Effect of the stirring speed on the struvite formation using the centrate from a WWTP

Efecto de la velocidad de agitación en la formación de estruvita usando el centrado de una planta de tratamiento de aguas residuales (PTAR)

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ABSTRACT: The formation of struvite (MgNH₄PO₄·6H₂O) for nutrient recovery in wastewater treatment plants has been widely investigated; however, little attention has been paid to the effect of stirring speeds on the resulting particle size, which could affect its agronomic value as a slow-release fertilizer. In this study, struvite formation from the centrate of sewage digestate was performed under six stirring speeds (0, 100, 200, 300, 400, 500 rpm). The resulting struvite crystals were characterised using X-ray diffraction and scanning electron microscopy with energy dispersive X-ray spectroscopy. The average particle size of struvite crystals increased from 55 µm at 0 rpm to 127 µm at 100 rpm and 128 µm at 200 rpm. Further increments in stirring speeds resulted in smaller crystal sizes. These results indicated that the largest particle size can be obtained at stirring speeds ranging from 100 to 200 rpm, equivalent to a velocity gradient between 79 and 188 s⁻¹, as there was no statistically significant difference between mean values (t-test, p<0.05). The optimum stirring speed range reported herein can be used to set operational conditions for struvite crystallisation with the benefit of producing large crystals and reducing energy consumption in stirring tanks.

RESUMEN: La formación de estruvita (MgNH₄PO₄·6H₂O) para la recuperación de nutrientes en plantas de tratamiento de aguas residuales ha sido ampliamente investigada; sin embargo, se ha prestado poca atención al efecto de las velocidades de agitación en el tamaño de partícula resultante, lo que podría afectar su valor agronómico como fertilizante de liberación lenta. En este estudio, la formación de estruvita del centrado se realizó a seis velocidades de agitación (0, 100, 200, 300, 400, 500 rpm). Los cristales de estruvita resultantes se caracterizaron mediante difracción de rayos X y microscopía electrónica de barrido con espectroscopía de rayos X con dispersión de energía. El tamaño de partícula promedio de los cristales de estruvita aumentó de 55 µm a 0 rpm a 127 µm a 100 rpm y 128 µm a 200 rpm. Incrementos adicionales en las velocidades de agitación resultaron en tamaños de cristales más pequeños. Estos resultados indicaron que el mayor tamaño de partícula se puede obtener a velocidades de agitación entre 100-200 rpm, equivalente a un gradiente de velocidad entre 79 y 188 s⁻¹, ya que no hubo diferencias estadísticamente significativas entre los valores medios (prueba t, p<0.05). El rango óptimo de velocidad de agitación descrito aquí se puede usar para establecer condiciones operativas para la cristalización de estruvita con el beneficio de producir cristales grandes y reducir el consumo de energía en tanques agitados.

1. Introduction

Phosphorus (P) and nitrogen (N) are the main nutrients causing eutrophication that lead to biodiversity loss and...
water quality problems \cite{1,2}. However, although an excess of phosphorus is problematic, modern agriculture is highly dependent on phosphorus fertilizer, which is derived from phosphate rock, a non-renewable resource with global reserves expected to be depleted within the following 50–100 years \cite{3}, making a strong problem for its contribution to agricultural and industrial development.

Both problems, phosphorus source depletion and eutrophication, can be tackled at the same time by the application of technologies aimed at recovering nutrients from wastewater streams \cite{4}. For example, the centrate from anaerobic digesters in Wastewater Treatment Plants (WWTP), which is a N- and P-rich stream produced after sewage digestate dewatering, with concentrations between 750–1,500 mg N L$^{-1}$ for nitrogen and 40–400 mg P L$^{-1}$ for phosphorus compounds. Those figures are typical from WWTP using enhanced biological nutrient removal (EBNMR) processes \cite{5–8}, which represents a good opportunity to recover nutrients from domestic wastewater. A technique proven to be a successful method to recover phosphorus from phosphorus-rich streams is the precipitation of struvite \cite{9} \([\text{MgNH}_4\text{PO}_4\cdot6\text{H}_2\text{O}]\). Struvite is an effective, slow-release fertilizer that can be used to close the nutrient loop in between waste production and agriculture, as it simultaneously recovers N and P from waste effluents \cite{10,11}.

Hence, struvite offers multiple benefits including limited negative impacts from agricultural runoffs into freshwater bodies, when compared with traditional chemical fertilizers. In fact, N losses have been reported to be considerably lower in struvite-treated soils than in soils receiving chemical fertilizers \cite{10}. Struvite can also reduce nitrous oxide emissions and the need for phosphate rock, thus creating a more sustainable environment in line with a wider circular economy approach.

To date, various kinds of reactors have been developed at laboratory, pilot and full scales showing a great potential in recovering struvite from waste streams, among them are: fluidized bed reactor (FBR), air lift reactor and continuously stirred tank reactor (CSTR) \cite{12,13}. FBR is the most commonly used due to its ability to produce struvite pellets of large size, high crushing strength and purity \cite{14}; however, its operation requires high flow rate and energy for mixing to ensure that the bed is continuously fluidized, so different efforts have been made to reduce energy demand and minimize losses of fine particles remaining in solution \cite{15}. Guadie \textit{et al.} \cite{1} used a novel cone-inserted fluidized bed reactor for reduce unwanted crystal loss, while Le Corre \cite{16} used stainless steel meshes to reduce the loses of fine particles in the effluent; Wang \cite{17} conducted an economic analysis in FBR reactor using different Mg sources. On the other hand, in air-lift reactors air increases the pH by CO$_2$ stripping and creates an internal recycle flow that allows crystals to grow until a critical particle size \cite{18}, however the costs of energy used in aeration can be considerably high. The stirred reactor or CSTR is one of the simplest and most effective technologies to produce struvite \cite{19,20}, although the mixing force and the reactor hydrodynamics are significant factors that contribute to the formation of struvite crystals \cite{21,22}.

In a stirred reactor, hydraulic shear forces can break struvite crystals into smaller fractions, which ultimately affect the resulting size of the precipitates \cite{21}. These fine precipitates are more likely to be washed out with the effluent, decreasing the recovery of precipitates. Therefore, in order to optimize P-recovery efficiency after crystallization, it is beneficial to aim for larger particle sizes \cite{20}. Additionally, larger struvite particles will have a longer effect on soils, increasing the absorption of nutrients from plants and crops \cite{18}.

The effect of stirring speed on struvite formation and on particle size has been little investigated. Liu \cite{23} showed that mixing rates and contact time increase phosphorus removal efficiency, while Cerrillo \cite{24} found that the maximum particle size decreased when the mixing speed was increased from 100 to 200 rpm. These results agree with the work reported by Kim \cite{25} indicating that in struvite precipitation processes, removal patterns of nitrogen and phosphorus from the solution increased logarithmically as a response to the increment of $Gt_d$ values – i.e., the product between mixing intensity/velocity gradient (G) and mixing duration (t$_d$); however, the individual effect of G on particle size at speeds between 0 and 500 rpm is yet unknown. Therefore, it is important to evaluate the effect of stirring speed and mixing intensity in a wide range of velocity gradients (G) in order to optimize the amount of energy needed in these reactors to obtain high percentages of P and N recovery and larger particles sizes of struvite precipitates.

Thus, the main objective of this study was to evaluate the effect of stirring speed on struvite precipitation using the centrate from a conventional municipal WWTP. The removal of nutrients and the particle size were assessed to obtain the optimum stirring range for a struvite mixing reactor.

\section*{2. Materials and methods}

\subsection*{2.1 WWTP}

Centrate samples were collected in the municipal WWTP of La Llagosta, located in the Catalonia autonomous community, province of Barcelona, Spain, at 41° 30'12.62
"N and 2° 12’5.12” E (Figure 1), which has a treatment capacity of 43,000 m³ d⁻¹ and a population equivalent (p.e) of 358,000 inhabitants. This WWTP has a conventional system for the treatment of a blend of domestic and industrial wastewaters, which comprises pretreatment, primary settlement, and activated sludge units followed by a final secondary settling with chemical precipitation (Figure 2). The process can remove 56% TKN and 86% TP for the raw municipal wastewater.

2.2 Analytical methods

Centrate samples were analyzed at IRTA facilities (Caldes de Montbui, Spain). Cations (Ca²⁺, Mg²⁺, K⁺) and anions (P-PO₄³⁻) concentrations were analyzed by ionic chromatography using a Metrohm 790 IC equipped with a Metrosep C4 150/4.0 column and a Metrohm 861 Advanced Compact IC equipped with a Metrosep A Supp 5-250 column and an 853 CO₂ suppressor, respectively. Total solids (TS), total suspended solids (TSS), volatile solids (VS), NH₄⁺-N, TKN, TP and total alkalinity concentrations were determined according to Standard Methods [26]. The appearance of the precipitates was analyzed by using a scanning electron microscope [SEM, ZEISS EVO®MA 10] coupled to an energy dispersive X-ray microanalysis detector (EDX-Oxford, Inca) at Autonomous University of Barcelona (Cerdanyola, Spain). Meanwhile, the crystalline nature of precipitates was determined by X-ray diffraction (XRD, X’Pert Powder of Panalytical), using CuK-alpha radiation. The XRD patterns were recorded in the scanning range of 2-theta from 10° to 60°. Data were collected and processed using High-score-plus software. Identification
of the phase peaks was accomplished by comparing the observed XRD patterns with a standard of struvite compiled by the the International Center for Diffraction Data (ICDD), the crystallography open database and the Powder Diffraction File (PDF) [27]. Crystal sizes were measured using ImageJ [1.51J8] software [28] by making 120 measurements of the crystals length for each sample.

2.3 Materials

Due to the characteristics of the centrate, MgCl₂·6H₂O and NaH₂PO₄ were added to balance the P: N: Mg molar ratio to 1:6:1.7 (P = 300 mgP·PO₄³⁻/L). This molar relationship was chosen because higher N:P molar ratio improves the precipitation of struvite instead of amorphous calcium and magnesium phosphates [29, 30], while Mg: Ca higher than 2:1 have the same effect [31]. Although struvite can precipitate at a wide range of pH values [7.0-11.5] [32], many investigations agree that the optimum pH value is around 9 [24, 33], for that reason NaOH in solution was added to reach pH 9. The principal conditions of the experiments are listed in Table 1.

<table>
<thead>
<tr>
<th>Parameter</th>
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<td>mg N-NH₄⁺·L⁻¹</td>
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</tr>
<tr>
<td>mg Ca²⁺·L⁻¹</td>
<td>261</td>
</tr>
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</table>

Molar ratios

| N-NH₄⁺ / P·PO₄³⁻ | 6.1     |
| Mg²⁺ / P·PO₄³⁻   | 1.7     |
| Mg²⁺ /Ca²⁺       | 2.5     |
| Stirred speed (rpm) | 0, 100, 200, 300, 400, 500 Replicates | 2 |

was calculated by multiplying the mmols of phosphate present in the centrate by the molecular weight of struvite.

2.4 Experimental setup

A standard jar tester with six paddles was used for struvite precipitation. Jars consist of 1L cylindrical glass vessels with 130 mm internal diameter. The flat paddles at the end of each stirrer shaft, which were made of stainless steel with a length of 7.6 cm and a height of 2.5 cm. The stirring device included a tachometer and a rev controller to adjust mixing rates ranging from 0 to 300 rpm. For the testing of mixing rates higher than 300 rpm, the same paddles were coupled to a laboratory stirrer. Mixing intensity (G) values tested in this study varied from 0 to 591 s⁻¹. The applied G value was calculated based on the interaction between mixing rates of flat paddles and the corresponding velocity gradient based on a work previously reported [34] which found that the velocity gradient has a linear correlation to the impeller speed.

The applied mixing duration (o) was 3 hours in the six-stirring speeds tested (0, 100, 200, 300, 400 and 500 rpm) followed by 16 hours without stirring (Figure 3). After 16 hours without stirring, the solutions were filtered using 1.2 µm membrane filters. In order to avoid struvite thermal decomposition that occurs above 55°C [35], the precipitates were air-dried at ambient temperature for 24 hours. Filters were weighed with and without the precipitate to calculate net struvite production; the TSS concentration from the centrate was also considered in order to calculate the total mass of struvite.

Considering that phosphate is the limiting reagent in this reaction, the maximum amount expected of struvite precipitates were air-dried at ambient temperature for thermal decomposition that occurs above 55°C. The applied mixing duration (t) was 3 hours in the six-stirring speeds tested (0, 100, 200, 300, 400 and 500 rpm) followed by 16 hours without stirring (Figure 3). After 16 hours without stirring, the solutions were filtered using 1.2 µm membrane filters. In order to avoid struvite thermal decomposition that occurs above 55°C, the precipitates were air-dried at ambient temperature for 24 hours. Filters were weighed with and without the precipitate to calculate net struvite production; the TSS concentration from the centrate was also considered in order to calculate the total mass of struvite.

3. Results and discussion

The characterization of centrate samples reported the following mean values: pH = 8.3; reactive phosphorus = 1 mgP·PO₄³⁻·L⁻¹; ammonium = 825 mgN-NH₄⁺·L⁻¹; calcium = 261 mgCa²⁺·L⁻¹; total suspended solids = 116 mg TSS·L⁻¹; and total alkalinity = 3.1 gCaCO₃·L⁻¹.

Struvite was obtained even under no stirring conditions, but the total amount recovered increased with the increment in stirring rates. Higher removal of NH₄⁺ was found as a result of increasing stirring speeds (Figure 4). In contrast, the Mg²⁺ removal increased between 0 and 100 rpm but later decreased as the stirring speeds increased, being slightly higher than the maximum that could be achieved by struvite precipitation (Figure 4). In this case, high Mg²⁺ removals at different speeds may be due to the precipitation of other Mg precipitates, such as Mg(OH)₂ since the excess mass of external magnesium source can rapidly react with OH⁻ ions in the solutions to produce these amorphous products [25].

The Ca²⁺, Mg²⁺ and N removal from the soluble phase was between 2.73 and 4.55 mmol Ca²⁺, between 11.0 and 13.8 mmol Mg²⁺ and between 9.9 and 16.9 mmol N-NH₄⁺. The Ca²⁺ removal may be due to the formation of calcium precipitation. Jars consist of 1L cylindrical glass vessels for that reason NaOH in solution was added to reach pH 9. The principal conditions of the experiments are listed in Table 1.
Figure 3 Experimental procedure for struvite precipitation

Table 2 Removal efficiencies (%), struvite similarity score XRD (%) and SEM-EDX results

| Stirring Speed (rpm) | Velocity gradient (s⁻¹) | Removal Efficiency (%)* | mass of crystals (g) | XRD Score (%) | Mg | Ca | N | P | SEM-EDX (%) |
|----------------------|--------------------------|-------------------------|----------------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
| 0                    | 0                        | 81.8                    | 69.7                 | 16.8           | >95            | 2.1            | 97%            | 14.9           | 18.5           | 0.6            | 14.9           | 18.5           | 0.6            |
| 100                  | 79                       | 85.1                    | 56.7                 | 19.0           | >95            | 2.4            | 88%            | 14.6           | 17.6           | 0.6            | 14.6           | 17.6           | 0.6            |
| 200                  | 188                      | 81.8                    | 69.7                 | 23.9           | >95            | 2.5            | 90%            | 13.8           | 16.4           | 1.6            | 13.8           | 16.4           | 1.6            |
| 300                  | 312                      | 73.5                    | 45.2                 | 24.5           | >95            | 2.5            | 97%            | 14.9           | 17.2           | 1.4            | 14.9           | 17.2           | 1.4            |
| 400                  | 447                      | 67.6                    | 63.6                 | 28.5           | >95            | 2.6            | 87%            | 14.3           | 15.4           | 1.8            | 14.3           | 15.4           | 1.8            |
| 500                  | 591                      | 68.1                    | 57.9                 | 28.7           | >95            | 2.6            | 82%            | 14.5           | 31.0           | 1.4            | 14.5           | 31.0           | 1.4            |

*regarding the initial content in the soluble phase; ** percentage of total atoms

Figure 4 Reduced Mg, Ca and N (mmol) and recovered crystals depending on the stirring speed-velocity gradient
carbonates like calcite [CaCO$_3$] and monohydrocalcite [CaCO$_3$·H$_2$O], more favoured in comparison with the formation of calcium phosphate [29, 36].

As reported in Figure 3, there was no direct relationship between the removal of Mg$^{2+}$ and Ca$^{2+}$. This is because the initial high magnesium content enhances the phosphate reaction with Mg$^{2+}$ rather than Ca$^{2+}$ [37], thereby reducing the interference due to the formation of Mg$^{2+}$ and Ca$^{2+}$ precipitates such as dolomite [CaMg(CO$_3$)$_2$] and favours the formation of struvite.

High stoichiometric NH$_4^+$: PO$_4^{3-}$ ratio (≈ 4.7) favor the struvite precipitation [36]; however, the mass of crystals above the maximum expected for struvite at stirred speeds between 200 rpm and 500 rpm (Figure 3) may be due to the formation of other precipitates not detected in the XRD, since the agitation possibly favours other reactions.

Despite finding greater removal of N and Mg$^{2+}$ than the maximum expected by struvite precipitation, it was found by X-ray diffraction that the main product obtained at all stirred speeds was struvite, finding similarities
between 82 and 97% with the standard of struvite (Score in Table 2 and Figure 5). The SEM-EDX analyses confirmed the presence of Mg and P in the precipitates, with a minimum presence of Ca (Table 2 and Figure 6).

Although Pastor [33] concluded that the degree of agitation of the system can exert a considerable influence on the crystalline habit (external appearance of the crystals), similar crystal shapes were observed in this work — i.e., stars, X shapes and needles (Figure 6). The last shape, needles, are the most reported in by researchers, including Crutchik et al. [22].

The attained data in this work confirmed the strong influence of stirring speeds on particle size distribution. The largest mean particle size was found at 200 rpm (128 µm; Figure 7) and the smallest size at 0 rpm (55 µm). According to t-student tests, there were no statistically significant differences between the particle size of 100 and 200 rpm (p >0.05); however, there were statistically significant differences between the particle size of 0 and 100 rpm, and 100 and 500 rpm (p<0.05). Therefore, the largest particle sizes were between 100 and 200 rpm, which corresponds to a G value between 79 and 188 s⁻¹. According to Capdevielle et al.[30], high mixing rates have a negative impact on struvite crystal sizes due to struvite dissolution and crystal break, thus, although the mixing effect enhances mass transfer rates [25] and therefore the particle size, stirring speeds higher than 200 rpm were detrimental to the struvite crystallisation process.

4. Conclusions

The struvite formation is affected by the stirring speed. The largest mean particle size of 128 µm was obtained at 200 rpm — i.e., equivalent to a G of 188 s⁻¹. The particle size of struvite crystals decreased with higher stirring speeds. Although the velocity gradient improves mass transfer rates of ions in the solution to support better crystal growth, high G values beyond the optimal can revert this effect, reducing the size of crystals formed or disfavoring their growth rates.

The P removal was not affected by stirring speeds, while the N removal increased with stirring speed achieving at 500 rpm a removal of 7 mmol more than the maximum possible by the sole formation of struvite. In this case, the agitation may favor the volatilization of N in the form of NH₃ [38]. Although this did not affect the formation of struvite by the high contents of N-NH₄⁺ in the centrate, it implied an increased emission and a potentially higher environmental risk.

Although the removal of Mg appears to be influenced by the stirring speed, the precipitation of other salts was minimal compared to the struvite precipitation, according to XRD. Therefore, a stirring speed range between 100 and 200 rpm (79 s⁻¹ < G < 188 s⁻¹) is recommended for struvite crystallisation with the benefit of producing large crystals, reducing energy consumption in stirring tanks, and reducing the volatilization of ammonia nitrogen.
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References


