Article

Precipitation of CaCO₃ polymorphs from aqueous solutions: the role of pH and sulphate groups

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Abstract: In this work we aim to experimentally study the nucleation and growth of CaCO₃ phases precipitated from supersaturated aqueous solutions in the presence of varying concentrations of sulphate oxyanion. The experiments were conducted under pH conditions close to neutral (7.6) and considering a wide range of initial (SO₄²⁻)/(CO₃²⁻) ratios (0 to ~ 68) in the aqueous solution. We paid special attention to the evolution of the precipitates during ageing within a time framework of 14 days. The mineralogy, morphology and composition of the precipitates were studied by X-ray diffraction, Fourier transform infrared spectroscopy, scanning electron microscopy and EDX microanalysis. The concentration of sulphate ions in the reacted aqueous solution was study by ICPs. The experimental results show that the mineral composition of the precipitate recovered in each run varied with the (SO₄²⁻)/(CO₃²⁻) ratio in the parental solution, which influences the mineral evolution of the precipitates during ageing. We observe that high concentrations of sulphate in the aqueous solution stabilize the vaterite precipitates and inhibit calcite formation. Furthermore, aragonite never precipitates directly form the solution and it is only formed via a dissolution-precipitation process in solutions with high (SO₄²⁻)/(CO₃²⁻) ratio after long reaction times. Finally, gypsum only precipitates after long ageing in those aqueous solutions with the highest concentration of sulphate. The reaction pathways during ageing, the morphology of the calcite crystals and the composition of vaterite and calcite are discussed considering both, kinetic and thermodynamic factors. These results show a considerably more complex behavior of the system than that observed in experiments conducted under higher pHs and supersaturation levels and lower (SO₄²⁻)/(CO₃²⁻) ratios in the aqueous phase.

Keywords: CaCO₃; polymorphs; sulphate; ageing process; aragonite; gypsum

1. Introduction

The presence of calcium carbonate mineral phases is ubiquitous in surface and subsurface environments in the Earth. Calcium carbonate phases in such environments range from amorphous to crystalline and from anhydrous (calcite, aragonite and vaterite, CaCO₃) to differently hydrated (monohydrocalcite, CaCO₃·H₂O, and ikaitie, CaCO₃·6H₂O) [1]. In these environments, the only stable phase of calcium carbonate is calcite, which appears as the main constituent of sedimentary carbonate rocks [1-2] and hard tissues in marine organisms [2-4]. However, all other phases can also form and often remain metastable over long periods due to kinetic factors and/or the influence of specific chemical species in a variety of bio-geological contexts [5-7]. Thus, amorphous calcium carbonate (ACC) plays a main role in the early stages of the formation of carbonate hard tissues, where it forms at far from equilibrium conditions and, following the Ostwald’s step rule, subsequently transforms into other calcium carbonate phases through different reaction pathways [8-10]. In carbonate hard
tissues ACC can also remain stabilised in specific locations under the influence of organic molecules and/or inorganic foreign ions like magnesium [11] as a sort of calcium carbonate storage which can eventually be disposable for hard tissue-damage reparation [12]. The appearance of the hydrated forms of calcium carbonate ikaite and monohydrocalcite is restricted to very specific environments like near-freezing conditions aqueous environments, where the presence of magnesium or phosphate ions has been proposed to promote their formation [13-14]. Kinetics effects facilitate the formation of vaterite in natural environments, where this phase represents an intermediate step in the reaction pathway that leads from ACC to calcite following Ostwald’s steps rule [15]. The seasonal formation and temporal stabilization of vaterite in evaporite-rich geological settings has also been linked to the influence of high concentrations of dissolved sulphate [16]. Finally, aragonite is the second most abundant calcium carbonate phase in surface and subsurface environments after calcite [17]. Aragonite also is a main component of carbonate biominerals, often coexisting with calcite in the same hard tissue [4,18-19]. The main role played by high Mg\(^{2+}/Ca^{2+}\) ratios in aqueous environments as the determining factor in the crystallization and stabilization of aragonite has been long known [5]. This role has been attributed to the inhibitory effect of dissolved magnesium on the growth of calcite [20-23]. Changes in Mg\(^{2+}/Ca^{2+}\) ratio in seawater have been proposed to explain recurrent calcite-aragonite switches in marine calcium carbonate biominerals along the Earth’s history [24]. Consequently, the influence of different foreign ions in the crystallization of calcium carbonate and, specifically, in its polymorph selection, has long been a main topic of research in carbonates mineralogy. Understandably, most research efforts have been focused on shedding light on the role of magnesium and, to a lesser extent, that of other cations like Sr\(^{2+}\) or Co\(^{2+}\) [5,23,25-27]. However, much less attention has been paid to the influence of different anions on calcium carbonate polymorph selection, even though there exist strong evidences that this influence, especially that of tetrahedral oxyanions, could also be most relevant [28]. Indeed, as mentioned above, phosphate oxyanions have been linked to the formation of ikaite [13]. A combined role of sulphate and Mg\(^{2+}\) in the switch from calcite to aragonite seas has been claimed on both, geochemical and experimental evidences [29]. The notion that sulphate (SO\(_4^{2-}\)), chromate (CrO\(_4^{2-}\)) and selenate (SeO\(_4^{2-}\)) promote the formation of vaterite and inhibit its transformation into calcite is supported by overwhelming experimental evidence [30-38]. Furthermore, this notion has been given theoretical support by computational studies that relate the effect of tetrahedral oxyanions on calcium carbonate polymorph selection to thermodynamic effects arising from the different impact that their incorporation substituting carbonate groups has in the structure of the three calcium carbonate polymorphs calcite, aragonite and vaterite [28,31]. This incorporation produces a most significant disruption of the local structure of aragonite. Although less so, this disruption still is very marked in calcite, while vaterite structure is the least disturbed by the isomorph incorporation of tetrahedral oxyanions.

In an earlier work, the specific influence of the presence of sulphate oxyanions in the crystallization medium on the formation and evolution upon ageing of calcium carbonate precipitates was studied at high pHs (10.9) by conducting batch experiment [31]. High pH conditions are restricted to limited geological settings such as hyper alkaline lakes in connection to tectonic rifts [39]. Even though other authors have studied the influence of sulphate ions on the precipitation of calcium carbonate under lower pHs [37-38], these studies only explored a narrow range of SO\(_4^{2-}/CO_3^{2-}\) ratio conditions and a comprehensive study of close to neutrality conditions is still missing. In this study we aim to improve our current understanding of the influence of sulphate oxyanions in calcium carbonate formation and evolution by filling this gap. We present here the results of precipitation experiments conducted under circa neutral conditions (initial pH = 7.6), closer to those commonly found in surface and subsurface waters, and considering a wide range of initial SO\(_4^{2-}/CO_3^{2-}\) ratios (0 to ∼68) in the aqueous solution. Furthermore, we study the evolution of the precipitates upon ageing within a time framework of 14 days, regarding its mineralogy and the morphology and composition of the different phases that constitute the precipitate at each stage. Different reaction pathways during ageing are unraveled depending on the initial (SO\(_4^{2-}\))/CO\(_3^{2-}\) ratio in the solution. These pathways are discussed considering both, kinetic and thermodynamic arguments. Our goal is to draw general
conclusions on the influence of sulphate oxyanions and the (SO₄²⁻)/(CO₃²⁻) ratio in aqueous environments in the polymorphic selection of calcium carbonate.

2. Materials and Methods

Stock solutions of CaCl₂ (Fluka, 1M), NaHCO₃ (99.7%, Panreac) and Na₂SO₄ (>99%, Sigma-Aldrich) were prepared with deionized water (Millipore, resistivity = 18.2 MΩ·cm). CaCO₃ precipitates were obtained by mixing 50 mL of a CaCl₂ solution with stirred 50 mL of a NaHCO₃ + Na₂SO₄ solutions (SO₄²⁻)/(CO₃²⁻) = 0 - 67.6) in Erlenmeyer flasks (Pyrex). The flasks were immediately closed and kept at 25 ± 0.5 °C while continuously stirred by means of a magnetic stirrer. PHREEQC [40] was used to calculate solution speciation and supersaturation with respect to the different solid phases that can form in the system using the phreeqc.dat database. This database includes a Ksp of 10⁻⁸.48 for calcite, of 10⁻⁸.34 for aragonite, of 10⁻⁴.58 for gypsum and of 10⁻⁴.38 for anhydrite. Ksp of 10⁻⁷.91 for vaterite [41] and of 10⁻⁷.0 for ACC [42] were manually added to this database.

Table 1 shows the initial concentration of the reactants and calculated pH, activities of the relevant species and saturation indexes (SI) with respect to calcite, aragonite, vaterite, gypsum and anhydrite. In all the experiments solutions were initially highly supersaturated with respect to the three CaCO₃ polymorphs. In order to maintain the supersaturation at an approximately constant value with respect to the three CaCO₃ polymorphs (SIcalc ~ 2.35, SIArg ~ 2.2 and SILTP ~ 1.8), CaCl₂ concentrations were adjusted. Only solutions in experiments A15 to A25 were supersaturated with respect to gypsum (SIgyp ≤ 0.25). In all experiments, solutions were undersaturated with respect to anhydrite. It is important to note that the mentioned SI values are those that would be reached if mixing were instantaneous and no crystallization occurred before the complete homogenization of the solution were reached after mixing. This is an unlikely situation and it can be assumed that some CaCO₃ precipitation will take place before complete mixing of the reactant solutions [43-45]. This might lead to precipitates that are chemically inhomogeneous regarding their concentration of impurities.

<table>
<thead>
<tr>
<th>Exp</th>
<th>Solution composition (M)</th>
<th>Relevant dissolved species</th>
<th>Saturation Indexes</th>
<th>(SO₄²⁻)/(CO₃²⁻)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[CaCl₂]</td>
<td>[NaHCO₃]</td>
<td>[Na₂SO₄]</td>
<td>pH</td>
</tr>
<tr>
<td>A0</td>
<td>0.05</td>
<td>0.05</td>
<td>---</td>
<td>7.620</td>
</tr>
<tr>
<td>A3</td>
<td>0.04</td>
<td>0.05</td>
<td>0.003</td>
<td>7.762</td>
</tr>
<tr>
<td>A5</td>
<td>0.04</td>
<td>0.05</td>
<td>0.005</td>
<td>7.765</td>
</tr>
<tr>
<td>A7</td>
<td>0.04</td>
<td>0.05</td>
<td>0.0007</td>
<td>7.768</td>
</tr>
<tr>
<td>A15</td>
<td>0.05</td>
<td>0.05</td>
<td>0.015</td>
<td>7.640</td>
</tr>
<tr>
<td>A20</td>
<td>0.05</td>
<td>0.05</td>
<td>0.020</td>
<td>7.645</td>
</tr>
<tr>
<td>A22</td>
<td>0.05</td>
<td>0.05</td>
<td>0.022</td>
<td>7.647</td>
</tr>
<tr>
<td>A25</td>
<td>0.05</td>
<td>0.05</td>
<td>0.025</td>
<td>7.650</td>
</tr>
</tbody>
</table>

Three independent runs were conducted for each experiment. Experimental runs were terminated after six different ageing times (5 minutes, 10 hours, 1, 3, 7 and 14 days). Precipitates were...
separated by filtering the aqueous phase through a 0.45-μm cellulose acetate membrane (Sartorius stedim; Albet), thoroughly rinsed with deionized water, subsequently rinsed with 10 mL of ethanol and left to dry overnight at 30 °C in a drying chamber. Filtered aqueous solutions were stored in closed Pyrex flasks at 4 °C in a refrigerator until they were prepared for analysis.

2.1. Sample characterization

The filtered solid phases recovered from each experiment were characterized by different techniques to determine their mineralogy, their chemical composition and the morphology and size of their crystalline constituents. The precipitates were first analyzed by powder X-ray diffraction (XRD) to identify the mineral phases by means of a Panalytical X'Pert PRO MRD microprocessor-controlled X-ray diffractometer. Most of the diffraction patterns were recorded between 20° and 60°, and few of them between 10° and 60° with a step size of 0.001° and using Cu Kα radiation. Diffractograms were processed with the X'Pert HighScore Plus (PANalytical B.V.) software. Diffraction patterns were compared to standard powder diffraction files from the ICDD-PDF2 data base (release 2007): 00-005-0586 for calcite, 01-080-2790 for aragonite, 04-017-8634 for vaterite, and 04-016-3025 for gypsum.

The precipitates were further analyzed by Fourier Transform Infrared (FTIR) spectroscopy using a BRUKER IFS 66v/S spectrophotometer equipped with a TGS detector. FTIR spectra were recorded on samples diluted in KBr by co-addition of 64 scans, with a precision of 0.2 cm⁻¹. The morphology of the mineral phases of the precipitate was imaged using a JEOL JSM 6400 Scanning Electron Microscope (SEM). Semi-quantitative chemical analyses of the precipitates were obtained using an Oxford Instrument Energy Dispersive X-ray (EDX) detector. KristalShaper software [46] was used to model the habit of calcite crystals.

In order to determine the content of S in the aqueous solution from each experiment, the solutions were analyzed immediately after filtering using inductively coupled plasma-optical emission spectrometry (ICP-OES) (SPECTRO Arcos).

3. Results

3.1. Mineralogical evolution of the precipitation during ageing

The mineralogical evolution of the precipitates during ageing is summarized in Table 2 and illustrated in Figures 1 to 4.

<table>
<thead>
<tr>
<th>Exp</th>
<th>(SO₄²⁻)/(CO₃²⁻)</th>
<th>5 minutes</th>
<th>10 hours</th>
<th>1 day</th>
<th>3 days</th>
<th>7 days</th>
<th>14 days</th>
</tr>
</thead>
<tbody>
<tr>
<td>A0</td>
<td>0</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Cal</td>
<td>Cal</td>
<td>Cal</td>
</tr>
<tr>
<td>A3</td>
<td>8.48</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Cal</td>
<td>Cal</td>
</tr>
<tr>
<td>A5</td>
<td>14.03</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Cal</td>
<td>Vtr</td>
</tr>
<tr>
<td>A7</td>
<td>19.51</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal</td>
<td>Vtr, Cal, Arg</td>
<td>Cal, Vtr, Arg</td>
</tr>
<tr>
<td>A15</td>
<td>41.78</td>
<td>Vtr, (Cal)</td>
<td>Vtr, (Cal)</td>
<td>Vtr, (Cal)</td>
<td>Vtr, (Cal)</td>
<td>Vtr, Arg, (Cal)</td>
<td>Vtr, Arg, Cal</td>
</tr>
<tr>
<td>A20</td>
<td>54.87</td>
<td>Vtr, (Cal)</td>
<td>Vtr, (Arg), (Cal)</td>
<td>Vtr, Arg, (Cal)</td>
<td>Vtr, Arg, (Cal)</td>
<td>Vtr, Arg, Cal, (Gp)</td>
<td></td>
</tr>
<tr>
<td>A22</td>
<td>60.00</td>
<td>Vtr</td>
<td>Vtr, (Arg)</td>
<td>Vtr, Arg</td>
<td>Vtr, Arg, Gp</td>
<td>Vtr, Arg, Gp</td>
<td></td>
</tr>
<tr>
<td>A25</td>
<td>67.61</td>
<td>Vtr</td>
<td>Vtr, Arg</td>
<td>Vtr, Arg</td>
<td>Vtr, Arg, Gp</td>
<td>Vtr, Arg, Gp</td>
<td></td>
</tr>
</tbody>
</table>

The XRD and FTIR analyses of the precipitates evidence that soon after their formation (ageing time = 5 minutes) they mainly consist of vaterite, with calcite present as a minor phase. The amount of calcite in the precipitate at 5 minutes reaction progressively decreases as the initial (SO₄²⁻)/(CO₃²⁻)
ratio in the aqueous solution increases. This is clearly evidenced by the increase in the intensity of the
peaks assigned to vaterite, and the decrease of those attributed to calcite in the diffraction patterns
recorded in the samples corresponding to this ageing time (Figure 1a). Thus, the amount of calcite is
significant in A0, where the initial (SO\text{2−})/(CO\text{3−}) ratio in the aqueous solution is 0, while no calcite is
detected in experiment A22, where the initial (SO\text{2−})/(CO\text{3−}) ratio is 60.0. FTIR spectra show
absorbance bands characteristics of vaterite and/or calcite, whereas the presence of aragonite is not
detected (Figure 1b). A very weak band at ~1130 cm\textsuperscript{−1} (found in the spectrum of precipitates from
experiments A5 – A22, but absent in that corresponding to the solids formed from sulphate-free
solutions, A0) can be assigned to sulphate ions incorporated in the structure of vaterite and/or calcite.

![Figure 1](https://www.preprints.org/Content/Preprint/1919020202v1/Figure1.png)

**Figure 1.** XRD patterns (a) and infrared spectra (b) of precipitates sampled after 5 minutes of reaction
in the case of experiments A0, A5, A15 and A22, with initial (SO\text{2−})/(CO\text{3−}) ratios of 0, 14.0, 41.8 and
60.0, respectively.

Furthermore, the mineralogical composition of the precipitates evolves with ageing time,
following trends whose characteristics vary depending on the initial (SO\text{2−})/(CO\text{3−}) ratio in the
solution. The composition of precipitates formed in experiments A0 to A5, where the initial (SO\text{2−})/
(CO\text{3−}) ratio in the aqueous solution varies between 0 and 14.03, show a similar evolution trend.
This trend is characterized by the progressive decrease of the amount of vaterite present in the
precipitate as ageing progresses (Figure 2). This decrease takes place at a slower rate with increasing
initial (SO\text{2−})/(CO\text{3−}) ratios in the aqueous solution, as confirmed by both, XRD (Figure 2a) and FTIR
spectroscopy (Figure 2b). Thus, in the absence of SO\text{2−} in the solution (A0 in Tables 1 and 2; (SO\text{2−})/
(CO\text{3−}) = 0), the precipitate exclusively consists of calcite after 3 days of ageing. In experiment A3
((SO\text{2−})/(CO\text{3−}) = 8.48), vaterite and calcite are detected after 3 days but calcite is the only CaCO\text{3} phase
present in the precipitate after 7 days of ageing. In contrast, vaterite is still present in the precipitate
formed in A5 ((SO\text{2−})/(CO\text{3−}) = 14.0) after 7 days of ageing (Figure 2; Table 2) but is undetected in the
solid phase recovered after 14 days of reaction. This trend is observed in all these three experiments,
i.e., vaterite can remain for longer ageing periods when the (SO\text{2−})/(CO\text{3−}) ratio increases.

![Figure 2](https://www.preprints.org/Content/Preprint/1919020202v1/Figure2.png)

**Figure 2.** XRD patterns (a) and infrared spectra (b) of precipitates sampled from aqueous solutions
with low (SO\text{2−})/(CO\text{3−}) ratios (A0 – A5) as a function of time.
A different mineralogical evolution trend is observed in experiments A7 to A20, where the initial 
(SO₄²⁻)/(CO₃²⁻) ratio in the aqueous solution varies between 19.51 and 54.87 (Figure 3). In this case, the 
formation of aragonite is also observed during ageing. This CaCO₃ polymorph appears as a 
constituent of the precipitate formed after ageing times that are shorter as the initial (SO₄²⁻)/(CO₃²⁻) 
ratio in the aqueous solution is higher. Thus, aragonite is detected in the precipitate formed in A7 
and A15 after 7 days of ageing, while this phase is found much earlier in the precipitate formed in 
A20, after only 1 day of ageing. This evolution trend is particularly evident in the FTIR spectra, which 
show weak bands at ~ 699 cm⁻¹ and 712 cm⁻¹ and at ~ 855 cm⁻¹, consistent with the positions of the υ₄ 
antisymmetric bending and υ₃ out-of-plane vibrations of carbonate ion in aragonite, respectively. This 
bands are absent in the precipitates formed after short times of ageing (72 h and shorter for A7 and 
A15, and 10 hours and shorter for A20). It is worthwhile to note that after 14 days of ageing these 
precipitates consist of the three CaCO₃ polymorphs, vaterite, aragonite and calcite, together with 
gypsum as a minor phase. Calcite is still the most abundant phase in A7, whereas vaterite is the most 
abundant one in A15 and A20 (Figure 3a). Bands at 1130, 628 and 608 cm⁻¹, particularly evident in the 
precipitate formed in A20 after 2 weeks of ageing can be assigned to sulphate ions incorporated into 
the lattice of the CaCO₃ polymorphs.

![Image](https://www.preprints.org/doi:10.20944/preprints201902.0202.v1)

**Figure 3.** XRD patterns (a) and infrared spectra (b) of precipitates sampled from aqueous solutions 
with a (SO₄²⁻)/(CO₃²⁻) ratios of 19.51 and 54.87 (A7 and A20) as a function of time.

Another mineralogical evolution trend is observed in experiments A22 and A25, where the 
initial (SO₄²⁻)/(CO₃²⁻) ratio in the aqueous solution is 60.00 and 67.61, respectively (Figure 4). In these 
experiments, vaterite is always the first and only phase that precipitates whereas calcite never forms 
part of the precipitate regardless the ageing time. It also is worthwhile to note that the formation of 
aragonite is unequivocally observed after 1 day of ageing in both experiments A25 and A22. After 7 
days gypsum precipitates and coexists with vaterite and aragonite, also in both experiments.

![Image](https://www.preprints.org/doi:10.20944/preprints201902.0202.v1)

**Figure 4.** XRD patterns (a) and infrared spectra (b) of precipitates sampled from aqueous solutions 
with the highest (SO₄²⁻)/(CO₃²⁻) ratio (67.6) (A25) as a function of time.

3.2. **Characterization of the precipitate and the aqueous solution**
SEM micrographs in Figure 5 show that vaterite appears as cauliflower-like aggregates (Figures 5a-b) or radial aggregates that consist of lens-shaped crystallites (Figure 5c-d). As is apparent, no significant evolution of vaterite morphology is observed as a function of the initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio in the aqueous solution nor the ageing time. In contrast, calcite appears as single crystals whose morphology and size are highly dependable on both, reaction time and initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio in the solution. Thus, in experiment A0, where there is no sulphate in the solution, calcite crystals are bounded by flat \(\{10\overline{4}\}\) rhombohedron faces regardless the reaction time and show sizes in the 10-25 \(\mu m\) range (Figures 5a, c, e). In experiments A3 to A7, where the initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio varies between 8.5 and 19.5, calcite crystals that form early after the mixing of the aqueous solution also show the typical rhombohedron-like shape with flat \(\{10\overline{4}\}\) faces (Figure 5f). However, their morphology undergoes an evolution as ageing progresses. This evolution is characterized by a progressively increasing elongation along [001] (Figures 5 f-h) as well as the emergence of rough pseudofacets that lead to the development of a new form, the more acute \(\{02\overline{1}\}\) rhombohedrum (see inset in Figure 5h). This evolution seems to be more marked in precipitates A7 and A5 than A3, which points to a positive relationship between higher initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio in the aqueous solution and more elongated and rougher-faced calcite crystals. Thus, after 14 days of ageing calcite crystals formed in experiment A3 show a length/width ratio around 2.4. This ratio is around 3.3 and 3.5 in calcite crystals formed in experiments A5 and A7, respectively (Figures 5i-j). As mentioned above, calcite crystals are hardly noticeable in experiments A15 to A20 and are absent in experiment A22 and A25 regardless the reaction time.
Figure 5. SEM images of precipitates recovered from solutions with different (SO$_4^{2-}$)/(CO$_3^{2-}$) ratios after various elapsed times ranging from 5 minutes to 14 days. Vaterite appears as cauliflower-like aggregates (a-b) or lens-shaped crystallites (c-d). When there is no sulphate in the solution, calcite crystals are bounded by flat [10-14] rhombohedron faces (a-c,e). In solutions with moderate concentration of sulphate calcite crystal habit undergoes an evolution characterized by progressive increase in elongation along [001] (f-j) and the emergence of the {02-21} form (see inset in h). Aragonite appears as crystals with the so-called “morning-star” morphologies (k). Gypsum crystals exhibit tabular habits with the [010] form as the main face (l).

Aragonite is observed in experiments A7-A25 ((SO$_4^{2-}$)/(CO$_3^{2-}$)=19.51-67.61). In all cases, aragonite appears as crystals bounded by curved prismatic faces. These crystals grow on the surface of vaterite aggregates, forming so-called “morning-star” aggregates (Figure 5k). Finally, gypsum crystals, which are observed in experiments A22 and A25, typically exhibit tabular habits with the [010] form as the main face with an accentuated elongation along [001] (Figure 5l).

EDX analyses conducted on vaterite and calcite crystals evidence that they contain S in all the cases, except for those vaterite and calcite crystals formed in the absence of SO$_4^{2-}$ (experiment A0) in the aqueous solution. For both, vaterite and calcite, a direct correlation between the [S] (% at) and the initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio in the aqueous solution is detected (Figure 6a). Thus, after 5 minutes reaction a [S] (% at) of 0.91 was measured in vaterite aggregates formed in experiment A3, while S/Ca
ratio is as high as 1.54 and 3.23 in vaterite formed in experiments A7 and A25, respectively. A similar trend is observed in calcite crystals, whose S/Ca ratio after 5 minutes reaction is 2.81 in experiment A3, 4.48 in experiment A5 and 5.76 in experiment A20. It is interesting to note that for both, calcite and vaterite, for a given (SO$_4^{2-}$)/(CO$_3^{2-}$) initial ratio, the highest S content is always detected in the precipitates with the shorter ageing (5 minutes reaction) and progressively decreases as ageing progresses. This evolution is depicted in Figure 6b for calcite formed in experiment A5. It is also striking that, regardless the initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio and the ageing time, calcite crystals always show higher S contents than vaterite aggregates. The small size of aragonite crystals formed in experiments A7 to A25 and gypsum crystals formed in experiments A22 and A25 prevented that EDX analyses could be conducted on them.

![Figure 6](image1.png)

Figure 6. (a) Concentration of S (% at) in calcite and vaterite sampled after 5 minutes of reaction as a function of the initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio. (b) Variation of [S] (% at) with ageing in calcite crystals sampled from aqueous solutions with a (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio of 14.03 (A5).

Finally, ICP-OES analyses of the S concentration in the aqueous solution show that at 5 minutes reaction, the concentration of S in the aqueous solution is progressively lower than the initial aqueous solution as the initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio in the aqueous solution increases (Figure 7a). Furthermore, the concentration of S in the aqueous solution varies with ageing time, following similar trends independently of the initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio in the solution (Figure 7b). Thus, the concentration of S in the aqueous solution is characterized by a rapid drop followed by a progressive increase of as ageing progresses.

![Figure 7](image2.png)

Figure 7. (a) Concentration of S (from ICP-OES analyses) in aqueous solutions sampled after 5 minutes of reaction as a function of the initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratios (dashed lines). The initial concentrations of S in the solutions are connected by a solid line. (b) Evolution of the concentration of S (from ICP-OES) of aqueous solutions with initial (SO$_4^{2-}$)/(CO$_3^{2-}$) ratios of 0 (A0), 14.3 (A5), 41.8 (A15) and 67.7 (A25), respectively.

4. Discussion
The experimental results described in the previous section (XRD and FTIR analyses) show that
the mineral composition of the precipitate recovered in each run varied as a function of two main
factors: the high supersaturation of the parental aqueous solution, which controls the phases that are
formed at the initial stages of crystallization, and the \((\text{SO}_4^{2-})/(\text{CO}_3^{2-})\) ratio in the parental solution,
which influences the mineral evolution of the precipitates during ageing. After the mixing of the
reactants, aqueous solutions are highly supersaturated with respect to all the crystalline \(\text{CaCO}_3\)
polymorphs (Table 1) and spontaneous precipitation takes place. According with the Ostwald step
rule [15] the formation of the most soluble and disorder phase, is kinetically favored [47]. However,
under the conditions used in these experiments the precipitation of the amorphous calcium
carbonate, the most soluble \(\text{CaCO}_3\) phase, can be discarded since the aqueous solution is always
undersaturated with respect to this phase \((\text{SI}_{\text{ACC}} \leq -0.1)\). Therefore, kinetics factors promote the
formation of vaterite as the first phase that precipitates immediately after the mixing of the parental
solutions. After vaterite nucleation, the system can further reduce its free energy by the subsequent
transformation of vaterite into calcite through a solvent mediated transformation [48]. In addition, a
reduction of supersaturation resulting from the growth of vaterite promotes the nucleation of calcite.

Therefore, as the system approach to equilibrium, the most stable \(\text{CaCO}_3\) phase, calcite, should
become the prevailing phase at the expense of vaterite. The mineralogical evolution observed in
experiments with the lower initial \((\text{SO}_4^{2-})/(\text{CO}_3^{2-})\) ratios (A0-A3) is in good agreement with this
expected evolution. Furthermore, it is in good consonance with the precipitate mineralogical
evolution upon ageing previously observed by, in similar mixing experiments conducted in more
highly supersaturated systems \((\text{SI}_{\text{CAL}} \sim 3.7)\) under higher pH (~10.9) [31]. Our current results are
further evidence that the presence of a high amount of sulphate ions in the aqueous media slows
down or even inhibits the direct precipitation of calcite and transformation of vaterite into calcite,
thereby contributing to the stabilization of vaterite with respect to calcite. The prolonged existence of
vaterite as a major constituent of the precipitate is observed in all the experiments. The stabilization
of vaterite is more evident the higher the \((\text{SO}_4^{2-})/(\text{CO}_3^{2-})\) ratio is in the aqueous solutions. Indeed, in
experiments A22 and A25, where this ratio is highest, vaterite does not transform into calcite even
after 2 weeks of ageing. These results are again consistent with those previously reported by
Fernández-Diaz et al. [31], who notice that vaterite can remain as a major constituent phase for long
ageing periods when \((\text{SO}_4^{2-})/(\text{CO}_3^{2-})\) ratios higher than one. The stabilization of vaterite and the
retardation of calcite crystallization in the presence of sulphate ions can be explained by a
combination of both thermodynamics and kinetics factors. Computational modelling of the
substitution of a small amount of carbonate groups by sulphate groups in the structures of vaterite
and calcite [28,31] showed that vaterite structure is much less disrupted by the isomorphic
incorporation of \(\text{AO}_3^{2-}\) groups, including sulphate, chromate and selenate, than calcite structure. In
the case of sulphate, the incorporation of this oxyanion results in an increment of calcite lattice energy,
while it has the opposite effect on the lattice energy of vaterite. This different impact of sulphate
isomorphic incorporation on the energetics of calcite and vaterite structures is especially relevant for
the S content range up to 3% molar fraction, where it translate into an effective reduction of the energy
difference between the two calcium carbonate polymorphs. The larger capability of vaterite to
accommodate sulphate oxyanions in its structure compared to calcite seems to be in contradiction
with the experimental evidence provided by the EDX analyses of calcite and vaterite grains in the
precipitates obtained in these experiments. These analyses steadily show a higher S content in calcite
crystals than in vaterite aggregates formed in the same experiment. This is so regardless the initial
\((\text{SO}_4^{2-})/(\text{CO}_3^{2-})\) ratio and the ageing time (see Figure 6a). Similar results were reported by Fernández-
Díaz et al. [31] for precipitates formed under higher supersaturations and pHs. These authors explain
the apparent contradiction between experimental results and energy-based expectations considering
that the higher S contents measured in calcite most likely were the consequence of this phase forming
later than vaterite. In such a scenario, calcite crystals would have grown from a \(\text{CO}_3^{2-}\) depleted
medium after vaterite nucleation and, consequently, in a solution with a \((\text{SO}_4^{2-})/(\text{CO}_3^{2-})\) ratio
significantly higher than the initial one.
Although the results of computer simulations of sulphate incorporation do not predict stability crossovers between vaterite and calcite, the approximation of the lattice energy of both polymorphs can explain the increasingly larger duration of vaterite as precipitate component with growing initial sulphate concentration in the solution.

While the incorporation of impurities from the aqueous solution can change the lattice energy of the bulk crystal, surface-related phenomena like impurity adsorption can modify the surface chemistry of the different polymorphs, thereby changing their surface energies [49]. The impact of this phenomenon is particularly relevant in the case of small particles, where the surface/volume ratio is very high, and the effect of lattice energy in the stability of the different polymorphs is comparatively much less important than that of surface energy. A small increase of the surface energy of calcite nanoclusters and nanoparticles due to sulphate adsorption could dramatically increase the barrier for its nucleation, since such a barrier is proportional to the square of the surface energy. Were this to happen, the main outcome would also be a progressively slower transformation of vaterite into calcite as the (SO₄²⁻)/(CO₃²⁻) ratio in the parent aqueous solutions increases. Kinetics arguments involving the adsorption of impurities have often been invoked to explain the stabilization of vaterite with respect to calcite and the retardation of calcite nucleation and growth. Phenomena observed during the precipitation of CaCO₃ from supersaturated aqueous solution bearing phosphate ions are an example [50]. In this system, the stabilization of vaterite was rather attributed to the blocking of active sites for dissolution on the surface of this phase due to the adsorption of phosphate ions. A similar phosphate adsorption onto surface sites-related explanation was claimed to explain calcite growth inhibition. Indeed, a similar reasoning can be applied to explain the observed effect of sulphate ions. Another alternative thermodynamic and mechanistic argument has been invoked to explain the inhibition of calcite growth by the effect of sulphate ions. In situ AFM experiments of calcite growth performed using a fluid cell and flowing supersaturated sulphate-bearing aqueous solutions showed that a low concentration of sulphate in the aqueous solution (5 mM) is enough to decrease the rate of step spread on calcite surface [51]. These nanoscale AFM observations supported the interpretation that the trapping of sulphate ions by nanometric growing layers resulted in the generation of interfacial strain energy. Strain relaxation perpendicular to the growing layer introduces local variations in bond lengths, giving rise to local departures from calcite surface ideal nanotopography. The direct consequence of this phenomenon is a decrease of the step advancement rate according to the so-called “template effect” model [20,25,52], which is more marked the larger the amount of sulphate ions trapped within calcite monolayers.

An interesting result observed in experiments A7 to A25, where the initial (SO₄²⁻)/(CO₃²⁻) ratio in the aqueous solution > 19.51, is the formation of aragonite. It takes place through the solvent-mediated transformation of vaterite and can also be explained invoking both, thermodynamic and kinetics arguments. For equivalent compositions of sulphate in the polymorphs, computer modelling of sulphate isomorphous incorporation predicts a significantly larger increment of aragonite lattice energy compared to that of both, calcite and vaterite. This result indicates that the substitution of carbonate groups by sulphate in aragonite structure is very unfavorable [28,31]. As a result, even when this polymorph forms from highly supersaturated solutions with very high (SO₄²⁻)/(CO₃²⁻) ratios, it grows relatively sulphate-free compared to calcite. The direct consequence is a much more marked impact of the presence of sulphate in the growth medium on calcite stability. Indeed, the solubility of a sulphate-bearing calcite can overcome that of sulphate-free aragonite, as pointed out by Busenberg and Plummer [53], who reported this to be the case for calcites with sulphate contents above 3 mole%. Such solubility crossover could explain the observation that, upon ageing of the vaterite precipitate, aragonite forms simultaneously to or, even, after the formation of the theoretically stable polymorph calcite when the concentration of sulphate in the aqueous solution is very high. This is the case in experiments A7 to A20. Unfortunately the small size of the aragonite crystals formed in these experiments has prevented to obtain EDX analysis evidence that could give full support to our interpretation, although the high concentration of sulphate measured in calcite crystals point in this direction (see Figure 6). On the other hand, the fact that sulphate incorporation into aragonite structure is very unfavorable, it can be concluded that an inhibitory effect of sulphate
ions during the growth of aragonite will arguably be less effective than in the case of calcite, and so will be a hypothetical “template effect” mechanism.

Beside the precipitation of CaCO$_3$ polymorphs, gypsum was unequivocally detected after 7 days of interaction in the experiments with the highest (SO$_4^{2-}$)/(CO$_3^{2-}$) ratios (A20-A25). The precipitation of this phase is not surprising since the initial solutions are supersaturated with respect to this phase (SL$_{Ca}$ -0.17-0.25). However, the early precipitation of CaCO$_3$ phases after mixing of the parent solutions necessarily depletes the solution in Ca$^{2+}$ (and to a much lesser extent in SO$_4^{2-}$), leading to a very important reduction of aqueous phase SL$_{Ca}$. Thus, systems initially close to equilibrium with respect to gypsum, as is the case of the solution in experiment A15, will become undersaturated soon after the precipitation of CaCO$_3$ starts, making the formation of gypsum impossible. Those systems initially more highly supersaturated with respect to gypsum will either approach equilibrium or significantly less supersaturated, allowing gypsum precipitation be prevented over large periods.

The presence of sulphate in the aqueous solutions and the ageing not only appears to be a critical factor controlling the mineralogy of the recovered precipitates, but it also affects the morphology of calcite single crystals. As mentioned in section 3.2, after short ageing time, calcite single crystals exhibit the typical rhombohedral idiomorphic shape (see Figures 5a,c,e), which progressively become blockier and elongated along the c axis as ageing progresses. This elongation is clearly more pronounced as the (SO$_4^{2-}$)/(CO$_3^{2-}$) ratio in the aqueous solution increases. Since there is a clear correlation between the concentration of sulphate in the aqueous solution and the amount of this anion that is incorporated into the calcite (and vaterite) structure, directly evidenced by EDX (see Figure 6a) and supported by ICP analysis (Figure 7a), we can conclude that the modification of calcite crystal habit is a direct result of the incorporation/sorption of this anion. The effect of several ions on calcite habit has been profusely discussed in the literature. In particular, the elongation of calcite crystals has been interpreted as resulting of a significant lowering of the surface energy of certain faces other than {10014} due to a preferential sorption of foreign ions on these faces [54]. It has also been attributed to a modification of calcite surface nanotopography due to step-specific impurity interactions [55]. These interactions contribute to the stabilization of new crystal faces that become prevalent during growth. In the case under consideration here, the habit of calcite crystals grown in presence of sulphate, we observe the stabilization of the {0221} form. The resulting habits strikingly resembles those of calcite crystals grown in the presence of small cations such as Mg$^{2+}$, Co$^{2+}$ or Mn$^{2+}$ [56].

Finally, the progressive decrease of S content in both the calcite and vaterite precipitates with reaction time (Figure 6b), which is in good agreement with a parallel progressive increase of sulphate concentration in the aqueous solution (Figure 7b), points to the precipitate undergoing of recrystallization processes through a dissolution-precipitation mechanism. The driving force for this recrystallization process could be the reduction of the extra energy associated to an excess of sulphate incorporated in calcite and vaterite structures under the influence of the high supersaturations dominating the early staged of the precipitation process. The progress of this recrystallization process leads to a chemical purification of the precipitates that further contributes to the stabilization of calcite as ageing progresses.

**Author Contributions:** Iris Cuesta Mayorga performed the laboratory experiments, the DRX analysis, and prepared the samples for FTIR, SEM images, EDX, and ICP analysis. José Manuel Astilleros and Lurdes Fernández-Díaz contributed to the conception of the study. All authors analyzed the data and were involved in manuscript preparation. All authors read and approved the manuscript.

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