. The result was explained, however, by assuming that the alka-nesulfonates adopt a U-shaped conformation and are in a bilayer arrangement. Thus, the diterently shaped alkanesul- fonate ions may pair up in such a way that a constant layer separation is obtained (see Fig. 3).

A series of ZnAl LDHs, each containing a diterent isomerof the naphthalenedisulfonate anion, have been prepared.81 The relationship between the physical size of the incorporated anion and the interlayer spacing was investigated by changing the positions of the two sulfonate groups on the naphthalene moiety. A small diterence in interlayer spacing was observed for the three isomers studied, which suggests a small dependence of the interlayer arrangement of the naphthalenedisul- fonate anion on the positions of the two sulfonate groups. readily converted to the divalent form within the LDH. The observed diterent interlayer spacings revealed that the guest molecule occupies two distinct conformations within the inter-layer, depending upon the charge of the anion. Furthermore, it was found that the divalent phase may be converted into a

15.4 Å phase, which is intermediate between the monovalent and divalent forms, by exposure of the LDH to water vapour, or by dispersion in low pH (pH B 6) solution (Fig. 4).

Cai *et al.*93 used atomic force microscopy to investigate the adsorption of the BHMBS anion onto the surface of a MgAl LDH. It was found that the packing arrangement and orienta- tion of the BHMBS anion, adsorbed on the surface of the LDH layers in aqueous solution, are governed by factors such as coulombic and hydrogen bonding interactions between the anion and the positively charged LDH surface.



Fig. 4 PXRD pattern of ZnAl LDHs: (a) ZnAl(NO), (b) 13.8 Å phase ZnAl(BHMBS), (c) 20 Å phase ZnAl(BHMBS) (Franklin et al., 58 reproduced by permission of the Royal Society of Chemistry)

The incorporation of quite large organic molecules contain- ing sulfonate groups into LDHs has been reported. An example is 5,10,15,20tetra(4-sulfonatophenyl)-porphyrin (TSPP), which has been anion-exchanged into a MgAl LDH.54 The interlayer spacing of 22.4 Å (along with IR and visible spectra) suggest that the molecule is inserted intact with the molecular planes perpendicular to the hydroxide layers (Fig. 5).

3.4 Phosphonates

The preparation of organo-LDHs containing aromatic phos- phonates has recently been reported. Anion exchange of the benzenediphosphonate anion, [O_PwC_H_wPO_1]4~, into a through the layer surface OH groups. For the products of the exchange reaction carried out above room temperature, or if any of the products were dried rigorously, the phase with the smaller interlayer spacing was found to predominate. Carlino *et al.*¹² have also reported the existence of a MgA1 LDH coh- taining the phenylphosphonate anion, in which it is proposed that the phosphonate group is connected directly to the matrix cations. In addition, Vichi and Alves94 have recently investigated the anion-exchange reactions of CdAl(NO) and CdAl(CO)

withphenylphosphonicacidand2-carboxyethylphosphonicacid. 3 3

3.5 Polymeric anions

The preparation of LDHs containing organopolymers has received growing attention in recent years. Tanaka et al.95 prepared a MgAl LDH containing polyacrylate anions, via the in situ polymerization of interlayer acrylate anions. The acry- late anions were incorporated via anion exchange and the sub-sequent polymerization was carried out by the addition of an initiator (potassium peroxodisulfate) to the undried product followed by heat treatment for 1 day at 80 ¡C. The poly-merization of the acrylate anion was inferred by the disappearance of the absorption band due to CxC double bond stretching in the infrared spectrum of the organo-LDH, fol- lowing the heat treatment. In addition, the interlayer spacing decreased from 13.8 to 13.4 Å upon polymerization of the acrylate anion. The PXRD pattern of the MgAl LDH contain-ing polyacrylate revealed a poorly crystalline material. Suga- hara et al.96 have reported the preparation of a MgAl LDH containing polyacrylonitrile using a similar procedure. An in situ polymerization route has also been used for the prep-aration of a CuCr LDH containing polyaniline.80 In this case, the oxidative polymerization of aniline intercalated in aCuCr(terephthalate) LDH is reported.

Oriakhi et al.97 describe the preparation and character-

ization of a series of LDHs containing a variety of matrix cations and three vinylic polymers: polyacrylate, polyvinylsul-fonate and polystyrenesulfonate. The organopolymer LDHs were coprecipitated in a basic solution containing the dis-solved polymer. Interlayer spacings of between 12.0 and 21.6 Å were determined for the reaction products, indicating

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MgAl LDH was found to depend upon the temperature at which the reaction was performed.37 If the exchange was per-formed at B0 _iC, PXRD revealed a single, crystallo- graphically well-ordered product LDH phase, with an interlayer spacing (14.7 Å) corresponding to a perpendicular monolayer arrangement of the diphosphonate anion. If the exchange was carried out at room temperature, a second phase with a smaller interlayer spacing (9.6 Å) was observed, in addition to the 14.7 Å phase. This additional phase was attributed to dehydration of the OH groups of the hydroxide layer and subsequent attachment of the phosphonate groups directly to the metal cations in the layers, *i.e.*, no longer



Fig. 5 Proposed interlayer arrangement of a MgAl LDH containing 5,10,15,20-tetra(4-sulfonatophenyl)-porphyrin (after Park et al.54)

incorporation of the polymer, although the broad basal reflec- tions observed indicate poor crystallinity of the products. A bilayer arrangement of the anionic polymers was proposed. A similar method was used by Messersmith and Stupp98,99 for intercalating water soluble organopolymers into a CaAl LDH. In these methods, a preformed polymer is incorporated between the hydroxide layers of the LDH.

Whilton et al.100 have recently prepared a MgAl(polyaspartate) LDH via two synthetic routes: in situ

thermal polycondensation of MgAl(aspartate) and direct syn- thesis in the presence of the preformed polymer. In both cases, however, the crystallinity of the LDH product is poor.

3.6 Thermal characteristics

The thermal stability of LDHs is an important consider- ation for many potential applications. Consequently, the thermal characteristics of LDHs containing various charge balancing anions have received considerable atten- tion.29,64,67,69,91,99,101h109 In general, the thermal decomposition of organo-LDHs can

be divided into three stages: (I) the removal of physisorbed water at the surface and between the hydroxide layers, from room temperature to approximately 300 $_{\rm i}$ C; (II) dehydroxyla- tion of the hydroxide layers in the range 300 \dot{E} 500 $_{\rm i}$ C and (III) elimination and combustion of the organic anion. The precise temperature at which each stage starts and finishes depends on many factors, including the nature of M2^{and M3^c}, the M2^(M3^c) ratio and the nature of the anion. In particular the



The thermal properties of a CaAl LDH intercalated with polyvinylalcohol have been studied by Messersmith and Stupp.99 The intercalate has an interlayer spacing of approx-

imately 18 Å at room temperature, which persists upon heating to at least 400 ¡C. The organopolymer intercalate heated to 1000 ¡C transforms into an inorganic solid that has a di†erent phase composition than the unintercalated LDH heated to the same temperature.

The e†ect of thermal treatment on the interlayer arrange- ment of a MgAl(terephthalate) LDH has recently been investi- gated by Kooli *et al.*64 using PXRD at elevated temperatures. The LDH was prepared *via* coprecipitation from an initial Mg/Al ratio of 2. At room temperature it was found that the terephthalate anion is in a vertical orientation between the hydroxide layers. Above 150 iC (a temperature at which water loss becomes important) the expanded structure collapses, as was determined by a shift in the position of the 003 reflection from 14.6 to 8.9 Å, suggesting a horizontal arrangement for the anion. At intermediate temperatures, *e.g.*, at 75 iC, reflec- tions at 23.2, 11.7 and 7.4 Å were observed (Fig. 6). The reflec-

tion at 23.2 Å was rationalized in terms of an ordered

Fig. 6 Temperature dependence of the PXRD patterns obtained for MgAl(terephthalate) prepared with a Mg/Al ratio of 2, after heat treatment at di†erent temperatures. The reflection at 23.2 Å is associ-ated with the creation of an interstratified arrangement of vertical andhorizontal terephthalate anions (after Kooli *et al.*64)

temperature at which stage III occurs can vary considerably and depends upon the identity of the organic anion. For a MgAl(terephthalate) LDH, for example, elimination of the organic anion occurs above 500 ;C.64 For a MgAl(dodecyl sulfate) LDH, however, elimination of the organic anion occurs at much lower temperature, in the range $150\dot{E}300$;C.91 The di†erence between the two systems may be attributed to the strength of the interaction between the organic anion and the hydroxide layers. Kuwahara *et al.*106 have investigated the thermal properties of a

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MgAl LDH containing p-toluenesulfonate anions. It was found that the organic anion contained within the LDH was easier to decompose than the anion in the form of its sodium salt. It was considered that this e[†]ect is due to the existence of a stronger interaction between sodium and p-toluenesulfonate than between the LDH and the anion.



Fig. 7 Proposed model for structural changes occurring during the heat treatment and loss of interlayer water of the LDH described in Fig. 6 at (*a*) 25, (*b*) 100 and (*c*) 200 ¡C (after Kooli *et al.*64)

interstratification of collapsed and expanded layers, with the *d* spacing of the lowest angle reflection close to the sum of the *d* spacings of the 003 peak of the vertical form at room tem- perature and the *d* spacing of the horizontal form at 150 ¡C (Fig. 7).64,67

A low temperature (150 ; C) thermal analysis of a

LiAl(myrisate) LDH has been performed by Borja and Dutta.107 Incorporation of the myrisate anion was achieved *via* exchange in ethanol. At room temperature, a monolayer arrangement of the myrisate anions is formed with an inter- layer spacing of 22 Å. The interlayer spacing expands to 43 Å at 100 ¡C, with removal of the solvent (ethanol), and the anion adopts a vertical bilayer arrangement. The non-expanded state was re-formed by cooling and redispersing the LDH in ethanol. A molecular explanation of this phenomenon is given by Dutta and Robins.108 Borja and Dutta107 also examined the low temperature thermal properties of a MgAl LDH inter- calated with myristic acid. It was found that the interlayer spacing decreases from 39 to 24 Å upon heating from room temperature to 140 ¡C, through the formation of irregularities in the all-*trans* bilayer conformation of the alkyl chains. The as-prepared form could not be reformed by cooling or treat- ment with ethanol, unlike the LiAl system. The high tem- perature (up to 450 ¡C) thermal properties of the LiAl(myrisate) LDH have been studied by Jakupca and Dutta.109

4 Applications of organo-LDHs

The applications of LDHs (and their thermal decomposition products) containing inorganic charge balancing anions are well documented.10 There is a wide range of applications for these materials, although the major application is in the field of catalysis. It will be seen in this Section that there is also a variety of potential applications for organo LDHs.

4.1 Catalytic applications

Reichle110 has investigated the catalytic behaviour of the thermal decomposition products of a number of LDHs con- taining a variety of matrix cations and charge balancing anions. In general, the products are fairly strong bases and useful catalysts for vapour-phase aldol condensations. It was found that the thermal decomposition products of LDH pre- cursors containing organic anions such as 1,10-decanedicarb- oxylate, 1,12-dodecanedicarboxylate or oxalate anions showed improved activities for the aldol condensation of acetone com-pared with the thermal decomposition products of carbonate-LDH precursors.

The incorporation of inorganic polyoxometalate anions, such as $[V_{10}O_{28}]6\sim$ and $[Mo_7O_{24}]6\sim$, into LDHs has received considerable attention.33,34,36,38,74,104,111h113 Such polyoxometalate LDHs may represent a class of pillared materials for selective adsorption and oxidation catalysis. Although the direct synthesis of an LDH containing polyoxo- metalate anions has recently been reported,36,111 earlier work by Drezdzon33 demonstrated that the incorporation of these inorganic anions could be achieved *via* an intermediate MgAl(terephthalate) LDH. Acidifying the organo-LDH in the presence of NaVO, for example, leads to the simultaneous polymerization of the monovanadate and displacement of the terephthalate from the interlayer, resulting in the incorpor- ation of the polyoxovanadate ([V O]6~) species. The use of a terephthalate LDH intermediate avoids the problems that may be associated with exchanging a small anion such as nitrate with a large polyoxometalate anion.

Evans et al.39 recently investigated the incorporation of

polyoxometalate anions into LDHs *via* directly synthesized LDH precursors containing terephthalate, *p*-methylbenzoate or *p*-hydroxybenzoate. The most suitable organic anion forexchange with the polyoxometalate was found to be *p*-methylbenzoate.

There has been increasing interest44,56,76,77,114 in support- ing metallophthalocyanines on LDHs, owing to the fact that these macrocyclic complexes can function as biomimetic catalysts for the autoxidation of organic molecules in aqueous solution. Cobalt(II) phthalocyaninetetrasulfonate ([CoPcTS]4 \approx), for example, is active for the autoxidation of a thiolate to a disulfide.77 Incorporation of the phthalocyanine complex into a MgAl LDH was found to significantly improve the catalyst reactivity (Fig. 8) and longevity for this reaction, compared to the homogeneous catalyst.77 In addi- tion, the use of an LDH support a†ords easy removal of the catalyst from the reaction medium. The same material is also active for the autoxidation of 2,6-di-*tert*-butylphenol to the corresponding diphenoquinone,76 and the reductive dehaloge- nation of certain halogenated organic compounds.114 In addi-tion, Shannon *et al.*46 have recently studied the oxidation of cyclohexene over a ZnAl LDH containing [CoPcTs]4 \sim . Metallophthalocyanine complexes supported on LDHs, there- fore, have a potential application in the treatment of waste water *via* the degradation of organic pollutants. Similar

3



Fig. 8 Dioxygen uptake plots for the autoxidation of 1.5] 10~2 M 1-decanethiol at 35 ¡C in the presence of [CoPcTs]4~ as catalyst: (*a*) homogeneous catalyst, (*b*) MgAl([CoPcTs]4~). The reactions were carried out at a substrate/cobalt ratio of 154 and pH 9.25 (after Per- ezbernal *et al.*77) materials based on the incorporation of metalloporhyrins into LDHs have also been investigated.44,53,54,115

The use of organo-LDHs as precursors to LDHs with enhanced thermal stability has recently been reported.116 Thelayer structure of a MgAl(CO) LDH prepared *via* reconstruction of a calcined organo-LDH precursor in Na CO was found to collapse at a temperature approximately 100 ¡C above that of the layer structure of a MgAl(CO) LDH pre-pared *via* direct synthesis. The improved thermal stability of the layer structure was attributed to the presence of carb- onaceous residues in the calcined organo-LDHs.

4.2 Photochemistry

The photophysical and photochemical properties of inter- calation compounds, including smectites, zirconium phos- phates and LDHs have recently been reviewed by Ogawa and Kuroda.7 The interlayer region of LDHs provides a novel environment for photochemical reactions of photoactive mol- ecules. The photochemistry of cinnamate anions between the hydroxide layers of a MgAl LDH, for example, has also been studied.117 It was found that both photodimerization and photoisomerization of the cinnamate anions occurs between the layers.

The controlled photodimerization of a variety of unsatu-rated carboxylates between the hydroxide layers of a MgAl LDH has been reported by Takagi *et al.*118 *Syn* head-to-head cyclodimers were selectively formed in the irradiation of sodium cinnamates between the hydroxide layers, whereas two isomers of *syn* head-to-head and *syn* head-to-tail cyclo- dimers were formed for the case of phenylethenylbenzoates. The product selectivity was shown to be controlled by the packing of the anions in the interlayer. Furthermore, the same authors have recently communicated that the photo-dimerization products can to some extent be controlled by changing the Mg/Al ratio, and hence the packing density of the incorporated monomer, of the host LDH.119

The incorporation of photochromic molecules into LDHs may yield useful photoresponsive materials. Tagaya and coworkers120,121 have studied the photoisomerization of sul- fonated indolinespirobenzopyran (SP-SO \sim) to merocyanine (MC) in the interlayer region of a MgAl LDH. The photoiso- merization was found to be irreversible for a MgAl LDH con- taining SP-SO \sim due to the stability of MC in the polar environment of the hydroxide layers. In the presence of *p*- toluenesulfonate (PTS), however, the reversible photoisomerization between SP-SO \sim and MC in the interlayer region of the MgAl LDH was observed. Fig. 9 shows the proposed mechanism for the reversible photoisomerization between SP-SO₃ \sim and MC in the MgAl(PTS) LDH.



Fig. 9 Possible mechanism for the reversible photoisomerization between SP-SO ~ and MC in the interlayer of a MgAl(PTS) LDH (after Tagaya et al. 121)

The incorporation of anionic dyes into LDHs may yield materials with potential applications as pigments.122 However, only a few preparations of LDHs containing anionic dyes have been reported.28,122

Second harmonic generation is a frequency-doubling non- linear optical process. Cooper and Dutta123 have prepared a LiAl LDH intercalated with 4-nitrohippuric acid, which exhibits second harmonic generation: 532 nm radiation from incident 1064 nm radiation. The ability of the intercalated material to exhibit nonlinear optical properties is due to aperpendicular monolayer packing of the acid molecules in the interlayer. This orientation leads to an ordered arrangement of dipoles, giving rise to a bulk dipole moment in the solid. Crystals of the pure acid exhibit no frequency-doubling char- acteristics due to a centrosymmetric packing in the crystal.

An examination of a LiAl LDH containing myrisate anions as a support for photochemical assemblies has recently been performed by Robins and Dutta.55 The goal of the study was to assemble a LDH containing TiO_x in which the TiO_x par-

ticles are able to interact with an anion-exchanged porphyrin.

The incorporation of TiO_v in the interlayer was achieved by taking advantage of the hydrophobic nature of the organo-

LDH, *i.e.*, titanium butoxide was partitioned into the inter-layer, followed by hydrolysis under ambient conditions. A suggested application for such a system is in the photo- degradation of pollutants.

4.3 Adsorption

The hydrophobic nature and accessibility of the interlayer region of organo-LDHs makes these materials candidates for the adsorption of other organic molecules. A ZnCr LDH con- taining alkyl sulfate anions, for example, undergoes swelling perpendicular to the layers under *n*-alkyl alcohols or *n*- alkylamines.51 In addition, Kopka *et al*.90 have found that ZnCr LDHs containing surfactant anions have the ability to take up a large variety of small organic molecules into the interlayer such as ethylene glycol, propanediol and glycerol.

Novel membrane-like materials are formed upon the incorporation of organic anions into LDHs.107 These materials are of relevance to separation and membrane tech- nologies. A LiAl LDH containing myrisate or hexanoate anions, for example, has the ability to partition pyrene from a methanolÈwater solution containing the polycyclic aromatic (Fig. 10).124 No sorption of pyrene was observed for a LiAl(succinate) LDH, however, and this was attributed to a sieving e†ect of this compound towards the pyrene molecule. The sorption of polyaromatic hydrocarbons, such as pyrene, is important from an environmental perspective. It should be noted that calcined LDHs may also be used as absorbents of organic molecules in environmental applications.125 In addi- tion, the potential application of a LiAl(myrisate) LDH as a stationary phase in gas chromatography has been investi- gated, owing to the partitioning e†ect exhibited by this material.109 The adsorption properties of organo-LDHs may be utilized

for the preparation of new materials. The intercalation of C molecules into the hydrophobic interlayer of a MgAl(dodecylsulfate) LDH has recently been reported.126 It was found that heating the resulting compound under vacuum to decompose the dodecyl sulfate left C molecules sandwiched between thehydroxide layers, although the crystallinity of the sample waspoor.

A MgAl(terephthalate) LDH has been used for a novel method of studying the initial stages of the emulsifier-freeemulsion polymerization of styrene.82 The method involved trapping the negatively charge polystyrene oligomers from the polymerization reaction mixture. It was found that the tereph-thalate anions were rapidly exchanged for the charged oligo- mers.



Fig. 10 Isotherm data for the uptake of pyrene in 50% methanol \dot{E} water for (a) LiAl(myrisate) LDH, (b) LiAl(hexanoate) LDH (after Dutta and Robins.124)

4.4 Electrochemistry

The potential application of organo-LDHs as new modified electrodes has been investigated.127h134 Electroactive materials derived from LDHs containing organic electroactive molecules, such as anthraquinonesulfonates and *m*- nitrobenzenesulfonate, have been prepared by Mousty *et al.*133 The electroactive anion was exchanged into a ZnCr LDH and a thin film of the product deposited on a glassy carbon electrode. The electrochemical behaviour of the sub- strate was then studied using cyclic voltammetry. It was found that these materials have potential applications for new clay- modified electrodes. The potential of ZnCr and ZnAl LDHs containing the electroactive 2,2l-azinobis-3-ethylbenzo- thiazoline-6-sulfonate anion as clay-modified electrodes has also been established by Therias and Mousty.134 The electro- chemical behaviour of manganese porphyrins incorporated into a MgAl LDH has also briefly been studied.132

4.5 Miscellaneous

The surface modification of LDHs with organic molecules has recently been reported.135 It was found that the product of thereaction of a ZnAl(OH) LDH with an organic oxychloride is an esterified LDH, with the organic molecules attached directly to the cationic layer (*i.e.*, not through the surface OH



Fig. 11 Schematic representation of the interlayer structure of (*a*) ZnAl(OH) LDH, (*b*) surface modified ZnAl LDH after reaction of (*a*) with an acid chloride (after Morioka *et al*.135)

groups) (Fig. 11). It has been suggested that these materials are candidates for new functional materials and catalysts.135 Similarly, the preparation of modified MgAl LDHs, in which organophosphonate anions are connected directly to the cationic layer, has recently been reported.37,72

Finally, it is worth noting that organo-LDHs have been

linked to prebiotic chemistry and may, therefore, have had a role to play in the origin of life.68,136,137

5 Concluding remarks

In this review we have concentrated on the host-guest supra- molecular arrangements that form using layered doublehydroxides. The observed chemistry complements that report-ed for cationic clay systems. In addition to organic guests it is also possible to incorporate, as alluded to in the text, complex polyoxoanions. Particularly important is the way that the dis- position of the anions within the interlayer can be modified by changing the charge on the layers, the degree of hydration and the method of synthesis. The studies described here point to the possibility of creating novel inorganic Eorganic supramo- lecular arrangements with desired structural and chemical properties.

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