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Aqueous phase synthesis and electronic spectroscopy of nanostructured layered double hydroxides **†**

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Herein, we proposed a green synthetic route based on coordination chemistry to produce nanostructured layered double hydroxides (LDHs). The synthesized NiAl-, NiFe- and ZnAl-LDHs exhibited similar morphologies and crystal structures. A possible mechanism for LDH formation was proposed based on pHpotentiometric titrations. Extensive electronic spectroscopy analyses, interpreted through the lens of ligand field theory, provided key insights into the formation mechanism of NiFe-LDH.

Introduction 1.

Lavered double hydroxides (LDHs), also known as hydrotalcitelike compounds, are a class of natural¹ or synthetic² two-dimensional materials^{3,4} that find many applications as ion exchangers, plastic additives, flame retardants and catalysts for steam reforming, since the last century.² Nowadays, transition-metal-based LDHs have garnered significant research attention due to their promising electrochemical properties for energy storage and conversion applications.⁵⁻⁷ Furthermore, recent hypotheses suggest their potential role in the origins of life on Earth.^{8,9} Layered double hydroxide compounds are structurally derived from layered divalent metal hydroxides (e.g., Mg(OH)₂) through the isomorphous substitution of a fraction of divalent metal species with trivalent ones.² The resulting lattice consists of stacked positively charged metal hydroxide layers separated by charge-counterbalancing anions and water molecules.² Hence, LDHs represent a broad family of layered materials with a common chemical composition, which can be summarized as follows:

$$\left[\mathbf{M}_{1-x}^{2+}\mathbf{M}_{x}^{3+}(\mathbf{OH})_{2}\right]^{x+}\mathbf{A}_{(x/n)}^{n-}\cdot \mathbf{y}\mathbf{H}_{2}\mathbf{O}$$
(1)

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Here, M^{2+} (e.g., Mg^{2+} , Ca^{2+} , Fe^{2+} , Ni^{2+} , ...) and M^{3+} (e.g., Al^{3+} , Cr^{3+} , Fe^{3+} , ...) represent divalent and trivalent metal species, respectively, which make up the hydroxide layers, while A^{n-1} refers to inorganic or organic anions (e.g., Cl⁻, NO₃⁻, CO₃²⁻, $R-COO^{-}$, ...) that are intercalated in the interlayer space.² The coefficient x is defined as $M^{3+}/(M^{2+} + M^{3+})$ and typically takes values in the range of 0.2 to 0.33.^{2,10} For simplicity, in this study, we refer to these compounds as M²⁺ M³⁺-LDHs. The significant flexibility in the chemical composition of LDHs has led to the development of several synthetic approaches.^{2,11-19} However, the most commonly exploited chemical pathway for the synthesis of LHDs involves the slow reaction between a trivalent metal hydroxide (i.e., M(OH)₃) and M²⁺ cations dissolved in an aqueous solution.^{11-14,19} Both the "urea hydrolysis" and co-precipitation methods are frequently used for this synthesis route.^{11-14,19} However, these procedures often require the use of autoclaves to achieve sufficiently high temperatures (typically around 150 °C) to ensure the complete conversion of M(OH)₃ into LDHs.¹¹⁻¹⁴ Moreover, the addition of chelating agents to the reaction mixture is pivotal for forming pure transition-metal LDH phases.¹² Another common method for LDH synthesis involves conducting the reaction at a constant pH (usually pH \approx 9 or 10) at which both metal ions precipitate simultaneously.² Nanostructured LDH materials (i.e., single- or few-layer nanosheets) can eventually be obtained through a top-down approach by delaminating the bulk LDH material synthetized using one of the aforementioned procedures.⁶ The top-down approach involves the liquid phase-exfoliation^{4,20,21} of bulk LDH materials into an appropriate solvent. The most suitable delaminating solvent for LDHs is the toxic formamide.^{6,22,23} Alternatively, several bottom-up approaches have been proposed for the direct synthesis of nanostructured LDH materials.^{15–18,24} More specifically, the



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[†]Electronic supplementary information (ESI) available: Chemicals used, synthesis procedure and experimental details, samples compositions, supplementary TEM images, supplementary XRD patterns and crystallographic data, supplementary pH-potentiometric titration profiles, speciation insights and affinity constants with references, ligand field analysis details and MATLAB code, comparison tables. See DOI: https://doi.org/10.1039/d4qi01475h

reverse microemulsion route has been used to synthesize nanostructured MgAl-LDHs, which can be easily incorporated into polymer matrices.¹⁵ However, the use of a water-isooctane emulsion raises concerns from both safety and environmental standpoints for practical applications. Tokudome et al. proposed a synthetic route for nanostructured NiAl- and NiFe-LDHs from an ethanol-water solution,^{17,18} but 1-chloro-2-propanol is produced as a byproduct,^{17,18,25} thus requiring a proper disposal being a halogenated compound. The aqueous miscible organic solvent treatment (AMOST) procedure yields nanostructured MgAl- and ZnAl-LDHs dispersible into nonpolar hydrocarbons.¹⁶ However, AMOST is only limited to tetraborate anions in the LDH interlayer space. Noteworthily, Hibino and Kobayashi reported the synthesis and delamination in the water of lactate-intercalated LDHs.²⁴ On the flip side, a massive amount of lactate (more than 20-fold the stoichiometric ratio) is required to achieve the synthesis of lactateintercalated LDHs.²⁴ Herein, we propose a fully aqueous and direct precipitation of nanostructured LDH compounds by exploiting the coordination chemistry of citric acid through an alternative synthetic route. Our synthetic approach is inspired by chemical bath deposition processes (CBD),^{26,27} which are widely utilized for metal oxide and sulphide film deposition for optoelectronic applications. Hence, our LDH synthesis process could be exploited for future production of mixed metal oxide films. The main advantage of our synthesis approach is that it is one-pot. In addition, different from other one-pot LDH synthesis approaches, the availability of metal cations, resulting from the hydrolysis of metal-citrate complexes, is the rate-limiting factor in the LDH formation process. This means that the stability constants $(\log \beta)$ of metal-citrate complexes dictate the availability of cations, influencing the nucleation of LDH nanosheets. Hence, this study focuses on the formation mechanism of citrate intercalated LDHs probed by pH-potentiometric titrations coupled to ligand field analysis of the electronic transitions. We demonstrate that our method is straightforwardly applicable to nickel-based LDHs, and can also be applied to ZnAl-LDHs, with some restrictions. So far, the formation mechanism of LDHs has been probed through pH titrations exclusively for all-inorganic syntheses.^{11,13,19} Jaśkaniec et al.¹² and Tokudome et al.¹⁷ performed pH vs. time characterizations of their LDH synthesis procedures, both of which exploit coordination chemistry. However, pH vs. time characterizations provide information only on the hydrolysis rate of the ammonium (or hydroxide) releasing agent used during the synthesis, and not on the actual formation mechanism of LDH materials. On the contrary, a pH titration allows external control of the equivalents of hydroxide ions supplied to the reaction environment, and therefore, provides insights into the reaction mechanism. These understandings can be useful in designing new or improving the existing synthetic processes. Herein, we report how citrate anions alter the formation mechanism of LDHs from the classical precipitation pathway to a hydrolytic polymerization of metal-citrate complexes. Lastly, by interpreting the electronic spectra of the reaction environment and the

final LDH product through the lens of the ligand field theory, we report the values of ligand field strength and Racah parameters of Ni²⁺ and Fe³⁺ in LDH crystal lattices. Then, we critically discuss the obtained results and possible implications by considering the current knowledge on electronic spectroscopy of nickel hydroxides, ferric oxides, and related materials.^{28–42}

2. Results and discussion

2.1. Morphology and crystal structure

We investigated the particle morphology of NiFe-, NiAl- and ZnAl-LDHs synthesized from metal-citrate complexes and then dispersed them in water. All dispersions were made up of lamellar particles (*i.e.*, nanosheets) that are \sim 5–20 nm wide in the case of NiFe-LDH (Fig. 1a), ~5 nm wide in the case of NiAl-LDH (Fig. S1a[†]) and ~200-500 nm wide in the case of ZnAl-LDH (Fig. S1b[†]). The nanosheets are rarely found isolated, since they form micron-sized structures also visible under scanning electron microscopy imaging (Fig. 1b and c). The presence of vertically aligned nanosheets NiFe-LDH and ZnAl-LDH in the HRTEM image (Fig. S1c[†]) enabled us to estimate their thickness, which is in the 1-2 nm range, in agreement with our previous work, in which we produced citrate-intercalated NiFe-LDH nanosheets by coprecipitation.43 On the contrary, the NiAl-LDH sample does not present vertically aligned nanosheets. The detailed study of the dependence of the morphology of LDHs on the synthetic conditions lies beyond the scope of this work. The X-ray diffraction patterns of the citrateintercalated NiAl- and NiFe-LDH powders are presented in Fig. 1d. Analogous diffraction patterns of NiAl- and NiFe-LDHs with variable metal ratios are presented in Fig. S2,† showing that our synthesis technique allows for a wide range of compositions in nickel-based LDH in the 0.14–0.33 range of x, being consistent with what was previously reported by B. Grégoire et al.¹⁰ On the flip side, the ZnAl-LDH phase appears to be feasible only for the x equal to 0.33, with some incidental ZnO contamination, as illustrated in Fig. 1e. While, as illustrated in Fig. 1f, the synthesis of ZnAl-LDH with an x value lower than 0.25, mainly leads to the formation of wurtzite type ZnO. For the sake of clarity, in LDH XRD patterns, we refer to the Miller indices of the hydrotalcite crystal structure (i.e., natural carbonate-intercalated MgAl-LDH, $R\bar{3}m$ space group). This indexing allows an easier comparison with the past literature patterns of the synthetic LDH materials and LDH minerals (Table S3[†]).^{1,44-46} The LDH diffraction patterns can be divided into three different regions. At scattering angles (2θ) below 30°, the "basal reflections" appear. These reflections are ascribed to the (00l) crystallographic planes, *i.e.*, planes parallel to the hydroxide layers and perpendicular to the unit cell c axis.⁴⁷ Thus, the 2θ position of the basal reflections reveals the unit cell c axis of LDHs (by the weighted mean of d_{00l} lattice parameters, as described in ESI section 4⁺). Our analyses provide the following c axis lengths for the citrate-intercalated LDHs: 3.669 nm for NiAl-LDH, 3.680 nm for NiFe-LDH, and 3.665 nm for ZnAl-LDH. The thickness of one hydroxide layer





Fig. 1 (a) High resolution transmission electron microscopy image of citrate-intercalated NiFe-LDH nanosheets. (b) Transmission electron microscopy image of the citrate-intercalated NiFe-LDH aggregates of nanosheets (a few isolated nanosheets are visible in the top left corner). (c) Scanning electron microscopy image of the lamellar micron-sized structures in a NiFe-LDH powder. (d) Powder X-ray diffraction pattern of the citrate-intercalated NiAl- and NiFe-LDH pellets. (e) Powder X-ray diffraction pattern of the citrate-intercalated ZnAl-LDH pellet synthesized with an initial Zn : Al ratio of 2 : 1. (f) Powder X-ray diffraction pattern of the citrate-intercalated ZnAl-LDH pellet synthesized with an initial Zn : Al ratio of 3 : 1. Reflection marker with an asterisk refers to wurtzite type ZnO (PANISC: 98-002-6170, $P6_3mc$ space group). The incident X-ray radiation used in XRD pattern is Cu K α .

plus one interlayered distance, called "basal spacing", is equal to one third the length of the *c* axis (in the, $R\bar{3}m$ space group).² These results are consistent with those from the past literature on citrate-intercalated MgAl-, NiAl- and ZnAl-LDHs synthetized by the anion exchange or the dissolution/precipitation routes.^{44–46} Instead, the second region, in the 2 θ range of 30° to 55°, displays reflections from (0*kl*) and (*h*0*l*) crystallographic planes.

The shape of these reflections is indicative of faults in the layer stacking.⁴⁸ In fact, sharp and well-defined reflections in this region are characteristics of highly crystalline LDHs.⁴⁸ On the contrary, our LDHs show an asymmetric broad reflection covering the whole $30-55^{\circ}$, 2θ region. This feature is indicative of a turbostratic disorder, *i.e.*, randomly slid and twisted layers,

also visible in Fig. 1a, in which two NiFe-LDH nanosheets are stacked on top of each other. Turbostraticity is a characteristic property of α -Ni(OH)₂,²⁹ LDHs intercalated with bulky anions,^{44,45,48–50} as well as other defective two-dimensional materials.^{51,52} The third region in the LDHs diffraction pattern is located at 2θ angles above 55°. This region contains the (110) reflection, from which the unit cell parameter *a* is obtained² (ESI section 4†). Crystallographic parameters of the citrate-intercalated LDHs are summarized in Table 1.

2.2. pH-potentiometric titrations

In this section, we describe the hydrolytic behaviour of monometallic and bimetallic solutions, with and without dissolved citrate, studied using pH-potentiometric titrations. Titration

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 Table 1
 Basal spacing and unit cell parameter dimensions of citrateintercalated LDHs calculated according to Bragg's law

Sample	x	Basal spacing (nm)	a axis (nm)	c axis (nm)
NiAl-LDH	0.25	1.223	0.3019	3.669
NiFe-LDH	0.25	1.227	0.3053	3.680
ZnAl-LDH	0.33	1.222	0.3056	3.665

curves are expressed as a function of the hydroxylation rate (R), *i.e.*, the added moles of hydroxide with respect to the total moles of metal cations. The total metal/citrate molar ratio is set to 1/1, whereas the ratio between metals of bimetallic solutions was chosen to reflect the stoichiometry of the corresponding hydrox-ide-carbonate minerals (*i.e.*, natural LDHs):¹ 1/3 for aluminium-nickel solutions, 1/3 for iron-nickel solutions and 1/2 for aluminium-zinc solutions, corresponding to takovite, reevesite and zaccagnaite minerals, respectively.¹ We also deliberately chose not to add any amount of strong acid, since it can bring an unnecessary source of error in the R determination from titration profiles, while also making less straightforward the comparison with previously published literature.^{11,13,53} Hence the initial pH in the titration profiles will be determined by the Lewis acidity of the metal cations involved.

2.2.1. Hydrolytic behaviour of Al³⁺, Fe³⁺, Ni²⁺, Zn²⁺ and their bimetallic solutions. The hydrolytic behaviour of monometallic and bimetallic solutions has been extensively characterized in the past.^{11,13,19} However, we decided to briefly report our results since they serve as baselines for titrations of metalcitrate solutions and as validation experiments. The titration profile of the $Al(NO_3)_3$ solution presents three equivalent points, located at 0.04, 2.65 and 2.97 R_{Al}^{3+} , related to the formation of $[Al_2(OH)_2]^{4+}$, Keggin's clusters, and the complete precipitation of Al(OH)₃, respectively.⁵⁴⁻⁵⁶ The titration profile of the Fe(NO₃)₃ solution presents three closely spaced equivalent points, averaged at 2.97 $R_{\rm Fe}^{3+}$, related to the complete precipitation of ferric oxyhydroxides. Furthermore, the three different equivalent points are located at 2.90, 2.97 and 3.05 $R_{\rm Fe}^{3+}$ and are related to β -FeOOH, ferrihydrite (*i.e.*, FeOOH xH_2O and α -FeOOH, respectively.^{13,57,58} The titration profile of the Ni(CH₃CO₂) solution presents two equivalent points located at 0.06 and 1.89 $R_{\rm Ni}^{2^+}$ related to the condensation of $[Ni(H_2O)_6]^{2+}$ cations into dimers and tetramers, and the precipitation α -Ni(OH)₂ phase, respectively.^{13,29,59,60} The titration profile of the Zn(NO₃)₂ solution presents one equivalent point located at 2.01 R_{Zn}^{2+} , confirming the precipitation of the stoichiometric Zn(OH)₂, probably present in its most insoluble polymorph (ε-Zn(OH)₂).⁶¹ All titration profiles of bimetallic solutions present two equivalent points, namely E_1 and E_2 , separated by a buffer region. As illustrated in Table 2, E_1 is always delayed compared to its expected R_M^{3+} value, whereas E_2 precedes its expected R_M^{2+} value, as previously described by Braterman's and Carteret's groups, regarding the formation mechanism of LDHs.^{11,13,53} Moreover, the more the LDH phase is favoured, the more E_1 is delayed and less sharp.11,13,53 Titration profiles of the monometallic and bi-

Table 2Hydroxylation ratios (R) of equivalent points observed inmetal-citrate solutions

Cations	Mole ratio	E_1	E_2
Al ³⁺	_	$2.97 R_{\rm Al}^{3+}$	_
Fe ³⁺	_	$2.97 R_{\rm Fe}^{3+}$	_
Ni ²⁺	_	_ 10	$1.89 R_{\rm Ni}^{2+}$
Zn ²⁺	_	_	$2.01 R_{\rm Zn}^{2+}$
Al ³⁺ , Ni ²⁺	1/3	~1.1 R	2.11 R
Normalized		$\sim 4.4 R_{Al}^{3+}$	$\sim 1.3 R_{\rm Ni}^{2+}$
Fe ³⁺ , Ni ²⁺	1/3	0.79 R	2.16 R
Normalized		$3.16 R_{\rm Fe}^{3+}$	$1.82 R_{\rm Ni}^{2+}$
Al^{3+}, Zn^{2+}	1/2	1.05 R	2.28 R
Normalized		$3.15 R_{Al}^{3+}$	$1.84 R_{\rm Zn}^{2+}$

metallic solutions without dissolved citrate are presented in Fig. S3. \dagger

2.2.2. Hydrolytic behaviour of metal-citrate solutions. When citrate anions are present, metal-citrate coordination compounds are formed. The nature of such metal-citrate complexes depends both on the solution pH and on the metal/citrate ratio.⁶¹⁻⁸⁵ In the following titrations and speciation models, citric acid (*i.e.*, $C_6H_8O_7$) is considered as tetraprotic (*i.e.*, H_4Cit), having the following acidity constants:⁸⁶ $pK_{a1} = 3.13$, $pK_{a2} = 4.76$, $pK_{a3} =$ 6.40 and $pK_{a4} = 14.4$. This means that the three carboxylic groups of citric acid are readily deprotonated in water, whereas the alcoholic group may only deprotonate in the complex formation or under concentrated alkali conditions. Hence, we denote free citrate anions originating from the dissolution of trisodium citrate (*i.e.*, Na₃HCit) as HCit³⁻. Fig. 2 reports the titration profiles of the trivalent metal-citrate solutions in the 1st column, divalent metal-citrate solutions in the 2nd column, and mixed metal-citrate solutions in the 3rd column. In the aluminium-citrate titration profile, the equivalent point, E_{L1} , at 1.39 R_{Al}^{3+} is related to the formation of $[Al_3Cit_3OH]^{4-}$, according to the following equilibrium:

$$3Al^{3+} + 3HCit^{3-} + 4OH^{-} \rightleftharpoons [Al_3Cit_3OH]^{4-} + 3H_2O, \qquad (2)$$

which predicts a 1.33 R_{Al}^{3+} value. Moreover, the asymmetric 1st derivative profile of E_{L1} is related to the $[Al_2Cit_2]^{2-}$ formation, as follows:

$$2Al^{3+} + 2HCit^{3-} + 2OH^{-} \rightleftharpoons \ \left[Al_2Cit_2\right]^{2-} + 2H_2O, \qquad (3)$$

in agreement with our speciation model presented in Fig. 3a. The titration profile of the iron-citrate solution is similar to the aluminium-citrate one. However, in contrast to aluminium, two equivalent points related to iron-citrate complexes, namely E_{L1a} and E_{L1b} , are clearly resolved and located at ~1.3 and $1.52 R_{Fe}^{3+}$, respectively. Our simulation is based on Silva *et al.*⁶⁵ iron-citrate stability constants, and predicts the existence of two iron-citrate complexes (*i.e.*, $[Fe_2Cit_2]^{2-}$ and $[FeCit]^-$) that are predominant at acidic and neutral pH values, as shown in Fig. 3b. However, the observed position of E_{L1a} and E_{L1b} in Fig. 2d is not accurately predicted, because Silva *et al.*⁶⁵ could not account for acid-base equilibria of the coordinated water molecules and hydroxide ions, due to constraints of the electrospray ionization mass spec-



Fig. 2 Potentiometric titration profiles and their respective 1st derivative of metal/citrate solutions with 1/1 molar ratio, as follows: (a) A^{3+} , (b) Ni^{2+} , and (c) $A^{3+}-Ni^{2+}$ with 1/3 ratio, (d) Fe^{3+} , (e) Zn^{2+} , and (f) $Fe^{3+}-Ni^{2+}$ with a 1/3 ratio, (i) $A^{3+}-Zn^{2+}$ with a 1/2 ratio. Comparison of titration curves in the presence of citrate (continuous line) and without citrate (dotted line) of: (g) trivalent metals, and (h) divalent metals. E_{L1} refers to A^{3+} and Fe^{3+} complex formations, E_{L2} refers to Ni^{2+} and Zn^{2+} complex formation. E_{Hyd} refers to the hydrolysis of complexes that leads to the formation of the respective hydroxides.

trometry used to determine the iron–citrate stability constants. Hence, oligomeric $[Fe_2Cit_2OH]_n^{3n-}$ species (with *n* of few units), produced by the condensation of $[Fe_2Cit_2]^{2-}$ and $[FeCit]^-$, represent a more plausible picture of the iron–citrate speciation, as previously proposed by Spiro *et al.*,^{68,69} but without providing stability constants. More in detail, the 1.52 R_{Fe}^{3+} value of E_{L1b} is consistent with the following formation equilibrium:

$$n \cdot \left(2\mathrm{Fe}^{3+} + 2\mathrm{HCit}^{3-} + 3\mathrm{OH}^{-}\right)$$

$$\approx \left[\mathrm{Fe}_{2}\mathrm{Cit}_{2}\mathrm{OH}\right]_{n}^{3n-} + n \cdot 3\mathrm{H}_{2}\mathrm{O},$$
(4)

which predicted a theoretical 1.5 $R_{\rm Fe}^{3+}$, whereas a theoretical 1.3 $R_{\rm Fe}^{3+}$ value of $E_{\rm L1a}$ is yielded from the formation equilibrium:

$$3Fe^{3+} + 3HCit^{3-} + 4OH^{-} \rightleftharpoons [Fe_3Cit_3OH]^{4-} + 3H_2O, \quad (5)$$

which is the iron analogous of (2). Lastly, E_{Hyd} corresponds to ferrihydrite nucleation from the partial hydrolysis of ironcitrate complexes.^{68,69} Since E_{Hyd} is located at ~2.7 R_{Fe}^{3+} it

means that less than three hydroxide ions are consumed per iron ion, being consistent with the observations of Spiro et al., concerning the progressive hydrolysis of $[Fe_2Cit_2OH]_n^{3n-}$ into ~8 nm ferrihydrite clusters capped with citrate anions.^{68,69} We also stress that E_{Hvd} appears as a broad peak in the 1st derivative profile, which indicates that the kinetics of the $[Fe_2Cit_2OH]_n^{3n-}$ hydrolysis into ferrihydrite is slower than the titration speed of 2.8 R h⁻¹. An analogous E_{Hyd} at ~2.5 R_{Al}^{3+} is also present in the titration profile of the aluminium citrate solution. Overall, as shown in Fig. 2g, the aluminium- and iron-citrate hydrolytic behaviour are similar. Both metals form 1/1 oligomeric citrate complexes that are stable in the 1.3–2.5 $R_{\rm M}^{3+}$ range and up to ~9.5 pH. The nickel–citrate titration profile presents one equivalent, namely $E_{1,2}$, located at 1.02 $R_{\rm Ni}^{2+}$ prior to hydroxide saturation. In addition to this, the solution remained clear of any precipitate even at the end of the titration, suggesting the stability of nickel-citrate complexes up to ~12.5 pH. In literature, several polynuclear nickel-citrate compounds have been reported.^{67,72-76} However,



Fig. 3 Speciation models of (a) Al-citrate, (b) Fe-citrate, (c) Ni-citrate and (d) Zn-citrate systems, with a metal/citrate molar ratio of 1/1. Speciation models are based on data from ref. 61–67, 81 and 86 We accounted for the solubility products of freshly formed precipitates; the solubility products of the aged precipitates are not considered.

to the best of our knowledge, speciation studies that account for the hydroxyl group deprotonation in nickel–citrate complexes are scarce.⁶⁷ For this reason, the speciation model presented in Fig. 3c predicts the formation of a solid phase that is not experimentally observed. As expected from Fig. 3c, at $0.00 R_{\rm Ni}^{2+}$ and neutral pH, the formation of the [NiHCit]⁻ complex is predominant, as follows:

$$Ni^{2+} + HCit^{3-} \rightleftharpoons [NiHCit]^{-}.$$
 (6)

Then, the consumption of one equivalent of hydroxide leads to the formation of $[Ni_2Cit_2]^{4-}$, as follows:

$$2[\text{NiHCit}]^{-} + 2\text{OH}^{-} \rightleftharpoons [\text{Ni}_2(\text{Cit})_2]^{4-} + 2\text{H}_2\text{O}.$$
(7)

However, according to E. Still *et al.*,⁶⁷ the formation of a hydroxylated tetramer is more plausible according to the equilibrium:

$$4[\text{NiHCit}]^{-}+4\text{OH}^{-} \rightleftharpoons [\text{Ni}_{4}\text{Cit}_{3}\text{OH}]^{5-}+\text{HCit}^{3-}+2\text{H}_{2}\text{O}.$$
 (8)

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In any case, according to both (7) and (8), an equivalent point at 1.00 $R_{\rm Ni}^{2+}$ is predicted; while considering the formation of $[{\rm Ni}_4{\rm Cit}_3{\rm OH}]^{5-}$ instead of $[{\rm Ni}_2({\rm Cit})_2]^{4-}$, has a negligible impact on our simulation results, since the aforementioned complexes have similar stability constants (Table S4†). To explain the absence of a precipitate, we refer to studies of Güdel and co-workers in which they reported the crystal structure of several different complexes containing 7, 8 and 21 nickel atoms coordinated with citrates, hydroxides, and water molecules,⁷³⁻⁷⁵ with the latter having the following chemical composition: $[{\rm Ni}_{21}({\rm Cit})_{12}({\rm OH})_{10}({\rm H}_2{\rm O})_{10}]^{16-}$. Moreover, by considering the overall formation equilibrium of Ni₂₁ as follows:

$$21[\text{NiHCit}]^{-}+22\text{OH}^{-} \approx [\text{Ni}_{21}(\text{Cit})_{12}(\text{OH})_{10}(H_2O)_{10}]^{16-}+9\text{HCit}^{3-}+2\text{H}_2\text{O}$$
(9)

a theoretical 1.05 $R_{\rm Ni}^{2+}$ is yielded, being in line with the experimental value of 1.02 $R_{\rm Ni}^{2+}$ for $E_{\rm L2}$ in Fig. 2b. Noteworthily, this Ni₂₁ complex contains a cluster of nickel atoms with a structure that resembles a Ni(OH)₂ nucleation seed.^{60,75} Reasonably, large nickel–citrate complexes, such as Ni₂₁, are the result of the hydrolytic polymerization of smaller species originating from either (7) or (8), while residual bound citrates hinder the further growth of the Ni(OH)₂ solid phase,⁷⁵ explaining the absence of a precipitate. The zinc–citrate titration profile shows similarities with the nickel–citrate one, displaying an $E_{\rm L2}$ at 1.06 $R_{\rm Zn}^{2+}$.

According to our speciation model, this equivalent point is related to the complete formation of $[Zn_2(HCit)_2(OH)_2]^{4-}$ as follows:

$$2[\text{ZnHCit}]^{-} + 2\text{OH}^{-} \rightleftharpoons [\text{Zn}_2(\text{HCit})_2(\text{OH})_2]^{4-}$$
(10)

that yields a theoretical 1.00 $R_{\rm Zn}^{2+}$. In addition, the zinc–citrate system shows an $E_{\rm Hyd}$ at 2.28 $R_{\rm Zn}^{2+}$ that corresponds to the slow hydrolysis of $[{\rm Zn}_2({\rm HCit})_2({\rm OH})_2]^{4-}$ as follows:

$$\left[Zn_2(HCit)_2(OH)_2 \right]^{4-} + 2OH^- \rightleftharpoons 2Zn(OH)_2 + 2HCit^{3-}$$
(11)

consuming a second hydroxide equivalent to produce $Zn(OH)_2$, which is clearly visible as a precipitate. In summary, divalent metal-citrate solutions behave similarly, according to the following generalized equilibria:

$$n \cdot (2\mathbf{M}^{2+} + 2\mathbf{H}\operatorname{Cit}^{3-} + 2\mathbf{O}\mathbf{H}^{-})$$

$$\approx n \cdot [\mathbf{M}_{2}\operatorname{Cit}_{2}]^{4-} + 2n \cdot \mathbf{H}_{2}\mathbf{O} \qquad (12a)$$

$$\approx [\mathbf{M}_{2}\operatorname{Cit}\mathbf{O}\mathbf{H}]_{n}^{n-} + n \cdot (\mathbf{H}\operatorname{Cit}^{3-} + \mathbf{H}_{2}\mathbf{O})$$

or

$$2\mathbf{M}^{2+} + 2\mathbf{H}\mathbf{Cit}^{3-} + 4\mathbf{OH}^{-}$$

$$\Leftrightarrow \left[\mathbf{M}_{2}(\mathbf{H}\mathbf{Cit})_{2}(\mathbf{OH})_{2}\right]^{4-} + 2\mathbf{OH}^{-}$$
(12b)

$$\rightleftharpoons 2\mathbf{M}(\mathbf{OH})_{2} + 2\mathbf{H}\mathbf{Cit}^{3-}.$$

From literature on nickel- and zinc-citrate complexes crystal structure^{73-75,79,80} and speciations,^{61,67,81-85} the nickel- citrate system follows (12a), whereas the zinc-citrate system

follows (12b) equilibria. The titration profiles of aluminiumnickel, iron-nickel, and aluminium-zinc solution, in the presence of citrates, are reported in the 3rd column of Fig. 2. All three titration profiles present two equivalent points related to the formation of complexes that are analogous to the respective monometallic-citrate solutions. However, by normalizing E_{L1} by its trivalent metal ratio, it follows that E_{L1} is always delayed compared to the corresponding monometallic solution (Table 3), with the aluminium-zinc-citrate system showing the largest E_{L1} delay of 0.35 R compared to the aluminium-citrate solution. We ascribe the delay of E_{L1} to the progressive formation of mixed metal-citrate complexes by consuming trivalent metal-citrate complexes as reactants. In the aluminiumand iron-nickel systems, E_{L2} is also delayed by 0.16 R_{Ni}^{2+} compared to the nickel-citrate system. On the contrary, E_{L2} of the aluminium-zinc-citrate solution precedes the equivalent point expected for the zinc-citrate solution by 0.16 R_{Zn}^{2+} . The reason behind this discrepancy is not yet clear to us and may be worth further investigation. However, it is reasonably related to the different speciation of nickel- and zinc-citrate systems as inferable from eqn (12). Eventually, the increased hydroxylation of citrate complexes leads to their hydrolysis, as apparent from the translucent aspect of the titrated bimetallic solutions beyond pH ~12.

2.3. Electronic spectroscopy

2.3.1. Nickel. This section describes the nature of Ni²⁺ related features observed in the electronic spectra of NiAl- and NiFe-LDHs, as shown in Fig. 4. The ${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{3}T_{2g}({}^{3}F)$, the ${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{3}T_{1g}({}^{3}F)$ and the ${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{3}T_{1g}({}^{3}F)$ transitions are characteristic features of the electronic spectra of nickel-based compounds;²⁸⁻³⁸ for the sake of simplicity, we labelled the aforementioned transitions as ν_{1} , ν_{3} and ν_{6} , respectively. We also point out that the ν_{3} is often accompanied by the ${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{1}E_{g}({}^{1}D)$ "spin-flip" transition, due to the state mixing between ${}^{3}T_{1g}$ and ${}^{1}E_{g}$, in the case of their energy proximity.³⁴⁻³⁶ We labelled ${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{1}E_{g}({}^{1}D)$ as ν_{2} ; nevertheless, several authors prefer to report a single average transition as ${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{3}T_{1g}({}^{3}F)$, ${}^{1}E_{g}({}^{1}D)$.^{32,37,38,87} Interestingly, ν_{2} results in enhancing the NiFe-LDH electronic spectra due to super-exchange interactions in Fe³⁺–O–Ni²⁺ bonds (see section 2.3.3).

 Table 3
 Hydroxylation
 ratios
 (R)
 of
 equivalent
 points
 observed
 in

 metal-citrate solutions

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Cations	Mole ratio	$E_{\rm L1}$	$E_{\rm L2}$	E _{Hyd}
Al ³⁺	_	$1.39 R_{Al}^{3+}$	_	$\sim 2.5 R_{Al}^{3+}$ $\sim 2.7 R_{Fe}^{3+}$
Fe ³⁺	_	$\sim 1.3 R_{\rm Fe}^{3+}$	_	$\sim 2.7 R_{\rm Fe}^{11^{-3+}}$
		$1.52 R_{\rm Fe}^{-3+}$		
Ni ²⁺	_		$1.02 R_{\rm Ni}^{2+}$	_
Zn^{2+}	_	_	$1.06 R_{\rm Zn}^{2+}$	$2.28 R_{\rm Zn}^{2+}$
Al ³⁺ , Ni ²⁺	1/3	0.38 R	1.28 R	_
Normalized	$1.53 R_{Al}^{3+}$	$1.19 R_{\rm Ni}^{2+}$		
Fe^{3+} , Ni ²⁺	1/3	0.40 R	1.28 R	~1.9 R
Normalized	$1.60 R_{\rm Fe}^{3+}$	$1.18 R_{\rm Ni}^{2+}$		
Al^{3+}, Zn^{2+}	1/2	0.58 R	1.18 R	2.10 R
Normalized	$1.74 R_{Al}^{3+}$	$0.90 R_{\rm Zn}^{2+}$		



Fig. 4 Electronic spectra of: (a) citrate intercalated NiAl- and NiFe-LDHs powders, (b) carbonate-intercalated NiFe-LDH dispersed in ethanol according to ref. 12 and taken as reference material, (c) second derivative of (b), (d) synthesis solution of citrate-intercalated NiFe-LDH at different *R* values, (e) second derivative of (d). Dotted vertical lines at 300 and 800 nm indicate the switch of lamp and detector, respectively. Inset to (b) and (d) is the magnification of the electronic spectra in the 600–800 nm range.

Eventually, we also detected the contribution to the absorbance of the ${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{1}T_{2g}({}^{1}D)$, ${}^{1}A_{1g}({}^{1}G)$ spin-forbidden transition, labelled as ν_{5} . The wavelength positions of the observed Ni²⁺ electronic transitions are reported in Table 4 and visualized in the Tanabe–Sugano diagram for d⁸ configuration (Scheme 1a). A straightforward comparison with other nickelbased minerals and synthetic compounds is shown in Table S5.†

2.3.2. Iron. In the NiFe-LDH electronic spectrum, we observed, the ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}E_{g}, {}^{4}A_{1g}({}^{4}G)$ and ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}E_{g}({}^{4}D)$ spin-forbidden transitions of Fe³⁺, labelled as ν_{4} and ν_{7} , at 448 and 348 nm (Fig. 4a–c) by means of second derivative spectrophotometry. Moreover, ν_{7} presents an asymmetric shape due to the absorbance contribution of ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{1g}({}^{4}P)$ at 338 nm (*i.e.*, ν_{7b} in Fig. 4c). Eventually, the two absorbance bands ascribed to ligand-to-metal charge transfer transitions (LMCT) are located at 250 and 210 nm and are labelled as ν_{8} and ν_{9} , respectively. For the sake of completeness, we also include in ν_{9} , the absorbance contribution of the n $\rightarrow \pi^{*}$ tran-

Band	Wavelength (nm)	Туре	Transition
ν_1	1120	dd (Ni ²⁺)	${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{3}T_{2g}({}^{3}F)$
ν_2	750	dd $(Ni^{2+})^a$	${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{1}E_{g}({}^{1}D)$
ν_3	665	dd (Ni^{2+})	${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{3}T_{1g}({}^{3}F)$
$ u_4$	448	dd $(\mathrm{Fe}^{3+})^a$	$Ni^{2+6}A_{1g}(^{6}S) \rightarrow {}^{4}E_{g}, {}^{4}A_{1g}(^{4}G)$
ν_5	420	dd $(Ni^{2+})^a$	${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{1}T_{2g}({}^{1}D), {}^{1}A_{1g}({}^{1}G)$
ν_6	386	$\frac{\mathrm{dd}(\mathrm{Ni}^{2^+})}{\mathrm{dd}(\mathrm{Fe}^{3^+})^a}$	${}^{3}A_{2g}({}^{3}F) \rightarrow {}^{3}T_{1g}({}^{3}P)$ ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{2g}({}^{4}D)$
ν_7	348 ^b , 338 ^c	dd $(\mathrm{Fe}^{3+})^a$	${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}E_{g}({}^{4}D), {}^{4}T_{1g}({}^{4}P)$
ν_8	250	LMCT	$t_{1u}^{\beta} \rightarrow t_{2g}^{\beta} \left(Fe^{(III)} \right)$
$ u_9$	210	LMCTs MOs of RCOO ⁻	$\begin{array}{l} t_{1u}^{\ \ \beta} \rightarrow e_{g}^{\ \ \beta} \left(Fe^{(III)} Ni^{(II)} \right) \\ n \rightarrow \pi^{*} \left(HOMO \rightarrow LUMO \right) \end{array}$

^{*a*} Spin-forbidden transitions. dd = Ligand Field Transitions (parity-forbidden). LMCT = Ligand-to-metal charge transfer. MO = molecular orbital. HOMO and LUMO = Highest occupied MO and lowest unoccupied MO. ${}^{b} {}^{6} {A}_{1g} ({}^{6} {S}) \rightarrow {}^{4} {E}_{g} ({}^{4} {D})$ contribution to the absorbance. ${}^{c} {}^{6} {A}_{1g} ({}^{6} {S}) \rightarrow {}^{4} {T}_{1g} ({}^{4} {P})$ contribution to the absorbance.



Scheme 1 Tanabe-Sugano diagrams of: (a) d^8 octahedral configuration, and (b) high-spin d^5 octahedral configuration. The abscissae report the ligand field strength (10*Dq*) normalized by the Racah parameter *B*, whereas the ordinates report the energies, normalized by *B*, of ligand field states. In each diagram, the abscissae axis always corresponds to the ground state. Here, energy states are colour-coded according to their spin–orbit terms which are reported on the left (*i.e.*, black for S, cyan for P, orange for D, violet for F and green for G). Moreover, energy states with the same spin multiplicity as the ground state are represented with a plain line, while energy states with a spin multiplicity different from the ground state are represented with a dashed line. The symmetry term of each energy state is reported on the top-right of the diagrams according to the Mulliken nomenclature. The red vertical lines indicate the 10Dq/B ratios we computed for NiAl- and NiFe-LDHs (Table 5). Racah parameters used: (a) $B = 887 \text{ cm}^{-1}$, $C = 4075 \text{ cm}^{-1}$; (b) $B = 916 \text{ cm}^{-1}$, $C = 2632 \text{ cm}^{-1}$. ESI section 6.3^{+} for computation details.

sition of the carboxylic groups that falls at ~210 nm. The wavelength positions of these Fe^{3+} electronic transitions are reported in Table 4 and visualized in the Tanabe–Sugano diagram for high-spin d^5 configuration (Scheme 1b). For a straightforward comparison with other ferric compounds see Table S5.†

2.3.3. Ligand field analysis. The electronic spectroscopy of transition-metal-based LDHs is a topic often disregarded in current literature. In fact, only a few studies analysed LDHs electronic spectra, merely assigning the ligand field transitions.^{28,43,88} Previously, we proved how second derivative spectrophotometry is effective in resolving multiple ligand field transitions in NiFe-LDH electronic spectra.⁴³ Here, by means of second derivative spectrophotometry, we computed the magnitude of the ligand field strength (10*Dq*), *B* and *C* Racah parameters related to Ni²⁺ and Fe³⁺ located within NiAl-and NiFe-LDH materials. From the Racah parameters, the nephelauxetic effect can be estimated. The nephelauxetic effect is the reduction of interelectronic repulsion of *d* electrons, with respect to the isolated ions, due to a coordination environment. This directly translates into an estimation of the

ionic degree of ligand-metal bonds,⁸⁹ given by the nephelauxetic ratio historically defined as:

$$\beta = \frac{B}{B_{\text{gas}}}.$$
 (13)

Here B_{gas} is the tabulated value of *B* for ions in the gas phase. During the years, the nephelauxetic effect has been further revisited.^{90–92} For our purposes, we used the (13) revisited version of Brik *et al.*,^{91,92} which also accounts for *C* and C_{gas} , as follows:

$$\beta_1 = \sqrt{\left(\frac{B}{B_{gas}}\right)^2 + \left(\frac{C}{C_{gas}}\right)^2}.$$
 (14)

We point out that a proper comparison between β values reported in the literature, β_1 should be obtained by normalizing the latter by $\sqrt{2}$. The ligand field analysis results are reported in Table 5. For Ni²⁺, located LDH crystal lattices, typical 10*Dq* values of ~9000 cm⁻¹, *B* values of ~890 cm⁻¹ and *C* values of ~4000 cm⁻¹ were computed, and these consistent

Ion	Material	$10Dq (cm^{-1})$	$B (\mathrm{cm}^{-1})$	$C \left(\mathrm{cm}^{-1} \right)$	10 Dq/ B	C/B	β	$\beta_1{}^a$
Ni ²⁺	NiAl-LDH	9009	892	4054	10.1	4.54	0.857	1.200
	NiFe-LDH	8929	887	4075	10.1	4.59	0.852	1.199
	Gas phase ⁸⁹	_	1041	4831	_	4.64	1	$\sqrt{2}$
Fe ³⁺	NiFe-LDH	7049^{b}	916	2632	7.7	2.87	0.903	1.056
	Gas phase ⁹⁰	_	1015	4800	_	4.73	1	$\sqrt{2}$
	1							

Table 5 Results of ligand field analysis of the electronic spectra of NiAl- and NiFe-LDHs

Ion	Material	$10Dq (\mathrm{cm}^{-1})$	$B(\mathrm{cm}^{-1})$	$C (\mathrm{cm}^{-1})$	10 Dq/ B	C/B	β	$\beta_1{}^a$
Ni ²⁺	NiAl-LDH	9009	892	4054	10.1	4.54	0.857	1.200
	NiFe-LDH	8929	887	4075	10.1	4.59	0.852	1.199
	Gas phase ⁸⁹	_	1041	4831	_	4.64	1	$\sqrt{2}$
Fe ³⁺	NiFe-LDH	7049^{b}	916	2632	7.7	2.87	0.903	1.056
	Gas phase ⁹⁰	—	1015	4800	—	4.73	1	$\sqrt{2}$
		. 2						

^{*a*} Defined as: $\beta_1 = \sqrt{\left(\frac{B}{B_0}\right)^2 + \left(\frac{C}{C_0}\right)^2}$, see ref. 91 for details. ^{*b*} Computed from the ν_7 contribution to the absorbance of ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{1g}({}^{4}P)$.

This article is licensed under a Creative Commons Attribution 3.0 Unported Licence. with those from the past literature concerning nickel hydroxides and related materials.^{28,37,42} This indicates that the nature of trivalent metals in nickel-based LDHs has a negligible influ-

ence on the ligand field parameters of Ni²⁺. In addition, Ni²⁺, in both NiAl- and NiFe-LDH crystal lattices, has a $\beta_1/\sqrt{2}$ of 0.849. By comparison, we performed the same ligand field analysis on the spectroscopic data published by Oliver-Tolentino et al.²⁸ yielding a $\beta_1/\sqrt{2}$ of 0.862 for carbonate-intercalated nickel-based LDHs, whereas the data published by Poul et al.42 yielded a $\beta_1/\sqrt{2}$ of 0.860 for nickel hydroxide-acetate (which is isomorphous to LDHs). Hence, we conclude that the nephelauxetic ratio in nickel-based LDHs is neither significantly influenced by the nature of the trivalent metal cation hosted in the hydroxide layers nor by the nature of the anions amid them. In high-spin Fe3+, all possible ligand field transitions are spin-forbidden. However, in correlated materials, the spin restrictions are lifted because of super-exchange interactions, making formally spin-forbidden transitions observable.^{33,40,41,93-97} The $^6A_{1g}(^6S) \rightarrow \, ^4E_g, \, ^4A_{1g}(^4G)$ and $\, ^6A_{1g}(^6S)$ \rightarrow ⁴E_g(⁴D) transitions, which are 10Dq-independent,³⁹⁻⁴¹ directly provide the Fe^{3+} values of *B* and *C*, which are 916 and 2632 cm⁻¹, respectively. To have a proper comparison, we also performed the same ligand field analysis on the spectroscopic data reported by Sherman et al.^{40,41} It follows that, according to (14), Fe³⁺ experiences a $\beta_1/\sqrt{2}$ value of 0.747 in NiFe-LDH. This means that the nephelauxetic ratio of Fe³⁺ in NiFe-LDH is comparable to the 0.75 value of $[Fe(H_2O)_6]^{3+}$ reported in literature,^{98,99} whereas the expected nephelauxetic ratio of Fe³⁺ in metal-oxide lattices ranges between 0.6 and 0.7, as shown in Fig. 5. In our opinion, these nephelauxetic similarities between Fe^{3+} in LDHs and $[\text{Fe}(H_2O)_6]^{3+}$ are related to the performance of the NiFe-LDHs as catalysts in water-splitting applications.100-102 In ferric compounds, the magnitude of 10Dq is usually obtained from the energy of ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{1g}({}^{4}G)$ and ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{2g}({}^{4}G)$ transitions. However, we have not detected any contribution to the absorbance spectrum ascribable to the aforementioned transitions. According to the Tanabe mechanism, the interaction between super-exchanged ions and incident light involves only orbitals that are partially filled,^{33,94,96,97} and is described by the following Hamiltonian:

$$H = \sum_{ij} \left(\mathbf{\Pi}_{\mathbf{a}_i, \mathbf{b}_j} \cdot \boldsymbol{E} \right) \left(\boldsymbol{s}_{\mathbf{a}_i} \cdot \boldsymbol{s}_{\mathbf{b}_j} \right), \tag{15}$$

where *i* and *j* are the half-filled d orbitals of a and b ions, respectively, s are the corresponding spin vectors, and E is the



Fig. 5 Scatter plot of the energy of the ${}^{6}A_{1q}({}^{6}S) \rightarrow {}^{4}E_{q}({}^{4}D)$ transitions in ferric compounds vs. their respective nephelauxetic ratios. Empty triangles indicate the classical nephelauxetic ratio, *i.e.*, $\beta = B/B_0$. Filled circles resent the new nephelauxetic ratio^{91,92} divided by $\sqrt{2}$, *i.e.*, $\beta_1/\sqrt{2}$. The dashed lines represent the expected range of $\beta_1/\sqrt{2}$ in ferric compounds. Apart from NiFe-LDH, the nephelauxetic ratios were computed from the spectroscopic data reported in ref. 40 and 41.

electric vector of the incident light. Π_{a_i,b_i} represents the partial derivative of exchange constants between a_i and b_j with respect to E coordinates. Since the t_{2g} orbitals of Ni²⁺ are full, they do not bring any contribution to (15), relegating the enhancement of ligand field transitions to the super-exchange interactions in Fe³⁺-O-Ni²⁺ bonds through e_g orbitals. This explains the observed intensity of ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}A_{1g}({}^{4}G)$ and ${}^{6}A_{1g}({}^{6}S) \rightarrow$ ${}^{4}E_{g}({}^{4}D)$, since the former transition is related to the spin canting of eg orbitals, whereas the latter is related to a spinflip in e_g orbitals of Fe^{3+} (see tables of the matrix elements of Coulomb interactions in ref. 103 and 104). On the contrary, ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{1g}({}^{4}G) \text{ and } {}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{2g}({}^{4}G) \text{ do not benefit from}$ (15) since both transitions involve an electron transfer from e_g to t_{2g} orbitals of Fe³⁺ with a concomitant spin-flip. The same phenomenon has been observed and described in the past for Ni2+-doped manganese halide perovskites.33,95 Given this, we could only rely on the energy of ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{1g}({}^{4}P)$ for the 10Dq of Fe³⁺, which turns out to be 7049 cm⁻¹. By comparison, this 10Dq value is approximatively half of the \sim 15 000 cm⁻¹ expected for Fe³⁺ in oxide crystal lattices,³⁹⁻⁴¹ being consistent with the linear relationship between Dq and the effective

charge of the ligands.¹⁰⁴ With known Dq, B and C values, the expected positions of the unobserved Fe³⁺ ligand field transitions are obtained from the Tanabe–Sugano diagram for high-spin d⁵ configurations (Scheme 1b). The ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{1g}({}^{4}G)$ and ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{2g}({}^{4}G)$ transitions are expected to fall at ~590 and ~480 nm, respectively.

However, it is evident from the second derivative electronic spectra in Fig. 4c and e that the contribution of those transitions, if any, is below our limit of detection, being consistent with the previous statement about super-exchange interactions in Fe³⁺–O–Ni²⁺ bonds. Eventually, the ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}T_{2g}({}^{4}D)$ transition is predicted to fall at ~370 nm, overlapping with the absorbance contribution of Ni²⁺ to ν_6 . We also stress that ν_7 has been assigned, in the past, to the ${}^{6}A_{1g}({}^{6}S) \rightarrow {}^{4}E_{g}, {}^{4}T_{2g}({}^{4}D)$ transition.^{28,43} We deem this assignment incorrect for two reasons. Firstly, the assignment is inconsistent with the electronic spectra of ferric oxides.^{40,41} Secondly, by considering the literature ν_7 assignment, our ligand field analysis yields improbable values of 10Dq close to zero. However, it is evident from the second derivative electronic spectra in Fig. 4c and e that the contribution of those transitions, if any, is below our limit of detection, being consistent with the previous statement about super-exchange interactions in Fe³⁺-O-Ni²⁺ bonds.

2.3.4. Formation mechanism of NiFe-LDH. The electronic spectra, and their 2nd derivative, of subsequent aliquots collected during the citrate-intercalated NiFe-LDH synthesis are displayed in Fig. 4d and e. Prior to the addition of any hydroxide equivalent (0.00 R), the solution appears green. This green colour is caused by three absorbance bands at 740, 660, and ~450 nm. As described earlier, the two bands at 740 and 660 nm are related to Ni²⁺ ligand field transitions, ν_2 and ν_3 , respectively, while the absorbance band at ~450 nm is related to Fe^{3+} , *i.e.*, ν_4 . However, we cannot exclude possible contributions from ν_5 and ν_6 . The LMCT absorbance edge begins at 400 nm saturating our detector at 290 nm. During the addition of NaOH, the solution turns from green to yellow after $E_{1,1}$ at 0.4 R is passed. Thereafter, a progressive increase in the absorbance intensity is observed in the 350-500 nm region. This observation confirms our interpretation of the potentiometric titration profiles. In fact, the progressive formation of bimetallic-citrate compounds implies an increasing number of possible super-exchange interactions in Fe³⁺-O-Ni²⁺ bonds, and therefore an increasing intensity of Fe³⁺ related spin-forbidden transitions (namely, ν_4 and ν_7) by the Tanabe mechanism. At the same time, as R increases, the LMCT absorbance edge shifts by ~100 nm to lower wavelengths. This shift can be explained by the hydrolysis of chemical bonds between transition metals and citrate carboxylic groups forming complexes. In fact, the increase of the photon energy required for a LMCT fits well with the alteration of the Fe³⁺ or Ni²⁺ coordination environment. In addition, it is consistent with a shift from a ligand electron-rich environment constituted by carboxylic groups to a ligand electron-poor environment constituted by hydroxide ions. After 1.88 R (ESI section 2⁺), the reaction mixture still appears clear of any precipitate, while its absorbance spectrum (red line in Fig. 4d) resembles that of carbonate-intercalated NiFe-LDH (Fig. 4b). Then, after performing the aging treatment at 90 °C, the reaction mixture became opaque due to suspended LDH particles, while its absorbance spectrum shows a further increase in the absorbance related to ligand field transitions (blue line in Fig. 4d).

Conclusions and perspectives

In summary, we developed an aqueous, coordination-chemistry-based synthetic procedure that yields nanostructured layered double hydroxides (LDHs) without using any anion exchange process, avoiding the use of organic solvents. We successfully synthetized nanostructured citrate-intercalated NiAl-, NiFe- and ZnAl-LDH materials, which possess a turbostratic disorder arising from the restacking of nanosheets ranging from 5 to 500 nm wide, depending on their chemical composition. The presence of vertically aligned nanosheets in the transmission electron microscopy images of NiFe- and ZnAl-LDH samples allowed us to estimate their thicknesses, which are in the 1-2 nm range. The LDH formation mechanism is a two-step process, as revealed by pH-potentiometric titrations and electronic spectroscopy investigations. The first step consists of forming bimetallic-citrate coordination compounds that occur by increasing the reaction environment pH. The second step consists of heating the reaction environment that causes the complete hydrolysis of the metal-citrate compounds, leading to the formation of LDH nanosheets. By means of electronic spectroscopy, we have shown a progressive enhancement of Fe³⁺ spin-forbidden ligand field transitions during the NiFe-LDH synthesis. This enhancement is due to super-exchange interactions in the increasing number of Fe³⁺-O-Ni²⁺ bonds formed during the LDH synthesis process. In addition, our ligand field analysis reveals that Ni²⁺ in NiAland NiFe-LDH crystal lattices behaves similarly to that in nickel hydroxide and other related materials. On the contrary, Fe^{3+} in NiFe-LDH behaves similarly to that in $[\text{Fe}(\text{H}_2\text{O})_6]^{3+}$, rather than in its ferric oxide counterparts, suggesting a correlation with the water-splitting electrocatalytic activity of NiFe-LDHs. Our approach and insights gained on the coordination chemistry of LDH synthesis may be useful to synthesize nanoparticles via coprecipitation and fabricate functional films for optoelectronics via electroplating or chemical bath deposition.^{26,27,105} Our ligand field analysis results may help to fully understand the electronic structure of transition metalbased LDHs, which constitute a valid option in the field of two-dimensional correlated insulators for future spintronic devices.106-108

Author contributions

The manuscript was written through the contributions of all authors. All authors have given approval to the final version of the manuscript.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

FB is the co-founder and Scientific Director of BeDimensional S.p.A., a company producing and commercializing 2D materials. The other authors have no conflicts to declare.

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