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1 Introduction

Ion-exchangeable layered metal oxides have received growing attention because they can be functionalized under soft conditions, providing access to a very wide variety of kinetic metastable phases, inaccessible via classical solid-state reactions.¹⁻³ Such new phases, very recently reviewed,⁴ range from complex new inorganic solids,⁵ organic-inorganic hybrid materials⁶ to 2D exfoliated nanosheets.⁷⁻⁹ They can present various interesting properties for instance in the field of energy (fuel cells, artificial photosynthesis, photovoltaics, batteries...) or in nanoelectronics (high-k dielectrics, ferroelectrics, multiferroics.).⁴

Among ion-exchangeable layered metal oxides, layered perovskites, Dion–Jacobson, Ruddlesden–Popper and Aurivillius phases, have probably been the most widely studied.^{1,10-15} Very recently, several reports have underlined the considerable progress offered by microwave activation for the functionalization of such layered perovskites, notably in terms of speed of the

Microwave-assisted functionalization of the Aurivillius phase $Bi₂SrTa₂O₉$: diol grafting and amine insertion vs. alcohol grafting†

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Microwave-assisted functionalization of the layered Aurivillius phase Bi₂SrTa₂O₉ by alcohols is thoroughly investigated. The grafting of linear aliphatic and bulky alcohols is studied as a function of the starting material, underlining the importance of the prefunctionalization of the layered perovskite, for instance by butylamine. In addition, the functionalization by α,ω -alkanediols is explored. α,ω -alkanediols bearing long alkyl chains ($n_c > 3$) adopt an unprecedented pillaring arrangement, whereas 1,3-propanediol and ethyleneglycol adopt a bilayer arrangement, only one out of the two hydroxyl groups being coordinated. Finally, the reactivities of alcohols and amines towards insertion are compared: the preferential reactivity of the two functional groups appears to be strongly dependent of the reaction conditions, and especially of the water content. This study is further extended to the case of amino-alcohol insertion. In this case, the amine group is preferentially bound, but it is possible to control the grafting of the alcohol moiety, thus going from a bilayer arrangement to a pillaring one. This work is of particular importance to be able to functionalize easily and rapidly layered oxides with elaborated molecules, bearing several different potentially reactive groups. EDGE ARTICLE
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reactions.6,16–¹⁸ In addition, apart from classical insertion or grafting reactions,¹⁶⁻¹⁸ microwave-assisted post-synthesis modification¹⁹ and exfoliation²⁰ have also been reported.

Nevertheless, despite the rapidity of the microwave approach compared to classical methods, the molecules which have been inserted up to now into the interlamellar space of layered perovskites remain essentially limited to relatively simple ones, which hence reduces the range of possible applications. As for amine insertion, we have recently shown that microwaveassisted reactions allowed to extend the variety of accessible amines, to aromatic or chiral ones for instance.¹⁸ As for grafting functionalization, either via classical heating, solvothermal or microwave-assisted reactions, a few Dion–Jacobson phases and Ruddlesden–Popper phases have been functionalized, essentially by simple alcohols (Table 1). To the best of our knowledge, only two examples describe the grafting of aliphatic linear polyols, namely (poly)ethylene glycols.^{13,21} In these reports, classical reactions are used, and only mono-grafting is observed. A more complicated alcohol (D-glucopyranose) has also been grafted into a Dion-Jacobson phase¹⁴ and into a Ruddlesden-Popper phase,²² and there again only one OH group (out of the five) reacts. Finally, only two examples of grafting with a molecule other than alcohol are reported (trifluoroacetic acid²³ and linear phosphonic acids¹⁵). Until now, there is one single report of grafting reaction in the protonated form of an Aurivillius phase.¹⁹

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In this article, we explore the microwave-assisted grafting of alcohols into a protonated Aurivillius phase $H_2Bi_{0.1}Sr_{0.85}Ta_2O_7$ (HST), derived from the Aurivillius phase $Bi₂SrTa₂O₉$ (BST). We more particularly investigate the influence of the starting reactant (HST or pre-functionalized HST) on the grafting of alcohols, including bulky alcohols. We report for the first time the bi-grafting of long alkyl diols in pillaring arrangement, whereas for short diols (ethylene glycol and 1,3-propanediol) only mono-grafting is observed. Finally, the comparison of the reactivity of $-NH₂$ and $-OH$ groups towards HST is also investigated, depending on the water content in the reaction, either for an intermolecular competition (using butylamine and ethanol) or for an intramolecular competition (using 5-aminopentan-1-ol). We describe in this article how the use of microwave-assisted reactions not only allows to speed up the insertion-grafting of aliphatic alcohols into a layered perovskite, but also provides new interesting tools to insert more

complicated molecules (diols and amino alcohols) and to control the preferential reactivity of the different functional groups.

2 Results and discussion

Details of the experimental procedures and methods, as well as results of the elemental analyses are given in ESI.†

2.1 Alcohol grafting and choice of the starting compound

It has been shown that some protonated forms of Dion–Jacobson phases, $HLaNb₂O₇·xH₂O$, $HCa₂Nb₃O₁₀·xH₂O$ and $HSr₂$ - $Nb₃O₁₀·xH₂O$, can react directly with *n*-alcohol to form grafted n -alkoxy derivatives.^{11,16,26} Yet, for Ruddlesden–Popper phases such a direct grafting of alcohol has been reported to be inefficient, either using classical heating^{10,27} or using microwave-

Table 1 Typical grafting reactions into layered perovskites

assisted reactions.¹⁷ HST $(H_2Bi_{0.1}Sr_{0.85}Ta_2O_7)$, the protonated phase derived from the Aurivillius phase $Bi₂SrTa₂O₉$ (BST) can react directly with n-alkylamines to form inorganic–organic hybrids either using classical heating²⁸ or using microwaveassisted reactions.¹⁸ Therefore it appeared worth trying to functionalize this phase by direct grafting of alcohol, but despite the use of drastic conditions in terms of temperature (up to 160 $^{\circ}$ C), duration (up to 3 days) and reaction conditions (classical heating, solvothermal or microwave activation), the direct grafting of ethanol into HST failed, providing only the unreacted starting compound. This result is not surprising since the same absence of reaction had already been reported for the structurally closely related Ruddlesden–Popper phases $\rm H_2$ CaTa \rm_2O_7 (ref. 17 and 27) and $\rm H_2L a_2Ti_3O_{10}.^{10}$

Yet, a solution to graft *n*-alcohols into the protonated doublelayered perovskite H_2 CaTa₂O₇, has been reported.^{17,27} It consists in using an intermediate pre-intercalation of H_2 CaTa₂O₇ with an n-alkylamine. Such a pre-intercalation strategy, already used for various lamellar hybrid materials²⁹⁻³⁶ has been used more scarcely for layered oxides.^{15,17,18,27} The easy accessibility to **HST** functionalized by *n*-alkylamines using microwave-activated reactions¹⁸ offers extended possibilities for the subsequent interlayer surface modification of this Ta-based layered perovskite with n -alcohols.

Three different intermediates have been chosen as starting materials: NH_3 -HST (HST intercalated by ammonium), C_2N -HST (HST intercalated by ethylamine) and C_4N -HST (HST intercalated by butylamine). The microwave-assisted $(130\,\mathrm{\degree C},\ 2\,$ h) grafting reactions were then performed using a large excess (more than 200 equivalents) of n -alcohols with respect to intermediates and $ca.$ 1.3 mass% of water with respect to *n*-alcohols.

First, the intermediate $NH₃$ -HST has been used as a starting material to react with *n*-alcohols (C_nOH, $n = 1, 2, 3$). Yet the XRD patterns and IR spectra of the reaction products show that no reaction occurs with ethanol and propanol (Fig. S1 and S2†). For methanol, the XRD pattern shows clearly a mixture of the starting material $NH₃$ -HST (interlamellar distance of 1.06 nm) and a new phase with an interlamellar distance of 1.23 nm, which corresponds to the effective grafting of methanol. This partial grafting is confirmed by IR spectroscopy (see ESI†). No improvement of these results could be obtained by changing the reaction conditions (temperature and duration), indicating that $NH₃$ -HST is not an ideal intermediate to perform grafting reactions with n-alcohols.

Fig. S3† shows the XRD patterns of the other intermediate tested, C_2N -HST, and its reaction products with *n*-alcohols $(C_nOH, n = 1, 2, 3, 4, 7, 12)$. For reaction with alcohols possessing relatively short carbon chains (C_nOH, $n = 1, 2, 3, 4$), the $(00l)$ reflections of C₂N-HST are no longer present in the final compounds, and new sets of $(00l)$ reflections appear. This is a positive signal indicating the occurrence of grafting reactions between *n*-alcohols (C_nOH, $n = 1, 2, 3, 4$) and **HST**. However, the compounds obtained after reactions with alcohols with longer carbon chains (C_nOH, $n = 7$, 12) are multiphasic and with a very low crystallinity, whatever the reaction conditions used.

Infrared spectroscopy (Fig. S4†) shows for all compounds the appearance of signals of asymmetric and symmetric stretching vibrations of CH_2 and CH_3 groups in the region 2800–

 3000 cm^{-1} . Yet, for the compounds obtained by reaction with C_7 OH and C_{12} OH, these signals are particularly weaker than expected for such long alkyl chains. In addition, the band in the region 1100–1150 cm^{-1} corresponding to the formation of the C-O-Ta motif³⁷ is visible for compounds with short alkyl chains but relatively weak for compound obtained with heptanol, or even absent for compound obtained with dodecanol.

It thus appears that the intermediate C_2N -HST can be used to graft alcohols with relatively short carbon chains (C_nOH , $n = 1$, 2, 3, 4), much better than the intermediate $NH₃$ -HST. Yet, for alcohols bearing longer carbon chain ($n = 7$, 12), no monophasic grafted products could be obtained.

Fig. 1a shows the XRD patterns of the last intermediate tested, C_4N -HST and its reaction products with *n*-alcohols $(C_nOH, n = 1, 2, 3, 4, 7, 12)$. The grafted products are denoted as

Fig. 1 (a) PXRD patterns of the starting material C_4N -HST and its reaction products with n-alcohols (C_nOH, $n = 1, 2, 3, 4, 7, 12$). (b) Left: Relationship between the number of carbon atoms in the aliphatic chain of n-alcohols and the interlayer distance of the corresponding grafted products (full line corresponds to the best linear fit). Right: Scheme of the bilayer arrangement of the mono-n-alkoxy-HST derivatives. (c) Corresponding IR spectra.

 C_n OH-HST (*n* represents the number of carbon atoms in *n*alcohols). After reactions, the XRD patterns show that the $(00l)$ reflections of C_4N -HST disappear, and new sets of $(00l)$ reflections appear. In contrast, the reflections, which are assigned to (100) and (110) in the XRD pattern of HST, are observed at the same positions (22.68° (3.92 Å) and 32.59° (2.74 Å) respectively, (CuK α 1 = 0.1540598 nm)) in all the obtained products, which indicates the retention of the inorganic perovskite-like slab structure.

Fig. 1b shows a linear relationship between the interlayer distance and the number of carbon atoms in n -alcohols. This relationship can be expressed as follows: $d_{001} = 0.24n + 1.04$. This linear relationship indicates that the conformation of nalkyl chains within interlayer spaces of the reaction products is similar. In addition, the slope, 0.24, is about twice the length of one $CH₂$ group in the case of an all-*trans* ordered *n*-alkyl chain (0.127 nm/carbon atom) which indicates a bilayer arrangement.²⁶ Similar results have been described for the intercalation of HST with *n*-alkylamines^{18,28} and for the grafting reactions of Ruddlesden–Popper phase H_2 CaTa₂O₇ (ref. 17 and 27) and of Dion–Jacobson phases $HLaNb₂O₇·xH₂O^{11,17}$ and $HCa₂Nb₃O₁₀$ - $\cdot xH_2O^{26}$ with *n*-alcohols.

Using the previous relationship, the tilt angle (with respect to the normal to the layers) can be estimated around 19 $^{\circ}$, which is the same as the one observed for the alkoxy derivatives of H_2 CaTa₂O₇.^{17,27} This tilt angle is larger than that of C_nOH- $\rm H_2 La_2 Ti_3 O_{10}$ (15°) which can be attributed to larger lattice parameters.10,27 Finally, there is a considerable difference between the tilt angle of alkoxy derivatives of HST (C_nOH-HST, 19 $^{\circ}$) and the one of the corresponding alkylamine intercalated **HST** (C_n **N-HST**, 40°).¹⁸ This can be attributed to the different bonding modes, electrostatic in the case of amines, covalent in the case of alcohols.

Fig. 1c shows the IR spectra of the starting material C_4N -HST and its reaction products with *n*-alcohols $(C_nOH, n = 1, 2, 3, 4, 7,$ 12). Comparing IR spectra of reaction products with that of the starting material, one notices the disappearance of the signals between 1530 and 1630 $\mathrm{cm}^{-1},$ which belong to the intermediate C4N-HST. ¹⁸ The appearance of asymmetric and symmetric stretching bonds of $CH₂$ or the stretching bonds of $CH₃$ groups $(2800-3000 \text{ cm}^{-1})$ is clearly observed, and one can also notice an intense absorption around 1145 cm^{-1} attributed to the C–O vibration in the C–O–Ta group. Comparing with the corresponding *n*-alcohols (Fig. S5†), there is an obvious blue shift (*ca.* 95 $\rm cm^{-1})$ of the C–O band with respect to the corresponding free

alcohol (ca.1050 cm⁻¹). This blue shift is attributed to the coordination of the alcohol.^{37,38} In addition, the disappearance of stretching bonds of OH groups around 3300 cm^{-1} in the obtained products also indicates the success of the grafting reactions. The above analyses are in accordance with the removal of butylamine and the formation of a covalent bond C– O–Ta.

The major IR features are collected in Table 2. The asymmetric stretching bond of the $CH₃$ groups is visible between 2950 and 2970 cm⁻¹ for C_nOH-HST ($n = 2, 3, 4, 7$ and 12) and hardly visible at around 2919 $\rm cm^{-1}$ for $\rm C_1OH\text{-}HST.^{\text{11,39}}$ For all compounds, the CH₃ symmetric stretching band is masked by the more intense CH_2 stretching vibrations, except for C_1OH -HST for which this CH_3 symmetric stretching band is 2817 cm⁻¹. The values of $v_{\rm as}$ and $v_{\rm s}$ for CH₂ group observed for C_7 OH-HST and C_{12} OH-HST correspond to all-trans conformation.⁴⁰ When the length of the alkyl chains decreases, the frequencies of these bands increase as expected in the case of the presence of some *gauche* conformations,⁴⁰ yet, the ratio of gauche conformation is too low to have an influence on the basal spacing, as evidenced by the linear variation of the basal spacing (Fig. 1b). The signals at around 1470 cm^{-1} and 1350 cm^{-1} are attributed to C-H bend and C-H rock, respectively. Ta–O elongation vibration is the dominant feature at 580 cm^{-1} . This band is slightly shifted depending on the samples, without any clear correlation with the functionalization. Finally, no explanation was found for the presence of two signals (591 and 524 cm^{-1}) in the Ta-O elongation vibration region for C_1 OH-HST. Edge Article

C_oOH-HSN (*a* represents the number of carbon atoms in n should (az 105 om⁻¹). In this included to the

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SEM observation for all compounds shows the same morphology of crystallites, inhomogeneous in size, with strati fication typical of lamellar compounds (see Fig. S6† for representative examples).

In order to further demonstrate the formation of covalent bond C–O–Ta, solid state NMR spectroscopy was employed. As illustrations, the solid-state ¹³C CP/MAS NMR spectra of C_1OH -HST and C_4 OH-HST are presented Fig. 2. The chemical shift of the α -carbon (-C–O–) of C₁OH-HST and C₄OH-HST are 60 and 74 ppm, respectively. Comparing with the chemical shift of the α -carbon of methanol and butanol in the liquid-state 13 C NMR spectra (50.0 and 62.3 ppm, respectively⁴¹), there is a downfield shift of about 10 ppm. This downfield shift indicates unambiguously the formation of the covalent bond C–O–Ta in the reaction products.10,13,19,27

Product	$v_{\rm as}$ (CH ₂) (cm ⁻¹)	$v_{\rm s}$ (CH ₂) (cm ⁻¹)	$v_{\rm as}$ (CH ₃) (cm ⁻¹)	$v_{\rm s}$ (C-O) (cm ⁻¹)	$v_{\rm s}$ (Ta-O) (cm ⁻¹)
C_1 OH-HST			2919	1147	591, 524
C_2 OH-HST	2926	2869	2968	1143	576
C_3 OH-HST	2933	2871	2960	1140	565
C_4 OH-HST	2931	2869	2957	1139	573
C_7 OH-HST	2919	2851	2955	1150	568
C_{12} OH-HST	2916	2848	2957	1144	579

Table 2 Positions of the IR absorption bands of the CH₂, CH₃, C–O and Ta–O groups in C_nOH-HST compounds

Fig. 2 Solid-state 13 C CP/MAS NMR spectra of C₁H-HST (red) and C_4 OH-HST (cyan)

The thermal behaviors of the grafted products have been studied with the help of thermo gravimetric and thermo differential analyses (TGA-TDA) (Fig. S7†). The TGA and TDA curves show small endothermic mass losses, of about 0.4–1.0%, before 120 $^{\circ}$ C, which are ascribed to the removal of water. It is worth noticing that there is no visible loss of free alcohol, which confirms the effective grafting of the alcohol molecules. Between 200 $^{\circ} \text{C}$ and 400 $^{\circ} \text{C},$ the obvious mass losses, associated to exothermic peaks are ascribed to the decomposition of the nalkoxy groups within the interlayer space of the layered perovskite. The XRD pattern of the compound obtained after heating at 450 °C (Fig. S8†) is very similar to the one of **HST**, suggesting that HST is recovered at this temperature. By analogy with what has been described on $\text{H}_2\text{SrTa}_2\text{O}_7$, the XRD pattern at 800 $^\circ\text{C}$ indicates the formation of a three-dimensional perovskite of formula ${\rm Bi}_{0.1}{\rm Sr}_{0.85}{\rm Ta}_2{\rm O}_6$.^{42,43} It appears that the layered perovskite structure of HST is more thermally stable when the perovskite has been grafted by alcohols. Operation Science

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Combining the above analyses and the results of elemental analyses, the formulae of the grafted products are collected in Table S1.[†] Amounts of *n*-alkoxyl groups per **HST** are around 1 per Ta₂ unit (except for methoxy), smaller than 2, the theoretical maximum value based on the chemical formula $H_{2,0}Bi_{0.1}Sr_{0.85}$ Ta₂O₇. This phenomenon has been well explained by steric reasons in a previous report concerning the insertion of amines into $HST²⁸$ and the grafting of alcohols into the Ruddlesden– Popper phase $H_2 La_2 Ti_3 O_{10}$.¹⁰ As a result, only half of the reaction sites can be occupied. However, the amount of $CH₃O$ group per HST in C_1O -HST is above this limit, around 1.5. It is noteworthy that this "excess" of methanol has already been reported for the grafting of alcohols into the Ruddlesden– Popper phase $\text{H}_{2}\text{CaTa}_{2}\text{O}_{7}$.²⁷ This dense arrangement of CH₃Ogroup on the interlayer surface of the perovskite-like slabs may be ascribed to the smaller cross-section area of methanol (0.126 nm²),⁴⁴ compared to the one all-trans aliphatic chains, estimated to be 0.186 nm².⁴⁵

The above study shows that the direct grafting reactions between HST and *n*-alcohols are totally inefficient and that the use of an intermediate is necessary. According to the work of van der Voort et al., primary amines are known as catalysts to promote the silylation of silica surface and formation of Si–O–Si linkage by acido-basic catalysis.⁴⁶ But the fact that the intermediate $NH₃$ -HST cannot react with *n*-alcohols (except partially with methanol) proves that the only presence of $\mathrm{NH}_3^{\ +}$ group is insufficient to promote the occurrence of grafting reactions in HST. *n*-alkylamine-HST hybrids are indeed efficient intermediates (provided the alkyl chain of the inserted amine is long enough). When using these intermediates (such as C_4N -HST), the microwave-assisted reactions are indeed much faster than the classical solvothermal ones (Table 1). For instance, the grafting of aliphatic mono-alcohols into a Ruddlesden-Popper phase H_2 CaTa₂O₇ (very similar to the protonated Aurivillius phase HST, $H_2Bi_{0.1}Sr_{0.85}Ta_2O_7$ lasts around 5 days using classical solvothermal approach.²⁷ The general sketch of the synthetic procedure is given in Scheme 1.

2.2 Alcohol exchange reaction

Various alkoxy derivatives of HLaNb₂O₇ xH₂O, HCa₂Nb₃O₁₀- $\cdot xH_2O$ and $H_2La_2Ti_3O_{10}$ can be prepared by alcohol-exchangetype reactions with an n -alkoxy derivative as starting material to react with various alcohols.10,13,26 Therefore, following the results described above, it appears worth trying to prepare *n*alkoxy derivatives of HST using microwave-assisted alcoholexchange-type reactions.

Among the obtained *n*-alkoxy derivatives, C_2 OH-HST was chosen as a starting material to react with some n -alcohols $(C_nOH, n = 3, 4, 7, 12)$ with the help of microwave irradiation $(130 \text{ °C}, 2 \text{ h})$. The grafting reactions were performed using a large excess (more than \times 200) of *n*-alcohols with respect to the intermediate C_2 OH-HST, using a small amount of water (addition of $ca.$ 1.3 mass% with respect to *n*-alcohols).

The XRD patterns and IR spectra of the reaction products of C₂OH-HST with *n*-alcohols (C_nOH, $n = 3, 4, 7, 12$) are identical to the ones obtained by using C_4N -HST as an intermediate (Fig. S9†). The insertion-grafting rates are also similar to the ones determined above. Therefore, we can safely ascertain that n -alkoxy derivatives of HST can be prepared by alcohol exchange from C_2 OH-HST, and that the obtained compounds are identical to the ones obtained by using C4N-HST as an intermediate. Considering the observed crucial role of water (without water the exchange reaction described above are inefficient), the relatively easiness to hydrolyse alcohol-functionalized layered perovskites,^{13,19} and by analogy with what has been described in the literature for alcohol exchange in other layered perovskites,^{10,13,15,26} we propose a hydrolysis-etherification mechanism to account for alcohol exchange reactions.

Scheme 1 Scheme of the general synthetic procedure.

2.3 Grafting of bulky alcohols

In order to further expand the scope of the microwave-assisted grafting reactions between HST and alcohols, we have tested some more bulky alcohols, other than linear aliphatic alcohols. As relatively bulky alcohols, propan-2-ol, tert-butanol and benzyl alcohol have been employed. The microwave assisted (130 $^{\circ} \mathrm{C},$ 2 h) grafting reactions were performed using a large excess (more than 200 times) of bulky alcohols, with respect to the intermediate C_4N -HST, and a small amount of water (addition of ca. 1.3) mass% with respect to alcohols).

After reaction with propan-2-ol, tert-butanol and benzyl alcohol, the interlayer distances change from that of C_4N -HST (2.05 nm) to 1.57 nm, 1.72 nm and 2.06 nm, respectively (Fig. 3). In contrast, the reflections assigned to (100) and (110) are observed at the same positions (22.68° (3.92 Å) and 32.59° (2.74 $\,$ A) respectively, $(CuK\alpha 1 = 0.1540598 \text{ nm})$.

Like for the insertion of aliphatic alcohols, IR spectra confirm the grafting of the bulky alcohols, essentially by the disappearance of the stretching OH vibration, and by the blue shift of the C–O vibration (see Fig. $S10\dagger$). The formulae of the obtained compounds, established from elemental analyses and Thermo-Gravimetric Analyses (TGA) (Fig. S11†) are presented on Table S2.† The thermal behavior of the last compound, tertbutanol-HST is not reported here, because the results of elemental analyses show the significant presence of nitrogen (which is not the case for the other two compounds). It proves that in this case, the grafting is not complete, even using harder conditions (130 °C, 48 h, classical oven; 160 °C, 2 h, microwave). We attribute this difficulty to the bulkiness of tert-butanol, close to the anchoring function. On the contrary, elemental analyses for propan-2-ol-HST and benzyl alcohol-HST show no trace of nitrogen, confirming the complete removal of C_4N . Edge Article

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It is noteworthy that, likely due to steric hindrance, the organic contents are 0.6 and 0.7 alcohol per Ta₂ unit for 2-propanol-HST and benzyl alcohol-HST respectively, smaller than the value obtained for aliphatic linear n -alcohol-HST (ca. 1.0).

2.4 Grafting of α , ω -alkanediols

The reactivity of HST with α , ω -alkyldiamines has been demonstrated, using either classical heating⁴⁷ or microwaveassisted reaction,¹⁸ but to the best of our knowledge, the

Fig. 3 Left: PXRD patterns of C_4N -HST and its reaction products with propan-2-ol, tert-butanol and benzyl alcohol. Right: As an example, proposition of arrangement of benzyl alcohol.

reactivity of a layered perovskite towards diols has been described only twice, with the grafting of (poly)ethylene-glycols into a Dion-Jacobson phase $HLaNb₂O₇$.^{13,21} We thus explored the possibility to functionalize HST by α, ω -alkanediols, using microwave activated reactions. C_4N -HST or C_2N -HST can be used indifferently as a starting material to react with α, ω -alkanediols ($HOC_nH_{2n}OH$, $n = 2, 3, 4, 7, 8, 9, 12$).

The microwave assisted (130 $^{\circ}$ C, 2 h) grafting reactions were performed using a large excess (more than \times 200) of α , ω -alkanediols with respect to the intermediate C_4N -HST, using a small amount of water (addition of ca . 13 mass% with respect to n alcohols).

Fig. 4a shows the XRD patterns of C_4N -HST and of the reaction products. After reacting with ethylene glycol, 1,3-propanediol and 1,4-butanediol, the interlamellar distances decrease from that of C_4N -HST (2.05 nm) to 1.52 nm, 1.56 nm and 1.59 nm, respectively. For longer diols, the interlamellar distances of the grafted products increase regularly with the increase of the number of carbon atoms in alkane-diols.

The relation between the interlayer distance and the number of carbon atoms in aliphatic linear α , ω -alkanediols is plotted in Fig. 4b. For the longer diols ($n = 4, 7, 8, 9$ and 12), there is a clear linear relationship between the interlayer distance and the number of carbon atoms in the aliphatic chain of α , ω -alkanediols. The relation can be expressed as follows: $d_{001} = 0.08n +$ 1.26. This linear relationship indicates that the conformation of n-alkyl chains in the interlayer space of the reaction products is identical for all compounds with $n \geq 4$. The slope provides an estimation of the tilt angle of the alkyl chains with respect to the normal to the inorganic layers around 51 $^{\circ}$ (Fig. 4b right). This tilt angle is much larger than the one obtained for aliphatic mono-alcohols $(19^{\circ}, \text{ see above})$ and larger than the one obtained for α , ω -alkanediamines (30°).^{18,47} Yet it is in line with the one observed for the grafting of aliphatic mono-alcohols into Nb-based layered Dion-Jacobson phases.^{11,17,26}

It is worth noticing that $C_2(OH)_2$ -HST and $C_3(OH)_2$ -HST present an interlayer distance similar to that of C_2 OH-HST $(1.52 \text{ nm} \text{ vs. } 1.55 \text{ nm})$ or slightly smaller than that of $C_3OH-HST$ (1.56 nm vs. 1.77 nm) respectively, whereas the interlamellar distance of hybrids with diols with longer alkyl chains ($n \ge 4$) is significantly smaller than that of the corresponding monoalcohols hybrids. For instance, the interlamellar distance $C_4(OH)_2$ -HST (1.59 nm) is much smaller than the one of C_4OH -HST (1.94 nm). Moreover, the interlamellar distance of $C_4(OH)_2$ -HST is similar to that of C_4N_2 -HST (1.56 nm), for which 1, 4butanediamine has a pillaring arrangement.^{18,47}

Fig. 4c shows IR spectra of ethylene glycol, 1,3-propanediol, 1,4-butanediol, $C_2(OH)_2$ -HST, $C_3(OH)_2$ -HST and $C_4(OH)_2$ -HST. The disappearance of the signal of $-CH_3$ group at 2962 cm⁻¹ from the starting material C_4N -HST is clearly observed, confirming the total removal of butylamine. The spectra of $C_2(OH)_2$ -HST and $C_3(OH)_2$ -HST present strong absorptions at the same positions as in ethylene glycol and propanediol respectively: 3296 cm^{-1} , ascribed to free -OH group and 1086 cm^{-1} ascribed to C–O group. This indicates the existence of free –OH groups in $C_2(OH)_2$ -HST and $C_3(OH)_2$ -HST. In addition, the appearance of a new signal at 1149 cm^{-1} indicates the formation of covalent bond C–O–Ta between ethylene glycol and layered perovskite.^{37,38} On the contrary, the spectrum of $C_4(OH)_2$ -HST shows the disappearance of the signal of the –OH group (visible at 3290 cm^{-1} in the spectrum of 1,4-butanediol). In addition, there is a blue shift of about 100 cm^{-1} of the signal of the C–O group $(1149 \text{ cm}^{-1} \text{ for the hybrid and } 1048 \text{ cm}^{-1} \text{ for } 1,4\text{-butanediol})$, without remaining signal at 1048 cm^{-1} .^{37,38} The spectra of $C_n(OH)_2$ -HST ($n = 7, 8, 9, 12$) present the same characteristics (Fig. S12†).

The above analyses strongly suggest that $C_2(OH)_2$ -HST and $C_3(OH)_2$ -HST present a bilayer arrangement of the guest

Fig. 4 (a) PXRD patterns of the starting material C_4N -HST and its reaction products with α, ω -alkanediols (HOC_nH_{2n}OH, $n = 2, 3, 4, 7, 8$) 9, 12). (b) Left: Relation between the number of carbon atoms in the aliphatic chain of α , ω -alkanediols and the interlayer distance of the corresponding grafted products (full line corresponds to the best linear fit for the longer diols (see text)). Right: Schemes of the pillaring arrangement of α , ω -alkanediols with long alkyl chains (top) and of the bilayer arrangement of ethyleneglycol or of 1,3-propanediol. (c) IR spectra of ethylene glycol, 1,3-propanediol, 1,4-butanediol, $C_2(OH)_2$ -HST, $C_3(OH)_2$ -HST and $C_4(OH)_2$ -HST

species, with only one side of the diol grafted to the perovskitelike slab, the other OH group remaining free. $C_n(OH)_2$ -HST ($n \ge$ 4) conversely, present a pillaring arrangement of guest species with both sides of the diols grafted to the perovskite-like slab.

This analysis is supported by solid state NMR spectroscopy (Fig. 5). The solid state ¹³C NMR spectrum of $C_2(OH)_2$ -HST shows two signals at 75 and 64 ppm whereas the liquid state 13 C NMR spectrum of ethylene glycol shows only one signal at 63.7 ppm.⁴¹ For $C_3(OH)_2$ -HST, three signals are observed at 69, 59 and 37 ppm, whereas 1,3-propanediol presents only two signals at 61.7 and 34.1 ppm.⁴¹ Therefore, for both compounds, there is a downfield shift about 10 ppm for one α -carbon, while the other remains unchanged. This confirms that one alcohol group is coordinated to the perovskite layer, while the other remains free. The small signals at 36 and 14 ppm (marked with asterisks) are ascribed to residual C_2N (for this peculiar experiment, C_2N -HST was used as starting material instead of C_4N -**HST**). For $C_4(OH)_2$ -HST, only one signal for the α -carbon is observed, at 73 ppm, downfield shifted by about 10 ppm with respect to 1,4-butanediol (62.6 ppm), whereas the other signals (B and B') remain unchanged.⁴¹ Therefore, both oxygen atoms of 1,4-butanediol are coordinated in $C_4(OH)_2$ -HST. The small signal around 62 ppm (marked with asterisk) is ascribed to free –OH groups of 1,4-butanediol, coming from residual solvent, defects or 1,4-butanediol mono-grafted onto the surface of the crystallites.

According to proposed formulae, established from elemental analyses and Thermo-Gravimetric Analyses (TGA) (Table S3†), it is worth noticing that $C_2(OH)_2$ -HST and $C_3(OH)_2$ -HST contain about 0.8 and 0.6 diol respectively per formula unit, which, considering that the theoretical maximum number of hydroxyl group per Ta₂ unit is $1,11,13$ constitutes a further argument in favor of the monografting of ethylene glycol and propanediol.

The bilayer arrangement of $C_2(OH)_2$ and $C_3(OH)_2$ in the interlayer space of HST is in good accordance with previous report of monografting of ethylene glycol on the Dion-Jacobson phase HLaNb₂O₇ xH_2O ¹³ or of 1,2- and 1,3-propanediols in Kaolinite.⁴⁸ In the latter example, when largely increasing the reaction times, the two OH groups of 1,3-propanediol are coordinated, but to the same inorganic layer. Yet, to the best of our knowledge, the pillaring arrangement of α , ω -alkanediols in the interlayer space of layered perovskite has never been

Fig. 5 Solid-state 13 C CP/MAS NMR spectra of $C_2(OH)_2$ -HST (green), $C_3(OH)_2$ -HST (blue) and of $C_4(OH)_2$ -HST (cyan).

described in literature. The modification of Kaolinite with butanediols shows no pillaring arrangement.⁴⁹ The grafting reactions between $HLaNb₂O₇$ and polyethylene glycols (PEGs) with various molecular masses lead only to mono-grafted arrangements.²¹ Finally, the functionalization of $HLaNb₂O₇$ with D-glucopyranose has also been described as a mono-grafted arrangement.³⁷

2.5 Alcohol grafting vs. amine insertion

Considering the above-described results and other reports in the literature concerning the microwave-assisted functionalization of layered perovskites by relatively simple amines and alcohols,6,16–¹⁹ it appears interesting to investigate the compared reactivity of amines and alcohols, and how the reaction conditions are able to change the preferential reactivity of HST towards $-NH₂$ group or -OH group. This study is of great importance when times come for the insertion of very complicated molecules, bearing several functional groups. We focus here on the role of water.

A series of reactions between an equimolar mixture of butylamine (5.1 mL, 3.7 g, 0.04 mol) and ethanol (2.9 mL, 2.3 g, 0.04 mol) and with HST (50 mg), using a variable amount of water were thus carried out (microwave, 130 $^{\circ} \mathrm{C},$ 2 h). The amount of distilled water is 0 mass%, 0.1 mass% and 1.0 mass% with respect to the total mass of the reactive mixture.

Fig. 6 shows the XRD patterns of HST and its reaction products depending on the reaction conditions. As expected, when no water is used, the reaction does not take place, neither with the amine, nor with ethanol. With a small amount of water (0.1 mass%), the interlamellar distance increases from 0.98 nm (HST) to 1.74 nm (which is slightly larger than that of $C_2OH-HST$ (1.54 nm)). When the amount of water increases further (1 mass%), the interlamellar distance increases to 2.07 nm, which corresponds to the one of C_4N -HST.¹⁸

The corresponding IR spectra are shown in Fig. S13.† In the absence of water, the spectrum of HST is observed, as expected and in accordance with the corresponding XRD pattern. With 0.1 mass% of water, in addition to the signal coming from the alcohol essentially in the 1100 cm^{-1} region, the spectrum shows the antisymmetric and symmetric stretching bands of the $CH₂$ group but also bands associated to the C–N group around

Finally, the solid-state 13 C CP/MAS NMR spectra of the reaction products with 0.1 mass% and 1.0 mass% of water further confirm the above observations (Fig. 7). When 1.0 mass% of water is used, only signals at 40, 30, 20 and 14 ppm were obtained, confirming the formation of C_4N -HST. When a smaller amount of water is used, 0.1 mass%, the solid-state ¹³C CP/MAS NMR shows the signals which corresponds to the insertion of butylamine at 40, 30, 20 and 14 ppm, but also a signal at 68 ppm. This signal is attributed to the α -carbon (-C-O–) of ethanol, which is downfield shifted with respect to the corresponding signal for free ethanol in the liquid-state (57.8 ppm).⁴¹ This indicates the formation of the covalent bond C–O– Ta and confirms the co-insertion of ethanol and butylamine when only 0.1 mass% of water is used. Edge Article

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As presented above, ethanol does not react with HST. Therefore, the fact that it is possible to graft ethanol into HST when a mixture of ethanol and amine is used indicates that butylamine serves as a catalyst. Depending on the conditions, C_4N -HST is formed first, and then the amine is replaced by ethanol as described when C_2N -HST or C_4N -HST are used as starting compounds. Yet, when the amine is present in excess (and not only in the form of $C_nN\text{-HST}$), the formation of a pure $C₂OH-HST$ phase is not possible. When the water content is high, the preferential reactivity of HST towards amines leads to the formation of a C_nN -HST phase and no insertion-grafting of alcohol. One hypothesis to explain the inhibition of the reactivity of the alcohol moiety when the amount of water is high, is the reduction of the nucleophilic character of the hydroxyl group by solvation with water.

2.6 Amino-alcohol insertion

The above study of the role of water on the preferential reactivity of HST towards –NH₂ group or –OH group, was further extended

Fig. 6 PXRD patterns of HST and its reaction products with an equimolar mixture of butylamine and ethanol and 0 mass%, 0.1 mass% and 1 mass% of distilled water.

Fig. 7 Solid-state 13 C CP/MAS NMR spectra of the products resulting from the reaction between HST and an equimolar mixture of butylamine and ethanol and 0.1 mass% (red) and 1 mass% (black) of distilled water

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to the study of the reactivity of HST and related compounds $(C_2N$ -HST, C_4N -HST and C_2OH -HST) towards an aliphatic amino-alcohol, 5-amino-pentan-1-ol.

Fig. 8 shows the XRD patterns of C_2N -HST and its reaction products with 5-amino-1-pentanol, using different quantities of water. The interlayer distance of the obtained product increases from that of C_2N -HST (1.57 nm) to 1.61 nm, when the volumetric ratio water/THF is 1%. This distance corresponds to the expected one in case of a pillaring arrangement (see Fig. 4a for pillaring diols and ref. 18 for pillaring diamines). When the volume of water is 2% or 3% the one of THF, the interlayer distance is around 1.75 nm. And when the volume of water is between 10% and 100% the volume of THF, the interlayer distance further increases to around 1.96 nm. The observed variation of the interlayer distances is attributed to the change of the arrangement of 5-amino-1-pentanol in the interlayer space, from pillaring arrangement to bilayer arrangement.

These different arrangements are further confirmed and precised by IR spectroscopy (Fig. S14†). First it is worth noticing that the sharp strong bands at 3332 cm^{-1} and 3286 cm^{-1} due to NH stretching of free -NH₂ group are greatly reduced in intensity in the spectra of the hybrid compounds compared to the spectrum of 5-amino-1-pentanol. In addition, new bands at 1578 cm⁻¹ and 1545 cm⁻¹ appear in the spectra of the hybrids, and can be attributed to $-NH_3^+$ groups deformation vibrations.50,51 This indicates that whatever the amount of water, the amino moiety is protonated and thus interacts with the inorganic negatively charged layers. Second, the signal coming from the stretching of the C–O bond appears at *ca*. 1060 cm⁻¹ when the volume of water is more than 4% the one of THF, which indicates that the alcohol moiety is not coordinated, whereas for low water content (below 0.02 times the volume of THF), this band shifts to *ca.* 1130 cm^{-1} , which indicates a grafting of the alcohol moiety to the inorganic layers and thus a pillaring arrangement of 5-amino-1-pentanol. When the volume of water is 2% and 4% the one of THF, the coexistence of signals around 1130 cm^{-1} and around 1060 cm^{-1} indicates the coexistence of the two arrangements. Unexpectedly, this coexistence of the two Operation Science (or the transition of the transition of COMMIC concerts are allowed to the members are the properties of the common Commute Commute

arrangements does not lead to a multiphasic compound (Fig. 8), but instead leads to the formation of compounds with an intermediate interlayer distance.

¹³C CP/MAS NMR spectra of different compounds (obtained with 1, 2 and 100% of water respectively) confirm the previous analyses (Fig. 9). Indeed, ¹³C NMR spectrum of free 5-aminopentan-1-ol in solution in CDCl₃ presents five signals at 61.6 , 42.0, 33.3, 32.6 and 23.2 ppm.^{41 13}C CP/MAS NMR spectrum of the 5-amino-pentan-1-ol-HST hybrid obtained with 100% of water presents a signal at 62 ppm, with no evidence of a signal at lower field, indicating that the –OH group is not coordinated. On the contrary, for compounds obtained with 2 or 1% of water the signal of the C atom bearing the hydroxyl group is clearly down-field shifted with respect to free 5-amino-pentan-1-ol, indicating the deprotonation and coordination of the hydroxyl group.10,13,19,27 Yet a residual signal at 62 ppm shows the presence of a small amount of free hydroxyl groups (less pronounced when the quantity of water is decreased). The other signals are essentially unchanged, only slightly upfield shifted as expected due to amine protonation.^{18,47}

Combining the above analyses and the results of elemental analyses, the formulae of the obtained products as a function of the amount of water used are collected in Table S4.†

The same results are obtained when C_4N -HST is used instead of C_2N -HST. With these starting materials, it is thus possible to finely control the interlayer spacing of the hybrids and the arrangement 5-amino-pentan-1-ol by playing with the water content in the reaction mixture.

On the contrary, when using C_2 OH-HST or HST are used as starting materials, only the bilayer arrangement, with an interlamellar distance around 1.96 nm, can be obtained. In addition, it is worth noticing that the insertion reaction of 5 amino-pentan-1-ol is far more difficult; complete disappearance of the starting materials C_2 OH-HST or HST requires a much larger quantity of water (Fig. S15–S18†).

2.7 Post-treatment: from bilayer to pillaring arrangement

Fig. 8 Left: XRD patterns of C_2N -HST and its reaction products with 5-amino-1-pentanol using different quantities of water. Right: Scheme of the pillaring arrangement (bottom), and of the bilayer arrangement (top).

The bilayer arrangement of 5-amino-pentan-1-ol leads to the presence of free –OH groups within the interlayer space of HST. These groups have the potential to undergo further grafting reaction to lead to pillaring arrangement. It is worth

Fig. 9 Solid-state 13 C CP/MAS NMR spectra of the products resulting from the reaction between HST and 5-amino-pentan-1-ol with 1 mass% (red), 2 mass% (green) and 100 mass% (blue) of distilled water.

underlining that 5-amino-pentan-1-ol-HST in its bilayer form is stable during at least 10 days in ambient conditions (in air, around 20 °C and about 50% humidity). Yet, after a moderate heating (70 °C) during one day, the interlayer distance decreases from 1.98 nm to 1.70 nm and further decreases to 1.62 nm when the heating time is prolonged to two days. Longer heating times do not lead to any further change of the interlamellar distance (Fig. 10).

It is noteworthy that the distance which is obtained after heating (1.62 nm) is the same as the one obtained by the reaction between C_2N -HST and 5-amino-pentan-1-ol using a low quantity of water with respect to THF (Fig. 8) which corresponds to a pillaring arrangement of 5-amino-pentan-1-ol. The corresponding IR spectra further support the above hypothesis, of a temperature induced pillaring of 5-amino-pentan-1-ol (Fig. S19†). Notably, the C–O stretching vibration undergoes a clear blue shift from 1058 cm^{-1} (C–OH) to 1120 cm^{-1} (ascribed to $C-OTa$). Finally, the down-field shift of the solidstate 13 C CP/MAS NMR signal attributed to the C bearing the hydroxyl group confirms the effective grafting of the alcohol moiety upon heating (Fig. 11). Nevertheless, a small peak at 61.7 ppm, corresponding to free –OH group, is still visible in the spectrum of the compound after heating. One hypothesis is that this peak is due to 5-amino-pentan-1-ol molecules linked to the surface of the crystallites via the ammonium group, and for which the terminal –OH group remains free. The broadening of the resonance lines is likely due to the more rigid structure induced by the pillaring arrangement of 5-amino-pentan-1-ol. Edge Article

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Fig. 10 XRD patterns of 5-amino-pentan-1-ol-HST "bilayer" and of the products obtained after heating.

Fig. 11 Solid-state ¹³C CP/MAS NMR spectra of 5-amino-pentan-1ol-HST "bilayer" (red) and the product obtained after heating (cyan).

3 Conclusions

The successful microwave-assisted interlayer surface modification of the protonated form of an Aurivillius phase, $Bi₂SrTa₂O₉$ (BST) via grafting reactions with various alcohols $(n$ -alcohols, linear diols and bulky alcohols) has been achieved. One marking result is the pillaring arrangement of α , ω -alkanediols with long alkyl chains in the interlamellar spacing of layered perovskite which is evidenced for the first time.

Taking advantage of the rapidity of microwave assisted reaction, the influence of the reaction conditions has been investigated more precisely. The importance of the choice of the pre-intercalated material for the preparation of alkoxy derivatives the Aurivillius phase has thus been underlined. In addition, it has been shown that water plays a key role not only to allow grafting of alcohols, but also to drive the preferential reactivity of the protonated perovskite with amines or alcohols. In the example chosen, 5-amino-pentan-1-ol, the amine group is preferentially bound, but it is possible to control the grafting of the alcohol moiety either by adjusting the quantity of water in the reaction medium, or by a post-reaction moderate heating.

These results will be most useful when times come for the insertion of more complicated molecules, bearing several functional groups, necessary for the precise design of functional hybrids with specific structures and properties. Such precise synthetic approaches can for instance lead to new luminescent activators, $52,53$ vapour sensors 54 or to a fine tuning of the ferroelectric properties of the starting **BST** phase.⁵⁵

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 R. E. Schaak and T. E. Mallouk, Chem. Mater., 2002, 14, 1455– 1471.
- 2 K. G. Sanjaya Ranmohotti, E. Josepha, J. Choi, J. Zhang and J. B. Wiley, Adv. Mater., 2011, 23, 442–460.
- 3 J. Gopalakrishnan, T. Sivakumar, K. Ramesha, V. Thangadurai and G. N. Subbanna, J. Am. Chem. Soc., 2000, 122, 6237–6241.
- 4 R. Uppuluri, A. S. Gupta, A. S. Rosas and T. E. Mallouk, Chem. Soc. Rev., 2018, 47, 2401–2430.
- 5 D. Montasserasadi, D. Mohanty, A. Huq, L. Heroux, E. A. Payzant and J. B. Wiley, Inorg. Chem., 2014, 53, 1773–1778.
- 6 S. Akbarian-Tefaghi and J. B. Wiley, Dalton Trans., 2018, 47, 2917–2924.
- 7 L. Wang and T. Sasaki, Chem. Rev., 2014, 114, 9455–9486.
- 8 B.-W. Li, M. Osada, Y. Ebina, S. Ueda and T. Sasaki, J. Am. Chem. Soc., 2016, 138, 7621–7625.
- 9 R. Ma and T. Sasaki, Adv. Mater., 2010, 22, 5082–5104.
- 10 S. Tahara, T. Ichikawa, G. Kajiwara and Y. Sugahara, Chem. Mater., 2007, 19, 2352–2358.
- 11 S. Takahashi, T. Nakato, S. Hayashi, Y. Sugahara and K. Kuroda, Inorg. Chem., 1995, 34, 5065–5069.
- 12 Y. Takeda, T. Momma, T. Osaka, K. Kuroda and Y. Sugahara, J. Mater. Chem., 2008, 18, 3581–3587.
- 13 H. Suzuki, K. Notsu, Y. Takeda, W. Sugimoto and Y. Sugahara, Chem. Mater., 2003, 15, 636–641.
- 14 C. Wang, K. Tang, D. Wang, Z. Liu, L. Wang, Y. Zhu and Y. Qian, J. Mater. Chem., 2012, 22, 11086–11092.
- 15 A. Shimada, Y. Yoneyama, S. Tahara, P. H. Mutin and Y. Sugahara, Chem. Mater., 2009, 21, 4155–4162.
- 16 J. R. Boykin and L. J. Smith, Inorg. Chem., 2015, 54, 4177– 4179.
- 17 S. Akbarian-Tefaghi, E. Teixeira Veiga, G. Amand and J. B. Wiley, Inorg. Chem., 2016, 55, 1604–1612.
- 18 Y. Wang, E. Delahaye, C. Leuvrey, F. Leroux, P. Rabu and G. Rogez, Inorg. Chem., 2016, 55, 4039–4046.
- 19 Y. Wang, E. Delahaye, C. Leuvrey, F. Leroux, P. Rabu and G. Rogez, Inorg. Chem., 2016, 55, 9790–9797.
- 20 S. Akbarian-Tefaghi, T. Rostamzadeh, T. T. Brown, C. Davis-Wheeler and J. B. Wiley, ChemNanoMat, 2017, 3, 538–550.
- 21 S. Hotta, N. Idota and Y. Sugahara, Key Eng. Mater., 2014, 616, 82–86.
- 22 Y. Wang, X. Zhu, X. Li, L. Wang, Y. Wang, Q. Hao and K. Tang, J. Mater. Chem. A, 2014, 2, 15590–15597.
- 23 S. Yoshioka, Y. Takeda, Y. Uchimaru and Y. Sugahara, J. Organomet. Chem., 2003, 686, 145–150.
- 24 Y. Asai, Y. Ariake, H. Saito, N. Idota, K. Matsukawa, T. Nishino and Y. Sugahara, RSC Adv., 2014, 4, 26932–26939.
- 25 Y. Takeda, H. Suzuki, K. Notsu, W. Sugimoto and Y. Sugahara, Mater. Res. Bull., 2006, 41, 834–841.
- 26 S. Tahara and Y. Sugahara, Langmuir, 2003, 19, 9473–9478.
- 27 Y. Wang, C. Wang, L. Wang, Q. Hao, X. Zhu, X. Chen and K. Tang, RSC Adv., 2014, 4, 4047–4054.
- 28 Y. Tsunoda, W. Sugimoto and Y. Sugahara, Chem. Mater., 2003, 15, 632–635.
- 29 T. Coradin, R. Clément, P. G. Lacroix and K. Nakatani, Chem. Mater., 1996, 8, 2153–2158.
- 30 P. G. Lacroix, R. Clément, K. Nakatani, J. Zyss and I. Ledoux, Science, 1994, 263, 658–660.
- 31 J. L. Colon, C. Y. Yang, A. Clearfield and C. R. Martin, J. Phys. Chem., 1988, 92, 5777–5781.
- 32 J. L. Colon, C. Y. Yang, A. Clearfield and C. R. Martin, J. Phys. Chem., 1990, 94, 874–882.
- 33 Q. Wang, D. Yu, Y. Wang, J. Sun and J. Shen, Langmuir, 2008, 24, 11684–11690.
- 34 G. Huang, S. Ma, X. Zhao, X. Yang and K. Ooi, Chem. Commun., 2009, 331–333.
- 35 E. Delahaye, S. Eyele-Mezui, J.-F. Bardeau, C. Leuvrey, ´ L. Mager, P. Rabu and G. Rogez, J. Mater. Chem., 2009, 19, 6106–6115.
- 36 E. Delahaye, S. Eyele-Mezui, M. Diop, C. Leuvrey, P. Rabu ´ and G. Rogez, Dalton Trans., 2010, 39, 10577–10580.
- 37 C. Wang, K. Tang, D. Wang, Z. Liu, L. Wang, Y. Zhu and Y. Qian, J. Mater. Chem., 2012, 22, 11086–11092.
- 38 M. V. Korolevich, R. G. Zhbankov and V. V. Sivchik, J. Mol. Struct., 1990, 220, 301–313.
- 39 F. C. Cruz, A. Scalabrin, D. Pereira, P. A. M. Vazquez, Y. Hase and F. Strumia, J. Mol. Spectrosc., 1992, 156, 22–38.
- 40 R. A. Vaia, R. K. Teukolsky and E. P. Giannelis, Chem. Mater., 1994, 6, 1017–1022.
- 41 AIST, Spectral Database for Organic Compounds, SDBS, http://sdbs.db.aist.go.jp/sdbs/cgi-bin/direct_frame_top.cgi, (accessed April 13, 2018). Openical Science

3. Copyright Content Article 2011, 1. Copyright Content Article 2012, 11 AM. This article is licensed und C. R. Martin, Phys.

4. Upyrightin, A. S. Guynn, A. S. Boston M. L. Milledon, S. Commons A. This a
	- 42 M.-P. Crosnier-Lopez, F. L. Berre and J.-L. Fourquet, J. Mater. Chem., 2001, 11, 1146–1151.
	- 43 P. J. Ollivier and T. E. Mallouk, Chem. Mater., 1998, 10, 2585– 2587.
	- 44 Y. Marcus, The properties of solvents, John Wiley & Sons Ltd., Chichester, 1998.
	- 45 A. Goñi, J. Rius, M. Insausti, L. M. Lezama, J. L. Pizarro, M. I. Arriortua and T. Rojo, Chem. Mater., 1996, 8, 1052–1060.
	- 46 P. van der Voort and E. F. Vansant, J. Liq. Chromatogr. Relat. Technol., 1996, 19, 2723–2752.
	- 47 S. Tahara, T. Yamashita, G. Kajiwara and Y. Sugahara, Chem. Lett., 2006, 35, 1292–1293.
	- 48 T. Itagaki and K. Kuroda, J. Mater. Chem., 2003, 13, 1064–1068.
	- 49 J. Murakami, T. Itagaki and K. Kuroda, Solid State Ionics, 2004, 172, 279–282.
	- 50 K. Nakanishi, T. Goto and M. Ohashi, Bull. Chem. Soc. Jpn., 1957, 30, 403–408.
	- 51 R. A. Heacock and L. Marion, Can. J. Chem., 1956, 34, 1782– 1795.
	- 52 T.-S. Chan, C.-L. Dong, Y.-H. Chen, Y.-R. Lu, S.-Y. Wu, Y.-R. Ma, C.-C. Lin, R.-S. Liu, J.-L. Chen, J. Guo, J.-F. Lee, H.-S. Sheu, C.-C. Yang and C.-L. Chen, J. Mater. Chem., 2011, 21, 17119–17127.
	- 53 S. Ida, C. Ogata, U. Unal, K. Izawa, T. Inoue, O. Altuntasoglu and Y. Matsumoto, J. Am. Chem. Soc., 2007, 129, 8956–8957.
	- 54 P. Ganter, L. M. Schoop, M. Däntl and B. V. Lotsch, Chem. Mater., 2018, 30, 2557–2565.
	- 55 N. C. Bristowe, J. Varignon, D. Fontaine, E. Bousquet and P. Ghosez, Nat. Commun., 2015, 6, 6677.