Model Validation of Nutrient Precipitation Using Bench-scale Reactor

Schneider, P., Pace, K., Kirsten, E. and Natividad, L.
College of Science, Technology and Engineering, James Cook University, Townsville, QLD 4811 AUSTRALIA

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Summary of key findings
A model for nutrient recovery by the precipitation of struvite is compared to the operation of a 12-litre bench precipitation reactor, using a defined medium as feedstock. The reactor was operated for durations up to 125 hours and its operation was mimicked by the model simulations. Both the model and the experimental data show the transition from a mass transfer limited system to a feed rate limited system. An apparent rate coefficient is adjusted using experimental data, including crystal mass fraction and orthophosphate concentration in solution. Recovery rates of 75% and 90% were observed at residence times of 2.5 and 4 hours, respectively. This model may be of utility to practitioners wishing to assess process designs and operating policies for their nutrient recovery systems.

Background and relevance
Large scale recovery of mineral phosphates through chemical precipitation will play a key role in society’s future nutrient recovery endeavours. Reducing potential risks resulting from uncertain process design requires deep process understanding. Furthermore, the operation of existing and future nutrient recovery plants would aim to minimise required instrumentation and control. Such process understanding can be gained through the co-application of process experimentation and process modelling.

The process in question is that of the precipitation of struvite, or magnesium ammonium phosphate, according to the following stoichiometry

$$Mg^{2+}_{(aq)} + NH_4^+_{(aq)} + PO_4^{3-}_{(aq)} + 6H_2O \rightleftharpoons MgNH_4PO_4 \cdot 6H_2O_{(s)}$$

While the past decade has seen a number of experimental contributions (Battistoni, Boccadoro, Fatone, & Pavan, 2005; Battistoni, De Angelis, Prisciandaro, Boccadoro, & Bolzonella, 2002; Battistoni, Pavan, Prisciandaro, & Cecchi, 2000; Britton et al., 2005; Fattah, Mavinic, Koch, & Jacob, 2008; Huang, Mavinic, Lo, & Koch, 2006), modelling studies are limited (S C Galbraith & Schneider, 2014; S.C. Galbraith, Schneider, & Flood, 2014; Hanhoun et al., 2013; Mehta & Batstone, 2013). This is somewhat paradoxical, since modelling has a faster development cycle and comes at much lower cost, compared to that of pilot and, most certainly, full-scale experimentation.

The process under study is presented schematically in Figure 1. This system is hybrid-mode; the solid phase is assumed to be retained within the reactor (i.e. batch mode) and the liquid phase flows through the well mixed volume (i.e. continuous mode). The inlet contained nitrogen, phosphorus and magnesium in a 1:1:1 elemental ratio. A base stream was employed to raise pH, driving the solution to a supersaturated state. Seed crystals (0.42 wt%) were added to a struvite saturated solution at the beginning of the campaign and recovered struvite is harvested at the end.

An model of this process was previously presented (Schneider, Wallace, & Tickle, 2013) and here modified to describe a simpler nutrient solution, while retaining all other features of that model. Of interest to this work is the reformulation of the molar production rate of struvite, which can be written

$$Rate = 3 \left( \frac{\rho \ \pi}{MW \ 6} \right)^{1/3} N^{1/3} (Mol)^{2/3} \frac{dL}{dt} = k_{app} (Mol)^{2/3} \ SI^n$$

The brace-bracketed term contains constants relating to particle geometry and conversion factors, $N$ is the total number of struvite particles, $Mol$ is the molar amount of struvite and $dL/dt$ is the average volume-equivalent spherical diameter growth rate of struvite particle population.
It is clear that the molar formation rate of struvite is driven by the struvite $SI$ and depends on the surface area of struvite available (i.e. $Mol^{2/3}$).

We operated a 12-litre, mixed-mode nutrient recovery reactor. In parallel we simulated the model and manually regressed the $k_{app}$ and growth rate order, $n$. To quantify the accumulation of solids in the reactor, the reactor contents very well mixed by speeding up the impellor and samples were taken at three points up the length of the side wall of the reactor. The filtrate was oven dried at 40°C and a mass balance was carried out to determine the weight fraction of particles. Dissolved orthophosphate concentration was determined using the ammonium molybdate spectrophotometric method.

**Results**

Our reactor was operated through several campaigns ranging from 30 to 125 hours. Figure 2 shows the reactor at several stages throughout a 70-hour campaign. It is apparent that some of the precipitated struvite is highly suspended, however the suspended particles eventually settled into a bed, which is clearly evident at 70 hours.

Figure 3 shows the dissolved orthophosphate concentration in the reactor liquid phase and the feed concentration throughout the 125-hour operating campaign. The combined feed rate to the reactor was reduced at the 49th hour, which is seen in the jump of reactor orthophosphate concentration in both experimental and simulated results. This reduction in feed rate increased the mean residence time of liquid from 2.5 hours to 4 hours.

Figure 4 shows the time course of struvite solids fraction in the reactor over the 70-hour campaign with the simulated results overlaid. Clearly the model and the experimental results were not in very good agreement.
The order of the power law rate equation was chosen as $n = 2$ on the basis of previously published work and since this gave reasonable agreement to the dynamic behaviour of the real process. The rate coefficient was regressed to a value of $k_{app} = 0.0285 ~ [mol/h]$.

Nutrient recovery rates were 75% at a residence time of 2.5 hours and over 90% at a residence time of 4 hours, which is also in line with the model predictions.

**Discussion**

It is clear that the 12-litre reactor operated in mixed mode, owing to the very clear supernatant present in the reactor, which supported the justification made in the process model.

The nutrient dynamics of orthophosphate concentration in the solution were well matched by the model, after the kinetic parameters were determined. Clearly, the mass retained in the reactor is significantly under predicted by the model. The experimental data shows the solids fraction appears to indicate that significant quantities of struvite may have been entrained in the exiting flow. This effect was reduced, since after about the 24th hour, the solids mass fraction increases as expected.

Of note is the linear response in the solids contained in the reactor over the processing time, which is mimicked by the process model. This indicates that mass transfer from liquid to solid phase was not limited by available surface area, but by the feed rate of nutrients to the reactor.
References


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Presenting Author

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<th>Dr Phil Schneider</th>
<th>James Cook University</th>
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Is the presenting author an IWA Young Water Professional? N (i.e. an IWA member under 35 years of age)

Bio: Phil Schneider is an academic at James Cook University who teaches into the Chemical Engineering program and carries out research in the area of nutrient recovery. His research goal is to apply chemical engineering principles to the issue of nutrient recovery, so that we can confidently design nutrient recovery systems that operate economically and, ideally, with a minimum of operator intervention. He enjoys teaching his undergraduate students and working with his research students and collaborators around the world.