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Catalytic Effect of Magnesium Ions on Polymorphic Crystallization of Calcium Carbonate

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Abstract

The effect of Mg²⁺ ions on the polymorphic crystallization of calcium carbonate has been studied by precipitating carbonate minerals from Ca(HCO₃)₂ (Ca²⁺, 400 mg/1) solutions containing various amount of Mg²⁺ ions (0 to 2000 mg/1) at the temperature of 2 to 80°C. The concentration of Mg²⁺ ions and temperature in a parent solution affect remarkably the aragonite formation. The precipitating fields of calcite and/or aragonite are systematically represented on log[Mg²⁺] vs. 1/T diagram. Energetic effect of Mg²⁺ ions on aragonite formation is estimated from an Ahrenius type relationship between required concentration of Mg²⁺ ions and absolute temperature to form aragonite; [Mg²⁺] = A·exp (—E/RT), where E = 13 Kcal/mol, which will be needed to form aragonite nucleous from solution.

The presence of calcite seeds favors the calcite formation in the presence of Mg²⁺ ions lower than 400 mg/l.

1. Introduction

It is important to reveal the role of Mg²⁺ ions and of the other divalent cations, anions and dissolved organic materials in parent solutions on the carbonate formation for the understanding of inorganic and biological precipitation of calcium carbonates. The biologically formed marine carbonate skeletons are composed of low Mg-calcite, high Mg-calcite and aragonite, where low Mg-calcite is thermodynamically stable under natural conditions (Jamieson¹, Macdonald²). The controlling factors of the polymorphic formation in marine carbonate skeletons are biological species, temperature, and salinity in the water environments (Lowenstam³, Chave⁴, Dodd⁵, Kitano et al.⁶, Milliman⁷).

It is evident that Mg²⁺ ions in a parent solution favor aragonite precipitation in marine environments (Kitano and Hood⁸, Simkiss⁹, Groot and Duyvis¹⁰). Mg²⁺ ions act as an inhibitor for nucleation and/or growth of calcite (Mclester et al.¹¹, Bischoff¹², Bischoff and Fyfe¹³, Pytkowicz¹⁴, Berner¹⁵,¹⁶). Katz¹⁷ and Mucci and Morse¹⁸ studied the incorporation of magnesium to calcite. Incorporation of magnesium makes calcite unstable thermodynamically (Kitano and Frutsu¹⁹, Jensen and Kitano²⁰, Winland²¹, Plummer and Mackenzie²²). And the incorporation of magnesium to carbonate decreases the surface tension of calcite (Mg poor and Mg rich) and aragonite, and decreases the nucleation energy of these minerals. The decrease in the nucleation energy is larger for aragonite than that for calcite (Moller and Lajacopalan²³).

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The purpose of this paper is to clarify the role of Mg\(^{2+}\) ions in a parent solution on the crystallization of calcium carbonates.

2. Experimental Procedure

Effects of Mg\(^{2+}\) ions and temperature in a parent solution on the formation of calcium carbonate polymorphs have been studied by precipitating carbonates from calcium bicarbonate solution containing various amounts of Mg\(^{2+}\) ions at various temperatures. The calcium bicarbonate solution was prepared by bubbling CO\(_2\) gas to calcium carbonate suspension and filtering with Toyo filter paper (No. 2). Then CO\(_2\) gas was bubbled again. Various amounts of MgCl\(_2\) solution were added to 1000 ml of the Ca(HCO\(_3\))\(_2\) solution. The solution were stirred continuously at 2, 10, 22, 40, 60 and 80°C. Also, sea water was used to have informations about carbonate polymorphism formed from it. The calcium and magnesium contents of solutions and precipitates were determined by EDTA titration and atomic absorption.

In order to get informations about the characterization of magnesium effect, the experiments on the effect of the presence of calcite seeds were also carried out in the presence of Mg\(^{2+}\) ions in a parent solution. The solution containing calcite seeds was prepared as follows: 2000 ml of calcium bicarbonate solution was stirred until CaCO\(_3\) separation started to come out. The formed precipitates were filtered off with Toyo filter paper (No. 2) during the presence of suspended carbonate in the solution, that is, before the formed carbonate deposited and the solution became transparent*. The prepared filtrates contain calcite seeds. Concentration of the seeds in a parent solution is around 5 mg/l or less. The various volumes of MgCl\(_2\) solution, saturated with calcite at P\(_{CO_2}\) = 1 atm., were added to the filtrates containing calcite seeds. When MgCl\(_2\) solutions not saturated with calcite were added, the seeds did not work. These solutions saturated with calcite were stirred for about 20 minutes and then kept still at 25°C. Carbonate precipitates appeared like snow fall after one day. For the complete precipitation, the solution was kept standing still further for one week at 25°C. The floating carbonates suspended in the surface were discarded carefully and the precipitates on the bottoms of the beakers were tested by X-ray diffraction, because the mineral composition of carbonates floating in the surface of the solution was usually complex.

*When the solution is filtered after the solution became transparent, the presence of seeds in the solution is negligible.

3. Results and Discussion

Crystallizing fields of calcium carbonate polymorphs are shown in Fig. 1. It is evident that the presence of Mg\(^{2+}\) ions and the high temperature in the parent solution favor the aragonite formation. The experimental result shows that only aragonite is precipitated from sea water under ordinary conditions.

The boundary of the precipitating fields of aragonite and calcite is shown by the strait line in the log[Mg\(^{2+}\)] vs. 1/T diagram, whereas the boundary line between calcite and calcite + aragonite deviates down from a straight line in a high temperature range (see Fig. 1). This may be caused by the mixing of complex carbonate formed in the surface of the solution at high temperatures. The boundary of the precipitating fields is considered to be expressed by an Ahrenius type relationship between
Precipitating fields of calcite and/or aragonite.

The required concentration of Mg$^{2+}$ ions and temperature to form aragonite is given by:

$$[\text{Mg}^{2+}] = A \exp \left( \frac{Q}{T} \right),$$

where $A$ and $Q$ are constant, and $Q$ contains an energy term. The above equation is rewritten as:

$$[\text{Mg}^{2+}] = A \exp \left( -\frac{E}{RT} \right),$$

where $A$ is constant and $R$ is gas constant. Mg$^{2+}$ ions in solution control the rate of calcium carbonate crystallization (Pytkowicz 14, Berner 16). So, $E$ is an energy term such as activation energy for aragonite formation from solution. From the experimental results shown in Fig. 1, the energetic effect of Mg$^{2+}$ ions is estimated to be shown as $E = 13$ Kcal/mol. Activation energies for calcite growth from solution without Mg$^{2+}$ ions and for transformation of aragonite to calcite were reported to be 11.0 Kcal/mol (Nancollas and Reddy 24) and 57.4 Kcal/mol (Bischoff 25), respectively. It seems that activation energy for calcite growth from Mg$^{2+}$ containing solution is larger than that from solutions without Mg$^{2+}$ ions.

In order to get informations about the characterization of magnesium role on the polymorphic crystallization of CaCO$_3$, the experiments on calcite seeds were carried out. The results are shown in
Fig. 2. Aragonite proportion of precipitates vs. added amount of Mg$^{2+}$ ions in solution.

Fig. 2 with the experimental results without the seeds. It is apparent that the presence of calcite seeds favor the calcite formation in the presence of Mg$^{2+}$ ions of lower than 400 mg/l. Aragonite is a dominant precipitates, when the parent solutions containing Mg$^{2+}$ ions of more than 200 mg/l without calcite seed were stirred at 25°C. But, aragonite was not formed from calcite seeded and non-stirred solution containing 240 mg/l of Mg$^{2+}$ ions. The seeds seem to exceed magnesium ions in the effect at moderate Mg$^{2+}$ concentration in a parent solution. The microscopic observations in the seed experiments recognized the growth of oolitic aragonite on the surface of calcite crystals. Magnesium ions may promote the formation of aragonite nucleous on the surface of calcite crystals. Nucleation stage is important for the formation of carbonate polymorphism. The energy estimated above is considered to be the energy needed to form aragonite nucleous from solutuions. Kitano et al.\textsuperscript{26} showed that the ions in a parent solution favorable for aragonite formation, such as copper and zinc, were much incorporated in carbonate precipitate at earlier stage of carbonate crystallization than at later stage, although the crystal form of those metal carbonate is not aragonitic but calcitic. Incorporation of magnesium to carbonate lattice sometimes causes the thermodynamical instabilization and the enhanced solubility of calcite (Berner\textsuperscript{16}, Plummer and Mackenzie\textsuperscript{22}). From the study on the kinetics for calcium carbonate formation from supersaturated solutions containing Mg$^{2+}$ ions, Möller and Lajacopalan\textsuperscript{23} reporated that the incorporation of magnesium to carbonate decreases the surface tension of carbonate minerals, and lowers the nucleation energy of calcite (Mg poor and Mg rich) and aragonite. The decrease in the nucleation energy for aragonite is maximum in the highly supersaturated solution having larger than 4 of Mg$^{2+}$/Ca$^{2+}$ molar ratio. The incorporation of magnesium may lower the energy needed for aragonite nucleous formation.
References


