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Synthesis of a Se$^0$/calcite composite using hydrothermal carbonation of Ca(OH)$_2$ coupled to a complex selenocystine fragmentation

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Abstract

In this study, the hydrothermal carbonation of calcium hydroxide under high CO$_2$-Ar pressure (90 bar) and high temperature (90 °C) coupled with a complex selenocystine fragmentation was carried out by using a batch system in order to synthesize an elemental selenium ($\text{Se}^0$/calcite composite. Under O$_2$-poor conditions, the composite was mainly characterized by red spherical selenium nanoparticles “non crystalline elemental selenium” (<500nm) deposited on the calcite matrix. In contrast, under O$_2$-rich conditions, the composite was characterized by grey rods selenium microparticles “crystalline elemental selenium” (<25µm) dispersed in the calcite matrix. The carbonate matrix was constituted by nano- and micro-rhombohedral crystals (<2µm) and micrometric agglomerates and/or aggregates (<5µm).

The results presented here demonstrate that $\text{Se}^0$/calcite composite, with spherical or rod morphologies for elemental selenium can be produced, this composite could possibly has a high potential for medical (ex. dietary supplement) or industrial (ex. pigments) applications. In addition, this study may have implications in the field of biomineralization.

Keywords: Elemental selenium, Calcite, Composite, Growth, Selenocystine, Nanoparticles, Hydrothermal method
1. Introduction

Selenium is well known for its photochemical and semiconductor properties and has been successfully used in solar cells, rectifiers, photographic exposure meters and xerography [1-3]. Selenium is also a key trace element required in small quantities in humans and animals for the function of a number of selenium-dependent enzymes, such as glutathione peroxidase (GPX) and thioredoxin reductase; however this element can also be toxic in larger doses. Both the beneficial and toxic effects of selenium are based on concentration ingested and on its chemical forms [4-10]. Inorganic and organic forms have been identified in the Nature. In addition, the selenium can easily form compounds with metals and occurs in about 50 minerals. It is present in four different oxidation states in aqueous and subsurface systems, namely -2, 0, +4 and +6 [11-13]. Obviously, the fate and transport of Se in contaminated sites are very much influenced by its chemical form and speciation. Recently, elemental selenium nanoparticles have been fabricated through various approaches, such as laser ablation, solution-phase approach, vapour-phase growth, electrochemical synthesis, photothermally assisted solution phase, ultrasonic, hydrothermal or solvothermal method and micelle-mediated synthesis [1]. Searching and designing novel methods to synthesize elemental selenium with controlled morphology is important from the viewpoint of fundamental issues and application. For example, recently, the Se\textsuperscript{0} nanoparticles are attracting more and more attention due to their excellent high biological activity and lower toxicity in animals and man. Calcium carbonate is an inorganic compound that has been widely studied due to its abundance in nature as a mineral and biomineral. Calcium carbonate particles are found in three polymorph structures, which are generally classified as rhombic calcite, needle-like aragonite and spherical vaterite. Calcite belonging to Trigonal-Hexagonal-Scalenohedral
crystallographic class is the most stable phase at room temperature under normal atmospheric conditions, while aragonite and vaterite belong to Orthorombic-Dipyramidal class and Hexagonal-Dihexagonal Dipyramidal class, respectively. The later are metastable polymorphs which readily transform into the stable calcite. The specific formation of one of the polymorphs of crystalline calcium carbonate particles depends mainly on the precipitation conditions, such as pH, temperature and supersaturation. Supersaturation is usually considered to be the main controlling factor [14]. Many experimental studies have been reported about the synthetic precipitation of the various forms of calcium carbonate and the conditions under which these may be produced, including the importance of initial supersaturation, temperature, pressure, pH and hydrodynamics. The effect of impurities and additives has also been well studied [15-29].

The above short description shows that the formation of calcite (CaCO$_3$) and elemental selenium (Se$^0$) are complex processes of considerable importance and numerous studies have been reported in the literature. The co-existence and the simultaneous precipitation/growth of these solid structures, however, have not been studied and/or reported to our knowledge. In this study, the hydrothermal carbonation of calcium hydroxide under high CO$_2$-Ar pressure (90 bar) coupled with a complex selenocystine fragmentation under O$_2$-poor and O$_2$-rich conditions was carried out by using a semi-batch system (sampling with time) in order to synthesize an elemental selenium (Se$^0$)/calcite composite.

Several analytical techniques were performed in order to characterize selected solid samples (SEM/EDS, TEM/EDS, XRD and XANES and EXAFS spectroscopy) and selected solution samples (ICP/AES and HPLC/ICPMS).
2. Materials and methods

2.1. Synthesis of Se⁰/calcite composite

(a) Under O₂-poor conditions or system with a purge step

One litre of high-purity water with electrical resistivity of 18.2 MΩ·cm, 3 g of commercial portlandite Ca(OH)₂ (calcium hydroxide provided by Sigma-Aldrich) with 96% chemical purity (3% CaCO₃ and 1% other impurities) and different quantities (0, 50, 100 or 200 mg) of seleno-L-cystine CO₂HCH(NH₂)CH₂(Se)₂CH₂CH(NH₂)CO₂H (provided by Sigma-Aldrich) with chemical purity (≥98.0%) were placed in a titanium reactor (Parr© autoclave with internal volume of two litres). The hydroxide and selenocystine particles were immediately dispersed with mechanical agitation (400 rpm). Then, at room temperature the gas argon with 99.999% chemical purity (provided by Linde Gas S.A.) was injected into the reaction cell in order to control the pressure at 90 bar during 30 minutes. After this time period, the suspension was heated to 90°C with a furnace adapted to the reactor. Obviously, during heating stage the pressure increased into the system, but it was kept constant at about 90 bar by using successive manual purge until the temperature was stabilized (about 90 minutes). Then, about 20 ml of suspension were sampled in the reactor (t=0) and immediately a flash purge was carried out until atmospheric pressure was reached. Theoretically, the argon adsorption in the suspension, the heating stage and the gas purge, allowed a partial removal of dissolved oxygen from the suspension (O₂-poor conditions). Unfortunately, the oxygen concentration was not monitored into the reaction cell. When the atmospheric pressure was reached in the reactor, 14.5 g of CO₂ with 99.995% chemical purity (provided by Linde Gas S.A.) were injected in the reactor and the total pressure in the system was immediately adjusted to 90 bar by argon injection. Under these T and P conditions, the vapour phase consists mainly of an Ar+CO₂ mixture with the CO₂ in a supercritical state (see Figure 1).
After this, a semi-batch system (sampling with time) was performed in order to measure the pH (using MA235 pH/ion analyzer) and, calcium and selenium concentration (using ICP Perkin Elmer Optima 3300 DV) in filtered solutions. For this case, about 20 ml of suspension were sampled in the reactor as a function of time (t=2, 6, 10, 30 and 60 minutes) during composite formation. Note that the pH measurement was carried out at 25 °C after filtration, cooling and degasification of the solutions. Obviously, this measurement is not representative of in situ pH behaviour, but, it can give a reasonable quantification of the solution saturation index with respect to the solid phases at standard conditions (25 °C and 1 atmosphere).

At the end of the experiment, the autoclave was removed from the heating system and immersed in cold water. The reaction cell was depressurized during the water cooling period. After water cooling at 35°C (about 15 minutes) the autoclave was disassembled, and the solid product was carefully recovered and separated by centrifugation (30 minutes at 12,000 rpm), decanting the supernatant solutions. Finally, the solid product was dried directly in the centrifugation flasks for 72 h at 65°C, manually recovered and stocked in plastic flasks.

(b) Under O2-rich conditions or system without a purge step

For these experiments, the dispersions (water-calcium hydroxide-selenocystine) cited above were directly heated to 90°C at atmospheric pressure. When the temperature was stabilized, about 20 ml of suspension were sampled in the reactor (t=0). Then, 14.5 g of CO2 were injected into the reactor and the total pressure in the system was immediately adjusted at 90 bar by argon injection. After the pressure setting, the sampling and composite recovery procedures were identical to the O2-poor system described above.

Height different experiments were carried out, four under O2-poor conditions and four under O2-rich conditions. For each experiment, six suspension samples were withdrawn from the
reactor. In addition, the experiments with 200 mg of selenocystine under O₂-poor and O₂-rich conditions were repeated three times in order to check their colour reproducibility.

2.2. Composite characterization

(a) Macroscopic observations

The coloration change with reaction time was registered by using a simple charge-coupled device camera (RICOH, Caplio R1v, 5.0M PIXELS 4.8X WIDE ZOOM).

(b) Microscopic observations and microstructure characterization

Morphological analyses of the five selected composite samples were performed by Scanning Electron Microscopy (SEM), with a HITACHI S-4800 microscope. Isolated fine particles (oriented on carbon Ni grids) of the same selected composite samples were also studied using a JEOL 3010 Transmission Electron Microscope (TEM) equipped with an energy dispersive X-ray analyzer (EDS) to image the morphology of the particles and to identify the precipitated phases.

The starting materials (Ca(OH)₂ and selenocystine) and four selected composite samples were characterized by X-ray powder diffraction using a Kristalloflex 810, SIEMENS diffractometer in Bragg-Brentano geometry. The XRD patterns were collected using Co kα₁ (λ_kα₁=1.7889 Å) and kα₂ (λ_kα₂=1.7928 Å) radiation in the range 2θ = 5 - 80° with a step size of 0.02° and a counting time of 8 seconds per step.

The selenocystine powder and three selected composite samples (water-saturated materials: pasties) were also characterized by XANES and EXAFS spectroscopy. X-ray Absorption Near-Edge (XANES) and Extended X-ray Absorption Fine-Structure (EXAFS) spectra were collected at the Rossendorf Beamline at ESRF (Grenoble, France) using a 13-element high-purity germanium detector (Canberra) together with a digital signal processing unit (XIA) for fluorescence detection. For energy calibration, a gold foil (K-edge at 11919 eV) was chosen.
because of its greater inertness and hence reliability in comparison to Se. With this approach, we determined an edge energy of 12656 eV for trigonal Se, instead of the tabulated value of 12658 eV for zerovalent Se. Dead time correction of the fluorescence signal, energy calibration and the averaging of single scans, as well as linear combination fits were performed with the software package SixPack. Shell fitting was done in WinXAS.

2.3. Physicochemical characterization of aqueous solutions

(a) Total element-concentration and pH

10 ml for all sampled suspensions were filtered through a 0.22µm pore-size filter and the obtained aqueous solutions were immediately acidified with a nitric acid solution and stored at 4°C for further measurement of [Ca] and [Se] by Inductively Coupled Plasma Atomic Emission Spectrometry (ICP Perkin Elmer Optima 3300 DV). The pH was also systematically measured at 25°C by using a MA235 pH/ion analyser in filtered solutions without acidification.

(b) Selenium speciation

10 ml for twelve sampled suspensions corresponding to two selected experiments were filtered through a 0.22µm pore-size filter and the obtained aqueous solutions were immediately stored at 4°C for further measurement (after one week) of selenocystine (SeCyst), selenomethionine (SeMet), selenite (Se(IV)) and selenate (Se(VI)) by liquid chromatography coupled with ICPMS detection (HPLC/ICPMS). The conditions for determination of SeCyst, SeMet, Se(IV) and Se(VI) species have been previously described [30]. Briefly, chromatographic separation was carried out using the Agilent 1100 series HPLC pump, equipped with an autosampler and variable volume sample loop. The analytical column was a Hamilton PRP-X-100, 10µm particle size, 25 cm length x 4.1 mm internal diameter.
The chromatographic separation of the four selenium species was obtained using a 5 mmol l\(^{-1}\) ammonium citrate buffer at pH 5.2. Injection volume was fixed at 100 µL. Low percentage of methanol (2% v/v) was added in the mobile phase to improve sensitivity. The mobile phase was delivered at 1ml min\(^{-1}\) isocratically. The HPLC-ICPMS interface consisted simply in a polyetheretherketone (PEEK) tube. Selenium selective detection was performed with an Agilent 7500ce ICPMS equipped with collision reaction cell. The instrumental and acquisition parameters were: rf power (1500 W); carrier gas (1 L/min); reaction cell gas (H\(_2\) at 5.0 mL/min). Integration time was 0.4 s for both \(m//z\) 77 and 78. Selenium species were quantified in diluted samples (1/10000) by external calibration with standard solutions of the respective pure compounds. Quantification was performed with selenium isotope \(m//z\) 78.

3. Results and discussion

3.1. Characterization of Se\(^0\)/calcite composite produced under O\(_2\)-poor conditions

Macroscopically, the typical colour of calcite is white, while the typical colours of Se\(^0\) are grey and red. In the current study, a red composite was produced under O\(_2\)-poor conditions. The red coloration depended on the selenocystine dose, for example, it was noticed that the red coloration of composite started to be observable when 50 mg/kg\(_{\text{water}}\) were used and the colour intensity increased with an increase of selenocystine dose (Figure 2). This coloration behaviour was reproducible at a constant calcium hydroxide dose (3 g/kg\(_{\text{water}}\) for this study). The coloration of composite was stable in its mother aqueous solution even after several weeks of storage at room conditions and without protection to the light.

Microscopically, the red composite was mainly characterized by spherical selenium nanoparticles (< 500 nm) deposited on the calcite matrix. The carbonate matrix was
constituted by nano- and micro rhombohedral crystals (< 2 µm) and micrometric agglomerates and/or aggregates (< 5 µm) (Figure 3).

Finally, the microstructure characterization of the red composite by using x-ray diffraction, suggests a complete Ca(OH)$_2$-calcite conversion, i.e. that the metastable crystalline phases of CaCO$_3$, such as vaterite and aragonite were not produced during the Ca(OH)$_2$ carbonation process in our experiments. Concerning the selenocystine, this crystalline organic material was not identified on the x-ray diffraction spectra for red Se$^0$/calcite composite (Figure 4). This suggests a complete chemical transformation or water dissolution of this starting solid material in our experiments and justifies the precipitation of red nano-particles of elemental selenium (observed by SEM and TEM microscopy), but, these results do not give any information on the selenocystine fragmentation mechanism. Note that red elemental selenium “spherical Se nanoparticles” was not identified by X-ray powder diffraction due to its poor crystalline property or amorphous property for this selenium form.

3.2. Characterization of Se$^0$/calcite composite produced under O$_2$-rich conditions

For these experiments, it was demonstrated that a grey composite can be produced under O$_2$-rich conditions. For this case, the grey intensity seems to be slightly dependent on the selenocystine dose from 50 to 200 mg/kg$_{water}$. The composite coloration was also stable in its mother aqueous solution even after several weeks of storage at room conditions and without protection to the light. Microscopic observations showed that the composite was characterized by rod hexagonal selenium microparticles (<25µm) dispersed in the calcite matrix (Figure 5). In addition, the X-ray diffraction measurements support these microscopic observations. It is well known that the trigonal or hexagonal morphologies are the most stable crystalline phases for Se$^0$. Here, a small proportion contained in a given composite can be identified by X-ray diffraction measurements (Figure 6). This figure also show a complete Ca(OH)$_2$-Calcite
conversion under O$_2$-rich conditions, revealing that the oxygenated conditions in the
suspension have not significant effect on the carbonation process efficiency. However, the
available quantity of oxygen in the suspension controls the fragmentation process of the
selenocystine, leading the precipitation/growth of elemental selenium with two different
morphologies and different sizes of particles. Consequently, it is demonstrated that the
fragmentation process of selenocystine under O$_2$-poor or O$_2$-rich conditions participates to the
precipitation/growth of calcite with unusual morphologies. This will be qualitatively
described in the following sub-section.

3.3. Precipitation/growth of calcite with unusual morphologies

For our experiments, the hydrothermal carbonation of calcium hydroxide described by the
global well-known reaction,

$$\text{Ca(OH)}_2(s) + \text{CO}_2(aq) \rightarrow \text{CaCO}_3(s) + \text{H}_2\text{O}$$

is an exothermic process that concerns simultaneously the dissolution of Ca(OH)$_2$,

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

and the dissociation of aqueous CO$_2$,

$$\text{CO}_2(aq) + \text{H}_2\text{O} \rightarrow \text{CO}_3^{2-} + 2\text{H}^+$$

These processes produce a fast supersaturation ($S_i$) of the solution with respect to calcite,

$$S_i = \frac{(\text{Ca}^{2+})(\text{CO}_3^{2-})}{K_{sp}} > 1$$

where (Ca$^{2+}$) and (CO$_3^{2-}$) are the activities of calcium and carbonate ions in the solution,
respectively, and $K_{sp}$ is the thermodynamic solubility product of calcite. Then, the nucleation
stage (formation of nuclei or critical cluster) takes place in the system,

$$\text{Ca}^{2+} + \text{CO}_3^{2-} \rightarrow \text{CaCO}_3(\text{nuclei})$$
The sampling with time of the suspensions in the reactor allowed the identification of the nucleation stage, characterized by the formation of an apparent stable emulsion after about two minutes of reaction time.

Finally, the crystal growth occurred spontaneously until the equilibrium calcite and the solution was reached,

\[ \text{CaCO}_3(\text{nuclei}) \rightarrow \text{CaCO}_3(\text{calcite}) \]

For this study, it was demonstrated that the selenocystine fragmentation during heating and carbonation process had insignificant effect on the Ca(OH)_2-calcite conversion, its effect on the calcite precipitation rate was not clear because the kinetic behaviour of ex-situ pH and Ca-concentration of sampled solutions were very similar for all experiments. However, several TEM micrographs revealed an unusual agglomeration/aggregation process of rhombohedral particles, this leading the unusual morphologies for calcite “ex. star-like morphologies” (see Figure 7). Recent microscopic observations show that the star-like morphologies in the solid product correspond to a physical aggregation of the crystalline nano-particles of calcite. This was attested by a simple physical treatment, i.e. by gently ground of composite product in a mortar in presence of ethanol. After this treatment, the star-like morphologies were not observed (unpublished data). For this case, the nanoparticles formation can be explained by the classic theory on the inhibition crystal growth, i.e. the surface poisoning of calcite nuclei by adsorption/incorporation of selenite (SeO_3^{2-}) and other unidentified impurities in-situ produced during selenocystine fragmentation process (see Table 1 and sub-section 3.4). In addition, recent studies in our research group shows that the incorporation of selenite (SeO_3^{2-}) into calcite retards the CO_2 transfer in the suspension during Ca(OH)_2 carbonation process (see Fig. A in the Supporting Information).
On the other hand, it is well known that impurities and additives in the solution can be modify crystal habits and could induce crystallographic twinning, which has been implicitly assumed to result from sorption of impurity/additive to the growing crystal surfaces, thus altering the growth kinetics. Recently, in-situ observations reveal that polyaspartate induces liquid-liquid phase separation of droplets of a mineral precursor. The droplets deposit on the substrate and coalesce to form a coating, which then solidifies into calcitic tablets and films. Transition bars form during the amorphous to crystalline transition, leading to sectorization of calcite tablets, and the defect textures and crystal morphologies are atypical of solution grown crystals [31-34].

3.4. Selenocystine fragmentation process

The powdered selenocystine is slightly soluble in pure water at atmospheric conditions, but a small dissolved amount increases the solution pH from 5.8 to about 8.5 due to a high proton (H\(^+\)) affinity of the two amine groups. The liquid chromatography coupled to ICPMS detection has shown that the selenocystine molecule can be conserved during several weeks in HCl acidic solution (pH=1) and preferentially at high selenocystine concentration (1 g(Se)/L) [35].

This same analytical technique revealed that selenocystine was rapidly fragmented in Ca(OH)\(_2\) alkaline solutions (pH ≅ 12.5 measured at 25°C). Here, about 25% of initial atomic selenium contained in the selenocystine (experiment with 200mg/kg\(_\text{water}\)) was oxidized to Se(IV) under O\(_2\)-poor conditions. Conversely, about 40% of initial atomic selenium contained in the selenocystine (experiment with 100mg/kg\(_\text{water}\)) was mainly oxidized to Se(IV) and a slight amount oxidized to (VI) under O\(_2\)-rich conditions (see Table 1). This partial oxidation of selenocystine taken place before the carbonation process; i.e. before the injection of CO\(_2\) in the reactor (\(t=0\)). This confirms the instability of selenocystine in alkaline solutions, in fact,
the SeCyst instability seems be controlled by the initial dose and the amount of dissolved oxygen.

Note also in Table 1 that before the CO$_2$ injection in the reactor ($t=0$), the selonocystine was slightly detected in the solution, but the kinetic data revealed a selenocystine desorption/liberation process from Ca(OH)$_2$ particles during carbonation process ($t\neq 0$), this effect being preferentially noticed under O$_2$-poor conditions. This suggests that a significant proportion of the initial SeCyst was adsorbed onto the Ca(OH)$_2$ particles. Consequently, at higher adsorbed quantity of SeCyst, i.e. under O$_2$-poor conditions, the precipitation/growth of spherical nanoparticles of red elemental selenium was favoured during the carbonation process. Conversely, at lower adsorbed quantity of SeCyst, i.e. under O$_2$-rich conditions, the precipitation/growth of hexagonal microparticles of gray elemental selenium was favoured during carbonation process.

The selenite-oxyanion produced during heating of Ca(OH)$_2$ suspension follows a complex path during carbonation process. Firstly, its solution concentration was drastically decreased by adsorption/incorporation on/in calcite matrix, leading the unusual morphologies for calcite “ex. star-like morphologies” (Figure 7). And secondly, a slight concentration increase was observed caused by a dissolution process of calcite fine particles due to a high molar excess of CO$_2$ in the system.

The adsorption and/or incorporation of selenite (SeO$_3^{2-}$) on/in the calcite were supported by XANES and EXAFS spectrometry. This powerful technique allowed also the identification and quantification of different selenium forms co-existed with calcite for three selected solid samples (labelled calcite 2, calcite 3 and calcite 5 in figures 8 and 9).

The Se K-edge XANES spectra of the three calcite samples are dominated by a white line at 12,656.5 eV, indicative of elemental Se (Figure 8.) [36-37]. Since the white line peak position of selenocystine is about 2 eV above that of elemental Se, we can discard that substantial
amounts of selenocystine is present in the samples. Samples 3 and 5 have an additional strong oscillation at 12,662 eV, which coincides with the white line of Se(IV), indicating that these samples contain tetravalent Se in addition to elemental Se.

The corresponding Se K-edge EXAFS are shown in Figure 9. The Fourier transform peaks at about 2.1 Å (uncorrected for phase shift) in the spectra of elemental Se and of selenocystine are due to Se-Se backscattering within the coordination sphere. This is also the strongest peak in the Fourier transform spectra of the three calcite samples. Red and gray elemental Se are discernable by the intensity of the 1st and 2nd shells, the ones for gray Se being much stronger due to the higher structural order. The spectrum of sample 5 is very similar to that of gray Se, while samples 2 and 3 are closer to red Se. The Se-O backscattering peak of the Se(IV) reference, Na₂SeO₃, occurs at a smaller distance of 1.4 Å. All three sample spectra show a contribution from this shell, although much lower in intensity than the pure Se(IV) compound. We can therefore conclude that all samples contain a small amount of Se(IV) adsorbed/incorporated onto/into calcite in addition to Se(0). This is in agreement with XANES results.

To further support this tentative phase identification and to derive quantitative information, we performed linear combination fits of the k³-weighted EXAFS spectra (Table 2). In confirmation of the visual interpretation of EXAFS spectra, sample calcite 5 contains a large amount of gray Se, while the other two samples are dominated by red Se. Note that satisfying fits were achieved only, when a reference spectrum of the aqueous selenite was added for samples calcite 2 and calcite 3, and the reference spectrum of a Se(IV) solid phase was added for sample calcite 5. The relative amount of Se(IV) corresponds to the relative height of the Se-O Fourier peak (Figure 9).

The sampling with time in the reactor and the use of multi-technique approach give relevant information on the selenocystine fragmentation process for our experiments, but, with this
information is still difficult to propose a clear chemical reaction mechanism. Note for example, that three transient selenium species were not identified and quantified by using HPLC/ICPMS measurements (Table 1) because only four selenium references were initially considered for this study.

4. Conclusion

In this study, the main purpose was to synthesize a Se\(^0\)/calcite composite. The synthesis was successfully performed by using the hydrothermal carbonation of calcium hydroxide under high CO\(_2\)-Ar pressure and high temperature coupled with a complex selenocystine fragmentation process under O\(_2\)-poor and O\(_2\)-rich conditions. In conclusion, under O\(_2\)-poor conditions (i.e. with purge stage), the composite was mainly characterized by spherical selenium nanoparticles (<500nm) deposited on the calcite matrix. The carbonate matrix was constituted by nano- and micro rhombohedral crystals (<2\(\mu\)m) and micrometric agglomerates and/or aggregates (<5\(\mu\)m). For this case, the spherical Se nanoparticles give a stable red coloration to the composite. In contrast, under O\(_2\)-rich conditions (i.e. without purge stage), the composite was characterized by hexagonal selenium microparticles (<25\(\mu\)m) dispersed in the calcite matrix. For this case, a gray coloration of composite was observed. It was also observed that the gas purge in the system and the selenocystine dose (mg/kg\(_{\text{water}}\)) play a crucial role on the selenocystine fragmentation mechanism during Ca(OH)\(_2\) suspension heating (at 90°C) and carbonation stages, this leading the precipitation/growth of elemental selenium with different morphologies and particle sizes. Consequently, it was noticed that the selenocystine fragmentation participates to the precipitation/growth of calcite with unusual morphologies “ex. star-like morphologies”.

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Finally, the results presented here demonstrate that Se$_0$/calcite composite, with spherical or hexagonal morphologies for elemental selenium can be produced, this composite possibly with a high potential for medical (ex. dietary supplement) or industrial (ex. pigments) applications.
Acknowledgements

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References


Table 1. The HPLC/ICPMS measurements of twelve solutions samples of two selected hydrothermal carbonation-selonocystine fragmentation experiments. Concentrations given in mg (Se) l⁻¹

<table>
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<tr>
<th>t (min)</th>
<th>SeCyst</th>
<th>SeIV</th>
<th>SeVI</th>
<th>t&lt;sub&gt;R&lt;/sub&gt; 2.80 min</th>
<th>t&lt;sub&gt;R&lt;/sub&gt; 5.91 min</th>
<th>t&lt;sub&gt;R&lt;/sub&gt; 7.03 min</th>
<th>pH&lt;sub&gt;25°C&lt;/sub&gt;</th>
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<tr>
<td>0</td>
<td>0.86 ± 0.08</td>
<td>24 ± 1</td>
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<td>ni</td>
<td>ni</td>
<td>12.60</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>nd</td>
<td>9.1 ± 0.7</td>
<td>nq</td>
<td>ni</td>
<td>12.43</td>
<td></td>
<td></td>
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<tr>
<td>6</td>
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<td>ni</td>
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<td>ni</td>
<td>7.30</td>
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<tr>
<td>30</td>
<td>4.0 ± 0.2</td>
<td>1.7 ± 0.2</td>
<td>nq</td>
<td>ni</td>
<td>7.34</td>
<td></td>
<td></td>
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<tr>
<td>60</td>
<td>nd</td>
<td>2.5 ± 0.1</td>
<td>nd</td>
<td></td>
<td>7.10</td>
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Under O₂-poor conditions and 200 mg/kg<sub>water</sub> of SeCyst dose

<table>
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<tr>
<th>t (min)</th>
<th>SeCyst</th>
<th>SeIV</th>
<th>SeVI</th>
<th>t&lt;sub&gt;R&lt;/sub&gt; 2.80 min</th>
<th>t&lt;sub&gt;R&lt;/sub&gt; 5.91 min</th>
<th>t&lt;sub&gt;R&lt;/sub&gt; 7.03 min</th>
<th>pH&lt;sub&gt;25°C&lt;/sub&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>nd</td>
<td>17 ± 1</td>
<td>0.21 ± 0.02</td>
<td>ni</td>
<td>ni</td>
<td>12.63</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>nd</td>
<td>4.7 ± 0.2</td>
<td>0.27 ± 0.01</td>
<td>ni</td>
<td>12.58</td>
<td></td>
<td></td>
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<tr>
<td>6</td>
<td>0.68 ± 0.05</td>
<td>1.5 ± 0.1</td>
<td>0.22 ± 0.2</td>
<td>8.03</td>
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<tr>
<td>10</td>
<td>0.65 ± 0.05</td>
<td>4.0 ± 0.3</td>
<td>0.27 ± 0.02</td>
<td>7.30</td>
<td></td>
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<tr>
<td>30</td>
<td>0.15 ± 0.03</td>
<td>5.6 ± 0.2</td>
<td>0.25 ± 0.02</td>
<td>7.28</td>
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<tr>
<td>60</td>
<td>nd</td>
<td>5.7 ± 0.3</td>
<td>0.24 ± 0.02</td>
<td>7.15</td>
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</tbody>
</table>

Under O₂-rich conditions and 100 mg/kg<sub>water</sub> of SeCyst dose

| t<sub>R</sub>: reaction time concerning the experiment (t=0 "before CO₂ injection in the reactor"); SeCyst: seleno-L-cystine; SeIV: selenite (SeO<sub>3</sub>²⁻); SeVI: selenate (SeO<sub>4</sub>²⁻); t<sub>R</sub>: retention time in the HPLC/ICPMS measurements; pH<sub>25°C</sub>: pH measured at 25°C; nd: not detected; nq: not quantifiable; ni: not identifiable.
Table 2. Linear combination fit of $k^3$-weighted EXAFS spectra

<table>
<thead>
<tr>
<th>Sample</th>
<th>Se grey</th>
<th>Se red</th>
<th>Se(IV)</th>
<th>Sum</th>
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<tr>
<td>Calcite 2</td>
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<td>0.87</td>
<td>0.12</td>
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<td>Calcite 3</td>
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<td>0.50</td>
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<td>0.93</td>
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<tr>
<td>Calcite 5</td>
<td>0.74</td>
<td>0.07</td>
<td>0.23</td>
<td>1.04</td>
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</tbody>
</table>

Calcite 2: experiment under O$_2$-poor conditions, 100 mg/kg$_{\text{water}}$ of selenocystine, 1 hour of reaction. Calcite 3: experiment under O$_2$-poor conditions, 50 mg/kg$_{\text{water}}$ of selenocystine, 1 hour of reaction. Calcite 5: experiment under O$_2$-rich conditions, 100 mg/kg$_{\text{water}}$ of selenocystine, 1 hour of reaction. Se(IV) correspond to Se(IV)$_{\text{aq}}$ and Na$_2$SeO$_3$ references in Charlet et al., 2007 [37].
Figure 1. Experimental P-T conditions represented on a pressure-temperature phase diagram for CO₂.
Figure 2. Coloration change of Se\(^0\)/calcite composite at different selenocystine doses and under O\(_2\)-poor conditions. Note the Se\(^0\)/calcite composite was manually dispersed to take these photographs.
Figure 3. Red Se\(^0\)/calcite composite synthesized by hydrothermal carbonation of calcium hydroxide coupled with a complex selenocystine fragmentation under O\(_2\)-poor conditions. (a) and (b) Backscattering SEM observations “without metal coating”; (c) and (d) TEM micrographs.
Figure 4. XRD measurements of starting solid materials (calcium hydroxide and selenocystine) and solid products (pure calcite and red Se⁰/calcite composite).

Hydrothermal carbonation of calcium hydroxide under O₂-poor conditions (P=200 bar; T=90 °C; Ca(OH)₂ dose=3 g/kg water; Selenocystine dose=200 mg/kg water; 1 hour of reaction time). P: Portlandite, C: Calcite.
Figure 5. Gray Se⁰/calcite composite synthesized by hydrothermal carbonation of calcium hydroxide coupled with a complex selenocystine fragmentation under O₂-rich conditions. (a) and (b) Backscattering SEM observations “without metal coating”.
Figure 6. XRD measurements of starting solid materials (calcium hydroxide and selenocystine) and solid products (pure calcite and gray Se\textsuperscript{0}/calcite composite). Hydrothermal carbonation of calcium hydroxide under O\textsubscript{2}-rich conditions (P=200 bar; T=90 °C; Ca(0H)\textsubscript{2} dose=3 g/kg\textsubscript{water}; selenocystine dose=200 mg/kg\textsubscript{water}; 1 hour of reaction time). P: portlandite, C: calcite.
Figure 7. (a), (b) and (c) Rhombohedral nano- and micro-particles of calcite; hydrothermal carbonation process without initial selenocystine dose. (d), (e), (f), (g), (h) and (i) Morphology for calcite “ex. star-like morphologies”; hydrothermal carbonation process with initial selenocystine dose (200 mg/kg$_{water}$)
Figure 8. Se K-edge XANES of three calcite samples in comparison to select references. Calcite 2: experiment under O$_2$-poor conditions, 100 mg/kg$_{\text{water}}$ of selenocystine, 1 hour of reaction. Calcite 3: experiment under O$_2$-poor conditions, 50 mg/kg$_{\text{water}}$ of selenocystine, 1 hour of reaction. Calcite 5: experiment under O$_2$-rich conditions, 100 mg/kg$_{\text{water}}$ of selenocystine, 1 hour of reaction. Se cystine correspond to the selenocystine reference.
Figure 9. Se K-edge EXAFS spectra (left) and corresponding Fourier transforms (right) of the three calcite samples and select references. Calcite 2: experiment under O₂-poor conditions, 100 mg/kg\textsubscript{water} of selenocystine, 1 hour of reaction. Calcite 3: experiment under O₂-poor conditions, 50 mg/kg\textsubscript{water} of selenocystine, 1 hour of reaction. Calcite 5: experiment under O₂-rich conditions, 100 mg/kg\textsubscript{water} of selenocystine, 1 hour of reaction. Se cystine correspond to the selenocystine reference.