PRELIMINARY RESULTS UPON CRYSTALLIZATION OF THE CALCITE-SPHEROcobALTITE SOLID SOLUTION

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Abstract

The (Ca,Co)CO₃·H₂O system was studied at 25 °C using both precipitation experiments and crystallization in silica gel in order to examine the thermodynamic properties and crystallization of the solid solution. Near the CaCO₃ endmember, unit cell parameters and volume of the solid phase show an almost linear variation, while near the CoCO₃ endmember, the growth of a phase of low crystallinity prevented those calculations. A detailed Fourier Transform Infrared Spectroscopy (FTIR) assigned this phase to a cobalt carbonate hydrate (CoCO₃·nH₂O). The fact that cell volumes are plotted above the line that corresponds to the theoretical cell volumes of the two endmembers, indicates positive value of excess volume of mixing (Vₑ), and consequently a non-ideal solid solution. Nucleation under conditions of high supersaturation in gels showed that in all the cases the degree of cobalt incorporation in crystals was low with an average value of cobalt mole fraction (XCoCO₃) around 0.03 while -even in the Co-rich zones- this value never exceeded the upper limit of 0.2. Crystal morphologies also exhibited a wide variety of forms as a function of aqueous phase composition.

Key words: solid solution, diffusion, crystal growth, free volume of mixing.

Περίληψη

Η μελέτη του στερεού διαλύματος (Σ.Δ.) ασβεστίτη-σφαφοκοβαλτίτη [(Ca,Co)CO₃] πραγματοποιήθηκε στους 25 °C μέσω πειραμάτων (i) χημικής καθίζησης από υπέρκορα υδατικά διαλύματα και (ii) κρυστάλλωσης σε gel πυριτίου, με κύριο σκοπό να εξεταστούν οι θερμοδυναμικές ιδιότητες καθώς και η κρυστάλλωση του συστήματος στη παραπάνω θερμοκρασία. Για συστάσεις του Σ.Δ. κοντά στο ακραίο μέλος του ασβεστίτη οι παράμετροι και ο όγκος (Vₑ) της μοναδιαίας κυψελίδας υπολογίστηκαν, ενώ κοντά στο ακραίο μέλος του σφαιροκοβαλτίτη η ανάπτυξη μιας φάσης χαμηλής κρυσταλλικότητας με σύσταση CoCO₃·nH₂O απέτρεψε τους παραπάνω υπολογισμούς. Επιπλέον, το γεγονός ότι οι πειραματικές τιμές των Vₑ βρίσκονται πάνω από την ευθεία που ενώνει τις αντίστοιχες θεωρητικές τιμές των ακραίων μέλων, υποδηλώνει πλεόνασμα ελεύθερου όγκου ανάμειξης (Vₑ) και συνεπώς ένα μη ιδανικό Σ.Δ.. Η κρυστάλλωση σε gel πυριτίου έδειξε ότι το κοβάλτιο συμμετείχε σε μικρό βαθμό στη σύσταση των σχηματισθέντων κρύσταλλων ενώ το μοριακό κλάσμα του σφαιροκοβαλτίτη (XCoCO₃) δεν υπερβαίνει σε καμία περίπτωση την τιμή 0.2. Τέλος, οι σχηματισθέντες κρύσταλλοι επιδεικνύουν μεγάλη ποικιλία μορφών ως συνάρτηση της συγκέντρωσης των υδατικών διαλυμάτων.

Λέξεις κλειδιά: στερεά διαλύματα, διάχυση, ανάπτυξη κρύσταλλων, ελεύθερος όγκος ανάμειξης.
1. Introduction

As calcite is one of the most abundant carbonate minerals on Earth, and cobalt a toxic metal (Lauwerys et al. 1994), the study of the (Ca,Co)CO₃ solid solution is of high importance. Although there are different works published, referring to morphological modification of calcite crystals doped by cobalt (Fernández-Díaz et al. 2006, Braybrook et al. 2002), or to calcite growth and dissolution in the presence of dissolved cobalt (Freij et al. 2004), the solid solution is not fully studied yet. The only available thermodynamic data comes from Goldsmith and Northop (Goldsmith et al. 1965) who studied the ternary system CaCO₃-MgCO₃-CoCO₃ by delimiting basic relations at 600-750 °C and 15 kilobars.

In this work, the interaction of Co(2+) with calcite is investigated, having as main intentions to characterize the solid solution (ideal or non-ideal), to evaluate the way in which composition at nucleation depends on supersaturation, and finally to examine the chemical and morphological evolution of crystals during growth.

2. Experimental Methods

2.1. Precipitation Experiments

Small crystals of the (Ca,Co)CO₃ solid solution were precipitated according to the following reaction:

\[ \text{Na}_2\text{CO}_3(\text{aq}) + x\text{CoCl}_2(\text{aq}) + (1-x)\text{CaCl}_2(\text{aq}) \rightleftharpoons \text{Co}_x\text{Ca}_{1-x}\text{CO}_3(s) + 2\text{Na}^+(\text{aq}) + 2\text{Cl}^-(\text{aq}) \]

In order to achieve precipitation of different compositions of the solid solution, 50ml of solutions with different ratios of CaCl₂ and CoCl₂ (Table 1) were used. The mother solutions were added in a vessel that contained 50 ml of a continuously stirred solution of 0.05M Na₂CO₃. The precipitates were then separated from the solution using a filter and dried in an oven at approximately 39 °C. By using adhesive carbon tape, a small quantity of crystals was taken from each precipitate and mounted on an aluminium sample holder. Finally all the samples were covered with carbon. The rest of the precipitates were powdered so they could be examined by X-ray powder diffraction.

<table>
<thead>
<tr>
<th>experiment</th>
<th>CaCl₂ (M)</th>
<th>CoCl₂ (M)</th>
<th>Ca²⁺/Co³⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td>P-1</td>
<td>0.05</td>
<td>0</td>
<td>Ca</td>
</tr>
<tr>
<td>P-2</td>
<td>0.05</td>
<td>0.001</td>
<td>50:1</td>
</tr>
<tr>
<td>P-3</td>
<td>0.05</td>
<td>0.003</td>
<td>17:1</td>
</tr>
<tr>
<td>P-4</td>
<td>0.05</td>
<td>0.005</td>
<td>10:1</td>
</tr>
<tr>
<td>P-5</td>
<td>0.05</td>
<td>0.007</td>
<td>7:1</td>
</tr>
<tr>
<td>P-6</td>
<td>0.05</td>
<td>0.01</td>
<td>5:1</td>
</tr>
<tr>
<td>P-7</td>
<td>0.05</td>
<td>0.02</td>
<td>5:2</td>
</tr>
<tr>
<td>P-8</td>
<td>0.05</td>
<td>0.03</td>
<td>5:3</td>
</tr>
<tr>
<td>P-9</td>
<td>0.05</td>
<td>0.05</td>
<td>1</td>
</tr>
<tr>
<td>P-10</td>
<td>0.03</td>
<td>0.05</td>
<td>3:5</td>
</tr>
<tr>
<td>P-11</td>
<td>0.01</td>
<td>0.05</td>
<td>1:5</td>
</tr>
<tr>
<td>P-12</td>
<td>0</td>
<td>0.05</td>
<td>Co</td>
</tr>
</tbody>
</table>

Crystal phases precipitated were characterized by X-ray powder diffraction analysis that was carried out on a Philips X’Pert Pro X-ray diffractometer in the 2θ angle range between 5° to 80° in...
steps of 0.02° using Cu Kα radiation. The study of the main reflections of diffractograms and the calculation of the unit cell parameters of each precipitate was made using the computer program X'Pert Plus. Characterization of low-crystallinity phases achieved by Fourier Transform Infrared Spectroscopy (FTIR) carried out on a Jobin-Yvon Horiba 800 UV LabRam-HR spectrometer equipped with a liquid nitrogen-cooled detector. A Golden Gate ATR Mk II modified attenuated total reflection system was also used. The spectral region spanned from 4000 to 600 cm⁻¹ with a resolution of 4 cm⁻¹. Experimental results were then compared to the spectra of (i) a natural calcite (Island Spar), and (ii) a commercial CoCO₃·Co(OH)₂·nH₂O substance.

2.2. Crystallization in Gels

In that group of experiments the reactants were introduced in two vertical reservoirs separated by a column of silica hydrogel, which occupied the horizontal branch between them, constituting in such a way a double diffusion system (U-tube), as shown in Figure 1. The silica gel was prepared by acidification of a Na₂SiO₃ solution (Merck, sp. gr.: 1.059 g/cm³) with HCl (IN) until a pH = 5.5 was obtained. The gel -due to polymerization- behaves as a porous medium (Dullien 1979), suppresses convection and advection of the reactants, which can only move through the gel, and eventually mix in a column profile by molecular diffusion. Because of the physical and chemical properties of the gel (micron-sized pores, 95.6 wt% “effective water” (Prieto et al. 2002)), the nucleation density decreases and consequently larger crystallite form. Three sets of experiments were carried out, using tubes with different lengths (Table 2), and parent solutions with different initial concentrations. In all cases one reactant reservoir contained 8 ml of 0.5M Na₂CO₃ and the other 8ml of mother solutions with different ratios of CaCl₂ and CoCl₂ (Table 3).

Table 2 - Length of tubes used in the experiments and length of gel column in the horizontal branch

<table>
<thead>
<tr>
<th>Exp.</th>
<th>tube characterization and length (cm)</th>
<th>gel column length (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1-S10</td>
<td>Small: 15</td>
<td>7-8</td>
</tr>
<tr>
<td>I1-I10</td>
<td>Intermediate: 25.7</td>
<td>16-17</td>
</tr>
<tr>
<td>L1-L10</td>
<td>Large: 35</td>
<td>26-28</td>
</tr>
</tbody>
</table>

Table 3 – Initial concentrations of reactants used in gel experiments. Na₂CO₃ concentration in all the experiments is 0.5 moles/l. Total waiting period until nucleation and distance of primary nucleation zone measured from the side of cations are also shown

<table>
<thead>
<tr>
<th>initial concentrations</th>
<th>small U-tubes set</th>
<th>int/dilate U-tubes set</th>
<th>large U-tubes set</th>
</tr>
</thead>
<tbody>
<tr>
<td>CaCl₂(M)</td>
<td>CoCl₂(M)</td>
<td>tₜ(h)</td>
<td>position(cm)</td>
</tr>
<tr>
<td>0.5</td>
<td>0</td>
<td>46</td>
<td>4</td>
</tr>
<tr>
<td>0.5</td>
<td>0.01</td>
<td>70</td>
<td>5</td>
</tr>
<tr>
<td>0.5</td>
<td>0.03</td>
<td>66</td>
<td>5</td>
</tr>
<tr>
<td>0.5</td>
<td>0.05</td>
<td>40</td>
<td>4</td>
</tr>
<tr>
<td>0.5</td>
<td>0.1</td>
<td>43</td>
<td>4</td>
</tr>
<tr>
<td>0.5</td>
<td>0.3</td>
<td>47</td>
<td>4.5</td>
</tr>
<tr>
<td>0.5</td>
<td>0.5</td>
<td>50</td>
<td>4.5</td>
</tr>
<tr>
<td>0.3</td>
<td>0.5</td>
<td>50</td>
<td>4.5</td>
</tr>
<tr>
<td>0.1</td>
<td>0.5</td>
<td>50</td>
<td>4</td>
</tr>
<tr>
<td>0</td>
<td>0.5</td>
<td>no crystal phase</td>
<td>no crystal phase</td>
</tr>
</tbody>
</table>
magnification) were then monitored. The total waiting period \( t_w \), from the beginning of the experiment, until the first crystallites appeared was noted, and the position of the nucleation area was measured (Table 3). The experiments were stopped two months after nucleation observed in the intermediate and small U-tubes, and one month after nucleation observed in the large U-tubes. The crystals were finally recovered by dissolving the gel in a 1M NaOH solution.

![Diagram of gel column with cations and anions](http://epublishing.ekt.gr)

**Figure 1** - A double diffusion system arrangement used in the gel experiments. The solutions that contained different ratios of Ca\(^{2+}\) and Co\(^{2+}\) were introduced in the “cations” side reservoir while the Na\(_2\)CO\(_3\) solutions in the “anions” side

The total concentrations of the reactants through different profiles of the gel column were calculated using a one-dimensional algorithm (Henisch et al. 1986) suitable for diffusion in finite systems:

**Equation 1** - Concentrations through the gel column

\[
C(d,t + \Delta t) = \frac{1}{6} \left[ C(d - \Delta d,t) + 4C(d,t) + C(d + \Delta d,t) \right]
\]

where \( C \) is the concentration, \( d \) is the position in the silica gel column, \( t \) is the diffusion time. This equation is valid when \( \Delta t \) is chosen to be related to \( \Delta d \) by:

**Equation 2** – Relation between time and position intervals in Equation 1

\[
\Delta t = \frac{1}{6D_w \phi \tau^2} \Delta d^2
\]

where \( D_w \) is the diffusion coefficient of the reactant in water, \( \phi \) is the porosity, and \( \tau \) the effective tortuosity of the gel (Lichtner et al. 1996). The characteristics of the silica gel are \( \phi = 0.969 \) and \( \tau = 1.19 \) (Andara 2004). The diffusion coefficients of the reactants Na\(_2\)CO\(_3\), CaCl\(_2\), and CoCl\(_2\) in water (\( D_w \)) were calculated and found 1.193x10\(^{-5}\), 1.336x10\(^{-5}\), and 1.227x10\(^{-5}\) cm\(^2\)s\(^{-1}\), respectively (Robinson et al. 1959). In that way, the total concentrations of the three reactants were calculated at \( t = t_w \) for the nucleation zone in the gel column, and the solution’s speciation was made using PHREEQC v2.12.5.669.
3. Results and Discussion

3.1. Precipitation Experiments

An X-ray powder diffraction analysis carried out for precipitation experiments showed the growth of a phase of low crystallinity from aqueous solutions high in cobalt (Fig. 2). This phase dominates in the precipitates preventing the definition of unit cell parameters from the corresponding diffractograms (Fig. 3). In order to reduce the spread of the low-crystallinity phase, all sequence of experiments repeated using NaHCO₃ instead of Na₂CO₃ as a reactant, but also in that case, the degree of crystallinity from aqueous solutions high in cobalt was proved very low. Moreover, precipitation experiments of the (Ca,Co)CO₃ solid solution were carried out in the atmosphere of N₂, using the same reactants, and initial concentrations as previous, but the diffraction analysis of those samples showed that the growth of the low-crystallinity phase could not be prevented. Therefore, all experimental procedures in the author’s knowledge are resulting to the suggestion that although it is achievable to obtain precipitation of spherocobaltite at hydrothermal conditions, mainly through autoclave techniques (Gamsjager et al. 1980, Egorov et al. 1976, Ikornikova et al. 1975), it is not possible to precipitate cobalt carbonate (CoCO₃) at room temperature (25 °C) and pressure (1 Atm).

Figure 2 - Two representative SEM images of two different precipitation experiments. (A) Low concentration of cobalt in initial solution (Ca²⁺:Co²⁺=17:1) where crystallinity is dominating on precipitates surface, and (B) precipitates from solution with Ca²⁺:Co²⁺=5:1, in which crystals are partially covered by the low-crystallinity phase

The low-crystallinity phase that precipitated in experiments within a wide Co²⁺/Ca²⁺ range of aqueous solutions was characterized by FTIR spectroscopy. In order to study the evolution and the chemical composition of this phase, a group of 6 different precipitates were analyzed. The analysis of the samples revealed a concurrent precipitation of crystalline (Ca,Co)CO₃ solid phases with a Co-rich matter of low crystallinity that was designated to a cobalt carbonate hydrate (CoCO₃·nH₂O).

The mole fraction of the crystal phase near the calcium carbonate endmember was determined using the INCA Energy 200 microanalysis system of the scanning electron microscope. For initial solutions high in calcium, the measured solid phase composition did not vary more than ±0.05 XCoCO₃. In experiments high in cobalt this value increased up to ±0.1 XCoCO₃ because of the presence of the low-crystallinity phase (Fig. 4).
Table 4 – Unit cell parameters of the precipitates obtained near the Ca-rich side of the solid solution. Numbers in parenthesis indicate the standard deviation

<table>
<thead>
<tr>
<th>experiment</th>
<th>XCoCO₃</th>
<th>α = b (Å)</th>
<th>c (Å)</th>
<th>Vcell (Å³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P-1</td>
<td>0</td>
<td>5.003(1)</td>
<td>17.078(6)</td>
<td>370.2(2)</td>
</tr>
<tr>
<td>P-2</td>
<td>0.17</td>
<td>5.002(1)</td>
<td>17.063(6)</td>
<td>369.8(2)</td>
</tr>
<tr>
<td>P-3</td>
<td>0.21</td>
<td>4.9888(9)</td>
<td>17.024(7)</td>
<td>366.9(2)</td>
</tr>
<tr>
<td>P-4</td>
<td>0.28</td>
<td>4.9876(4)</td>
<td>17.017(3)</td>
<td>366.6(1)</td>
</tr>
<tr>
<td>P-5</td>
<td>0.31</td>
<td>4.9824(6)</td>
<td>16.980(5)</td>
<td>365.1(2)</td>
</tr>
<tr>
<td>P-6</td>
<td>0.37</td>
<td>4.972(1)</td>
<td>16.886(6)</td>
<td>361.5(2)</td>
</tr>
<tr>
<td>P-7</td>
<td>0.46</td>
<td>4.9691(1)</td>
<td>16.866(6)</td>
<td>360.6(2)</td>
</tr>
</tbody>
</table>

For solid phase compositions near the CaCO₃ endmember, the X-ray diffraction patterns were indexed in the R₃c space group of the Hexagonal-Scalenohedral class and the cell parameters were calculated (Table 4) using the computer program X’Pert Plus version 1.0. The solid phase precipitated as pure calcium carbonate clearly corresponds to calcite (P-1 experiment), while the rest of the patterns exhibit a continuous shift towards higher 2θ angles as cobalt mole fraction increases. As expected, for different solid solution compositions, cell parameters and volume decrease with increasing Co content (Figure 5). This tendency is in agreement with the smaller radius of Co²⁺ (0.72Å) compared to that of calcium Ca²⁺ (0.99Å).
3.2. Gel Experiments

One of the main features of silica hydrogel experiments is the high supersaturation obtained, allowing the study of system behaviour under nonequilibrium conditions. In relation to precipitation experiments, larger crystals were formed allowing (1) the definition of crystal’s chemical evolution and (2) the description of different crystal forms that could be possibly developed.
Figure 6 - A SEM-BSE image showing a solid extracted from a gel experiment with Ca\(^{2+}\):Co\(^{2+}\)=1:1.7 in initial solution. As shown in the compositional profile at the right, along the line A-B, crystals exhibit a clear compositional zoning with two cobalt-rich cores surrounded by a calcium-rich area.

Figure 7 - Scanning electron microscope images showing the different morphologies obtained for different initial concentration of calcium and cobalt. Ratio of Ca\(^{2+}\)/Co\(^{2+}\) in initial solution decreases from 10:1 in (A), to 1:1 in (B), and finally to 1:5 in (C).

In all the experimental sets followed, crystals are enriched in Ca with the mole fraction of calcium carbonate (X\(\text{CaCO}_3\)) always above 0.95, as calculated using the SEM-EDS analysis. Referring to experiments with initial concentration of Ca equals to zero, no crystals were observed in gel column at the time at which the experiments stopped. As cobalt incorporates in a small proportion into the lattice of formed CaCO\(_3\), even if it is present in high concentrations in aqueous solutions, the existence of a miscibility gap between the end-members of the solid solution is of high possibility. This comes in agreement with previous studies of this system. According to Palache (Palache et al. 1951), in natural solids the range of Co\(^{2+}\) miscibility in calcite varies around 2.7% at 50 °C. In the same work, the excess free energy of mixing parameter was estimated to be around 4.12, which indicates a positive excess free energy of mixing and suggests the presence of a
miscibility gap in the system. Also, Goldsmith et al. (1965), studying the ternary system CaCO$_3$-
MgCO$_3$-CoCO$_3$ found that at 600 °C, about 5 mol % CoCO$_3$ is soluble in calcite.

Despite the fact that the composition of the solid solution is very close to pure calcite, some of the
crystals are revealing a compositional profile with a clear compositional zoning: a CoCO$_3$-rich
core surrounded by CaCO$_3$-rich regions (Fig. 6). This fact can be easily explained by the solubility
product constants of the two endmembers. Spherocobaltite (logKsp = -9.98) is about one and a
half order of magnitude less soluble than calcite (logKsp = -8.47), thus cobalt-rich carbonate
crystallizes before CaCO$_3$ occupying the central part of the crystals. In any case, mole fraction of
CoCO$_3$ oscillates around 0.05, and although it increases in Co-rich sections, it never exceeds the
value of 0.2 supporting the hypothesis of a wide miscibility gap in the system.

Examination of the crystals grown in the gel experiments through a scanning electron microscope
shows a wide variety of crystal morphology as a function of different Ca$^{2+}$/Co$^{2+}$ ratios in initial
solutions. Blocky hexagonal prismatic and needle-like crystals are forming spherulites with the
presence of rhombohedra for high ratios of Ca$^{2+}$/Co$^{2+}$ (Fig. 7A). For equal concentrations of Ca$^{2+}$
and Co$^{2+}$ in the initial solution, crystal morphologies are dominated by rose-like multicrystals (Fig.
7B). Crystallization from solution high in Co$^{2+}$ and low in Ca$^{2+}$, produces also spherulites of
broken or not very well formed crystals (Fig. 7C).

4. Conclusions

Precipitation experiments carried out in room temperature and pressure only produced crystals
near the CaCO$_3$ endmember of the solid solution. For Ca$^{2+}$/Co$^{2+}$ values below 17:1 a cobalt-rich
phase of low crystallinity was precipitating, and near the CoCO$_3$ endmember was dominating in
precipitates preventing the definition of mole fraction and unit cell parameters at this side of the
solid solution. An FTIR spectroscopy assigned the low-crystallinity matter to a cobalt carbonate
hydrate (CoCO$_3$·nH$_2$O). Crystals of the solid solution that were grown under conditions of
supersaturation in gels were Ca-rich (XCoCO$_3$=0.03) with a Co-rich core (XCoCO$_3$=0.15-0.2).
The crystal morphologies developed in those experiments exhibit a wide variety of forms
(spherulites, rose-like multicrystals etc.) depending mainly on initial solution concentration. No
crystals were extracted from the gel experiments referring to the pure CoCO$_3$ endmember of the
solid solution.

The presence of a miscibility gap in the system is the key factor in order to characterize the solid
solution. Therefore, it is obligatory to study the thermodynamic properties of the system, and this
can be achieved only by eliminating the presence of the low-crystallinity phase near the Co-rich
side of the solid solution. Crystals that have grown in gel exhibit various morphologies that
possibly correspond to the other two polymorphs of CaCO$_3$, that is to say aragonite and vaterite.

5. Acknowledgments

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Sustitución Aniónica. Sistemas BaSeO$_4$ - BaSO$_4$ - H$_2$O, CaSeO$_4$ - CaSO$_4$ - H$_2$O y CaMoO$_4$
- CaWO$_4$ - H$_2$O, Ph.D. Thesis, Universidad de Oviedo.


