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1 **Advocating circular economy in wastewater treatment: Struvite formation**  
2 **and drinking water reclamation from real municipal effluents**

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## 18 **Abstract**

19 In this pilot study, the circular economy concept in wastewater treatment was examined,  
20 through a zero liquid discharge (ZLD) process where struvite was recovered and drinking  
21 water was reclaimed. A stage wise approach was used for struvite formation and the  
22 subsequent reclamation of drinking water. Specifically, the early stages of treatment entail the  
23 synthesis of struvite via the chemical precipitation of nutrients (phosphate and ammonia),  
24 using thermally activated cryptocrystalline magnesite. Thence, reverse osmosis (RO) was  
25 employed for drinking water reclamation. With this dual approach, 3.5 m<sup>3</sup> of municipal  
26 wastewater were successfully treated at a pilot plant in South Africa, producing ~52.5 kg of  
27 struvite and ~3.4 m<sup>3</sup> of drinking water. The operating parameters were 30 min of residence  
28 time, 0.5 g : 500 mL solid to liquid (S/L) ratio, using ambient temperature and pH. X-ray  
29 diffraction (XRD) and High Resolution Scanning Electron Microscopy (HR-SEM) coupled  
30 with electron dispersion spectroscopy (EDS) confirmed the synthesis of struvite and the  
31 presence of notable Mg/P ratios. Fourier Transform Infrared Spectrometer (FTIR) further  
32 ascertained the obtained results. Moreover, it was identified that the reclaimed water meets  
33 the South African National Standard (SANS) 241 and the world health organisation (WHO)  
34 standards for drinking water. An economic analysis revealed the viability of the process,  
35 suggesting that the system could be self-sustainable. Therefore, the results of this work  
36 indicate that introducing the concept of circular economy in wastewater treatment can  
37 promote the sustainable management of the ever-increasing quantities of municipal  
38 wastewater and at the same time address problems of emerging concern, such as water  
39 scarcity and phosphate shortage.

40 **Keywords:** Municipal wastewater; struvite synthesis; reclamation of drinking water;  
41 reverse osmosis (RO); circular economy

## 42 1 Introduction

43 Municipal wastewater influents and poorly treated effluents comprise a heterogeneous  
44 mixture of microbial, organic, and inorganic contents [1-5]. Of prime concern is the  
45 ubiquitous presence of ammonia, nitrate, and phosphate, which can pose a major  
46 environmental threat to the receiving ecosystem if not managed properly. In particular,  
47 nutrient enrichment can render the environment conducive to aquatic plant a rapid growth.  
48 However, when aquatic plants die, they deplete the dissolved oxygen in the water through  
49 aerobic decomposition. This create anoxic conditions, thus depriving aquatic organisms of  
50 oxygen. A vicious cycle begins, where the death of more aquatic organisms will ensue, which  
51 reduces the aquatic biodiversity and its ability to foster life, including benthic organisms and  
52 habitat [6-10]. In light of that, techno-viable wastewater management techniques need to be  
53 developed and implemented, particularly in low- and middle-income countries (LMIC) such  
54 as South Africa, to address this problem. According to various standards and water quality  
55 limits, the proposed level for ammonia and phosphate should be  $\leq 1.5 \text{ mg L}^{-1}$  and  $\leq 10 \text{ mg L}^{-1}$ ,  
56 respectively, when discharged to the receiving environment, albeit ammonia can be converted  
57 into nitrate or nitrite with prescribed limits of  $\leq 11$  and  $\leq 0.90 \text{ mg L}^{-1}$ , respectively [11-14].

58 Nonetheless, the presence of ammonia and phosphate in municipal influents can also render  
59 them a viable option for their recovery through the synthesis of struvite [15-18]. In addition,  
60 the treated water can be further purified and reclaimed to address the alarming rate of water  
61 crisis, in South Africa and further afield. Specifically, according to the United Nations (UN)  
62 sustainable development report, by 2030 the water crisis will be very high in developing  
63 countries, primarily due to overpopulation and pollution [19].

64 This pilot study examines the performance of an actual struvite recovery system, scaled up at  
65 village-level and operating under the South African setting. Noteworthy, previous works  
66 focused on optimization of the operational parameters [1, 20], which were used for the  
67 sustainable and effective operation of this pilot unit. Furthermore, it was identified that  
68 ammonia removal was dependent on phosphate and magnesium concentration, with  
69 decreasing phosphate concentration largely affecting the system's efficacy in terms of  
70 ammonia removal [1]. In light of that, a fixed system under optimised conditions, which will  
71 yield high efficacy for ammonia removal, was examined as will be demonstrated. In addition  
72 to struvite, drinking water was also recovered by means of reverse osmosis (RO) filtration.  
73 Specifically, in recent decades, RO has emerged as a promising technique for pollution  
74 abatement and drinking water production. This is owed to its excellent separation process,  
75 which exhibits a high rejection rate of water contaminants, including a broad range of organic  
76 and inorganic pollutants as well as micropollutants and microorganisms and pathogens [21-  
77 23]. When RO is used for municipal wastewater reclamation, it has been observed that Ca  
78 and Mg in wastewater can lead to membrane scaling, thus an antiscalant should be injected  
79 [24]. Several studies have used lime, lime/soda ash,  $Al_2(SO_4)_3$ , and  $Na_2CO_3$ , among others, to  
80 protect the membrane from fouling, scaling and metal poisoning [21, 23]. However, in this  
81 work MgO is used towards struvite formation, which suggests that through Mg scavenging  
82 from the wastewater its concentration can be minimized, thus largely avoiding the need for  
83 antiscalants.

84 Overall, in this industrially orientated study, a problematic liquid waste (municipal  
85 wastewater) is valorised in the context of circular economy and through a zero liquid  
86 discharge (ZLD) processes to synthesize struvite and reclaim drinking water. Furthermore,

87 struvite can be used as slow release fertiliser [25, 26] or for the production of phosphoric acid  
88 [27-29], among others. This means that virgin materials that would be consumed for the  
89 manufacturing of these products are protected, while struvite can also address, at least partly,  
90 the problem of the dwindling reserves of phosphate rock. In addition, the RO reclaimed  
91 drinking water can also address water scarcity concerns, in South Africa and further afield.  
92 Future works of our group will also focus on cost effective alternatives to RO filtration, such  
93 as nanofiltration (NF) and ultrafiltration (UF) as well as on photocatalysis, given the high  
94 number of annual sunshine hours in South Africa. Overall, the system is versatile, amenable  
95 and retrofittable, hence denoting that it can be directly installed in different South African  
96 set-ups and can be further scaled up to industrial systems. Finally, by being a ZLD process  
97 the environmental impacts from discharging poorly treated or untreated wastewater, a  
98 problem in South Africa and other LMICs that are currently struggling to manage the ever-  
99 increasing wastewater volumes due to rapid population growth, is also addressed. Therefore,  
100 this is one of the first few studies in design and execution of a closed loop system for  
101 valorisation and beneficiation of real municipal influents in the circular economy context.

## 102 **2 Materials and methods**

### 103 **2.1 System description**

104 The stage wise treatment process, starting from raw municipal wastewater to struvite  
105 recovery and drinking water reclamation, is shown in **Figure 1**. This process was  
106 implemented at a pilot system, constructed at the premises of the Council for Scientific and  
107 Industrial Research (CSIR) in Pretoria, with a treatment capacity of 3.5 m<sup>3</sup> per run (around  
108 one hour). The pilot plant was found to be able to effectively treat raw municipal wastewater,  
109 collected from a typical treatment plant in Pretoria. It should be noted that after collection,

110 the raw wastewater was simply screened before being fed into the pilot unit. Furthermore,  
111 struvite was recovered and water that meets the South African National Standard (SANS)  
112 241-2:2015 and the world health organisation (WHO) standards for drinking water was  
113 reclaimed. The treatment process consists of five discrete stages, shown in **Figure 1**, while  
114 their functions are described in **Table 1**.

115 *Figure 1 here*

116 As shown in **Figure 1**, raw municipal influent is first introduced into a 3.5 m<sup>3</sup> clarifier, where  
117 an overhead stirrer is in place. Then, the treatment process is carried out in five (5) stages,  
118 which are: (1) struvite synthesis, (2) storage of the treated effluent, (3) treated effluent  
119 purification by means of RO, (4) purified water storage, and finally (5) struvite storage. It  
120 should be noted that the produced retentate is send back to the feed stream, i.e. to the raw  
121 municipal effluent stream, while the filtrate from the filter press is send to the storage tank  
122 (**Figure 1**). A discussion on the treatment process is given below.

123 *Table 1 here*

## 124 **2.2 Synthesis of struvite**

125 The first stage of the treatment process is the synthesis of struvite by means of thermally  
126 activated cryptocrystalline magnesite (calcined magnesite thereafter). The raw  
127 cryptocrystalline magnesite activation process, along with the main characteristics of the  
128 activated cryptocrystalline magnesite can be found in [1]. The conditions that optimize  
129 struvite synthesis, both in terms of pollutants removal and environmental sustainability, were  
130 adopted using the results of previous works of our group [1, 20]. Specifically, struvite was  
131 synthesized at 30 minutes of equilibration time, using a 0.5 g calcined magnesite to 500 mL

132 wastewater ratio (i.e.  $1 \text{ g L}^{-1}$ ), at ambient wastewater temperature and pH. After struvite  
133 formation, the treated effluent was left to settle for another 30 min and then the supernatant  
134 was moved to the holding tank (**Stage 2**). The produced sludge, which contains struvite, was  
135 dewatered using a filter press (**Stage 5**) and the filtrate was then directed a holding tank,  
136 where the supernatant is stored (**Figure 1**). The dewatered struvite was stored in a drying bed.  
137 The physicochemical and microbial parameters of the raw wastewater, the treated effluent,  
138 the reclaimed drinking water, and the recovered struvite were determined using state-of-the-  
139 art analytical techniques and are discussed below.

### 140 **2.3 Drinking water reclamation**

141 Drinking water was reclaimed by diverting the treated effluent from the holding tank to a RO  
142 unit for further polishing. Before the RO treatment, the calcined magnesite treated effluent  
143 was disinfected using chlorine, to remove biological contaminants such as microbes and  
144 *E.coli* and other coliform bacteria.

145 For RO membrane protection, the effluent was also dosed with 0.5 L diluted HCl (0.01M), to  
146 correct the pH to  $\sim 7.5$ , and 5 L of antiscalant solution (2 ppm concentration) per run ( $3.5 \text{ m}^3$   
147 supernatant). The treated effluent was then diverted to the RO system for further purification  
148 (**Stage 3**). After treatment, drinking water was reclaimed and retentate was also produced.  
149 The drinking water was further purified by chlorination and moved to the purified water tank  
150 (**Stage 4**). The residual limit was maintained at  $<5 \text{ mgL}^{-1}$  to ensure that the water remains  
151 disinfected throughout the supply chain until the end-user (Table 2). This was also  
152 conforming to the SANS 241-2:2015 and the WHO specifications for drinking water. The  
153 retentate was then diverted to the raw municipal wastewater stream for treatment (**Stage 1**).



## 154 **2.4 Materials and chemical reagents**

155 The raw municipal wastewater was collected from a treatment plant situated in Gauteng  
156 Province, South Africa, by means of a 10 m<sup>3</sup> vacuum truck. As such, seasonal variations in  
157 the chemical components of municipal wastewater were not considered. The source of  
158 magnesium oxide (MgO), which is required for nutrients precipitation from the wastewater,  
159 was prepared using raw cryptocrystalline magnesite, which was mined from an abandoned  
160 magnesite mine in Folovhodwe, South Africa. The raw magnesite was milled, then calcined  
161 at 1000 °C, as to activate it, and then re-milled to fine powder [1]. Finally, commercially  
162 available and off-the-shelf chemicals, i.e. chlorine, HCl, and antiscalant, were used in the  
163 pilot system.

## 164 **2.5 Reverse osmosis properties: membrane specifications and operating limits**

165 In this study, a FilmTec™ BW30-4040 membrane was used, which is typically employed in  
166 light industrial applications, including for wastewater treatment [30]. The selected membrane  
167 is able to provide a reliable output of high quality water, which is required in order for the  
168 reclaimed water to meet SANS 241-2:2015 and the WHO specifications for drinking water.  
169 The properties and specifications of the reverse osmosis (RO) system are shown in **Table 2**.

170 *Table 2 here*

## 171 **2.6 Characterisation**

### 172 **2.6.1 Aqueous characterisation**

173 In-situ analyses for pH, electrical conductivity (EC), and total dissolved solids (TDS) were  
174 carried out using a multi-parameter probe (HANNA instrument, HI9828). Ex-situ analyses

175 were carried out in a South African National Standard (SANAS) accredited laboratory,  
176 located at CSIR, Pretoria, South Africa.

### 177 **2.6.2 Solids characterisation**

178 The analytical pieces of equipment that were used to identify the elemental composition, the  
179 morphological and chemical properties, and the metal functional groups of the raw  
180 and calcined magnesite, as well as of the synthesized struvite are shown in **Table 3**. It should  
181 be noted that both the high-resolution scanning electron microscope (HR - SEM) and the  
182 high-resolution transmission electron microscope (HR - TEM) used in this work were  
183 coupled with an energy dispersive X-Ray spectroscopy (EDS) system.

184 *Table 3 here*

### 185 **2.7 Economic analysis**

186 A preliminary economic analysis of the assembled pilot plant was carried out to shed light on  
187 the estimated cost of the proposed technology and to quantify the possible economic gains of  
188 the recovered process co-products (struvite and drinking water). Specifically, both co-  
189 products have a commercial value, which only for the case of struvite can be as high as 200  
190 USD (or R3 500) per ton, as is suggested in different chemical supply industries catalogues.  
191 Furthermore, in South Africa, there is no rural-urban tariff for potable water, since the cost is  
192 determined by consumption, with penalties in place for heavy consumers. Municipalities  
193 purchase water, in bulk supply, from commercial companies and typically sell it to end-users  
194 at a higher rate. In the context of this pilot study, a mean cost of 0.54 USD (or R9.10) per m<sup>3</sup>  
195 of drinking water was considered. Here, the economic analysis was focused on the direct field  
196 cost (DFC) and operational expenditure (OPEX), taking into account the following  
197 assumptions and limitations:

- 198 • The plant will operate 24 hours, 7 days a week (24/7), 360 days per year (to account  
199 for maintenance activities) with a minimal of 95% availability on an annual basis.
- 200 • Extraordinary conditions, such as flooding and climate extremes, which could hamper  
201 the plant's operation, are not included in the analysis.
- 202 • The supply and transportation cost of the municipal influent is deemed negligible as  
203 conventional treatment plants could be retrofitted with the proposed technology.
- 204 • In the economic analysis only the DFC and OPEX were considered, and thus labor  
205 costs and other possible expenditures are outside of this analysis.
- 206 • The electricity input required for the machinery operation is included in the analysis.

### 207 **3 Results and discussion**

#### 208 **3.1 Chemical characteristics of the raw wastewater and treated effluent**

209 Table 4 list the chemical composition of the raw wastewater and of the effluent of each  
210 treatment stage, i.e. calcined magnesite treated effluent and RO water output. A discussion on  
211 the chemical characteristics of the effluent of each main stage of this technology, including  
212 the reclaimed drinking water is given below.

213 *Table 4 here*

##### 214 **3.1.1 Raw wastewater influent**

215 As expected, the raw wastewater is rich in organic, inorganic, and biological contaminants  
216 and particularly in ammonia, phosphate, organic carbon, and bacteria content (**Table 4**). This  
217 can be attributed to decomposition of organic waste, which lead to the release of pathogenic  
218 bacteria, ammonia, and phosphate, among others. Specifically, *E. coli*, total coliform, and the

219 colony counts of the heterotrophic bacteria (i.e. total plate count) were very high, as was  
220 expected. This is also the case for ammonia, phosphate, colour, turbidity and conductivity  
221 parameters. The reported values are typical for raw municipal wastewater. Finally, it should  
222 be mentioned that the retentate, i.e. the reject water stream of the RO process, is diverted to  
223 the raw wastewater stream for treatment. This ensures that no wastewater is discharged to the  
224 environment and that a ZLD processes is carried out.

### 225 **3.1.2 Calcined magnesite treated effluent**

226 As shown in **Table 4**, calcined magnesite treatment has a high efficacy in the removal of both  
227 biological and particularly the nutrient content of raw wastewater. The pH of the raw  
228 wastewater was observed to be within the SANS 241-2: 2015 specifications. However, after  
229 treatment with calcined magnesite, it was observed that the pH of the effluent grossly  
230 increases. This is attributed to the addition of hydroxyl groups from the reaction of calcined  
231 magnesite with the raw wastewater. The reactions that govern the observed increase in the pH  
232 value of the treated effluent are shown in equation (1) and (2):



235 Furthermore, the high value of EC in the raw wastewater was observed to decrease after the  
236 treatment. This could be explained by the reaction of the calcined magnesite with the raw  
237 wastewater to form struvite. Specifically, struvite is sinking certain contaminants that are  
238 associated with the observed high EC value in the raw wastewater. Hence, their removal  
239 attenuates the EC value in the treated effluent, albeit the EC value is still above the drinking  
240 water limit. The same pattern was observed in the TDS value, which also decreased after  
241 treatment, but its level was still above the drinking water limit. Mn and Fe values were below

242 the SANS 241:2015-2 limits for drinking water in the raw wastewater, nonetheless calcined  
243 magnesite treatment further reduced them to minuscule levels. Fe is a common inorganic  
244 pollutant causing membrane fouling [31, 32], and hence its precipitation along with struvite is  
245 beneficial for the membrane fouling inhibition. In addition, the results shown in Table 4,  
246 indicate that the concentration of *E. coli*, total coliform, total plate counts, conductivity,  
247 dissolved solids, and ammonia has been reduced but they are still above the SANS 241-  
248 2:2015-2 limits. On the other hand, alkalinity, pH, Mg, Ca, colour and turbidity were  
249 observed to increase after treatment. This may be attributed to the dissolution of Mg and Ca  
250 from the matrices of calcined magnesite, hence leading to an increase in pH and alkalinity.  
251 The existence of Mg and Ca was confirmed by FTIR, SEM-EDS and XRD techniques, as  
252 reported below. This was expected since the thermally activated magnesite is rich in MgO,  
253 while it also contains insignificant amounts of Ca. However, after RO treatment, high purity  
254 potable water was obtained that meets the SANS 241-2:2015-2 specifications (Table 4).  
255 Regarding the ammonia removal efficacy, this was observed to be dependent on phosphate  
256 concentration, as the struvite formation is dependent on the molar ratio of the two. In this  
257 work, approximately 87% and 98.8% removal efficacy for ammonia and phosphate  
258 respectively, was observed. However, in this context one notable drawback is the system's  
259 deficiency in phosphate, which is required to form a co-ordination of Mg, NH<sub>3</sub> and PO<sub>4</sub>  
260 reaction towards the synthesis of struvite, since a molar ratio of 1:1:1.1 is required. This  
261 claim was also noted in a study by Tansel, et al. [33]. To this end, future research of our  
262 group will focus on supplementing the phosphate level to enhance struvite synthesis. This  
263 will also minimize the Mg concentration in the treated effluent, thus limiting the need of an  
264 antiscalant.

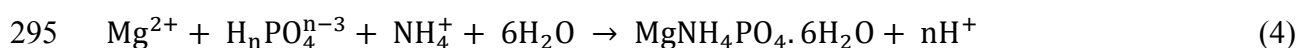
265 Overall, it was found that calcined magnesite treatment can effectively remove ammonia and  
266 phosphate, and possibly heavy metals such as zinc (Table 4), from municipal wastewater.  
267 Attenuation of biological contaminants in the raw wastewater was also observed. This  
268 suggests that this technology has some disinfection capabilities, which enable it to remove  
269 biological contaminants. In a nutshell, struvite was successfully synthesized and the nutrient  
270 load of the treated effluent was substantially attenuated. However, as was expected, the  
271 chemical characteristics of the treatment effluent did not meet those for drinking water, as  
272 specified in SANS 241-2:2015-2 (Table 4). Therefore, since in the context of this work  
273 drinking water reclamation is also anticipated, the calcined magnesite treated effluent was  
274 purified using RO and chlorination, as described below.

### 275 3.1.3 Reverse osmosis purified water

276 As shown in Figure 1, after the calcined magnesite treatment, the effluent was further  
277 purified by a combination of chlorination and RO, as to reclaim drinking water. Specifically,  
278 to disinfect the water and remove biological contamination first chlorine was added to the  
279 calcined magnesite treated effluent. Hydrochloric acid was also added to drop the pH, as to  
280 protect the RO membrane. This also contributed to the reduction of alkalinity (Table 4).  
281 Then, the effluent was passed through the RO system. As shown in Table 4, the use of the  
282 filtration technology efficiently reduced all examined parameters to meet the SANS 241-  
283 2:2015-2 specifications for drinking water. Among others, the colour, turbidity, magnesium,  
284 and calcium concentrations were drastically reduced. The RO treated water was then  
285 chlorinated again, keeping the residual limit  $<5 \text{ mg L}^{-1}$ , as to ensure that the water remains  
286 disinfected throughout the supply chain until the end-user. Finally, the generated retentate is  
287 sent back to the raw wastewater stream for treatment (Figure 1).

## 288 3.2 Struvite synthesis chemistry

289 It is well established that the removal of phosphate and ammonia from wastewater is mainly  
290 attributed to a combination of adsorption, crystallization, and precipitation. Adsorption  
291 precedes precipitation and then crystallization leads to the formation of struvite and the  
292 synthesis of other minerals [34]. Equation (3) and (4) show the possible routes for the  
293 formation of a number of product minerals during the treatment of municipal wastewater:



296 where  $n = 0, 1, 2$ , etc., and it corresponds to the pH of the solution [35].

297 Furthermore, the presence of MgO and CaO within the matrices of the thermally activated  
298 cryptocrystalline magnesite will enable it to scavenge ammonia and phosphate via the  
299 crystallization of calcium, magnesium, phosphate and ammonia, as depicted in equations (3)  
300 – (6). This pattern was also reported by Peng, et al. [35]. Furthermore, this system cannot be  
301 purely defined by adsorption models, since the process is not reversible and it is not utterly a  
302 surface phenomenon. This suggests that it can be classified as a precipitation process with  
303 subsequent crystallization and it is within the branch of co-ordination chemistry. This allows  
304 the EC, TDS, and other contaminants to be reduced to acceptable limits.

## 305 3.3 Characterization of the raw and calcined cryptocrystalline magnesite and 306 product minerals

307 According to the mass balance of this pilot unit approximately 52.5 kg of sludge per 3.5 m<sup>3</sup>  
308 will be produced. The product minerals contained in the produced sludge are also analyzed  
309 below, using different analytical techniques.

### 310 3.3.1 Mineralogical characterization

311 In **Figure 2**, the XRD patterns of the raw cryptocrystalline magnesite, the calcined magnesite,  
312 and the synthesized struvite are shown. Furthermore, Table 5 lists the identified mineral  
313 phases, along with the measured XRD peaks (2 theta degrees) and relevant studies that  
314 underpin the presented results.

315 *Figure 2 here*

316 Specifically, the identified 2 theta degrees of the raw cryptocrystalline magnesite (**Figure 2**)  
317 suggest that it is rich in magnesite, which is further corroborated by studies cited in Table 5.  
318 However, after calcination, new mineral phases, i.e. periclase and brucite, were observed in  
319 the calcined magnesite. This also indicates that CO<sub>2</sub> was released, as an airborne emission,  
320 from the matrices of cryptocrystalline magnesite, as shown in equation (5):



322 Finally, the recovered struvite matrix was observed to contain, apart from struvite, quartz,  
323 brucite, periclase, and magnesite. These results not only confirm the synthesis of struvite  
324 through the proposed technology, but also the presence of other Mg-based minerals. The  
325 latter could be beneficial in cases where struvite is going to be used for agricultural soil  
326 amelioration and fertility enhancement.

327 *Table 5 here*

### 328 3.3.2 Metal functional groups

329 The metal functional groups of the raw cryptocrystalline magnesite, the calcined magnesite,  
330 and the synthesized struvite were identified using the FTIR technique. In **Figure 3** the FTIR



331 spectrum of raw and calcined magnesite and of the synthesized struvite are shown, while  
332 **Table 6** lists the identified metal functional groups, their wavenumber ( $\text{cm}^{-1}$ ) and citations for  
333 relevant studies dealing with the identified metal functional groups.

334 *Figure 3 here*

335 As shown in **Figure 3** and **Table 6**, the metal functional groups of raw cryptocrystalline  
336 magnesite include carbonate and silicates, while water is also present. However, after  
337 calcination only the carbonates were still present and a new functional group of MgO was  
338 observed, which will be instrumental towards struvite crystallization. This confirms that  $\text{CO}_2$   
339 was volatilized and MgO was formed in the calcined magnesite matrix. The FTIR results also  
340 corroborate the ones obtained by the XRD technique. Furthermore, the presence of hydroxide  
341 (-OH) in both the calcined magnesite and the recovered struvite matrices suggest the  
342 possibility of brucite being present, which was confirmed in the XRD results. Regarding the  
343 recovered struvite, it was observed to contain  $\text{PO}_4^{3-}$ , N-H and MgO (**Table 6**), which  
344 confirms that indeed through the proposed technology struvite is formed. Hydroxide, along  
345 with carbonates were also observed. As mentioned above, hydroxide suggest the presence of  
346 brucite, while the carbonates could be attribute to the cryptocrystalline magnesite, as denoted  
347 by the XRD results. Finally, apart from suggesting that brucite might be present, prevalence  
348 of hydroxide also suggests that the synthesized struvite is hydrated, as shown in equation (6)  
349 [1]:



351 *Table 6 here*

### 352 **3.3.3 Morphological characteristics**

353 The morphology of the raw cryptocrystalline magnesite, the calcined magnesite, and the  
354 synthesized struvite are shown in **Figure 4**. As mentioned in **Table 3** a high-resolution  
355 Focused Ion Beam Scanning Electron Microscope (FIB FESEM) instrument was employed to  
356 identify the morphological characteristics of the matrices under study. Specifically, an Auriga  
357 Cobra FIB FESEM was used in this work, which can be also used for SEM imaging. Using  
358 the FESEM function clear, ultra-high resolution, and low electrostatically distorted images  
359 were obtained that revealed the morphology of the minerals under study (**Figure 4**).

360 ***Figure 4 here***

361 As shown in **Figure 4**, the raw cryptocrystalline magnesite has irregular sheets, which are  
362 homogenously distributed across its surface. Furthermore, regardless of the different  
363 magnification used, i.e. from 1  $\mu\text{m}$  to 100 nm, no changes in the morphological properties of  
364 the raw cryptocrystalline magnesite surface were identified. The FESEM results also suggest  
365 that cryptocrystalline magnesite is a nanomaterial with octagonal structures. However, after  
366 calcination the morphology of the raw cryptocrystalline magnesite was observed to have  
367 changed. This could be attributed to the fact that during calcination mass loss was observed,  
368 attributed to  $\text{CO}_2$  volatilization, which suggests that a reaction took place to form MgO. As a  
369 result, after calcination the morphology of the calcined magnesite was observed to have  
370 changed to round nanosheets, homogeneously distributed across its surface (**Figure 4**).  
371 Finally, after the calcined magnesite comes into contact with the municipal effluent struvite is  
372 synthesized, which was found to be rich in round nanosheets as well. Our group [1] and  
373 various other groups have reported similar findings. For example, Herald, et al. [36]  
374 reported that struvite can be present in different morphologies, such as coffin shaped,  
375 pyramid type, prismatic type, needle type, and feather shaped, amongst others, and this is

376 strongly dependent on growth parameters and synthesis conditions Herald, et al. [36].  
377 Similar findings were reported by Chauhan and Joshi [37]. In addition, other Mg based  
378 minerals, apart from MgO, could have influenced the morphology of the synthesized struvite,  
379 along with impurities associated with the calcined magnesite matrix and micelles that would  
380 have been co-deposited with struvite during the synthesis process.

### 381 **3.3.4 Spot analysis using HR-SEM**

382 To further corroborate the elemental composition of the raw and calcined cryptocrystalline  
383 magnesite and of the synthesized struvite, spot-analysis, by means of Auriga Cobra FIB-  
384 FESEM instrument which includes an EDS detector, was employed. The morphological spot-  
385 analysis results of the raw cryptocrystalline magnesite, calcined magnesite, and synthesized  
386 struvite are shown in **Figure 5** to **Figure 7**, respectively.

#### 387 *Figure 5 here*

388 As shown in **Figure 5a**, raw cryptocrystalline magnesite is predominated by octagonal  
389 structures on its surface. Elemental analysis further revealed that the predominant elements  
390 are C, Mg, and O, as well as traces (< 1%) of Ca and Si (**Figure 5b-f**). During the calcination  
391 process of the raw cryptocrystalline magnesite, C is the element that will be released in the  
392 environment through CO<sub>2</sub> volatilization. The EDS spot analysis results are also in agreement  
393 with the XRD and FTIR results for raw cryptocrystalline magnesite. Furthermore, the  
394 identified Ca and Si concentrations would be instrumental in increasing the pH of the raw  
395 wastewater effluent, if they would not be affected by the calcination process. Regarding the  
396 morphological properties of the calcined magnesite, these are shown in **Figure 6**.

#### 397 *Figure 6 here*

398 **Figure 6a** portrays a nanosheet like morphology for the calcined cryptocrystalline magnesite.  
399 Furthermore, C, Mg, and O are the predominant elements in the calcined magnesite, while  
400 also traces of Ca, Si, Al, and Fe were observed. However, it should be noted that in spectrum  
401 no.156 a large Ca concentration (18.8%) was observed. These results are also in agreement  
402 with the XRD and FTIR results. Furthermore, it appears that the calcination process does not  
403 affect the Ca and Si levels that were observed in the raw cryptocrystalline magnesite matrix.  
404 These concentrations are pivotal for the chemical precipitation of nutrients from wastewater,  
405 since they will aid in increasing the pH of the raw wastewater. The pH increase will create  
406 the conditions that are suitable for the synthesis of struvite, since a defined alkaline pH range  
407 is required for struvite crystallization, as is well established in the literature [1, 15, 16, 38-40].  
408 Finally, the morphological properties (SEM-EDS spot analysis) of the synthesized struvite  
409 are shown in **Figure 7**.

410 *Figure 7 here*

411 As shown in **Figure 7**, spherical and homogenous crystals were observed in the synthesized  
412 struvite. Furthermore, it was identified that the synthesized struvite is predominated by C,  
413 Mg, and O, while traces of Ca and Si were also observed. More importantly, the existence of  
414 P was identified, which confirms the synthesis of struvite. It should be noted that only the  
415 soluble component of calcined magnesite contributes to the synthesis of struvite and the non-  
416 soluble will settle. Moreover, the Mg and P content confirm struvite formation. Finally, the  
417 identified Si and Ca traces suggest the co-precipitation of struvite with those impurities. The  
418 obtained results are aligned with the ones obtained by the XRD and FTIR techniques.

419 **3.3.5 Morphological features obtained by the HR-TEM analysis**

420 The results regarding the morphological properties of raw cryptocrystalline magnesite, as  
421 well as the observed changes in the morphological properties of the calcined magnesite and  
422 the synthesized struvite are shown in **Figure 8**. In total, four different magnifications, i.e.  
423 1 $\mu$ m, 500 nm, 200 nm, and a high-magnification of 100 nm, were used.  
424 Specifically, as shown in **Figure 8** the raw cryptocrystalline magnesite has irregular  
425 nanosheets homogenously distributed across its surface, as was also identified in the FESEM  
426 analysis. Furthermore, regardless of the magnification the morphological properties of the  
427 raw cryptocrystalline magnesite were observed to remain the same. Moreover, **Figure 8**  
428 confirms that cryptocrystalline magnesite is a nanomaterial with octagonal to irregular  
429 structures. This corroborates, at least partially, the FESEM results. However, after  
430 calcination, the morphology of raw cryptocrystalline magnesite was observed to have  
431 changed. This could be attributed to the large CO<sub>2</sub> airborne emissions, during its calcination,  
432 to the atmosphere. More specifically, after calcination, the raw cryptocrystalline magnesite's  
433 morphology was observed to have changed to round nanosheets, (**Figure 8**). Furthermore, it  
434 was observed that the nanosheets are distributed homogeneously throughout the calcined  
435 magnesite's surface. Finally, after the calcined cryptocrystalline magnesite comes into  
436 contact with the raw municipal wastewater, this results to struvite synthesis. When using the  
437 four different magnifications, i.e. 1 $\mu$ m, 500 nm, 200 nm, and 100 nm, it is shown that the  
438 synthesized struvite is a nanomaterial with rich round nanosheets. This was also identified in  
439 the FESEM analysis.

440 *Figure 8 here*

#### 441 **3.4 Preliminary economic analysis of the struvite precipitation pilot unit**

442 Overall, the results of this pilot study confirmed the feasibility, from the technical  
443 perspective, of struvite precipitation and recovery, as well as of drinking water reclamation  
444 from real municipal wastewater. However, the economic feasibility of the proposed  
445 technology remains largely unknown for the South African setting and thus was examined  
446 here through a preliminary cost-benefit analysis. The analysis includes the operational  
447 expenditure (OPEX) of the process, but not the capital expenditure (CAPEX) required for the  
448 construction of the pilot unit. The reason that the CAPEX was not included in the analysis is  
449 twofold. First and foremost, this study deals with a pilot unit and as such when the process is  
450 scaled up to more comprehensive systems, economies of scale would be achieved. Therefore,  
451 including the CAPEX of the pilot unit could skew the results and make them case specific.  
452 However, this is not the case for the OPEX, since the same amounts of chemical reagents  
453 would be consumed in scaled up systems, while a relative lower electricity input per unit of  
454 volume of wastewater might be required in scaled up system. However it was identified that  
455 the cost of electricity does not significantly affect the total cost of the process (Table 7).  
456 Second, this technology is versatile and could be even incorporated into existing wastewater  
457 treatment facility, thus largely avoiding CAPEX. Furthermore, it is a prerequisite to treat the  
458 raw wastewater before being discharged to the environment, therefore its treatment will be  
459 associated with a cost regardless of the technology used. This suggest that a mean avoided  
460 cost would be required for conventional wastewater treatment, which could be ascribed to the  
461 system under study thus affecting the CAPEX of this technology. For the above reasons,  
462 CAPEX is not included in the economic analysis and a zero value was ascribed to the  
463 wastewater. Finally, the analysis was carried out in South African Rand (R) and then results  
464 were expressed using United States Dollar (USD). To provide context, the mean exchange  
465 rate of US dollar to rand was around R17 at the time of writing.

466 **Table 7 here**

467 Using the estimates shown in Table 7, a preliminary cost-benefit analysis, not including the  
468 CAPEX, was carried out for the 3.5 m<sup>3</sup> capacity pilot plant that is already constructed at the  
469 CSIR premises, in Pretoria, South Africa. By doing so, both the operating cost of treating  
470 municipal wastewater and the economic gains from the recovered struvite and the reclaimed  
471 drinking water were quantified and shown in **Table 7**. It was found that the OPEX to treat  
472 municipal wastewater treatment per run was calculated to be 0.8 USD per m<sup>3</sup> of wastewater.  
473 On the other hand, the economic gains from selling the generated products would amount to a  
474 maximum of 2.8 USD per m<sup>3</sup> of wastewater. It should be noted that the economic gains are  
475 just an estimate of the possible economic benefits of the process co-products, which does not  
476 include the cost of their further treatment, such as in the case of struvite, or their  
477 transportation cost. However, the results provide insight on the process economic  
478 sustainability and suggests that the process could be self-sustainable, from the economic  
479 perspective, providing a stable operating revenue to local communities. Furthermore, the  
480 results of this work appear to align with the literature, where this technology has been found  
481 economic viable, having a six year return of investment for larger plants with daily capacity  
482 in the range of few hundreds to few thousands m<sup>3</sup> [41].

#### 483 **4 Conclusions and recommendations**

484 The findings from the 3.5 m<sup>3</sup> pilot plant that was examined in this work suggest that through  
485 a zero liquid discharge (ZLD) processes is feasible and economic viable to recover struvite  
486 and reclaim drinking water from municipal wastewater. The pilot plant under study was  
487 found to effectively treat real municipal wastewater and at the same time i) recover struvite  
488 (52.5 kg per run) and other mineral products, which can be particularly useful in the

489 agricultural industry, and ii) reclaim drinking water (3.4 m<sup>3</sup> per run), which meets the SANS  
490 241:2015-2 specifications for drinking water. Future research will focus on the processing of  
491 struvite and of the Ca-based minerals towards the production of phosphoric acid.

492 Overall, this technology can provide the important water resource in water scarce regions and  
493 particularly in remote and insular communities in South Africa and further afield. This  
494 technology could be applied directly at village-level or further scaled up to industrial  
495 treatment systems. An additional stream of revenue will be also generated via the selling of  
496 the recovered struvite and the reclaimed drinking water. Specifically, through a preliminary  
497 economic analysis, the operational expenditure (OPEX) of the pilot unit was lower than the  
498 economic gains from the recovered struvite and the reclaimed water, suggesting that it can  
499 provide a stable operating revenue to local communities. Therefore, this technology can  
500 potentially be a game-changing approach in the endeavour to protect the environment and its  
501 precious resources at village-level in LMIC, where infrastructure is weak or non-existence,  
502 and possibly at more comprehensive scales for industrial wastewater treatment applications.

503 Finally, future works of our group will focus on introducing cost effective alternatives to  
504 reverse osmosis (RO) filtration, such as the use of other more cost effective membrane  
505 filtration process, e.g. nanofiltration (NF) and ultrafiltration (UF) as well as on advanced  
506 oxidation processes (AOPs), e.g. photocatalysis, given the high number of annual sunshine  
507 hours in South Africa.

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## Tables' captions

**Table 1:** The five treatment stages for the synthesis of struvite and drinking water reclamation.

**Table 2:** The operating limits and membrane specification of the FilmTec™ RO membrane system [42].

**Table 3:** The equipment used to determine the parameters of the raw and the thermally activated magnesite and of the synthesized minerals (struvite).

**Table 4:** The chemical composition of wastewater and of the Mg and RO treated water, along with the South African National Standard (SANS) 241-2:2015 for drinking water.

**Table 5:** The measured XRD peaks (2 theta degrees) and the identified mineral phases of raw magnesite, calcined magnesite, and synthesized struvite, along with relevant studies for each mineral phase.

**Table 6:** The identified metal functional groups of the raw magnesite, calcined magnesite, and synthesized struvite.

**Table 7:** A preliminary cost-benefit analysis for the 3.5 m<sup>3</sup> pilot plant, in Pretoria, South Africa.

## List of tables

**Table 1:** The five treatment stages for the synthesis of struvite and drinking water reclamation.

Number of stage	Unit process	Function
1	Struvite synthesis	To recover struvite
2	Holding tank	To store the treated water
3	Reverse osmosis (RO)	To purify the water
4	Storage tank	To store the purified water
5	Struvite storage tank	To store the dewatered struvite

**Table 2:** The operating limits and membrane specification of the FilmTec™ RO membrane system [42].

<b>Parameter</b>	<b>Values</b>
Product	BW30-4040
Part Number	80783
Surface Area m <sup>2</sup> (ft <sup>2</sup> )	7.2 (78)
Feed Spacer Thickness (mil)	34
Permeate Flow Rate gpd (m <sup>3</sup> /d)	2,400 (9.1)
Stabilized Salt Rejection (%)	99.5
<b>Membrane Type</b>	<b>Polyamide Thin-Film Composite</b>
Maximum Operating Temperature	45°C (113°F)
Maximum Operating Pressure	4.1 MPa ( 41 bar or 600 psi)
Feed Flow Rate: 4040	16 gpm (3.6 m <sup>3</sup> /h)
Maximum Pressure Drop	15 psig (0.1 MPa or 1.0 bar)
Length in cm	101.6
Weight in kg	17.0
pH Range, Continuous Operation	2 - 11
pH Range, Short-Term Cleaning	1 - 13
Maximum Feed Silt Density Index	SDI 5
Free Chlorine Tolerance	<0.1 mg L <sup>-1</sup>



**Table 3:** The equipment used to determine the parameters of the raw and the thermally activated magnesite and of the synthesized minerals (struvite).

Parameter	Equipment	Model
Mineralogical composition	XRD	PANalytical Aeris powder diffractometer with a PIXcel detector and fixed divergence- and receiving slits with Fe filtered Co-K $\alpha$ radiation ( $\lambda=1.789\text{\AA}$ ). Phases were identified using X'Pert Highscore plus software.
Elemental composition	XRF	The Thermo Fisher ARL Perform'X Sequential XRF instrument with Uniquant software was used for analyses.
Morphology properties and spot-analysis	SEM-EDS	Auriga Cobra FIB-FESEM (Model: Sigma VP FE-SEM with Oxford EDS Sputtering System, Make: Carl Zeiss, Supplier: Carl Zeiss, USA)
Morphology properties	HR-TEM	High-Resolution Scanning Electron Microscope (HR- TEM) (JEM – 2100 electron microscope, Angus Crescent, Netherland)
Functional groups	FTIR	Perkin-Elmer Spectrum 100 Fourier Transform Infrared Spectrometer (FTIR) equipped with a Perkin-Elmer Precisely Universal Attenuated Total Reflectance (ATR) sampling accessory equipped with a diamond crystal

**Table 4:** The chemical composition of wastewater and of the Mg and RO treated water, along with the South African National Standard (SANS) 241-2:2015 for drinking water.

Analysis	Units	SANS 241:2015-	Raw effluent	Mg treated	RO treated
		2 Limits			
E. coli	count/100mL	≤ 0	3,000	1,118	<0.001
Nitrate	mg L <sup>-1</sup> N	≤ 10	1.5	<0.16	<0.16
Nitrite	mg L <sup>-1</sup> N	≤ 0.90	0.20	<0.010	<0.010
Sulphate	mg L <sup>-1</sup> SO <sub>4</sub> <sup>2-</sup>	≤ 500	50	13	<10
Total Chromium	µg L <sup>-1</sup> Cr	≤ 50	70	<0.81	<0.5
Total Iron	µg L <sup>-1</sup> Fe	≤ 2,000	67	54	<0.01
Total Manganese	µg L <sup>-1</sup> Mn	≤ 400	50	<0.05	<0.01
Total Coliform	count per100mL	≤ 10	4,900	3,000	<0.001
Total Plate Count	count per1mL	≤ 1,000	570,000	24,000	<0.001
pH @ 25°C	-	≥ 5.00 to ≤ 9.70	7.76	10.25	7.5
Turbidity	NTU	≤ 1.00	3.32	5.37	<0.1
Colour	mg L <sup>-1</sup> Pt-Co	≤ 15	31	33	<0.5
Electrical Conductivity	mS m <sup>-1</sup>	≤ 170	327	211	20
Total Dissolved Solids	mg L <sup>-1</sup>	≤ 1,200	1,458	1,062	258
Ammonia	mg L <sup>-1</sup> NH <sub>3</sub>	≤ 1.50	300	40	0.10
Total Sodium	mg L <sup>-1</sup> Na	≤ 240	26	22	0.5
Total Zinc	mg L <sup>-1</sup> Zn	≤ 5.0	<0.01	<0.01	<0.01
Temperature	°C	≤ 30	25	26	25
Total Alkalinity	mg L <sup>-1</sup> CaCO <sub>3</sub>	≥ 250 to ≤ 300	138	350	<0.5
Total Calcium	mg L <sup>-1</sup> Ca	≤ 300	16	19	0.9
Total Magnesium	mg L <sup>-1</sup> Mg	≤ 100	15	278	0.5

Total Potassium	mg L <sup>-1</sup> K	≤ 100	19	17	<1
Total Phosphate	mg L <sup>-1</sup> P	≥ 1 to ≤ 10	180	2.19	<0.08

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**Table 5:** The measured XRD peaks (2 theta degrees) and the identified mineral phases of raw magnesite, calcined magnesite, and synthesized struvite, along with relevant studies for each mineral phase.

Mineral	2 theta degree	Mineral phase	References
Raw magnesite	39	Magnesite	[1, 43, 44]
	41		
	42.5		
	50.5		
	52.5		
	60.5		
	62		
	72		
	73		
	79		
	81.5		
82			
Calcined magnesite	21	Brucite	[16, 43, 45-47]
	42.5	Periclase	
	60	Brucite	
	69	Brucite	
	72	Periclase	
	81	Brucite	
Struvite	21	Brucite	[36, 38, 46, 48-52]

22	Struvite
31	Quartz
33	Calcite
39	Magnesite
44	Brucite
45	Struvite
50	Periclase/magnesite/struvite
60	Brucite
69	Quartz
72	Periclase
79	Struvite
82	Magnesite/Struvite

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**Table 6:** The identified metal functional groups of the raw magnesite, calcined magnesite, and synthesized struvite.

Mineral	Wavenumber (cm <sup>-1</sup> )	Metal functional	References
Raw magnesite	750	CO <sub>3</sub> out-of-plane bending vibration	[53, 54]
	905		
	1010		
	1050		
	1100		
	1200	Si-O-Si stretching	
	1400	CO <sub>3</sub>	
	3000	H <sub>2</sub> O	
Calcined magnesite	850	MgO	[1, 54]
	1400	CO <sub>3</sub>	
	3700	-OH (Water)	
Struvite	850	MgO	[1, 16, 36, 50, 55,
	1005	PO <sub>4</sub> <sup>3-</sup>	56]
	1200	PO <sub>4</sub> <sup>3-</sup>	
	1400	CO <sub>3</sub>	
	1690	N-H	
	3000	-OH (Water)	
	3700	-OH (Water)	

**Table 7:** A preliminary cost-benefit analysis for the 3.5 m<sup>3</sup> pilot plant, in Pretoria, South Africa.

<b>Raw materials and electricity</b>	<b>Unit cost (USD)</b>	<b>Input</b>	<b>Total cost (USD)/3.5 m<sup>3</sup></b>
Wastewater	0	3500	0
<b>Materials</b>			
Calcined magnesite	205.8823529	0.0035	USD
Chlorine	235.2941176	0.00001	0.002353
HCl	705.8823529	0.00005	0.035294
Antiscalant	5294.117647	0.0001	0.529412
CIP chemicals	1058.823529	0.0001	0.105882
<b>Total cost</b>			<b>0.672941</b>
<b>Electricity</b>			
Wastewater feed pump	0.066470588	0.167	0.011101
Transfer pump	0.066470588	0.167	0.011101
RO feed pump	0.066470588	0.333	0.022135
Mixers	0.066470588	0.5	0.033235
Reagent mixers	0.066470588	0.25	0.016618
Reagent dosing	0.066470588	0.167	0.011101
RO dosing pumps	0.066470588	0.167	0.011101
RO membranes	0.066470588	0.167	0.011101
<b>Total cost</b>			<b>0.127491</b>
<b>Overall cost</b>			<b>0.800432</b>
<b>Valuables</b>			
Struvite	200	0.01	2
Clean water	0.535294118	3	1.605882
<b>Total return</b>			<b>3.605882</b>
<b>Total commission for 3.5 KL</b>			<b>2.805451</b>

The convention rate of 1 USD = 17 South African Rand on an average exchange rate.

## Figures caption

**Figure 1:** The multi-stage treatment process for the synthesis of struvite and drinking water reclamation.

**Figure 2:** The X-ray diffractogram and mineralogical composition of raw magnesite, calcined magnesite and the synthesized struvite.

**Figure 3:** The metal functional groups of the raw magnesite, calcined magnesite, and synthesized struvite.

**Figure 4:** The FESEM images showing the changes in morphological properties of raw magnesite, calcined magnesite, and the synthesized struvite from lower to higher magnification, i.e. a) 1  $\mu\text{m}$ , b) 200 nm, and c) 100 nm.

**Figure 5:** The SEM-EDS results of the raw magnesite showing (a) the spot analysis image and (b) to (f) the relevant EDS spectra.

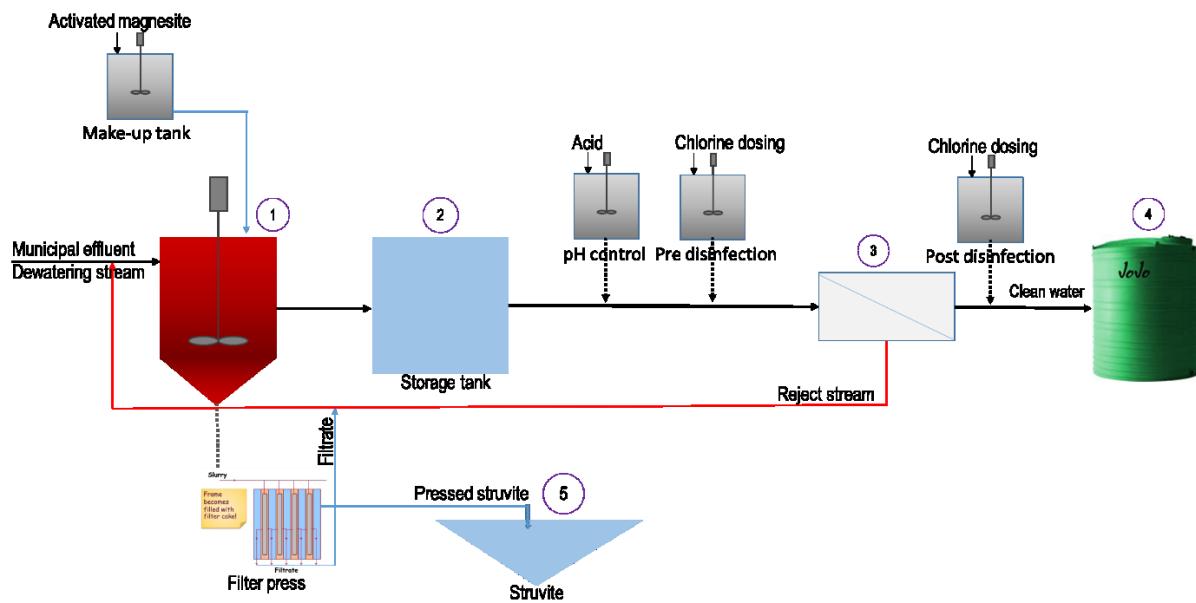
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**Figure 7:** The SEM-EDS results of the synthesized struvite showing (a) the spot analysis image and (b) to (f) the relevant EDS spectra.

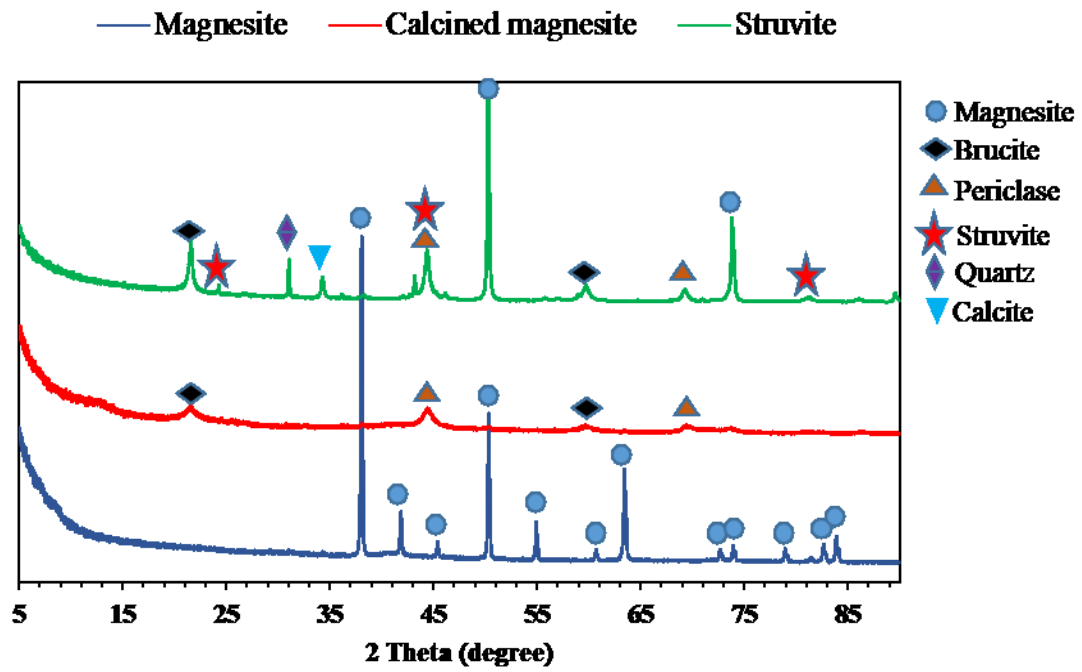
**Figure 8:** The morphological properties of raw magnesite, calcined magnesite and the synthesized struvite using four magnifications, i.e. a) 1  $\mu\text{m}$ , b) 500 nm, c) 200 nm, and d) 100 nm.



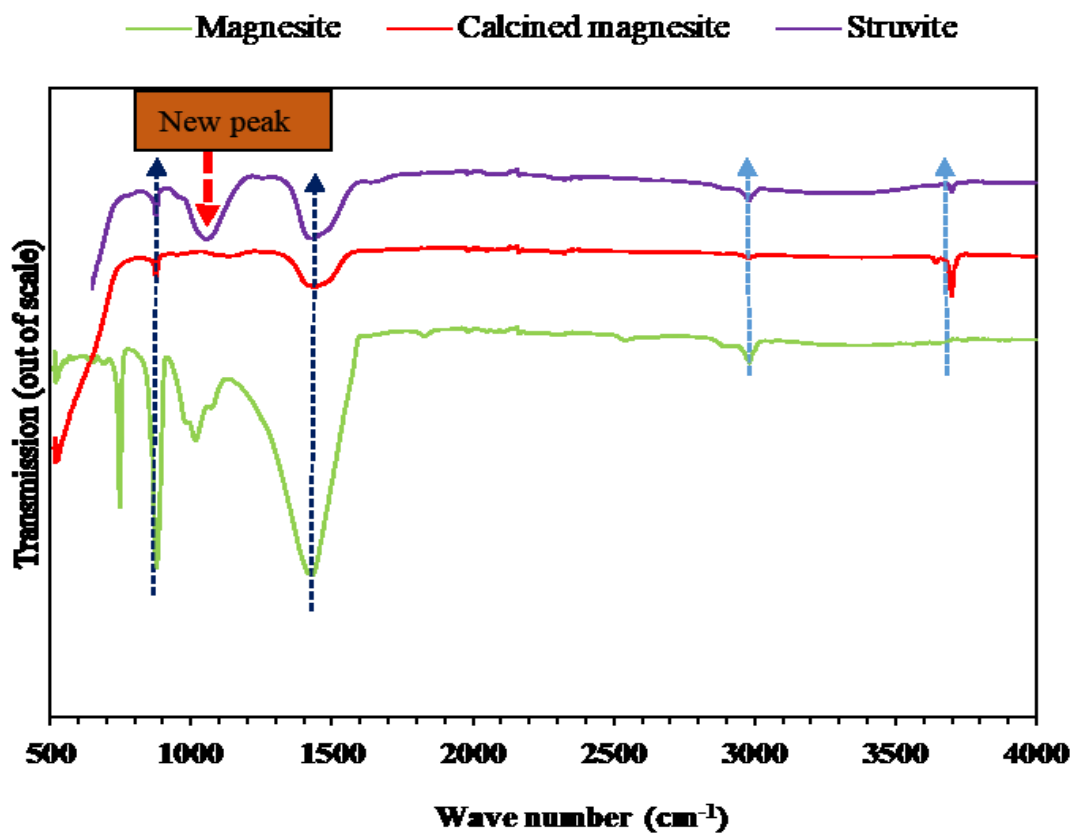
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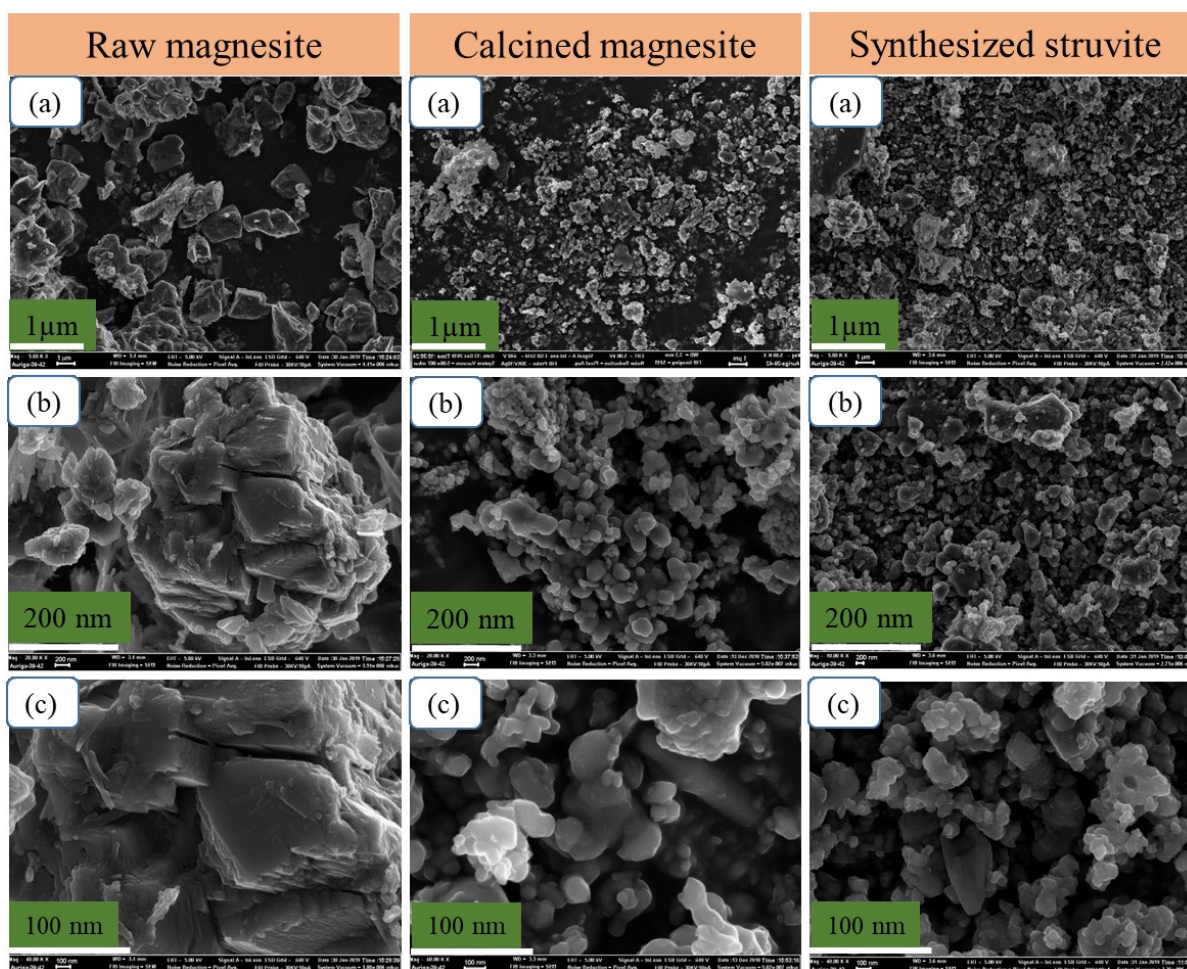
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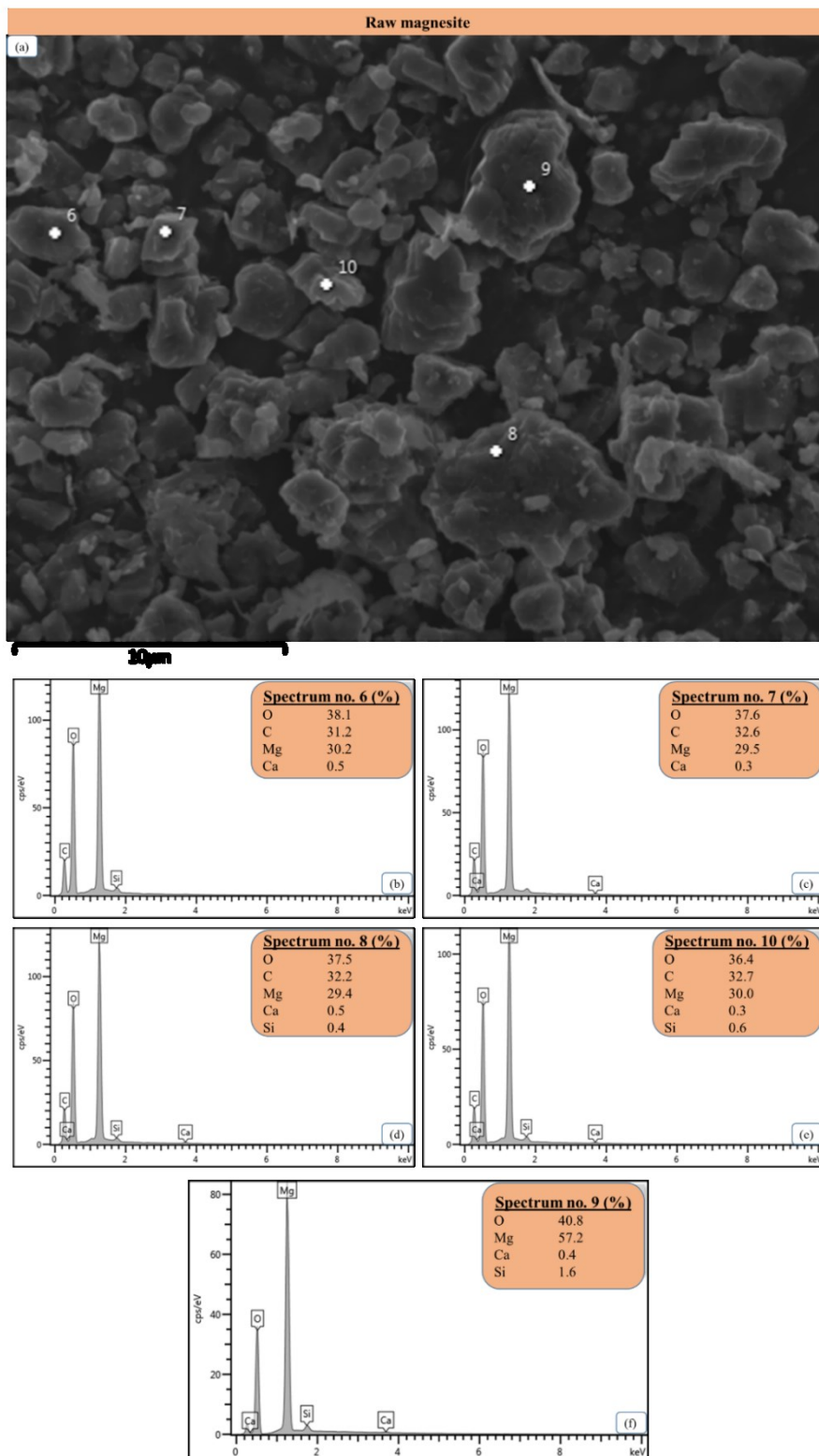
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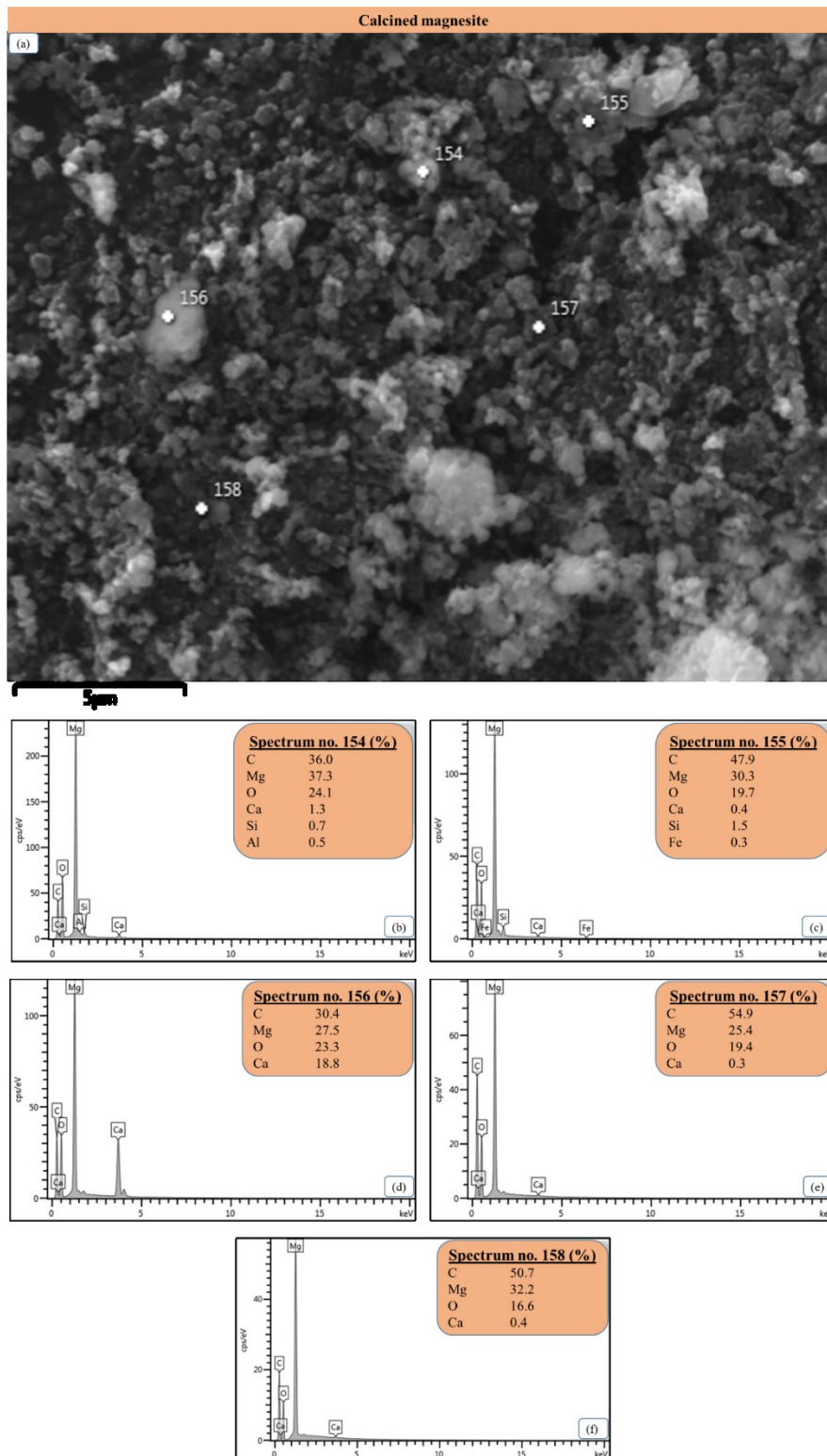
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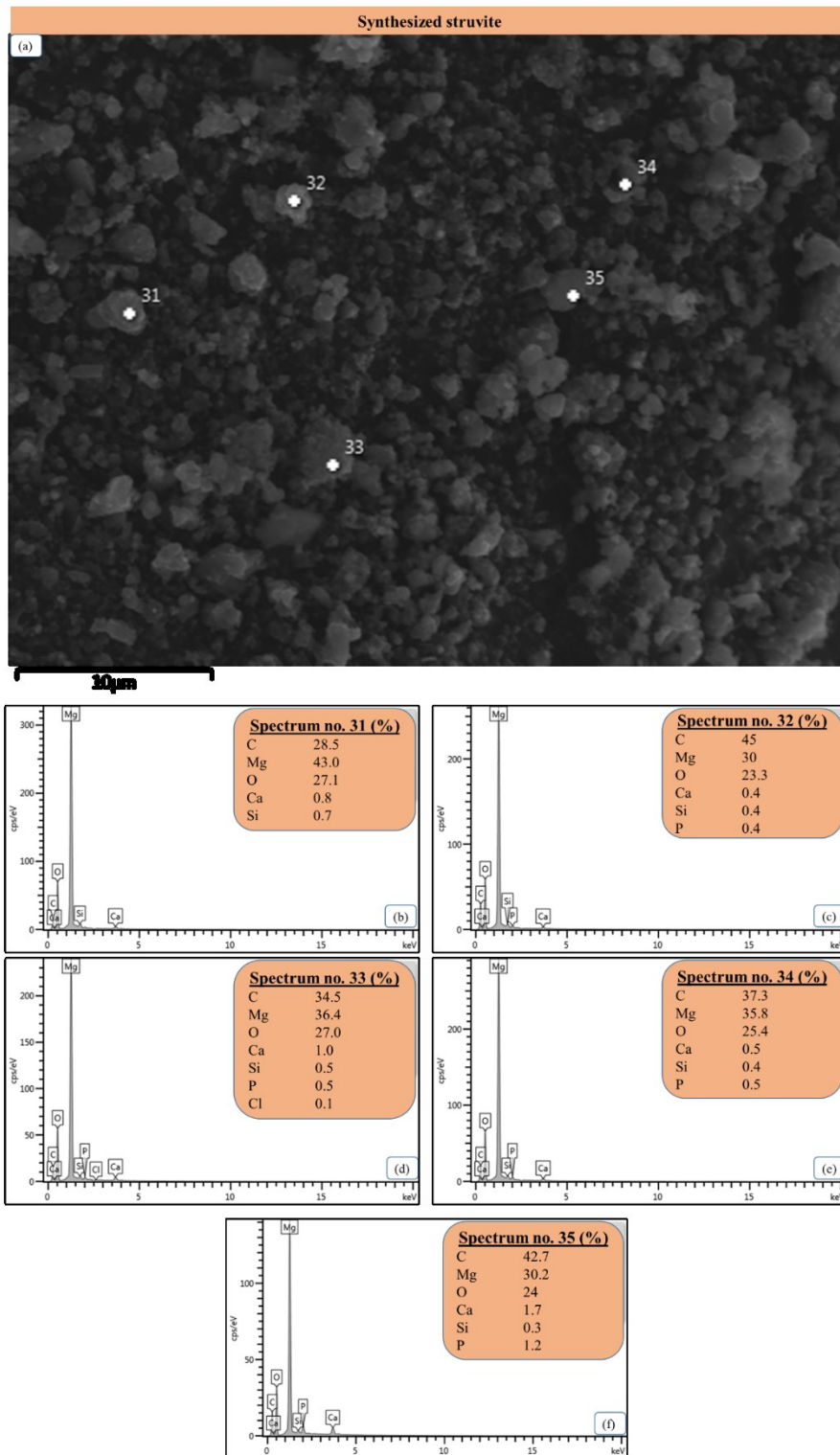


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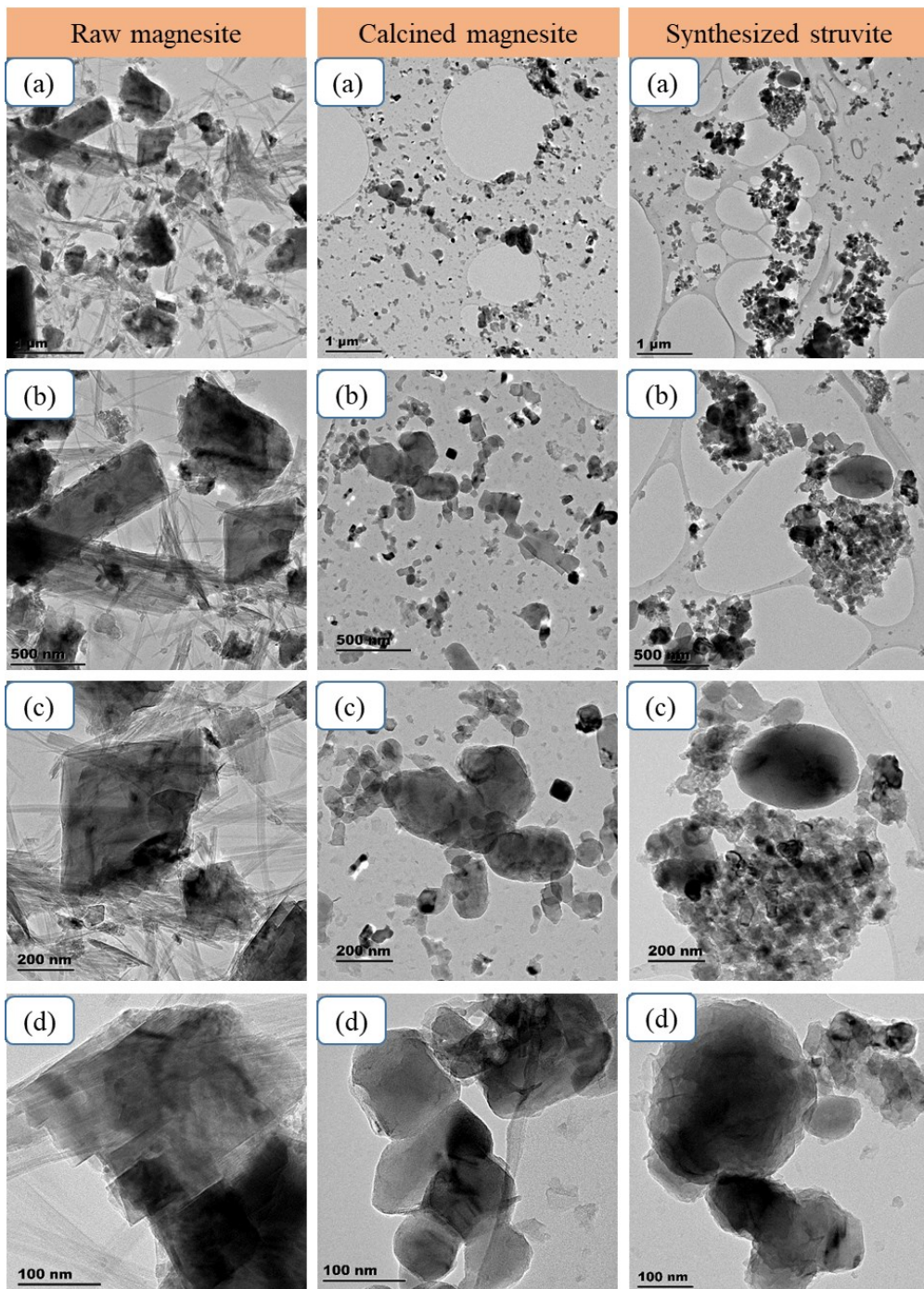


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