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Low-Temperature Continuous Flow Synthesis of Metal Ammonium Phosphates

Alistair F. Holdsworth^{1,2}, Harry Eccles¹, Alice M. Halman¹, Runjie Mao¹ & Gary Bond¹

The synthesis of the high performance inorganic materials essential to the quality of modern day life is hindered by traditionalist attitudes and reliance on outdated methods such as batch syntheses. While continuous flow methods have been extensively adopted in pharmaceutical circles, they remain largely unexplored for the preparation of inorganic compounds, despite higher efficiency, safety and versatility. In this publication, we demonstrate a step-change for the synthesis of metal ammonium phosphates through conversion of the extant batch process to a low-temperature continuous regime, exhibiting a tenfold increase in throughput combined with a significant decrease in particle size.

Although inorganic materials are essential to life in the 21st century, advancements in their preparation are underutilised as traditionalist attitudes in academia and particularly industry mean batch processes predominate. Continuous flow synthesis (CFS) is one such alternative technique: it is prevalent in pharmaceutical research^{1,2}, yet has received lesser attention for inorganic synthesis outside of nanomaterials research³. This is largely due to the typically high capital costs associated with flow systems⁴; despite the technique's inherent flexibility^{1,2,4}, safety^{1,2}, scalability^{1,3,5,6}, and controllability^{1-3,7,8}. Low temperature (<100 °C), ambient pressure flow syntheses presents an affordable preparative route to a plethora of inorganic materials⁸⁻¹⁵, without the high capital cost of proprietary systems⁴. In this communication, we demonstrate a step-change, low-cost continuous preparation of versatile metal ammonium phosphates and derivatives. We compare CFS with the commonly used batch method¹⁶ and demonstrate a tenfold increase in throughput combined with a significant reduction in particle size for continuously synthesised particles.

Metal ammonium phosphates (MAPs) and other compounds of the form $AMPO_4 \cdot xH_2O$ (where A^+ is Na, K, NH_4 and M^{2+} is a divalent metal such as Mg, Fe or Zn) were first explored in detail by Bassett and Bedwell in the 1930s¹⁶. These compounds have since been investigated for a myriad of uses including catalysis¹⁷⁻¹⁹, pigments¹⁷, fertilisers²⁰, energy storage²¹, biomaterials⁸⁻¹¹, flame retardants¹⁷ and as ion exchange materials²²⁻²⁸. Their preparative route, however, has changed little since the initial research¹⁶: an inefficient batch precipitation process requiring vast excesses of phosphorus reagent, and extended reaction times (>3 h) at high temperatures (>80 °C) to fully crystallise the product. Hydrothermal and solvothermal routes to MAPs and related compounds have been explored²⁹⁻³³, though these are often as, if not more time, resource and energy intensive than the traditional precipitation route.

Based upon a novel low temperature (<100 °C) ambient pressure continuous flow synthesis of the biomaterials brushite and hydroxyapatite⁸⁻¹¹, we adapted the synthesis of MAPs to a continuous mode. Bassett noted that an additional source of NH_4^+ or performing the precipitation at an elevated pH, can serve to increase the rate of crystallisation of some MAPs¹⁶. Through a combination of these factors and the rapid mixing of a continuous flow regime, we are able to reduce the reaction time for preparation of MAPs from a typical 180 minutes down to 7 minutes. An excess of phosphate relative to the metal is required to prevent formation of the undesired $M_3(PO_4)_2$ phase^{16,34}.

We prepared six MAPs ($M = Mg, Mn, Fe, Co, Ni$ and Zn) and one related hydrogen phosphate ($M = Sn$) using this technique in a variety of hydration states. Our CFS process provides two feeds: one containing the metal (M^{2+} , 0.05 M) typically as the nitrate salt, and the other a mixture of triammonium phosphate (TAP, 0.25 M) and ammonium nitrate (AN, 0.5 M)⁸⁻¹¹. These feeds mix continuously before heating the flow, which serves to crystallise the initially amorphous product⁸. For some metals (notably Fe and Co), this crystallisation results in a clear colour change. The characterisation data of our MAPs are presented in Table 1, with comparative particle sizes,

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M	XRD Phase		Yield		Particle Size (CFS, μm)			Particle Size (Bx, μm)			Throughput		BET
	(CFS)	(Bx)	% (CFS)	% (Bx)	D ₁₀	D ₅₀	D ₉₀	D ₁₀	D ₅₀	D ₉₀	CFS	Bx	m ² /g (CFS)
MgAP*	Mon/Hex	Mon	94	99	2.5	13.6	44.3	14.6	24.6	66.0	6.6	1.3	15.7
MnAP	Mon	Mon	99	99	6.4	22.8	65.1	7.4	18.4	42.2	8.4	1.5	6.8
FeAP	Mon	Mon	99	99	2.9	9.5	23.9	15.4	52.8	108.3	8.4	1.6	49.8
CoAP	Mon	Mon	97	99	4.9	23.5	51.3	15.8	65.0	135.9	8.4	1.6	6.5
NiAP	Hex	Mon	83	99	1.3	7.7	26.6	2.6	12.1	81.4	7.1	1.6	9.5
ZnAP	Anhyd	Anhyd	98	99	2.0	9.7	20.9	6.3	19.9	45.5	7.9	0.5	3.6
SnHP	Anhyd	Anhyd	79	78	2.7	6.5	22.8	2.3	19.4	40.3	7.5	1.5	4.7

Table 1. Characterisation data for MAPs: reaction yield (%), particle size for flow (CFS) and batch (Bx) produced material as 10th, 50th and 90th percentiles (D₁₀, D₅₀ and D₉₀, μm), BET surface area (m²/g) and observed XRD crystal hydration phase. MgAP (*) required the addition NH₃ to the Mg²⁺ feed in flow to precipitate the desired product⁸.



Figure 1. SEM images of FeAP platelets, ZnAP flowers and SnHP rods.

yields, and throughputs for material produced using the common batch process¹⁶. Reaction yields are expressed as a percentage of the theoretical maximum. Throughputs are expressed in grams per litre of liquor per hour, accounting for the reaction time.

MgAP forms a mixed-phase mono- and hexa-hydrate in CFS, and a monohydrate in batch (Fig. S1)^{19,35–37}. MnAP, FeAP and CoAP are all prepared in the monohydrate dittmarite phase regardless of synthesis method (Figs S2–S4)^{19,35–37}, forming plate-like crystallites (Fig. 1a). NiAP forms the hexahydrate (Fig. S5) struvite-like phase in flow, and the monohydrate in batch³⁴. ZnAP forms an anhydrous phase regardless of synthesis method (Fig. S6)¹⁶, forming flower-like crystallites (Fig. 1b). SnHP forms anhydrous rod-like particles (Figs 1c and S7).

MgAP can interchange between mono- and hexahydrated phases through hydrolysis and dehydration respectively^{35,38}, though our continuous flow process forms a mix of the two phases, though a longer or hotter reaction time fully dehydrates the product. NH₃ was added to the Mg²⁺ feed to prevent formation of MgHPO₄⁸. Mn, Fe and CoAP all favour the monohydrated phase and were prepared in this form in both batch and flow^{16,19,36,37}. NiAP can, like MgAP, interconvert between the two phases, though the monohydrate phase is metastable and the synthesis is sensitive to the reaction conditions used³⁴. The NiAP batch sample was of the monohydrate phase as the crystals have sufficient time to dehydrate. No stable ZnAP hydrate has been prepared as the system favours the anhydrous form¹⁶. We were unable to synthesise SnAP as hydrolysis of the acidic Sn²⁺ ion takes precedence over the formation of the desired phase. SnHP is prepared by exchanging the triammonium phosphate for phosphoric acid and omitting the ammonium nitrate. The phases of the batch produced materials match those of the flow produced materials except for NiAP³⁴, which is synthesised as the monohydrate in batch¹⁶. Neither CaAP or CuAP can be synthesised via our continuous flow route, as hydroxyapatite³⁹ (for Ca) or a hydroxyphosphate⁴⁰ (for Cu) form preferentially instead. Bassett noted the difficulty in synthesising CuAP relative to other divalent metals¹⁶.

The yields of all our flow process were excellent (>94%) except for NiAP (83%) and SnHP (79%) due to the marginal solubilities of the metals in aqueous ammonia and acid respectively^{41–43}. The yields can be increased through process optimisation. Yields for the batch produced material were quantitative, except for SnHP, where the yield matched the flow produced example. All of our flow produced materials, except MnAP, were at least half the size than the batch produced materials with narrow size distributions. Flow-produced FeAP is less than one fifth the size of equivalent batch material. The reduction in particle size is due to the increased rates of mixing and turbulence within the flow process and our adapted reaction conditions which promote a significant reduction in crystallisation time. For MnAP, the larger sized particles produced in flow are due to increased Ostwald ripening during the crystallisation process⁴⁴. The observed surface areas of our flow-produced compounds vary with the nature of the crystalline product. FeAP shows a BET high surface area as the plate-like crystals are significantly

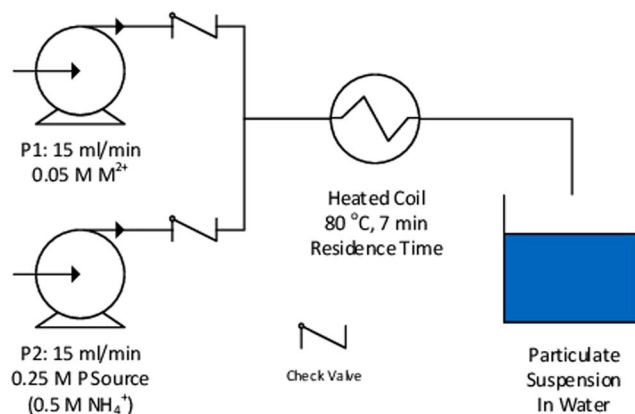


Figure 2. Schematic of CFR.

thinner (Fig. 1a) than those of the similarly plate-like MnAP and CoAP. Throughputs for the CFS process are at least five times that of the batch process, and up to sixteen times for ZnAP due to the lengthy reaction times required in batch¹⁶.

The efficiency of the CFS process can be further improved by the feasible regeneration of the effluent liquor. Concentration, replacement of precipitated phosphate and addition of the required NH_3 allows for use in further syntheses. Recovery of process heat by pre-heating precursor streams would further increase efficiency. Changing reagent stoichiometry, overall concentrations, and process parameters such as temperature, flow rate and mixing regime would allow for control of particle sizes produced within the CFR⁴⁵. Use of specialist mixers such as confined impinging jet mixers may also allow for preparation of MAPs on the nanoscale due to more turbulent mixing during the precipitation step⁴⁵. The flows within our demonstrated system are all laminar.

Conclusions

We have developed an efficient continuous flow synthesis of metal ammonium phosphates and compared this to traditional batch methods. We have demonstrated a step change in the preparation of these compounds as synthesised particles are smaller, more evenly sized and produced with far greater efficiency than previous syntheses. In comparison to the traditional batch methods, we observe up to a tenfold increase in throughput (in terms of mass per unit volume per unit time) with our flow process. This value can be increased further with process optimisation. For many applications, smaller particles are preferable as higher surface areas provide higher activities. Continuous flow synthesis provides this benefit with the option of greater control over particle size and morphology. Through the application of appropriate engineering and chemistry, the syntheses of other key inorganic materials could be improved in a similar manner as demonstrated here.

Experimental

$\text{Mg}(\text{NO}_3)_2$, $\text{Mn}(\text{NO}_3)_2$, $(\text{NH}_4)_2\text{Fe}(\text{SO}_4)_2$, $\text{Co}(\text{NO}_3)_2$, $\text{Ni}(\text{NO}_3)_2$, $\text{Zn}(\text{NO}_3)_2$, SnCl_2 , NH_4NO_3 , H_3PO_4 , NH_4OH , and $(\text{NH}_4)_2\text{HPO}_4$ were obtained from Fisher Scientific as reagent grade (>98%) quality and used as acquired with no further purification. $(\text{NH}_4)_3\text{PO}_4$ was prepared *in situ* by the addition of excess ammonia to a concentrated solution of $(\text{NH}_4)_2\text{HPO}_4$ and gently heating the resultant mixture to 65 °C so as to remove the excess ammonia. The solution was then cooled and diluted to the required concentration. Deionised water (>18 M Ω /cm) was used for all syntheses.

Our continuous flow reactor (CFR) was constructed based on previous literature references^{8–11}. Simply, this consists of two peristaltic pumps (Watson Marlow SciQ 323), a simple Y-mixer and a coil of PVC tubing (16 m length, 4 mm internal diameter, wall thickness 1 mm, total volume 210 cm³) contained within a thermostatically-controlled water bath (Grant JB Aqua 18 Plus). This is illustrated in Fig. 2.

The rotational rate and thus flow rate of each pump can be varied from 5–40 ml/min, corresponding to a total flow rate of 10 to 80 ml/min and thus a residence time of between 3 and 20 minutes, though a total flow rate of 30 ml/min and thus a residence time of 7 minutes was used in this work. Both pumps were run at the same flow rate in all cases, with one providing the metal salt feed, and the other the phosphorus source and any ancillary reagents. The temperature of the water bath could be varied between ambient and 100 °C. All reactions presented here were performed at 80 °C. A narrow aperture (c.a. 1 mm) at the outlet of the CFR served to even the flow within the system, and check valves after each pump ensured a continuous unidirectional flow. Products were purified by repeated washing with deionised water, and were then dried overnight at 80 °C. The batch syntheses of MAPs were adapted from the work of Bassett and Bedwell¹⁶, using the same procedure but scaled to give the same volume of product as in our flow process.

The rate of heating within the process is rapid, as a total flow rate of greater than 120 ml/min emerges from the reactor at 80 °C with the same tubing setup utilised for these syntheses, suggesting that temperature equilibrium is attained within 2 minutes and thus 5 minutes are spent at temperature to allow completion of reaction. Further modelling of this factor is beyond the scope of the research presented here.

Powder X-Ray Diffraction (PXRD) analysis was conducted using a Bruker D2 Diffractometer with a copper $\text{K}\alpha$ radiation source with data collected between 5 and 80 degrees 2θ . XRD patterns were converted from

proprietary formats using PowDLL⁴⁶. Scanning Electron Microscopy (SEM) and X-Ray Energy Dispersive elemental analysis was conducted under high vacuum using a FEI Quanta 200 scanning electron microscope equipped with an EDAX Sapphire Energy Dispersive X-Ray detector. Surface areas were determined using the Brunauer–Emmett–Teller (BET) model at 77 K using nitrogen absorption (Micrometrics ASAP2010), with accuracy checked against an alumina standard. Particle size distribution analysis (PSD) was conducted using a Malvern Mastersizer 2000. Samples were sonicated in deionised water before analysis to disperse any agglomerates. Results are expressed as D₅₀, the 50th percentile of particle size.

Data Availability Statement

The data underpinning this publication can be found on <https://clock.uclan.ac.uk/>.

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Author Contributions

Principal Investigator of this research was G. Bond, the manuscript was co-written by H. Eccles and A. Holdsworth. Experiments were carried out by A. Holdsworth, with BET experiments carried out by A. Halman. Results were analysed and interpreted by A. Holdsworth and R. Mao.

Additional Information

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