Green Synthesis of Spherical Calcium Hydroxide Nanoparticles in the Presence of Tannic Acid

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1. Introduction

Calcium hydroxide (CH) nanoparticle has gained extensive attentions as an eco-friendly material [1, 2] with numerous applications in the natural stone, wood, and cultural heritage conservation [2–6], wall painting [7, 8], and consolidation of artworks [9], as well as for the treatment of catalysts and adsorbents [10, 11]. Although great efforts have been made on the synthesis approaches of such nanoparticles, conventional precipitation process hardly works due to its moderate solubility [1, 7]. The assistance of high temperature and high pressure or organic solvents is to some extent essential to synthesis desired CH nanoparticles [1]. Many methods have been proposed to synthesize CH nanoparticles, including the homogeneous hydrolytic reactions [12], water-in-oil microemulsion [13], hydrogen plasma-metal reaction [7], sonochemical method, and wet chemical approaches [14, 15]. Nevertheless, more environmentally friendly and less expensive method is always more desirable for CH nanoparticle production [2].

Recently, we found that tannic acid (TA), an inexpensive plant-derived polyphenol, can be used as an effective surfactant to stabilize and facilitate the production of CH nanoparticles through a co-precipitation process at ambient temperature and pressure. TA was noticed because it not only is inexpensive and available worldwide unlimited [16] but also possesses the prominent functional properties in terms of antioxidant, anti-mutagenic, anti-inflammatorily, biodegradability, and anti-tumour activity [17, 18]. TA is molecularly composed of a central glucose core and several gallic acid monomers. The benzene hydroxyl groups within TA molecular structure give TA the inherent bonding capacity with metal cations, such as Ca$^{2+}$ through complexion reaction [19, 20]. Therefore, the nucleation and growth of CH crystals in the co-precipitation system is assumed to be influenced, decelerating the growth speed of CH crystals, which may help achieve CH precipitates in nanosizes.

Therefore, this paper was conducted to investigate the effect of TA on the morphology and crystal form of CH nanoparticles in a co-precipitation process using CaCl$_2$ and NaOH. To this end, Ca$^{2+}$ was firstly mixed with TA solution to produce sufficient complexion. Then, NaOH was added as the precipitator to produce CH precipitates. Field-emission scanning electron microscopy (SEM), transmission electron microscope (TEM), and Brunauer, Emmett, and Teller (BET) surface area analyzer were applied to characterize the morphological and geometric properties of the CH precipitates. X-ray diffractometer (XRD) and Fourier transform
**Figure 1:** XRD patterns of CH precipitates with different dosages of TA.

**Figure 2:** FTIR spectra of CH precipitates and TA.

**Figure 3:** Continued.
infrared spectrometer (FTIR) were used to study the morphological and crystal features of CH. Mechanism regarding the formation of CH nanoparticles was discussed to understand the effect of TA on the nucleation and growth of CH particles.

2. Experimental Study

2.1. Materials and Methods. CaCl$_2$, NaOH, and isopropanol of analytical purity were purchased from Sinopharm Chemical Reagent Co., Ltd. and were used as received without further purification. Analytically, pure TA was provided by Aladdin Industrial Corporation. Double deionized water was used throughout this study. The proposed synthesis route of CH was simply complied with a conventional co-precipitation process expressed in

$$\text{Ca}^{2+} + 2\text{OH}^- \rightarrow \text{Ca(OH)}_2(s)$$  \hspace{1cm} (1)

The dosage of TA was examined as the experimental variable to understand its effect on the morphology and the crystal form of CH precipitates. To this end, TA solutions with mass concentration of 0%, 0.01%, 0.03%, 0.05%, 0.1%, 0.3%, and 0.5% were made by adding TA with 100 ml double deionized water. These solutions were sealed by plastics immediately after which TA had been agitated by glass rods to be completely dissolved. Then, 2.2 g CaCl$_2$ was added into TA solutions and was vigorously stirred. These solutions were sealed and statically placed for 1 hour to ensure a sufficient complexation reaction to take place between TA and Ca$^{2+}$. Subsequently, 100 ml 0.2 M NaOH was added into the aged solutions slowly. The mixed solutions were then sealed and stirred for another 1 hour by a magnetic stirrer at the speed of 50 rpm. The final concentrations of both the CaCl$_2$ and NaOH in the solutions were the same at 0.1 M. At the predetermined time, about 300 ml isopropanol was poured into the solutions aiming to cease the nucleation or growth.

![Figure 3: Micrograms of CH precipitates. SEM images of (a) TA0, (b) TA01, (c) TA03, (d) TA05, (e) TA1, (f) TA3, and (g) TA5 and (h) TEM image of TA05.](image-url)
of CH crystals. The precipitates were vacuum filtrated and washed by isopropanol two times and double deionized water three times. Finally, the precipitates were vacuum dried for 24 hours and stored in vacuum plastic bags prior to the characterization analysis. These synthesized CH precipitates were coded as TA0, TA01, TA03, TA05, TA1, TA3, and TA5, respectively, based on the percentage concentration of the used TA solutions.

2.2. Characterization Methods. The morphology of the synthesized CH particles was studied by a FlexSEM 1000 SEM. Platinum particles sputtering for 120 s was performed to gain well electrical conductivity of the samples. A JEM-2100F TEM was applied to further examine the morphology of the produced CH nanoparticles. XRD was conducted to investigate the crystalline forms of the CH samples by using a TTR-III h/h rotating anode X-ray diffractometer with Cu_Ka radiation at 40kV and 200 mA. The scan speed and scope were, respectively, 8°/min and 2θ = 5°–55° for each specimen. Nicolet iS50 FTIR was used to further determine the composition of the CH samples. BET surface area of CH samples was also analyzed using a Micromeritics Tristar II 3020 M.

3. Results and Discussion

The effect of TA on the crystal form of CH precipitates was investigated by XRD, as presented in Figure 1. Both crystalline and amorphous CH have been identified along with a small amount of calcite, which was produced by the slight carbonation of CH samples. In general, the nucleation and growth of CH crystals have been prominently affected by TA molecules. The characteristic peak heights of CH precipitates at 2θ = 34° are found to decrease or even disappear in TA5 with increasing dosages of TA. In addition to that the residual peaks are hardly detected in TA3 and completely disappear in TA5. This suggests that the existence of TA interfered the crystallization of CH, resulting CH with poor crystallinity when more TA has been added, especially amorphous CH when 0.5% TA was used.

FTIR was applied to further reveal the role of TA in formation of CH, as shown in Figure 2. The absorption band at 3643 cm\(^{-1}\) belongs to the O-H stretching in the CH crystal [7, 21]. The minor absorption bands at 872 cm\(^{-1}\) and 1411–1439 cm\(^{-1}\) correspond to the symmetric and asymmetric stretching of CO\(_3\) group of calcite [7], indicating the slight carbonation of CH, which agrees with the XRD result. The absorption bands at 3438 and 1604 cm\(^{-1}\) are attributed to the O-H stretching and bending vibrations, respectively, implying that physisorbed H\(_2\)O molecules are linked to CH crystals [2]. Moreover, the absorption bands of TA at 1713, 1613, 1535, 1195, and 1030 cm\(^{-1}\) correspond to C=O stretch of benzene ring (1713 cm\(^{-1}\)), C–C stretch and C–H deformation in plane of benzene ring (1613 cm\(^{-1}\)), C=O stretch of carboxylic acid (1535 cm\(^{-1}\)), (O–H) deformation in plane of phenolic and carboxylic acid, (C–C) stretch and (C–H) deformation in plane of benzene ring (1195 cm\(^{-1}\)), and (C=O) stretch of carboxylic acid (1030 cm\(^{-1}\)) [22]. The absorption bands at 1195 and 1030 cm\(^{-1}\) were detected in CH samples of TA01 to TA05, suggesting the existence of residual TA in CH samples.

With the increasing dosage of TA, the absorption band at 3643 cm\(^{-1}\) of the CH precipitate decreases or disappears in TA3 and TA5, displaying a similar tendency to that of XRD results. This further confirms the impeding effect of TA on crystallization of CH. In addition, more distinct absorption peaks at 1604 cm\(^{-1}\) can be found when more TA has been used. This may indicate that CH with poor crystallinity contains physisorbed H\(_2\)O molecules in its structure.

Figure 3 shows the micrographs of CH precipitates by SEM and TEM. These figures clearly demonstrate that TA also exerted significant influence on the morphology of CH particles. The control group TA0 shown in Figure 3(a) displays prismatic microscale particles with much dense structure. This is a kind of classic morphology of the hexagonal CH crystal. When TA has been added, the CH particle becomes porous in structure. Both nanosized fibrous and spherical particles are observed, which acts as a coating layer of the dense matrix, as can be found in Figure 3(b). This is consistent with the results of XRD and FTIR as CH is in poor crystallization form when TA was presented and therefore no hexagonal CH was observed. When more TA has been added (Figures 3(c)–3(g), spherical CH nanoparticles are formed instead of the fibrous CH nanoparticles or dense micron-scale CH particles. These spherical CH nanoparticles have fairly homogeneous sizes of about 50 nm, some of which adhere to each other, which can be further confirmed by the TEM image of TA05 in Figure 3(h) as the representative.

One of the direct benefits of producing CH nanoparticles with TA solution is the enlarged BET area, as charted in Figure 4. The development tendency of the BET areas of all CH samples generally complies to a parabolic-like trend. The BET area of TA0 is just 0.487 m\(^2\)/g; nevertheless, it increases sharply to the maximum 94.087 m\(^2\)/g of TA05 samples, which is about 193 times larger than that of TA0. Slight reduced BET areas as compared to that of TA05 are found in CH samples incorporated with more TA than 0.05%.

**Figure 4**: BET area of CH precipitates.
Based on the above analysis, the formation of CH nanoparticles in the presence of TA can be viewed as a four-step process. Firstly, the phenolic hydroxyl groups in the TA molecule capture Ca\(^{2+}\) through a complexion reaction. Ca\(^{2+}\) is bound to two adjacent oxygen atoms. This process is crucial for the formation of CH nanoparticles. Normally, without TA, Ca\(^{2+}\) and OH\(^{-}\) in a co-precipitation system react freely to nucleate and grow into insoluble CH particles, as shown in Figure 5. Nevertheless, with the presence of TA, Ca\(^{2+}\) are bound to TA molecules. Consequently, the reaction between Ca\(^{2+}\) and OH\(^{-}\) is hindered due to the competitive binding between TA and OH\(^{-}\) to Ca\(^{2+}\). The binding of Ca\(^{2+}\) and OH\(^{-}\) thus become more difficult due to the bonding of Ca\(^{2+}\) by TA. In this way, the newly formed CH nucleus is much more stable, decelerating the nucleation and growth of CH particles. As a result, the growth of CH nanoparticles is inhibited, leading to nanosized CH particles with poor crystallinity.

It is of great importance to control the crystallinity and morphology of the CH precipitates since it is a major hydration products of Portland cement. Not only compacted microstructure can be gained but also a series of synergistic binding effect between CH and other hydration products, i.e., calcium silicate hydrate (CSH) gel may be triggered due to the poor crystallinity of CH in the presence of TA. This will be further investigated in our ongoing studies.

4. Conclusions

A green, low-cost, robust, and simple synthesis method for CH nanoparticles has been proposed. This method uses TA as the surfactant through a co-precipitation method of Ca\(^{2+}\) and OH\(^{-}\) under ambient temperature and pressure. It was found that TA played a pivotal role in controlling both the morphology and crystallization of the produced CH nanoparticles. With the increasing dosage of TA, the crystallinity of CH tended to be poorer to even the amorphous phase. Meanwhile, CH nanoparticles with sizes of about 50 nm have been successfully synthesized in groups containing 0.3% TA or more. This is because TA in the co-precipitation system not only acted as an ions capturer to hinder the reaction between Ca\(^{2+}\) and OH\(^{-}\) but also functioned as a layer of protective coating, leading to decelerated nucleation and growth of CH particles. This study may provide a useful hint on the controlling of the morphology and crystallinity of the CH during cement hydration so that the properties of hardened cementitious composites can be tuned.

Data Availability

Data used to support the findings of the study are available from the corresponding author upon request.

Conflicts of Interest

The authors declare no conflicts of interest.

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