Research article

Struvite precipitation within wastewater treatment: A problem or a circular economy opportunity?

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HIGHLIGHTS

- Biological P removal at wastewater treatment plants causes precipitation problems.
- We evaluated if problem can be overcome by adopting circular economy thinking.
- Mass balance estimates & struvite precipitation profiles used to identify hotspots.
- Struvite recovery via a crystallization reactor demonstrated as economically viable.
- Struvite recovery from wastewater treatment is viable circular economy opportunity.

GRAPHICAL ABSTRACT

Circular Economy from struvite recovery
- Recovery of nutrients.
- Displacement of fossil fuel derived fertiliser (Ammonia).
- Displacement of mined phosphate.
- Potentially reduced leaching/ eutrophication rate.
- Developing market emerging.
- Revenue stream from the fertiliser.

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ABSTRACT

Enhanced biological phosphorus removal at wastewater treatment plants that use anaerobic digesters for sludge treatment have historically encountered phosphate precipitation problems in the form of struvite. Literature on struvite is thin which is surprising given it can foul/block the sludge return lines and associated pumps and valves, causing significant operational problems. This study has evaluated if a typical large wastewater treatment plant can overcome this problem by adopting circular economy thinking. The struvite profile based on the supersaturation ratio of \((\text{Mg}:\text{NH}_4:\text{PO}_4^2)/\text{CO}_3\) \(^{2-}\), pH and temperature demonstrates the potential operational hotspots that can present uncontrolled struvite formation. Based on current struvite monitoring technologies and a cost-benefit analysis, the controlled struvite recovery via an Ostara crystallization reactor has been demonstrated to be economically viable with a pay-back period of less than a decade. An integrated evaluation illustrates the positive environmental impact arising from the utilisation of the recovered product. Economic viability and payback periods will vary according to circumstances, but we recommend that WWTP operators globally consider fitting a crystallisation reactor to appropriate plants. The outcomes and recommendation from this study are particularly timely given the global fertiliser shortage (2022) that is driving up food prices and reducing crop sizes.

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1. Introduction

Wastewater treatment plants (WWTPs) incorporating secondary treatment and anaerobic sludge digestion facilities have historically encountered phosphate precipitates in the form of struvite (Barr and Münch, 2001; Jaffer et al., 2002). These agglomerates can foul and block the sludge return lines and associated pumps and valves (Ohlinger et al., 1999; Jaffer et al., 2002; Mudragada et al., 2014). Uncontrolled struvite precipitation increases pumping and maintenance costs, as well as reducing the overall capacity of the plant piping system via hydraulic loss capacity. This also lowers the biological treatment efficiency (Battistoni et al., 1997; Barr and Münch, 2001; Doyle and Parsons, 2002; Jaffer et al., 2002; Mudragada et al., 2014).

However, it is possible that controlled production and recovery of struvite could promote circular economy (CE) opportunities for the wastewater treatment industry (Barr and Münch, 2001; Jaffer et al., 2002; Molinos-Senante et al., 2011; Siciliano et al., 2020), especially in coastal areas (Roberts et al., 2021). The precipitated struvite can be recovered and sold as fertiliser, reducing resource depletion by replacing fossil-fuel based P (Jaffer et al., 2002; Zhang and Mo, 2013). Mavhungu et al. (2021) report that struvite precipitation could act as a fast, efficient, and environmentally friendly pre-treatment step to remove P and reduce N from wastewater. However, a potential concern is that the environmental impacts of struvite recovery via the use of additional chemicals and energy are not offset by its benefits (Sena et al., 2021). Nevertheless, removal of struvite can significantly decrease the operational and maintenance costs of WWTPs, promoting the optimisation of operational and environmental performance (Jaffer et al., 2002; Pastor et al., 2008).

Within WWTPs, enhanced biological phosphorus removal (EBPR) processes produce a P-rich activated sludge, causing struvite deposit problems in anaerobic sludge digestion (Barr and Münch, 2001). Generally, struvite (MgNH₄PO₄) is a white crystalline mineral compound that forms under conditions of elemental supersaturation within liquids (Barr and Münch, 2001). It can be found in WWTP systems when Mg²⁺, NH₄⁺ and PO₄³⁻ (magnesium, ammonium and phosphate – MAP) concentrations exceed solubility levels, supersaturation occurs and minerals combine and precipitate into solid form (Wu and Bishop, 2004).

WWTP design generally incorporates flow and partial pressure reduction through pump impellers and pipe bends, forcing the removal of dissolved CO₂ and increasing the pH of the liquor (Battistoni et al., 1997; Fattah et al., 2010; Xavier et al., 2014). High pH levels (~7.5–10) are essential for the struvite precipitation (Ohlinger et al., 1999; Barr and Münch, 2001; Wu and Bishop, 2004). Reduction of the partial pressure of reject waters from 0.5 atm to 0.05 atm within the WWTP contributes to the release of dissolved CO₂ and thus increases pH levels from 7.0 to 8.0, increasing the chance of struvite precipitation (Fattah et al., 2010).

The most common P removal technologies include biological P removal, crystallization, chemical precipitation via ferric chloride, tertiary filtration and ion exchange (Morse et al., 1998). Crystallization of struvite is probably the most sustainable solution due to the minimal waste production that needs to be managed (Marti et al., 2008).

The primary aim of this study was to determine and evaluate the economic and environmental impacts arising from the uncontrolled struvite formation within a typical large WWTP. For this aim, the objectives were to:

i. Establish a mass balance for the nutrients present in the system. This is to enable the specific locations within the operational system that present higher possibility for struvite precipitation to be identified (Jaffer et al., 2002; Marti et al., 2008).

ii. Create a profile for struvite that incorporates the potential operational hotspots for uncontrolled struvite precipitation and other factors that can affect struvite formation.

iii. Identify the operational and maintenance costs and the environmental impacts arising from struvite precipitation at the WWTP.

The secondary aim of this study was select the most appropriate monitoring pathway for the uncontrolled struvite precipitation based on the characteristics of the studied WWTP. For this aim, the objectives were to:

i. Identify and evaluate current technologies regarding nutrient removal and/or controlled struvite formation.

ii. Evaluate the economic costs and benefits of the controlled struvite formation.

iii. Develop an integrated cost benefit analysis for the proposed struvite monitoring technology.

We critically discuss findings in the context of enabling wastewater treatment processes to recover more value from wastewater (resources), reducing treatment and maintenance costs and enabling circular economy (CE) thinking to be operationalized.

2. Methodology

2.1. WWTP characteristics

Budds Farm wastewater treatment plant (BF-WWTP), owned by Southern Water (SW) in England, treats the domestic wastewater of Portsmouth, Havant, Hayling Island, Cosham, Paulsgrove, Waterlooville, Horndean and Hambledon. It has a general capacity of 109,000 m³ of wastewater per day. Before its establishment, untreated wastewater arising from Hampshire was discharging to Langstone Harbour causing significant environmental impact at local and national level (Smith et al., 1999; Doyle and Parsons, 2002). Indeed, SW has admitted deliberately and illegally dumping untreated wastewater into this harbour, resulting in a huge fine (BBC News, 2021a) and damaging publicity (BBC News, 2021b).

BF-WWTP is responsible for a population equivalent (PE) of approximately 410,000. It presents an outfall of an average 2000 L/s and it operates for 24/7 for 365 days per year. It has one of the biggest BNR facilities in UK with 3 main operational units. BF-WWTP handles indigenous and imported sludge with its anaerobic digesters to treat 800 m³ of activated sludge on 14 day cycle. The site is under tight nutrient consents, restricting discharges into the harbour to 10 mg/L total nitrogen or less. Figure 1 illustrates the main operational units associated with the wastewater treatment process within the BF-WWTP.

2.2. Sampling and analysis

Operational units and locations (hot spots) within WWTPs that have historically presented P precipitation in the form of struvite are summarised in Table 1 (Le Corre et al., 2009). A sampling strategy was created to determine concentrations for the three struvite ions (Mg²⁺NH₄⁺PO₄⁻²) or MAP. External factors such temperature (T) and pH were determined to evaluate struvite behaviour within the system (Bouroupolous and Koutsoukos, 2000; Doyle and Parsons, 2002; Jaffer et al., 2002; Fattah, 2012).

Wastewater samples were professionally collected by an operational engineer employed at BF-WWTP. The samples were processed and analysed on-site at BF-WWTP’s engineering laboratory. All samples were filtered via Advantec Grade No.1 filter paper or cellulose acetate syringe filter. Because of the high density of the sludge samples, filtration was necessary in order to remove coarse particles and prepare samples for qualitative analysis. Temperature (T) and pH were measured potentiometrically in the undiluted liquid wastewater sample after filtration and in less than one hour from sample collection. The DO700 analyser provided automatic measurements for dissolved oxygen, T, pH, conductivity and total dissolved solids (TDS) (Pass, Extech DO700 Portable Dissolved Oxygen Meter).

Ammonia concentrations were determined via a QUANTOFOX test kit that provides a detection range between 10 and 400 mg/L NH₄
Phosphate concentrations were determined via a QUANTOFIX test kit that only detected ortho-phosphate. This method provides a detection range between 3 and 100 mg/L PO₄³⁻. Magnesium concentration was determined via a viscolar ECO total hardness test kit that provides a detection range between 10-100 mg/L CaO. Dilutions of 1:4 or 1:10 (based upon previous experience) of wastewater samples to double-distilled deionised water were required based on the colour indication of the test kits.

2.3. Creation of the struvite precipitation profile within BF-WWTP

Struvite precipitation generally correlates with the degree of supersaturation (Ohlinger et al., 1999; Bouropoulos and Koutsoukos, 2000; Kofina and Koutsoukos, 2005), magnesium to phosphorus molar ratio (Mg:P) (Barr and Münch, 2001), pH (Barr and Münch, 2001; Doyle and Parsons, 2002), temperature (Doyle and Parsons, 2002), crystal retention time, recycle ratio (Ohlinger et al., 1999; Bouropoulos and Koutsoukos, 2000), turbulence and mixing (Ohlinger et al., 1999; Bhuiyan, 2007). Eq. (1) summarises the kinetics of struvite formation:

\[
\text{Mg}^{2+} + \text{NH}_4^+ + \text{PO}_4^{3-} + 6\text{H}_2\text{O} \rightarrow \text{MgNH}_4\text{PO}_4 \cdot 6\text{H}_2\text{O} \tag{1}
\]

The struvite profile (behaviour within the WWTP system) arose from the concentrations of the main chemical components of MAP and the struvite supersaturation rate (SSR) for each sampling point in correlation with pH and T. This correlation indicated which locations present higher potential for uncontrolled struvite precipitation (Doyle and Parsons, 2002; Le Corre et al., 2009; Fattah et al., 2010; Fattah, 2012). In general, uncontrolled formation can occur when the concentrations of Mg²⁺, NH₄⁺ and PO₄³⁻ exceed the Ksp of struvite and precipitates in a 1:1:1 molar ratio following Eq. (1) (Le Corre et al., 2009). If the general equation of soluble salt in water is assumed then Eq. (2) follows:

\[
\text{AaBb(s)} \rightarrow a\text{Az}^{+} + b\text{Bz}^{2-} \tag{2}
\]

and the constant solubility product Ksp can be expressed as:

\[
\text{Ksp} = [\text{A}^{z+}]^a[\text{B}^{z-}]^b \tag{3}
\]

where:

- \([A^{z+}]\) and \([B^{z-}]\) are the total concentrations of ions in solution
- \(z^+\) and \(z^-\) are the valencies of the considered ions.

By applying Eq. (3) on Equation 1, Equation 4 is produced (Michalowski and Pietrzyk, 2006):
### 2.4. Cost benefit analysis of P recovery via struvite precipitation

An economic feasibility study of P recovery was established using a cost-benefit analysis (CBA). The net profit (NP) is the difference between benefits and costs (i.e., NP = IB + EB where net profit [NP = total income - total costs]; internal benefit [IB = internal income - internal costs]; and external benefit [EB = positive impact – negative impact]). Through a CBA, a project is only economically feasible only if NP > 0 (Molinos-Senante et al., 2011).

The term internal income includes the sale of recovered P and treated water, as well as savings from reduced WWTP operating costs arising from chemical utilisation for the chemical precipitation of phosphorus; reduced sludge generation and associated management cost and reduced pipe and tube cleaning due to less uncontrolled struvite scaling (Molinos-Senante et al., 2011).

Internal costs include investment cost, civil works (equipment, machinery, and auxiliary facilities); operating and maintenance costs (reagents for chemical precipitation and pH value maintenance); and financial costs (Molinos-Senante et al., 2011).

Investment costs depend largely on the size of the WWTP (Molinos-Senante et al., 2011). Operating and maintenance cost proposed values depend on the concentration of P in the waste streams and the reagents used to operate the plant. The cost depends on the people equivalent covered from the plant and the operational streams the P is be extracted. Thus, considering these incomes and costs, the internal benefit can be expressed as Eq. (6) (Molinos-Senante et al., 2011):

\[
IB = \sum_{t=0}^{T} \left[ (ASR \times SPP) + (ASR \times CR) + (ARP \times CR) - (IC + OMC + FC) \right]
\]

where:

\[ IB = \text{internal benefit (}$\text{)} \]
\[ ASR = \text{annual volume of struvite recovered (}$\text{)} \]
\[ SPP = \text{present selling price of struvite ($}/$\text{kg}) \]
\[ ASR = \text{annual volume of sludge generation reduction (}$\text{)} \]
\[ CSM = \text{cost of sludge management ($}/$\text{kg}) \]
\[ ARR = \text{annual volume reduction of reagents (}$\text{)} \]
\[ CR = \text{cost of reagents ($}/$\text{kg}) \]
\[ APR = \text{annual volume reduction of uncontrolled struvite precipitation (}$\text{)} \]
\[ CDD = \text{cost of cleaning struvite deposit ($}/$\text{kg}) \]
\[ IC = \text{investment cost (}$\text{)} \]
\[ OMC = \text{operational and maintenance cost (}$\text{)} \]
\[ FC = \text{financial costs (}$\text{)} \]

External benefits (EB) refer to any positive or negative impact that derives from a proposal and effects on people without economic value. P recovery from waste streams is regarded as having positive EB, such as an increase in the availability of a non-renewable resource and important environmental benefits, because if phosphorus discharge is prevented, then its level in water bodies is reduced and, consequently, there are fewer eutrophication problems. The external benefits can be expressed as Eq. (7) (Molinos-Senante et al., 2011):

\[
EB = \sum_{t=0}^{T} (PI - NI)
\]

where:

\[ PI = \text{positive impact (}$\text{)} \]
\[ NI = \text{negative impact (}$\text{)} \]

In general, IB can be estimated directly as monetary units while the EB is difficult to estimate as marketable price. In order to give a comparable value for the EB, the term shadow price may be used (Molinos-Senante et al., 2011). Given the volatility of markets during the COVID19 pandemic, pre-COVID monetary values have been used to establish order-of-magnitude indicative costs/prices.

### 2.5. Integrated evaluation of chosen struvite monitoring technology

Except for the financial elements, the integrated evaluation (IE) took into consideration the environmental (shadow indicator) and social impacts that a proposal can cause at a national and international level. The CBA determined if a technology can be viable from an economic perspective. The IE demonstrated the sustainable trends of the modern industrial models and aimed to present a holistic approach regarding the P monitoring pathway (Molinos-Senante et al., 2011; Bird, 2015).

### 3. Results

#### 3.1. Operational diagram and struvite precipitation

Based on the BF-WWTP operational diagram as well as the locations shown in Table 1, the sampling map was established in consultation with the BF-WWTP management team. Figure 2 illustrates the sludge
treatment process within the plant focusing on the areas that present higher possibility for uncontrolled struvite precipitation.

Table 2 presents data arising from the determination of the main factors that affect the struvite formation at the different operational hot-spots.

Several struvite Ksp values have been proposed in the literature. The most widely accepted value was found to be $10^{-12.6}$ (Snoeyink and Jenkins, 1980; Michałowski and Pietrzyk, 2006). At full-scale wastewater treatment processes, the Ksp value was determined as $10^{-13.26}$ (Ohlinger et al., 1999), thus struvite estimated to be less soluble than presented to be based on other researches. Table 3 presents the concentrations of the tested locations of BF-WWTP in M (mol/L) and the SSR.

The SSR for the selected locations indicates that the influent that reaches BF-WWTP presents extremely high concentrations of the three ions of MAP. Thus there is high possibility of uncontrolled struvite formation in the majority of the operational system; this is regularly observed on-site. However, a positive SSR does not ensure that struvite is necessarily formed (Fattah, 2012). This is mainly because the kinetics of P precipitation (how fast P is precipitated) and the competing reactions must be taken into consideration. Furthermore, ions such as $\text{Ca}^{2+}$, $\text{K}^+$, $\text{CO}_3^{2-}$, etc., can influence the saturation of struvite by reacting with $\text{Mg}^{2+}$, $\text{PO}_4^{3-}$, and $\text{NH}_4^+$. It is therefore necessary to evaluate the availability of free ions (i.e. the ionic activity) for a given pH in order to accurately estimate the precipitation rates (Snoeyink and Jenkins, 1980; Ohlinger et al., 1999; Barr and Münch, 2001). Figure 3 presents the variation of pH in the different operational hot spots at BF-WWTP.

Struvite can potentially be formed at locations that present positive SSR and a pH range between 7 and 10.7 (Tables 2 and 3, Figure 3) (Snoeyink and Jenkins, 1980; Ohlinger et al., 1999; Barr and Münch, 2001). In general, the digester recirculation pumps, PDST, centrifuge feed, centrate and SAS storage tank demonstrate suitable conditions for P precipitation in the form of struvite. The correlation between Ksp and pH indicates that struvite solubility decreases with increasing pH which in turn leads to an increase in the precipitation potential of a solution (Booram et al., 1975; Ohlinger et al., 1999; Musvoto et al., 1999b; Doyle and Parsons, 2002).

Generally, in digestion of BNR sludge, the stabilising counter ions of polyphosphate produce Mg in a molar ratio of ~0.3 mol Mg/mol P. Thus, it can be assumed that all the removed Mg potentially can be precipitated as struvite (Doyle and Parsons, 2002). Moreover, in the presence of high influent P concentrations (>200 mg/L) struvite crystals may demonstrate a growth rate up to 0.177 mm/d (Abe, 1995). If it is assumed that BF-WWTP uses 3.94 in (100 mm) sludge lines, then based on these struvite growth rates, an operational hot spot in the

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1. Screened sludge storage tank.
2. Thickened sludge storage tank.
3. SAS storage tank – Surplus Activated Sludge (SAS).
4. Blended cake.
5. Digester feed.
6. Digester recirculation pump.
7. PDST (digested sludge) – Post digested sludge treatment.
8. Centrifuge feed.
9. SAS filtrate.
10. Primary thickening filtrate.
11. Centrate.
12. SAS after thickened.
Table 2. Data arising from the samples analysis of the most potential locations regarding the struvite formation within the operational system of Budds Farm WWTP. The locations that present the symbol * or ** indicate locations with the same sludge characteristics due to operational design. In addition, the brackets indicate the dilution of wastewater samples: double-distilled deionised water in order to reach the detection limits.

| Hot-spots of struvite precipitation | pH     | T (°C) | PO₄³⁻ (mg/L) | NH₄⁺ (mg/L) | Mg²⁺ (mg/L) |
|-------------------------------------|--------|--------|--------------|-------------|-------------|
| Screened sludge storage tank        | 6.5    | 20.8   | 100 (1:4)    | 400         | 390         |
| Thickened sludge storage tank*      | 5.4    | 20.8   | 100 (1:4)    | 100         | 390         |
| SAS storage tank                    | 7.2    | 21.1   | 100          | 10          | 350         |
| Blended cake                        | 6.4    | 22.0   | 100          | 100         | 190         |
| Digester feed*                      | 5.9    | 20.8   | 100          | 100         | 390         |
| Digester recirculation pump         | 8.4    | 21.2   | 40           | 100         | 150         |
| PDST (digested sludge) **           | 8.4    | 20.8   | 200 (1:4)    | 200 (1:4)   | 190         |
| Centrifuge feed**                   | 8.4    | 20.8   | 200          | 200         | 190         |
| Thickened sludge storage tank       | 7.2    | 21.1   | 100 (1:4)    | 100         | 390         |
| Screened sludge storage tank        | 7.1    | 20.3   | 100 (1:4)    | 10          | 380 (1:4)   |
| Primary thickening filtrate         | 5.6    | 20.7   | 200 (1:4)    | 50          | 340         |
| Centrate                            | 8.1    | 21.5   | 40           | 200 (1:4)   | 230         |
| SAS after thickened                 | 7.0    | 20.5   | 100          | 10          | 260         |

Table 3. Concentrations of struvite ions in mol/L and the respective SSR.

| Hot-spots of struvite precipitation | PO₄³⁻ (mol/L) | NH₄⁺ (mol/L) | Mg²⁺ (mol/L) | Ksp          | SSR |
|-------------------------------------|---------------|--------------|--------------|--------------|-----|
| Screened sludge storage tank        | 0.0010        | 0.0200       | 0.0163       | 3.26 x 10⁻⁷  | >1  |
| Thickened sludge storage tank*      | 0.0010        | 0.0050       | 0.0163       | 8.15 x 10⁻⁸  | >1  |
| SAS storage tank                    | 0.0010        | 0.0005       | 0.0146       | 7.3 x 10⁻⁹   | >1  |
| Blended cake                        | 0.0010        | 0.0050       | 0.0079       | 3.95 x 10⁻⁸  | >1  |
| Digester feed*                      | 0.0010        | 0.0050       | 0.0163       | 8.15 x 10⁻⁸  | >1  |
| Digester recirculation pump         | 0.0004        | 0.0050       | 0.0063       | 1.26 x 10⁻⁸  | >1  |
| PDST (digested sludge)**            | 0.0021        | 0.0100       | 0.0079       | 1.65 x 10⁻⁷  | >1  |
| Centrifuge feed**                   | 0.0021        | 0.0100       | 0.0079       | 1.65 x 10⁻⁷  | >1  |
| PDST filtrate                       | 0.0010        | 0.0005       | 0.0158       | 7.9 x 10⁻⁹   | >1  |
| Primary thickening filtrate         | 0.0021        | 0.0027       | 0.0142       | 8.05 x 10⁻⁸  | >1  |
| Centrate                            | 0.0004        | 0.0100       | 0.0096       | 3.84 x 10⁻⁸  | >1  |
| SAS after thickened                 | 0.0010        | 0.0005       | 0.0110       | 5.5 x 10⁻⁹   | >1  |

3.2. Current and upcoming phosphorus removal techniques

There are a wide range of technologies to remove and recover P from wastewater, including chemical precipitation, biological phosphorus removal, crystallisation, novel chemical precipitation approaches. Phosphorus in wastewater represents a significant renewable resource and there is no environmental or technical reason why phosphorus cannot be recycled (Morse et al., 1998; De-Bashan and Bashan, 2004; Xie et al., 2016). Morse et al. (1998) summarises applied technologies focusing on the main inputs, auxiliary inputs, main output as well as the form that the P is recovered, industrial and agriculture value, associated advantages and limitations.

It was reported that chemical precipitation can achieve an average of 0.6 mg/L of total P in the effluent with an average Alum dose of 45 g/L (Patoczka, 2005). In real operational conditions, crystallisation via chemical precipitation can remove 45 mg/L of P to 6 mg/L of P within a pH of 8.7. BNR can reduce mean influent total P concentration from 6 mg/L to 1.5 mg/L. Egle et al. (2016) categorised the most industrial applied and robust technologies in three categories: aqueous phase (digester supernatant, dissolved P in anaerobically digested sludge and effluent), sewage sludge and sewage sludge ash.

4. Discussion

4.1. Decision-making for struvite monitoring

Based on the finding regarding the availability of the main ions that constitute struvite as well as the external factors that can affect the uncontrolled formation, large plants such as BF-WWTP demonstrate a high potential for the application of an economically-viable struvite controlled-recovery technology. Within a typical operational system, locations such as digester feed, digester recirculation pump, PDST and cent rate are likely to be highly suitable for struvite formation and hence extraction.

In general, monitoring technologies target: the disruption of solubility or alteration of the growth mechanisms of crystals or promotion of controlled struvite formation for recovery (Barr and Münch, 2001; Le Corre et al., 2009; Fattah, 2012; Egle et al., 2016). Since struvite precipitation is mainly based upon the equilibria of the main chemical components, most previous case studies tried to overcome this problem by either forming other phosphate based salts or by reducing the pH under the optimum formation range (Egle et al., 2016). This is mainly because once one of the P, Mg or NH₃ ions are removed, this automatically leads inhibition of struvite formation (Le Corre et al., 2009).
Acid washing with mainly CH₃COOH or H₂SO₄ has been used by many WWTPs in order to overcome blockage problems within their operating systems (Williams, 1999; Doyle and Parsons, 2002). The use of FeCl₃ is the most dominant option for struvite inhibition. However, it can produce large quantities of sludge mainly because of the poor molar removal of P per mole of Fe added. A ratio between 0.38 and 0.48 molar removal of phosphate per mole Fe added has been reported (Mamais et al., 1994). Na₂HPO₄ has also been described as a struvite inhibitor. The REM-NUT®, Ostara®, PRISA, Gifhorn, Stuttgart and LOPROX technologies produce struvite as final product with P content in the range of 10–12%. This final product typically presents similar plant uptake efficiency as commercial fertilizers in acidic soils and partially in alkaline soils (Kratz et al., 2010). In general, except from MgNH₄PO₄.6H₂O (the most dominant form), P can be precipitated as magnesium hydrogen phosphate trihydrate or newberyite (MgHPO₄.3H₂O) at pH < 6 or trimagnesium phosphate (Mg₃(PO₄)₂.22H₂O/Mg₃(PO₄)₂.8H₂O) at pH between 6 - 9 (Musvoto et al., 1999b; Michałowski and Pietrzyk, 2006).

The REM-NUT technology is the most expensive (2017) with >€42 per kg P produced at full operational scale (pre-COVID value), because of the tremendous requirements for resins and chemicals (Egle et al., 2016). Ostara®, DHV®, PRISA and P-RoC®, present an annual cost of approximately €6–10 per kg P produced or €0.8–2 per PE per year (for BFWWTP = 0.8 + 2 = €2.8/PE/2 = €1.4/PE * 410,000 PE/year = €574,000/year). The cost of 1 kg of P recovered via wet-chemical processes is €9–16 (Gifhorn and Stuttgart processes). The high requirements regarding chemicals, including acids, caustics and precipitation agents set the recovery costs at relatively high levels. The cost of wet-oxidation processes, such as Aqua Reci®, PHOXNAN and MEPHREC®, is approximately

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**Figure 3.** Operational pH variations for the proposed hot spots. The red lines indicate the thresholds for struvite precipitation. Values lower than 7 and higher than 10.7 present high solubility of struvite and thus less danger for operational problems arising from scaling.

**Figure 4.** Temperature variations for the different operational hot-spots. T reduction is an indicator for uncontrolled struvite formation.
between £23–27 per kg Prec, which makes them economically unattractive (Egle et al., 2016).

DHV Crystalactor® is not economically viable, even with maximum revenues, maximum savings and an up-scaling of the plant to 500,000 PE due to the high operational resources demand (Egle et al., 2016). High annual costs have also been reported for the Aqua Reci® and the MEPHREC® processes.

A P controlled crystallization process appears to be a suitable method to recover P in the form of struvite (Battistoni et al., 1997, 2000, 2005; Barr and Münch, 2001; Doyle and Parsons, 2002; Pastor et al., 2008). Thus an Ostara fluidised bed crystallisation reactor seems to be a very promising option regarding the sustainable management of the struvite precipitation problem (Merlo, 2011; Bird, 2015; Egle et al., 2016).

4.2. P recovery and recycling through struvite crystallisation

The operational design of BF-WWTP is likely to enhance struvite formation at specific operational hotspots when the optimum conditions are met. Consequently, a controlled struvite recovery can potentially be viable. Controlled struvite precipitation process via crystallisation can recover struvite at rates of 80–90% from reject waters (Shu et al., 2006; Xavier et al., 2014). This system can reduce ammonia concentrations by 29% (Shu et al., 2006; Xavier et al., 2014).

A struvite crystallizer reactor incorporates reverse gravitational flows of reject waters from anaerobic digesters and solids dewatering facilities (Britton et al., 2005; Fattah et al., 2010; Cullen et al., 2013). Struvite crystals are separated by density and size as result of the upward flow through increasingly larger reactor chambers. Spherical pellets or beads (known as prills) remain in the upper chamber until enough minerals accumulate, increasing the size and density while the smaller particles drop down into the smaller reactor chamber in the lower section. Each reactor zone has a reduced liquid retention time. The larger diameter area at the top of the reactor has the smallest struvite particles with higher retention time to allow for crystal growth (Cullen et al., 2013). The largest prills present in the lowest and smallest reactor chamber are harvested, dried and bagged.

Precipitation of struvite requires a molar ratio of 1(Mg2+:1(NH4+)**:1(PO43-) (De-Bashan and Bashan, 2004; Le Corre et al., 2009). It has been estimated that 95% of P can be precipitated as struvite from the centrifuge supernatant, by the addition of 1.05–1.3 or 1:1 molar ratio of Mg to P, i.e. a Mg dose of about 210 mg/L (if P concentrations 200 mg/L is assumed) (Fujimoto et al., 1991).

 Supernatants and reject water pH levels are often unsuitable for struvite formation. Thus, pH levels must be regulated. Sodium hydroxide is mainly used for pH regulation within a crystallizer reactor (Jaffer et al., 2002; Britton et al., 2005; Xavier et al., 2014). In general, recovery cannot be viable in mainstream activated sludge, trickling filter and anaerobic processes mainly because of low of P and NH4 concentrations (Williams, 1999). However, based on the results arising from this study, plants such as BF-WWTP produce high concentrations of MgNH4PO4 at a variety of operational locations. Anaerobic digestion supernatants as well as centrate from sludge filter can be potential locations, because of the high decomposition of microbial mass due the microbial bio-activities and the release of optimum P and NH4 concentrations (Bhuiyan, 2007).

4.3. Cost benefit analysis

For the CBA of a WWTP the following (pre-COVID) data are considered (Molinos-Senante et al., 2010, 2011):

For a given data set of 22 WWTP with treated water volume between 1,000,000 and 8,000,000 m³/year the average operational cost is as follow: energy = 0.0392 €/m³, staff = 0.0712 €/m³, regents = 0.0301 €/m³, waste management = −0.0342 €/m³, maintenance = 0.0453 €/m³ (Molinos-Senante et al., 2010, Molinos-Senante et al., 2011). Based on these data, the overall (pre-COVID) operating cost of WWTPs are in the range 0.1158 €/m³ and 0.7491 €/m³. Based on the treating capacity the weighted average is approximately 0.2200 €/m³. Considering that Budds Farm WWTP presents treating capacity (109,000 m³/day * 365 days/year = 39,785,000 m³/year = 4.97 times higher than the proposed, then the overall operational cost can be estimated approximately as (4.97 * 0.22 €/m³ =) 1.09 €/m³. Because of the incorporation of BNR in the BFWWTP operational system, the annual cost for chemical struvite precipitation is estimated as (1.09 €/m³ * 109,000 m³/day * 365 days/year =) 43,365,650 €/year.

Environmental benefits reflect the value of the environmental damage resulting from the uncontrolled management and dumping of the undesirable outputs (known as shadow price). By taking into account the volume of pollutants removed in the treatment process (kg/year), the volume of treated wastewater (m³/year), and the prices obtained for each pollutant, the overall environmental benefit resulting from wastewater treatment is calculated between 0.0099 €/m³ and 1.0039 €/m³ (Molinos-Senante et al., 2010, 2011). Based on the treating capacity of the WWTPs that have been examined the weighted average is 0.3609 €/m³. For Budds Farm WWTP this value can be expressed as (0.3609 €/m³ * 4.97 =) 1.79 €/m³ * 109,000 m³/day * 365 days/year = 71,215,150 €/year.

The price of treated water is estimated to be 0.345 €/m³ (Molinos-Senante et al., 2010). If the WWTP proceeds to sale of the treated water, then assuming 50% and 100% of the treated water is sold, the mean net profit is increased by ~55% (=0.3138 €/m³) and 71% (=0.4863 €/m³), respectively (Molinos-Senante et al., 2010). If the 50% sale water scenario is applied to BF-WWTP, the (pre-COVID) income is estimated as (0.3138 €/m³ * 109,000 m³/day * 365 days/year =) 12,484,533 €/year.

Regarding a full-scale pilot plant with treating capacity 400 m³/d of homogeneous liquids, the NaOH addition costs range from 0.0014 to 0.51 €/m³, which can be responsible for the 97% of the chemical expenses (Barr and Münch, 2001; Jaffer et al., 2002; Battistoni et al., 2005). For BF-WWTP this estimation can be calculated as (0.51 + 0.014 = 0.5114/2 = 0.2557) * 109,000 m³/day = 27,871 €/day. An alternative to NaOH addition was examined by Battistoni et al. (2005) with use of air stripping to adjust the pH of struvite precipitation. The addition of Mg(OH)2, which is cheaper than MgCl2 and simultaneously helps to increase the pH, is a possible option (Barr and Münch, 2001). A further potential option is the incorporation of sea water as an alternative source of magnesium. This proposal can reach 95% P removal compared to 97% removal with MgCl2. BF-WWTP could utilise saline water from the nearby Langstone Harbour and because of its location the transportation cost associated will be minimal. Furthermore, Battistoni et al. (2005) indicated that the cost associated with struvite recovery can be reduced from 0.28€ to 0.19€ per m³ (0.28–0.19 = 0.09 *100 = 9%). For BF-WWTW this reduction can be estimated as (1.09 €/m³ * 0.90 = 0.9981 €/m³, 1.09 €/m³ - 0.0981 €/m³ = 0.99 €/m³ * 109,000 m³/day * 365 days/year =) 39,462,741.2 €/year by using sand as auto-nucleation media.

The feasibility of controlled recovery of struvite via crystallisation is heavily dependent on the profits generated from struvite sales as fertilizer. The revenue produced from the struvite sales is difficult to estimate because of the differences in regional demand and rates of production (Le Corre et al., 2009). It should be noted that currently (2022) there are concerns about the increasing costs of energy used for ammonification process of making ammonia is not a green process as it is most commonly made from methane, water and air, using steam methane reforming and the Haber process. In Japan, struvite was historically sold as fertiliser for agricultural purposes for approximately 250 €/t. For a process treating 400 m³/d of centrate liquors incorporating an activated sludge handling system and a BNR, the potential revenue regarding the struvite recovery and sale has historically been estimated as ~25,000 €/month (= 300,000 €/year) with a 90% P removal (Barr and Münch, 2001; Jaffer et al., 2002). It reported that 1 kg of struvite can be recovered from 100 m³. Based on the Budds Farm outfall [(2000 L/s * 1
kg)/100,000 L = 0.01 kg s⁻¹3600 s/h * 24 h/day * 365 day/year = 315 tonnes/year can be produced (Barr and Münch, 2001). Based on historical data, the average marketable price of struvite is estimated as 200–1,885 S/tonne. Assuming an average price of 1042.5 S/tonne (Barr and Münch, 2001; Doyle and Parsons, 2002) then an annual income of (1042.5 S/tonne * 315 tonnes/year = ) $328,387/year can be estimated. The income from this activity will cover only ~33% of the chemical cost requirements (amount of NaOH and Mg needed for 90% P recovery). For BF-WWTP this is (109,000/400 = 272.05/25,000 = 6,812,500 (€/year) with a chemical cost 20,643,939.39 €/year; thus based on historical data and assuming all values change over time at the same rate, the proposal is financially unviable (~20,643,939.39 €/year+6,812,500 €/year = – 13,831,439.39) (Nethling and Benisch, 2004; Le Corre et al., 2009).

Currently the fertiliser market is dominated by phosphate rock, with 2020 price ~65 €/t (i.e. much cheaper than the assumption used). Hence rock P fertiliser is by far more economic. However, if the sludge handling optimisation is taken into consideration, the reduction of sludge disposal can significantly impact the feasibility of controlled struvite recovery. The recovery of phosphorus can reduce the volume of sludge generated by 49%. For WWTPs treating 100 m³/d, 1,000 m³/d, and 55,000 m³/d of wastewaters, the sludge disposal cost reduction would be significant (Shu et al., 2006). For BF-WWTP, overall cost of cake disposal pre-COVID was ~880 k with a total sludge production of 48,000 tonnes. Based on historical data, an annual saving of (374/€d * 2 = 748 €d * 365 days/year = ) 273,020 €/year could be achieved.

An Ostara reactor presents an annual operational cost of 574,000 €/year (Egle et al., 2016). In general, the cost arising from struvite precipitation problems within a medium size WWTP (25 MGD = 25 × 10⁶ gallon/day * 4.54 L/gallon = 113,500,000 L/day) has historically exceeded US $100,000 or 83,366.68 €/year. By taking into account the savings arising from the reduction of operational and maintenance cost due to struvite scaling, including chemical addition for chemical struvite precipitation, manpower, and maintenance costs, the overall saving could range from 1,470 € to 7,335 € per 4,540 m³ depending on the size of the treatment plant (Nethling and Benisch, 2004; Le Corre et al., 2009). If a mean price is assumed then the cost saving is estimated as 0.97 €/m³ of waste water. For Budds Farm WWTP cost saving can be estimated as (0.97 €/m³×109,000 m³/day × 365 days/year = ) 38,591,450 €/year.

The investment cost for a controlled crystallisation reactor depends on the size of the treatment facilities. Investment cost for the recovery of P from the effluent and sludge can be 3,732,549 € and 1,417,739 € respectively, for a population of 100,000. BF-WWTP covers a total of 410,000 PE thus, investment cost can be up to 15,303,450.9 € and 5,812,729.9 € respectively (mean price = 10,558,089.95 €) (Montag et al., 2009).

For the determination of the environmental benefits, according to the results that Molinos-Senante et al. (2011) presented, the average value of the P shadow price is approximately 42.74 €/kg (meaning that for every kg of phosphorus that is not dumped into the environment, the damage prevented, or the environmental benefit generated equals €42.74). Base on the data set tested, the weighted average shadow price, depending on the volume of treated wastewater and it is approximately 0.218 €/m³. If it is considered that BF-WWTP presents an outfall of an average 2000 L/s and it operates for 24/7 for 365 days per year, then the shadow price arising from its treating capacity is (0.218 €/m³ × 2000 L/s × 3600 s/min × 60 min/h × 24 h/day × 365 days/year = ) 824,981,760,000 €/year.

Table 4 summarises the integrated evaluation of the 3 proposed technologies by taking into account all the financial and environmental associated aspects.

Based on the data extracted from the CBA, struvite recovery via chemical precipitation is not viable. This also applies for chemical precipitation in combination with saline water and sand as auto-nucleation media. However, a crystallisation reactor is potentially more attractive. Considering a WWTP with treatment capacity 410,000 PE that recovers P in the form of struvite from digested sludge and its wastewater volume is 110,000 m³/day with a phosphorus concentration in the influent of 200 mg/L, this proposal is potentially economically viable with pay-back period of less than 10 years (assuming interest rate 6% and discount rate 3.5%).

However, when the average value of the EB is taken into account, phosphorus recovery is economically feasible even for the chemical struvite precipitation. Note that for the estimation of EB, only environmental benefits have been estimated, and not the increase in resource availability; if this impact was incorporated into the feasibility analysis, the results would be even more favourable.

5. Conclusions

Uncontrolled struvite formation is a significant operational problem that many WWTPs, especially those incorporating secondary treatment and anaerobic sludge digestion facilities, are required to overcome. Although uncontrolled formation of struvite can be a nuisance, controlled production and recovery of struvite can be beneficial for WWTPs. This study has demonstrated for the first time that struvite recovery from wastewater treatment is an economically viable circular economy opportunity.

Plants such as BF-WWTP provide an example that could potentially deliver a sustainable solution to this problem around the world due to its particular design and operational characteristics. Operational hotspots such as digester recirculation pumps, PDST, centrifuge feed, centrate and SAS storage tank demonstrate suitable conditions for precipitation in the form of struvite. These locations present a positive SSR, optimum pH ran between 7 and 10.7 and a significant T variation indicating, based on the literature, potential struvite precipitation.

Controlled crystallisation process is one of the most promising P recovery technologies in the form of struvite. Using a fluidised bed reactor,
this technology discharges the WWTP operational system from Mg:NH₄:PO₄²⁻ by creating optimum formation conditions. A crystallisation reactor seems to be the most robust technology for this process. Considering the great environmental and recovery performance of this proposal, a CBA based on the characteristics of BF-WWTP indicates that the pay-back period would be less than 10 years, especially given the likely increasing future energy costs and greenhouse gas emissions associated with fertiliser production. Of course, economic viability and payback periods will vary according to circumstances, but we recommend that WWTP operators generally consider fitting a crystallisation reactor to appropriate plants. The outcomes and recommendation from this study are particularly timely given the global fertiliser shortage (2022) that is driving up food prices and reducing crop sizes.

Declarations

Author contribution statement

Panayiotis Achilleos: Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Ian D. Williams: Conceived and designed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Keiron Roberts: Conceived and designed the experiments; Analyzed and interpreted the data; Wrote the paper.

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Data will be made available on request.

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The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

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