Continuous synthesis of doped layered double hydroxides in a meso-scale flow reactor

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Abstract
Layered double hydroxides is a class of low cost structured nanomaterials with many potential applications in environmental catalysis and sustainable technologies. Their large scale use is hindered by the challenge of reproducible synthesis at scale. Here we report a general, readily scalable process for the reproducible synthesis of transition metal doped hydrotalcites using a two-step process: co-precipitation in a mm-scale (meso-scale) continuous flow reactor, followed by aging. We have shown that co-precipitation in flow at a residence time close to the micromixing time affords good control of particle formation. Reproducible synthesis allowed us, for the first time, to investigate the formation of the pore morphology of hydrotalcites and their thermal stability as a function of metal doping. The obtained samples exhibited surface areas (80-150 m² g⁻¹) higher than those typically attained in batch syntheses, with very low standard deviation between the samples, a high degree of crystallinity and small crystallite sizes, in the range of 9.5-11.9 nm, depending on composition. A systematic characterization allowed us to elucidate the mechanism of the pore morphology formation: the crystallites were found to agglomerate into disk-like platelets, whereas the pore structure of the hydrotalcites is formed by agglomeration of the platelets.

Keywords: hydrotalcites; flow chemistry; continuous processes; pore morphology

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Introduction

Anionic clays or layered double hydroxides (LDHs) form a broad class of materials, defined by the characteristic lamellar structure with two kinds of metal cations in the layers and anionic species in the interlayer region [1]. Despite LDHs being known since 19th century and first synthetic LDHs developed in 1940s [2], there is recent significant growth in scientific and practical interest in these inexpensive structured materials. Due to the relative ease of modification of their composition, a significant variation in functional performance can be attained, which is highly attractive in many application areas, especially in environmental catalysis, sustainable technologies, including healthcare technologies [3-7]. Modulation of properties is typically achieved by modification of the feed composition in the controlled co-precipitation batch synthesis methods. Batch synthesis results in materials with broad variability in their functional properties and very poor reproducibility [8]. Even minute changes to synthesis parameters (pH, concentrations, rates of addition, duration, temperature and amount of water during aging, method of washing) from one preparation to the next result in vast differences in the final material’s properties: for example, Wang et al [9] demonstrated that morphology, pore structure and chemical composition are highly sensitive to pH of co-precipitation; Climent et al [10] showed that crystal size of similar materials is highly sensitive to temperature of the solutions during precipitation. More generally in the liquid phase synthesis of nanomaterials, the impact of process conditions on the outcomes of synthesis of crystalline materials is not well characterised due to the fast rates of precipitation and complex mechanisms of crystal growth and aggregation. As a result, it is very challenging to develop detailed structure-property relationships of LDHs that would enable rational design of new LDH-based functional nanomaterials, and applications of these materials in novel sustainable technologies are hindered by lack of reproducibility of material properties at scale.

This limitation of the co-precipitation synthesis at scale stems from the fundamental mismatch of the rates of nucleation and crystal growth and aggregation, and the rates of mixing in stirred vessels. It is impossible to attain a uniform distribution of concentrations within a reaction vessel under such conditions; this automatically leads to a broad distribution in the properties of the obtained materials and poor reproducibility in manufacture. We hypothesised that achieving reproducible outcome in the co-precipitation step is critically important for attaining a narrow distribution of properties of the final LDHs in the two-step synthesis method. Although there are several reports of LDH synthesis under continuous flow conditions using micro-scale [11-13] and meso-scale reactors [14-18], there are no studies detailing systematic characterizations of morphological properties and thermal stability of the produced materials. In most of the earlier studies the produced materials are characterized by broad particle size distribution.
Thus, we developed a process of co-precipitation under continuous flow conditions in a meso-scale reactor, as a two-step (precipitation-aging) process of synthesis of doped catalysts precursors, where the step of aging can also be realised as a continuous process. This synthesis modification was sufficient to significantly reduce variance in the structural properties of the synthesised materials, and this allowed us, for the first time, to carefully examine the mechanism of formation of the pore structure of the doped Mg-Al hydrotalcites as a function of the composition change. We focused on the systematic study of the synthesis of natural Mg-Al LDHs doped with a range of transition metals, which is an important system in the field of heterogeneous catalysis. To the best of our knowledge there are no reports of such systematic studies for natural Mg-Al LDHs that allow mechanistic insights into the formation of the pore structure of these materials. This synthesis method is likely to have a significant impact on development of new catalysts: an example is the very recent work published by our collaborating group, who adopted our synthesis method for preparation of new active catalysts for a cleaner energy solution [19].

2. Experimental
2.1 Materials
Aluminium(III) nitrate nonahydrate (98%), magnesium(II) nitrate tetrahydrate (98%), iron(III) nitrate nonahydrate (98+%), copper(II) nitrate 2.5-hydrate (98%), nickel(II) nitrate hexahydrate (99%), zinc(II) nitrate hexahydrate (98%) from ACROS Organics; chromium(III) nitrate nonahydrate (98.5%) from Alfa Aesar; sodium hydroxide (97%) from VWR AMRESCO® Life Sciences and sodium carbonate (99.5%) from Fisher were used in synthesis of doped hydrotalcites.

2.2 Synthesis protocol
Batch synthesis: 300 mL of a solution of Mg(NO$_3$)$_2$·6H$_2$O (0.15 mol) and Al(NO$_3$)$_3$·9H$_2$O (0.05 mol) was added dropwise over 2 h to 300 mL of 0.083 M Na$_2$CO$_3$. The pH was maintained between 9.5 and 10.5 by addition of 1 M NaOH. After completing the addition, the solution was heated to 65 °C for 2 h. The resulting solid was filtered and washed with DI water until the pH of the filtrate was neutral.

Flow synthesis: hydrotalcites with a general formula [Me$^{2+}$$_{1-x}$Me$^{3+}$$_{x}$(OH)$_2$]$^{x+}$(CO$_3$)$_{x/2}$·mH$_2$O were synthesized by the following procedure: two 60 mL plastic syringes were filled with metal precursor solution (solution A) and base solution (solution B), placed on a syringe pump (Cole Parmer EW-74900-20) and connected via a Y connector and 2-mm internal diameter polyethylene tubing (see reactor scheme in Figure 1). The syringe pump was set with a constant flow rate of 4 mL min$^{-1}$ and was positioned to drip into
a beaker containing 100 mL of DI water. The mixture in the beaker was stirred at 200 rpm at the set temperature in the range 65-95 °C for 2 h. Subsequently the mixture was cooled to ambient temperature, filtered and washed with DI water until the pH of the filtrate was neutral. All the samples were dried at 110 °C overnight.

This procedure was consistent for all doped HTs except the copper-doped sample. In this case we found it is important to wash the precipitate formed in the continuous synthesis from residual salts and base, as the reaction appears to be incomplete. Failure to do so results in formation of additional phases. The formed precipitate was thus filtered, washed with DI water until the pH of the filtrate was neutral, collected and placed into a beaker containing 100 mL of DI water. The following procedure was identical to all other flow synthesized samples.

![Flow Diagram](image)

**Figure 1.** Schematic diagram of the flow synthesis method.

**Solutions composition.** For standard MgAl HT composition with Mg/Al = 3, solution A was prepared using a mixture of Mg(NO$_3$)$_2$·6H$_2$O (0.15 mol) and Al(NO$_3$)$_3$·9H$_2$O (0.05 mol) dissolved in 700 mL of deionized (DI) water, while solution B was prepared using a mixture of NaOH (0.40 mol) and Na$_2$CO$_3$ (0.025 mol) dissolved into 700 mL of DI water.

The composition of the doped HTs was set to be Mg:Al:Me = 2.4:1:0.6, when doped with Me$^{2+}$, and Mg:Al:Me = 3:0.4:0.6, when doped with Me$^{3+}$. The HTs doped with Fe$^{3+}$, Co$^{2+}$, Ni$^{2+}$, Cu$^{2+}$, Zn$^{2+}$, targeting a 15% mole fraction over the sum of all metal species, were prepared in an analogous synthetic protocol by adding the respective metal nitrate salt to solution A (details of compositions are given in Table S1 in ESI). Table 1 shows designations used subsequently in the text and compositions for all the synthesized...
hydrotalcites, as well as the results of their ICP-AES analyses. To distinguish between samples prepared at different aging conditions, temperature will be added after sample designation, for example “Fe HT80” corresponds to the Fe-doped hydrotalcite aged at 80 °C.

2.3 Materials characterization
Elemental analysis was carried out with atomic absorption spectroscopy (AAS) performed on a Perkin Elmer AAnalyst 200.

Powder X-Ray Diffraction (PXRD) was performed on a Rigaku MultiflexII Desktop X-ray Diffractometer (Cu Kα, 40 kV, 30 mA) instrument in the stepscan mode with diffraction angle (2θ) from 5 to 80°. The crystallite size was calculated from Debay-Shrerrer equation, using the full-width at half-maximum (FWHM) value of the (003) reflection.

Nitrogen adsorption isotherms were measured on a Micromeritics TriStar surface analyzer at liquid nitrogen temperature. Samples were degassed under vacuum at 110 °C for three h prior measurements. Surface areas were calculated using the BET method in the relative pressure range of 0.05-0.30. Pore size distribution was calculated using the BJH method using the desorption branch of the isotherm, following the literature procedure [20].

Table 1. Theoretical composition and the results of elemental analysis for the flow synthesized hydrotalcites.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Metals Ratio</th>
<th>Designation</th>
<th>Theoretical Molar Ratio</th>
<th>Molar Ratio by ICP-AES 65 °C</th>
<th>Molar Ratio by ICP-AES 80 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>MgAl</td>
<td>3:1</td>
<td>HT</td>
<td>75:25</td>
<td>75:25</td>
<td>76:24</td>
</tr>
<tr>
<td>MgAlFe</td>
<td>3:0.4:0.6</td>
<td>Fe HT</td>
<td>75:10:15</td>
<td>77:7:16</td>
<td>75:7:18</td>
</tr>
<tr>
<td>MgAlCo</td>
<td>2.4:1:0.6</td>
<td>Co HT</td>
<td>60:25:15</td>
<td>62:23:15</td>
<td>60:26:14</td>
</tr>
<tr>
<td>MgAlNi</td>
<td>2.4:1:0.6</td>
<td>Ni HT</td>
<td>60:25:15</td>
<td>60:23:17</td>
<td>60:25:15</td>
</tr>
<tr>
<td>MgAlCu</td>
<td>2.4:1:0.6</td>
<td>Cu HT</td>
<td>60:25:15</td>
<td>62:24:14</td>
<td>60:25:15</td>
</tr>
<tr>
<td>MgAlZn</td>
<td>2.4:1:0.6</td>
<td>Zn HT</td>
<td>60:25:15</td>
<td>64:20:16</td>
<td>62:24:14</td>
</tr>
</tbody>
</table>

TEM images were obtained using a JEOL JEM 2100F FETEM at 200 kV. Dried samples were dispersed in deionized water under ultrasonication for 30 s at ambient conditions, prior to being deposited on copper
grids coated with holey carbon or silicon monoxide films (Agar Scientific). Most imaging work was done with silicon monoxide coated grids due to better dispersion of the samples, although at the expense of the resolution.

Thermogravimetric analysis (TGA) was performed on a Perkin Elmer Pyris 1 TGA. Samples were heated from 30-650 °C at 20 °C min⁻¹ in nitrogen.

Raman spectra of hydrotalcites powders were obtained using a commercial spectrometer (Horiba Jobin-Yvon LabRAM HR) equipped with a CCD detector. The 532 nm line from an Argon ion laser was used to illuminate the samples at 5 mW. For in situ Raman spectroscopy, the transitions during thermal decomposition of Mg:Fe:Al HT were studied in air (50 mL(STP) min⁻¹) in a high temperature cell (Linkam CCR1000). The spectra were collected in the temperature range of 25-600 °C at intervals of 20-50 °C at a heating rate 20 °C min⁻¹.

3. Results and Discussion

3.1. Reactor characteristics and reproducibility

For the reactor developed in this work, given its ID and the flow rates used, the space time, also equal to mean residence time, is estimated to be 0.34 s. The reactor’ Re number is approximately 2,857. For the composition of the reaction mixture and the conditions used in this study the theoretical limit of the micromixing time is 0.4 s. This is calculated using the engulfment mixing model [21] and the estimated energy dissipation rate (0.0032 W kg⁻¹) [22]. Thus, we operate the reactor at residence time close to the micromixing time. This is significantly faster than a cylindrical flask with a magnetic stir bar (~9.3 s) [23] and is responsible for attaining smaller particles with narrow particle size distributions. Further dilution of the product mixture in the large volume of the aging vessel should reduce growth of primary particles and limit their aggregation.

In order to quantify the reproducibility of the flow method described here and compare it to that of batch synthesis, we focused on the property that showed greatest variability in syntheses: the BET surface area (see discussion below). The surface areas for the materials synthesized in flow are substantially more narrowly distributed than those from the syntheses in batch, as shown in Table 2. Standard deviations for the HTs synthesized was less than 6.05 m² g⁻¹, which shows good control of precipitation and aging for the flow method. Greatest variability in batch synthesis was observed for the HT aged at 65 °C, which ranged from 27 – 110 m² g⁻¹ in six replicates. In comparison, four replicates of flow syntheses of the same
HT had surface areas between 139 and 148 m² g⁻¹. Similar trends were observed for the Fe HT and the Ni HT aged at 65 °C.

Table 2. Standard deviation of BET surface areas based on multiple replicates of HTs synthesized using flow procedure and by co-precipitation in batch.

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Ageing temperature, °C</th>
<th>Standard deviation of S_BET, m² g⁻¹ (# replicates)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Flow</td>
</tr>
<tr>
<td>HT</td>
<td>65</td>
<td>1.82 (4)</td>
</tr>
<tr>
<td></td>
<td>80</td>
<td>3.05 (4)</td>
</tr>
<tr>
<td>Fe HT</td>
<td>65</td>
<td>6.05 (3)</td>
</tr>
<tr>
<td></td>
<td>80</td>
<td>2.55 (3)</td>
</tr>
<tr>
<td>Ni HT</td>
<td>65</td>
<td>4.15 (2)</td>
</tr>
<tr>
<td></td>
<td>80</td>
<td>2.35 (2)</td>
</tr>
</tbody>
</table>

3.2. Structural characterization of the HTs synthesized in a meso-scale flow reactor

ICP-AES analysis of the HTs samples prepared in flow shows consistent elemental composition, closely matching the expected theoretical ratios, as shown in Table 1.

The presence of the hydrotalcites phase was confirmed by PXRD, as all doped HTs exhibited characteristic reflections of LDHs, Figure 2. The lower angle [00l] reflections ((003) at 2θ ≈ 11.4 °, (006) at 22.9° and (009) at 34.6° for all examined samples indicate formation of a layered structure with 3R symmetry and can be used to calculate the interlayer spacing parameter, c, which varies from 23.1 to 23.6 nm for the doped samples aged at different temperatures, Table 3. The value of c depends on multiple factors, including the strength of Coulombic attraction between the positively charged cation layers and the negatively charged interlayer anions [24], as well the relative quantity of interlayer anions and water [25]. For all studied HTs the c value increased when aging temperature was increased from 65 to 80 °C.

The unit cell parameter a was calculated from (110) reflection at ~60.6°. Its value depends on the nature of the doping metal, as it reflects cation-cation distance within the hydrotalcite layer, Table 3. In the HTs doped with M²⁺ metals, the ionic radii of the divalent ions isomorphically substituting Mg²⁺, are within close range to that of Mg²⁺ (0.72 Å), see Table 3. As a result, the cell parameter a remains unchanged (3.06 Å). On the contrary, in the HTs doped with M³⁺ metals, such as Fe³⁺, there is a more substantial difference in the M³⁺ ionic radius relative to that of Al³⁺ (0.54 Å vs 0.65 Å for Al³⁺ vs Fe³⁺). This difference
manifests itself in structural changes, reflected in the value of $a$: in the sample doped with 15% Fe$^{3+}$ the parameter $a$ increases to 3.09 Å compared to the neat HT sample.

For all the compositions the crystallite sizes range from 9.5 to 12.9 nm. The smallest crystallites are observed for the Ni HT samples aged at 65 and 80 °C: 9.5 and 9.7 nm respectively. Changing the aging temperature from 65 to 80 °C has little or no effect on the crystallite size, with the largest effect observed in the Fe HT sample: from 11.9 to 12.9 nm.

For the HT samples aged at 65 °C very similar PXRD patterns are obtained to those of the samples aged at 80 °C (compare Figure 2 with Figure S1 in ESI). On this basis we infer that there are minimal structural differences in the hydrotalcites aged at these two temperatures.

Raman spectra of the flow-synthesized HTs aged at 80 °C are shown in Figure 3. The spectra exhibit characteristic bands in the region of 100-600 cm$^{-1}$, which arise from M-O normal modes in the LDHs, and bands at 1040-1060 cm$^{-1}$ assigned to CO$_3^{2-}$ stretches. Raman spectra of the hydrotalcites aged at 65 °C are similar to these aged at 80 °C and are shown in Figure S2 in ESI.

Assignment of the main 548 and 474 cm$^{-1}$ bands was carried out on the basis of the observed trends in the shift of bands with substitution of M$^{2+}$ or M$^{3+}$. A comparison of the line at ~548 cm$^{-1}$ for the HT80 and the Fe HT80 samples shows a shift to lower wavenumbers with increasing iron content, Figure 3, which suggests that Fe$^{3+}$ affects this normal mode. Thus, the band at 548 cm$^{-1}$ is unlikely to be assigned to Mg-O-Mg stretching only, as in that case little or no shift would be observed with increasing M$^{3+}$ content. If the band would be assigned to M$^{3+}$-O-M$^{3+}$ stretching, no shift should be observed when substituting Mg$^{2+}$ for another M$^{2+}$, while if it were assigned to M$^{2+}$-O-M$^{3+}$ stretching, it would be sensitive to both M$^{2+}$ and M$^{3+}$ composition. Given that we do observe shifts of this band with both M$^{2+}$ and M$^{3+}$ substitution, we propose that this band can be assigned to M$^{3+}$-O-M$^{2+}$ stretching. However, it should be noted that the shift of the 550 cm$^{-1}$ band is small when M$^{2+}$ cation has comparable radius to that of Mg$^{2+}$ (e.g. Cu, Zn, Ni), as the change in bond lengths is relatively small.
Table 3. Unit cell parameters and crystallite sizes of flow synthesized hydrotalcites, as calculated from PXRD patterns.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Cell Unit a, Å</th>
<th>Cell Unit c, Å</th>
<th>Crystallite size, nm</th>
<th>Ionic radius of dopant, Å [26]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>65 °C</td>
<td>80 °C</td>
<td>65 °C</td>
<td>80 °C</td>
</tr>
<tr>
<td>HT</td>
<td>3.06</td>
<td>3.06</td>
<td>23.31</td>
<td>23.55</td>
</tr>
<tr>
<td>Fe HT</td>
<td>3.09</td>
<td>3.09</td>
<td>23.37</td>
<td>23.58</td>
</tr>
<tr>
<td>Co HT</td>
<td>3.06</td>
<td>3.06</td>
<td>23.06</td>
<td>23.23</td>
</tr>
<tr>
<td>Ni HT</td>
<td>3.05</td>
<td>3.05</td>
<td>23.44</td>
<td>23.46</td>
</tr>
<tr>
<td>Cu HT</td>
<td>3.06</td>
<td>3.06</td>
<td>23.17</td>
<td>23.23</td>
</tr>
<tr>
<td>Zn HT</td>
<td>3.06</td>
<td>3.06</td>
<td>23.14</td>
<td>23.35</td>
</tr>
</tbody>
</table>

Figure 2. PXRD patterns of flow synthesized hydrotalcites aged at 80° C: a) HT, b) Fe HT, c) Co HT, d) Ni HT, e) Cu HT, f) Zn HT.

This assignment is in agreement with a study of cation ordering in hydrotalcite layers carried out by Sideris et al [27], who examined Mg-Al HTs with different Al content by multinuclear NMR spectroscopy. They showed that for the HT with Mg/Al ratio of 3 the two most abundant cation environments are Mg₂AlOH (80%) and Mg₃OH (20%), with almost no MgAl₂OH or Al₃OH environments. Thus, the most intense Raman band of Mg-Al HT (~548 cm⁻¹) is most likely due to Mg-O-Al or Mg-O-Mg vibrations (the bands appear symmetrical, with no apparent shoulder peaks).
The band at ~470 cm\(^{-1}\) is assigned to M\(^{3+}\)-O-M\(^{3+}\), as this band shifts only with substitution of M\(^{3+}\). The even weaker band at 300 cm\(^{-1}\) can be assigned to an M-O vibration, which seems to be more profound for the samples with iron and nickel. Bands around 1040 cm\(^{-1}\) and 1060 cm\(^{-1}\) are both assigned to carbonates anions in different environments [28].

Figure 3. Raman spectra of the substituted hydrotalcites aged at 80 °C. a) HT, b) Fe HT, c) Co HT, d) Ni HT, e) Cu HT, f) Zn HT.

3.3 Thermal properties

All hydrotalcites synthesized in flow exhibited the characteristic thermal decomposition profiles of hydrotalcites, with three endothermic transitions: (I) at low temperature, <100 °C, representing the loss of physisorbed interlayer water, (II) at ~ 150-250 °C, corresponding to the loss of interlayer water and carbonates, and (III) the high temperature transition, at 350-550 °C, corresponding to dehydroxylation and decarbonation processes [29]. Depending on composition and aging temperature the shapes and positions of these transitions vary significantly (Table 4; Figures 4 and 5; ESI Figure S3a-l).

For all the flow synthesized samples, apart from the Fe HT, increasing aging temperature causes transitions II and III to shift to the lower temperatures, indicating a decrease in thermal stability. Thermal stability is an indicator of strength of Coulomb interactions between cationic layers and anions in-between these layers. We have observed the increase in parameter c upon the increase in the aging temperature, Table 3. This will reduce the electrostatic interaction between the layers and, thus, would reduce thermal stability for the loss of interlayer anions and water. Hence, we observe the lower temperature shift of
transitions II and III in DTG profiles. Depending on composition, the degree of the increase in parameter $c$ and the degree of loss of thermal stability are different, indicating differences in layer charge density and the strength of Coulomb attraction.

The order of stability of HT samples is the following, for aging at 65 °C: HT > Ni HT > Cu HT > Co HT > Fe HT > Zn HT, for aging at 80 °C: Ni HT > HT > Fe HT > Cu HT > Co HT > Zn HT. Knowledge of thermal decomposition profiles is important for controlled calcination of HTs to oxides used as catalytic materials. It has been shown that the number of basic sites of hydrotalcite-based oxides depends on the calcination temperature and goes through the maximum [30]. The phase composition of the sample determines the number of basic sites: mixed oxide phase provides higher basicity, whilst spinel phase has fewer and weaker basic sites.

The fact that the “HT” hydrotalcite with no dopant is the most stable is logical, as substitution within the structure, especially with different size cations, creates disorder and structural changes, as seen from structural analysis of the Fe HT sample by PXRD and Raman. However, thermal stability also depends on the nature of the incorporated cation. For instance, it has been reported that in the cobalt-containing HTs, Co$^{2+}$ in the octahedral environment is easily oxidized to Co$^{3+}$ during aging or drying stage of preparation, resulting in decreased stability [31].

Table 4. A summary of thermal properties of the flow synthesized HTs aged at 65 and 80 °C.

<table>
<thead>
<tr>
<th>Sample, 65 °C</th>
<th>HT</th>
<th>Fe HT</th>
<th>Co HT</th>
<th>Ni HT</th>
<th>Cu HT</th>
<th>Zn HT</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>57</td>
<td>56</td>
<td>54</td>
<td>56</td>
<td>56</td>
<td>60</td>
</tr>
<tr>
<td>DTG</td>
<td>II</td>
<td>211</td>
<td>182</td>
<td>208</td>
<td>208</td>
<td>191</td>
</tr>
<tr>
<td>(°C)</td>
<td>III</td>
<td>450</td>
<td>406</td>
<td>409</td>
<td>425</td>
<td>415</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sample, 80 °C</th>
<th>HT</th>
<th>Fe HT</th>
<th>Co HT</th>
<th>Ni HT</th>
<th>Cu HT</th>
<th>Zn HT</th>
</tr>
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<tbody>
<tr>
<td>I</td>
<td>51</td>
<td>58</td>
<td>67</td>
<td>59</td>
<td>51</td>
<td>45</td>
</tr>
<tr>
<td>DTG</td>
<td>II</td>
<td>201</td>
<td>192</td>
<td>155</td>
<td>206</td>
<td>177</td>
</tr>
<tr>
<td>(°C)</td>
<td>III</td>
<td>421</td>
<td>416</td>
<td>363</td>
<td>423</td>
<td>381</td>
</tr>
</tbody>
</table>

*I, II and III are first, second and third minima in 1st differential of TGA.*
To gain additional insight into the evolution of structure of the synthesized HTs during thermal treatment we carried out in situ Raman studies for the Fe HT 80 sample. The choice of the Fe-containing sample was due to the lower fluorescence from this material, which allowed to monitor the variation in Raman bands better, than for the other materials in the series. In the range of 150-250 °C, which also corresponds to region (II) of the thermogravimetric analysis, the main characteristic bands at 289, 462 and 531 cm\(^{-1}\) for \(\text{M-OH}\) gradually disappear, Figure 6, indicating collapse of the hydrotalcite structure. Concurrently, the 1042 cm\(^{-1}\) and 1057 cm\(^{-1}\) bands for carbonate ions (high symmetry) shift to 1033 cm\(^{-1}\) and gradually diminish to zero as temperature is increased, see Figure S4 in ESI. This is in agreement with DTG analysis, where the endothermic minimum attributed to carbonate loss is observed at 192 °C, Figure 5.

Broad bands around 3500-3600 cm\(^{-1}\), assigned to stretching vibrations of water molecules in the interlayer and the physically adsorbed surface water, also disappear in the temperature range of 150-250 °C. Bands at ~3650-3700 cm\(^{-1}\) are attributed to OH-stretching of hydroxyl groups. Concurrent decrease of 3500-3600 and 1033 cm\(^{-1}\) bands, Figure 6, is evidence of decomposition and of dissipation of the interlayer water and carbonates ions.
Figure 5. TGA-DTG curves for the hydrotalcites aged at 80 °C.

Already at ~150 °C we observe the evolution of a band at ~690 cm\(^{-1}\), which develops into a strong band at 640 cm\(^{-1}\) with further temperature increase up to 600 °C. After heating the Fe HT80 sample to 600 °C three bands in the 200-1000 cm\(^{-1}\) region can be observed: at 320, 404 and 640 cm\(^{-1}\). For comparison, the Raman spectrum of the same sample calcined \textit{ex situ} at 800 °C shows strong bands at 205, 316, 470 and 696 cm\(^{-1}\), with a shoulder at 640 cm\(^{-1}\) (ESI Figure S5). This evolving phase is a mixed oxide \textit{pre}-spinel phase, which likely changes its composition during the thermal treatment. The bands observed for the \textit{ex situ} calcined sample are generally consistent with the bands characteristic of MgFe\(_2\)O\(_4\) spinel [32]. The spinel-like phase begins to form as hydrotalcite layers lose hydroxyls at higher temperatures. The second endothermic peak in DTG with the minimum at 416 °C, Table 4, represents this dehydroxylation / decarboxylation process. The \textit{in situ} Raman data is in agreement with the DTG data, but provides finer detail on the evolution of the structure.
Figure 6. *In situ* temperature-controlled Raman spectroscopic study of thermal decomposition of the Fe HT80.

### 3.4 Morphological properties of the flow synthesized HTs

TEM images of HT samples aged at 65 and 80 °C are shown in Figure 7a,b. In both cases hexagonal platelets with diameter of 20-30 nm are observed. The average crystallites size measured by XRD is in the range of 10-12 nm, Table 3; furthermore, both PXRD and Raman spectra are characteristic of well-crystalline samples. This leads to the conclusion that the platelets observed in the TEM images are agglomerates of primary crystallites. The platelets are further loosely aggregated into larger structures. The level of aggregation seems to be visibly higher for the sample treated at the higher temperature: the samples were prepared identically for the TEM measurements, but we see more of individual platelets in the image corresponding to the HT 65 sample, compared to that in the image of the HT 80 sample. Based on the literature, an increase in aging temperature results in a higher rate of crystal growth [10]. For the
conditions used in this study there is a minor increase in the size of crystallites based on PXRD data, which does not contradict the literature. There is, however, a more pronounced aggregation of the particles upon aging, which should result in the more pronounced changes in the morphology of the materials, specifically their pore structure.

To investigate the morphology of hydrotalcites synthesized in flow and examine the effect of aging temperature on surface area and pore size distribution, nitrogen adsorption-desorption isotherms were obtained for all the samples under study. All samples exhibit type IV isotherm with an H3 hysteresis loop (ESI Figure S6), which is consistent with the presence of a mesoporous solid.

**Figure 7.** TEM images of HT samples aged at (a) 65 and (b) 80 °C.
LDHs synthesized by conventional batch co-precipitation in our laboratory have relatively low surface areas of 27-110 m² g⁻¹ with mean value of 73±28 m² g⁻¹. The BET surface areas of the flow synthesized hydrotalcites, reported in Table 5, are much higher, in the range of 80-150 m² g⁻¹. Such a direct comparison is, however, somewhat arbitrary, since $S_{BET}$ value of LDHs depends on the sample’s chemical composition and its aging temperature, and as we argue here, on the method of synthesis: batch vs flow co-precipitation. As there are few reports that examine pore structure of uncalcined hydrotalcites, we systematically considered the effect of composition and aging temperature for the flow-synthesized materials.

For the flow-synthesized HTs reported here, the surface area ($S_{BET}$) does not change significantly when increasing the aging temperature from 65 to 80 ºC, which is in agreement with results from Climent et al for the batch-synthesized samples [10]. The biggest difference in $S_{BET}$ is observed for the Ni HT and the Co HT samples, showing an increase by 23 and 19 m² g⁻¹ respectively for the samples aged at 80ºC. However, we observe significant differences in the surface area as a function of HT composition. The largest surface areas were observed for the HT65, the Ni HT80 and the Cu HT65 samples, being 148, 149 and 140 m² g⁻¹ respectively, Table 5. The Co HT65 sample exhibited the lowest specific surface area amongst the HT samples, 79 m² g⁻¹.

**Table 5.** Characterization of morphology of the flow-synthesized HTs aged at two temperatures.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Ageing Temp, ºC</th>
<th>$S_{BET}$, m² g⁻¹</th>
<th>BJH pore volume, cm³ g⁻¹</th>
<th>Pore width, nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>HT 65</td>
<td>148</td>
<td>0.26</td>
<td>3.6</td>
<td></td>
</tr>
<tr>
<td>HT 80</td>
<td>142</td>
<td>0.29</td>
<td>5.8</td>
<td></td>
</tr>
<tr>
<td>Fe HT 65</td>
<td>126</td>
<td>0.34</td>
<td>6.6</td>
<td></td>
</tr>
<tr>
<td>Fe HT 80</td>
<td>120</td>
<td>0.37</td>
<td>8.3</td>
<td></td>
</tr>
<tr>
<td>Co HT 65</td>
<td>79</td>
<td>0.21</td>
<td>7.3</td>
<td></td>
</tr>
<tr>
<td>Co HT 80</td>
<td>98</td>
<td>0.33</td>
<td>10.5</td>
<td></td>
</tr>
<tr>
<td>Ni HT 65</td>
<td>126</td>
<td>0.17</td>
<td>3.5</td>
<td></td>
</tr>
<tr>
<td>Ni HT 80</td>
<td>149</td>
<td>0.21</td>
<td>4.5</td>
<td></td>
</tr>
<tr>
<td>Cu HT 65</td>
<td>140</td>
<td>0.27</td>
<td>3.6</td>
<td></td>
</tr>
<tr>
<td>Cu HT 80</td>
<td>126</td>
<td>0.28</td>
<td>6.2</td>
<td></td>
</tr>
</tbody>
</table>
The more pronounced influence of both composition and the aging temperature is observed in the pore size distributions, calculated using the BJH method. Results are shown in Figures 7a-f. Despite having little effect on $S_{BET}$, aging temperature has a profound effect on the morphology of hydrotalcites. For the standard sample “HT”, aging at 65 °C results in the formation of narrowly distributed pores with the mean diameter of 3.6 nm. The increase in aging temperature for the HT sample to 80 ºC, and subsequently to 95 °C, clearly results in larger pores, Figure 7a, with the average pore diameters of 5.8 and 7 nm respectively.

The interlayer spacing between hydrotalcite sheets, as determined by PXRD, is smaller - 2.3 nm, Table 3. This implies that $N_2$ adsorption is measuring pores between particles and not the interlayer space. Using simplistic geometrical considerations, if we assume that pores are spaces between aggregated particles and that our particles are disks, as we observe in the TEM images in Figure 5, then the 3.6 nm diameter pore would translate into a 23.2 nm diameter disk. Details of the calculation are given in SI. This corresponds surprisingly well to the TEM images in Figure 5, where we note that the shape of the agglomerated particles is disk-like and the diameters of the agglomerates are in the order of 20-30 nm. We conclude that the measured pores are between the agglomerated particles within the larger aggregates.

In the case of the Fe HT samples the measured pore diameters are 6.6 and 8.3 nm respectively for the two aging temperatures, Table 5. Following the same argument as for the case of the HT sample, this implies that the rate of the agglomerated particle growth is faster in the case of the Fe HT sample, than that for the HT sample: the crystallite sizes, based on PXRD data, do not vary significantly between the HT samples as a function of composition. Therefore, the large variation in the pore sizes could only be attributed to the differences in the sizes of the agglomerates of the crystallites.

The cobalt-doped sample forms large agglomerates of crystallites already when aged at 65 °C. The porous structure is formed in-between the agglomerates, with an average pore size of 7.3 nm, which increases to 10.5 nm when aged at 80 °C, Figure 7c. The BJH pore volume increases from 0.21 to 0.33 cm$^3$ g$^{-1}$ when the aging temperature is increased, indicating that the size and the number of pores are both increasing, Table 5.
The nickel HT sample is somewhat different from all the other doped samples. Its surface area and the number of pores increase when the aging temperature is increased from 65 to 80 °C, as the BJH pore volume increases from 0.17 to 0.21 cm$^3$ g$^{-1}$, Table 5. Yet the pore size changes little with the increase in the aging temperature and is similar to the HT65 sample. This suggests that the rate of agglomeration of primary crystallites in the Ni HT sample does not depend so much on temperature.

The copper-doped hydrotalcite samples show a very similar behavior to the undoped HT samples, with a narrow pore size distribution when aged at 65 °C (mean pore diameter of 3.6 nm), which broadens when aged at 80 °C (average pore size of 6.2 nm, Figure 7e). However, the number of pores seems to stay the same as the BJH pore volume does not change and the surface area even decreases.

The Zn HT samples demonstrate a principally different morphology compared to the other doped samples. The Zn HTs have two pore size distribution maxima when aged at either 65 or 80 °C, Figure 7f, suggesting the formation of several types of agglomerates of different sizes. Both Zn HT65 and Zn HT80 have pores of average size of ~ 10 nm, as well as a second set of much larger pores, especially for the sample aged at 80 °C.

There appears to be a correlation between pore size and thermal stability of hydrotalcites samples doped with M$^{2+}$ metals: all samples with small pores, which we attribute to small agglomerates of primary crystallites (HT 65, Ni HT65, Ni HT80, Cu HT 65) also exhibit superior thermal stability for loss of interlayer carbonates. This may be a result of similar physical phenomena controlling both, thermal stability and the crystallites agglomeration rate.

**3.5 Comparison with other flow synthesis methods**

We can compare performance of the reactor developed in this work and the obtained materials with the reported to-date precipitation syntheses of LDHs in flow in the open literature. There is a very small number of reports of continuous synthesis of LDHs in the open literature. Thus, a study of the effect of different mixing regimes attained in a microchannel reactor compared with a stirred tank baffled reactor on particle size distribution of hydrotalcites demonstrated that hydrotalcites particles of smaller mean size are obtained in flow with a lower power input, compared to the stirred tank [11]. The particle sizes obtained were in the order of 10 µm, which are claimed to be non-agglomerated. This, however, was not supported by any structural characterization of the synthesized materials; particles of hydrotalcites in this range of sizes are certainly agglomerates. As we have shown above, the primary crystallites of hydrotalcites are in the order of 9-12 nm and any observed larger structures are aggregates.
Narrow particle size distributions and particles in the range of 20-50 nm were obtained in microreactors in two studies. Thus, plate-like particles with the mean size of 50 nm were obtained following different durations of hydrothermal aging of the materials precipitated in the microreactor at Reynolds numbers of approximately 6,200 [12]. These are much higher Re numbers than those used in the present study, whereas the size of the obtained particles is twice larger, possibly due to the overall residence time or the impact of high sheer in the microreactor. Plate-like particles of a similar size to those obtained in this work, approx. 20-30 nm, where obtained in another microreactor synthesis of hydrotalcites [13]. Microreactors do allow to obtain small particles with a narrow distribution of particle sizes. However, the product yield of microreactors (space-time-yield) is rather small for industrial production.

In a study using a meso-scale reactor with embedded static mixer elements, particles with the mean size of ~ 40-45 nm were obtained, similar to some of micro-reactor studies, but with a very broad size distribution [14]. There is no data in the paper to calculate residence time or micromixing time for their specific system. It is therefore most likely that the choice of conditions for synthesis in the static-mixer fitted reactor was much larger than the micromixing time.

In other studies, a continuous version of the hydrothermal synthesis of LDH-type materials was explored. The one-step continuous method combines co-precipitation with aging into a single process, which reduces the overall time to obtain crystalline LDH materials compared to the two-step processes. However, in some cases the resulting materials were highly heterogeneous, contained impurities of other inorganic phases and had low surface areas of around 6 m² g⁻¹ [15, 16]. With more precise control of the reactants concentration in the hydrothermal continuous synthesis of functionalized LDHs, materials with delaminated structures were obtained [17, 18]. This method is potentially suitable for scaling the manufacture of high-surface area functionalized LDH-based materials, typically obtained via a hydrothermal synthesis in very low yields. However, there is little further work reported to date.

There are other possibilities to affect mixing and hence nucleation step in the hydrotalcite synthesis, apart from performing the reaction in flow. For example, synthesis was performed in batch under high sheer mixing, affording primary crystallite sizes of ~ 14-30 nm depending on Al: Mg ratio, and resulting in the disk-like particles with a relatively narrow size distribution of the aggregates with the mean of ~0.1 µm [33]. In this work the rates of mixing were significantly enhanced due to high sheer in the colloidal mixer, but the operation was batch-wise with 60 s reaction time, followed by aging. Improvement in mixing time
is a significant advance over conventional batch syntheses of HTs, but the broad residence time distribution of the batch reactor will favor, by its physical nature, broad distribution of compositions and hence of primary crystals and then the aggregates. High sheer mixer was later implemented as a continuous stirred tank reactor (CSTR) with a space time of 18 s [34]. The residence time distribution in CSTR is still very broad, which necessarily results in a broad distribution of compositions.

Thus, analyzing the available literature on the flow syntheses of LDHs by co-precipitation we note that plate-like particles with the aggregate sizes in the range of 20-50 nm could be obtained, compared to large 3D particles typically synthesized in batch. Furthermore, although narrow particle size distribution could be attained in micro-reactors, we show that it is sufficient to reduce the size to a mm-scale reactor, easier for scaling, but to carefully select the conditions such that mixing time is in the order of the micromixing characteristic time. This results in the narrow distribution of not only particle sizes, but also the key morphological properties of the obtained materials.

4. Conclusions
In this study we thoroughly investigated the structure and morphology of the doped hydrotalcites synthesized under continuous co-precipitation in a meso-scale flow reactor. The new synthesis method is proposed to overcome the limitations of the widely used batch co-precipitation: irreproducibility of the synthesis and significant variance of properties obtained in the batch co-precipitation due to lack of control over mixing in the process. The hypothesis was that the mixing time in the reactor should be commensurable to micromixing characteristic time. This was proven by obtaining a significantly more narrow distribution of the key property of the flow-synthesized materials – specific surface area, measured as BET surface area – as well as by higher degree of crystallinity and smaller particle sizes obtained in flow.

Within the range of the flow synthesized doped hydrotalcites we observed high sample crystallinity, high surface area and pore volumes. We obtained trends of thermal stability and proposed a simple geometrical explanation for the formation of the pore structure of the hydrotalcites materials. Crystallite size, shape and pore structure are all important technical characteristics of the final materials for most applications, such as electronics, ion-exchange and catalysis. The flow synthesis method allows to obtain materials with a high degree of crystallinity for various compositions. The crystallite sizes vary within 2.5 nm for all the different compositions. However, the pore sizes and, hence, the agglomerates of the crystallites vary significantly. For the conditions of flow synthesis used in this study, we conclude that the formation of the secondary structure is strongly dependent on the composition and conditions of post-synthesis (aging) when conditions of the flow co-precipitation are fixed at the characteristic time of micromixing.
Overall, the flow synthesis method affords HT materials with a narrow distribution of properties. This, in turn, allows one to perform a systematic study of the effects of composition and synthesis conditions on the material properties of interest, opening the way to rational design of LDHs with targeted properties and scale up of their manufacture.
Figure 8. Pore size distributions for the flow synthesized HTs. (a) undoped hydrotalcites, (b) Co-doped hydrotalcites, (c) Fe-doped hydrotalcites, (d) Ni-doped hydrotalcites, (e) Cu-doped hydrotalcites, (f) Zn-doped hydrotalcites. The aging temperatures for the samples are given in the sample labels in the graphs.
Supporting Information

Supporting information includes detailed synthesis composition, additional XRD and Raman data, full set of DTG-TGA data, details of the in situ Raman spectra, nitrogen sorption isotherms and details of the geometrical model used to estimate pore sizes.

Acknowledgements

This research was in part supported by the National Research Foundation, Prime Minister’s Office, Singapore under its CREATE programme. AV thanks the NSF Chemistry Division for funding through the CAREER program (award 1554963).

References


Continuous synthesis of doped layered double hydroxides in a meso-scale flow reactor

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Supporting Information

Content

1. Synthesis compositions
2. Additional XRD data
3. Additional Raman data
4. Full set of DTG-TGA data
5. Details of in situ Raman spectra
6. Nitrogen sorption isotherms
7. Details of geometrical model for pore size – particle size calculation

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1. Synthesis compositions

Table S1. Synthetic protocol.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Metals ratio</th>
<th>C Mg(NO$_3$)$_2$, M</th>
<th>C Al(NO$_3$)$_3$, M</th>
<th>C dopant nitrate, M</th>
</tr>
</thead>
<tbody>
<tr>
<td>HT</td>
<td>3:1</td>
<td>0.21</td>
<td>0.07</td>
<td>-</td>
</tr>
<tr>
<td>Fe HT</td>
<td>3:0.4:0.6</td>
<td>0.21</td>
<td>0.03</td>
<td>0.04</td>
</tr>
<tr>
<td>Co HT</td>
<td>2.4:1:0.6</td>
<td>0.17</td>
<td>0.07</td>
<td>0.04</td>
</tr>
<tr>
<td>Ni HT</td>
<td>2.4:1:0.6</td>
<td>0.17</td>
<td>0.07</td>
<td>0.04</td>
</tr>
<tr>
<td>Cu HT</td>
<td>2.4:1:0.6</td>
<td>0.17</td>
<td>0.07</td>
<td>0.04</td>
</tr>
<tr>
<td>Zn HT</td>
<td>2.4:1:0.6</td>
<td>0.17</td>
<td>0.07</td>
<td>0.04</td>
</tr>
<tr>
<td>MgFe HT</td>
<td>3:1</td>
<td>0.21</td>
<td>-</td>
<td>0.07</td>
</tr>
</tbody>
</table>

2. Additional XRD Data

Fig S1. XRD patterns of substituted hydrotalcites aged at 65° C. a) HT, b) Fe HT, c) Co HT, d) Ni HT, e) Cu HT, f) Zn HT.
3. Additional Raman data

Figure S2. Raman spectra of substituted hydrotalcites aged at 65 °C. a) HT, b) Fe HT, c) Ni HT, d) Zn HT, e) Cu HT, f) Co HT.
4. Full set of TGA-DTG data

![Graph a) HT 65](image)

![Graph b) HT 80](image)
**Figure S3.** a-l) TGA-DTG analysis of flow synthesized HTs.
5. Details of in situ Raman data

**Figure S4.** *In situ* Raman spectra of Fe HT 80, enlarged 900-1150 cm$^{-1}$ region.
Figure S5. Comparison of Fe HT sample calcined in situ at 600°C (black) and ex-situ at 800°C (blue).
6. Nitrogen sorption Isotherms

- **a)**
- **b)**
- **c)**
- **d)**
- **e)**
- **f)**
Figure S6. N$_2$ BET adsorption isotherms for doped HTs aged at 65 - 80 °C.
7. Details of geometrical calculation

**Figure S7.** Illustration of geometry for calculation of pore diameter formed between circular plates.

\[
\cos(\theta) = \frac{R}{R + r}
\]

\[
R = \frac{r}{1 - \frac{1}{\cos(\theta) - 1}}
\]

For pore diameter 3.6 nm, \( r = 1.8 \) nm, \( R = 11.6 \) nm, which gives particle diameter 23.2 nm.