Molecular-scale surface processes during the growth of calcite in the presence of manganese

J. M. Astilleros,1,* C. M. Pina,1,2 L. Fernández-Díaz2 and A. Putnis1
1Institut für Mineralogie, Universität Münster, Corrensstrasse 24, D-48149, Münster Germany
2Dpto. Cristalografía y Mineralogía, Universidad Complutense, 28040 Madrid, Spain

Abstract—This paper deals with the growth behaviour of the Mn-Ca-CO3-H2O solid solution-aqueous solution system on calcite (1014) surfaces. This system represents a model example, which allows us to study the effect of a number of controlling factors on the crystallisation: (1) the supersaturation function, \( \beta(x) \), and nucleation rate function, \( J(x) \), for the Mn-Ca-CO3-H2O system, (2) the relationship of such functions to the molecular scale growth mechanisms operating on growing surfaces, and (3) the surface structure of the calcite (1014) faces. In situ atomic force microscopy (AFM) growth experiments revealed a wide variety of surface phenomena, such as the transition between growth mechanisms, anisotropic changes in the step rates, and the influence of the Mn-bearing newly formed surface on subsequent growth (step stoppage followed by the formation of two-dimensional nuclei and the reproduction of the original calcite (1014) surface microtopography). These phenomena result from the interplay between the controlling parameters and are explained in those terms.

1. INTRODUCTION

Calcite, because of its reactive surfaces and ubiquity in natural aqueous environments, is the most important carbonate controlling divalent metal mobility. Among the different surface metal uptake mechanisms, coprecipitation reactions and subsequent formation of (M,Ca)CO3 solid solutions (where M is a divalent cation) have been found to be significant in geochemical systems (Reeder, 1996; Tesoriero and Pankow, 1996; Böttcher, 1997; Temmam et al., 2000). Cations incorporated into the calcite structure from supersaturated solutions include a wide range of environmental contaminants (e.g., Cd2+, Pb2+, Co2+, Zn2+2+) and diagenetic tracers (e.g., Mg2+, Mn2+). The incorporation of such cations not only affects the composition of the growing carbonate, but also plays an important role in the crystallization process, evidenced by changes in growth rates, shape, and thickness of growth layers, etc. (Sangwal, 1993). In addition, the variability of morphologies and textures usually exhibited by natural and synthetic calcite is frequently attributed to the substitution of Ca2+ by other cations in the calcite structure (Parker et al., 1993; Fernández-Díaz et al., 1996; Temmam et al., 2000). The study of the relationships between crystal morphology and chemical composition is, therefore, of special interest for understanding the formation of calcite in both geologic and biogenic environments.

Recently, considerable efforts have been made to understand the effect of different cations on the growth behaviour of calcite at a molecular scale. AFM observations have revealed a number of nanometric processes occurring on the calcite (1014) surface, including the modification of the morphology of spirals and step dynamics, the anomalous thickening of certain growth steps, and the selective incorporation of cations in crystallographic non-equivalent sites on the surfaces (Paquette and Reeder, 1995; Davis et al., 2000; Astilleros et al., 2000, 2002). It seems clear that all these phenomena are the result of the interplay between physicochemical conditions (i.e., composition of the aqueous solutions and their saturation states with respect to the crystallizing calcitic solid solution), the kinetics of the possible growth mechanisms (spiral growth, two dimensional nucleation, and continuous growth) operating at a molecular scale on the different faces of the crystal, and the surface structure of the growing face.

In this paper we present a study of the effect of Mn2+ on calcite (1014) surfaces during growth. The specific thermodynamics of the Mn-Ca-CO3-H2O SS-AS (solid solution-aqueous solution) system, i.e., the non-ideality of the solid solution and the existence of miscibility gaps close to the endmembers (McBeath et al., 1998), together with the clear structural control exerted by calcite surfaces on Mn incorporation, makes this system a very suitable example to evaluate the influence of the factors that determine the actual growth behaviour of the MnCa1-xCO3 solid solution. Our study includes (1) the evaluation of the supersaturation, \( \beta(x) \), and nucleation rate, \( J(x) \), functions for the Mn-Ca-CO3-H2O system on the basis of solid solution-aqueous solution theory, (2) the relationship of such functions to the operating growth mechanisms, and (3) the present knowledge about the surface structure of the calcite (1014) faces. By considering these controlling parameters, we can interpret a number of molecular-scale surface phenomena observed using in situ atomic force microscopy (AFM) (e.g., anisotropic modification of step rates, changes in growth mechanisms, and an inhomogeneous covering of the surface, reproducing the original microtopographic features).

2. CALCITE (1014) SURFACES GROWING FROM Ca-Mn-CO3 AQUEOUS SOLUTIONS

As a starting point for studying the growth of the calcite (1014) face in the presence of Mn, it is necessary to consider the thermodynamic properties of the Mn-Ca-CO3-H2O SS-AS
system as well as the interrelation between the calcite (1014) surface structure and the incorporation and adsorption behaviour of Mn. In the following subsections, we will present: (1) a general study of the thermodynamics and kinetics of the Mn-Ca-CO₃-H₂O SS-AS system at 25°C, including how to evaluate supersaturated states and nucleation rates within such a system, and (2) a description of the interaction of calcite (1014) surfaces with manganese, based on the structural model proposed by Reader (1996). As we will see, AFM observations of growth phenomena at a molecular scale can be explained as a result of the interaction of thermodynamic, kinetic, structural, and microtopographic factors.

2.1. The Thermodynamics of the Mn-Ca-CO₃-H₂O SS-AS System and the Evaluation of Supersaturation and Nucleation Rates

2.1.1. Thermodynamic background

In the Mn-Ca-CO₃-H₂O SS-AS system at 25°C, a limited rhombohedral solid solution can be formed between the endmembers calcite (CaCO₃) and rhodochrosite (MnCO₃). In addition, there exists a double salt with the dolomite structure called kumarahite (MnCaCO₃). Although this SS-AS system seems to be simple, neither the extent and non-ideality of the solid solution nor the stability field for kumarahite have been unequivocally established. Several studies on natural and synthetic Mn-Ca-CO₃ solid solution series have reported compositional ranges from total miscibility between endmembers (Goldschmidt and Graf, 1957; Fubini and Stone, 1983) to limited solid solubility with different ranges for miscibility gaps (Brand and Veizer, 1980; Ten Have and Heijnen, 1985). Furthermore, there is no agreement for the distribution coefficient values (Lorens, 1981; Mucci, 1988; Pingitore et al., 1988; Dromgoole and Walter, 1990; Böttcher, 1997), even though all the researchers found that, in comparison to Ca, Mn is strongly incorporated into the calcite structure (i.e., Kₐ > 1). Finally, the determination of the solubility product for the endmember rhodochrosite, KₘnC₀₃, seems to be problematic, and a number of values for this parameter can be found in the literature. Some of the KₘnC₀₃ values reported are: 10⁻¹⁰⁻⁹⁹ (Garrels et al., 1960), 10⁻¹⁰⁻⁶² (Wagman et al., 1982), 10⁻¹⁰⁻⁵⁵ (Johnson, 1982) and 10⁻⁹⁻⁴³ (McBeath et al., 1998). It is clear that this lack of agreement with respect to the precipitation behaviour in the Mn-Ca-CO₃-H₂O SS-AS system makes it difficult to propose a consistent thermodynamic model. However, McBeath et al. (1998) have recently published new experimental data for the Mnₑ₋ₓCaₓC₀₃ solid solution, obtained using electrochemical cells. These data allowed them to derive the excess Gibbs energy as a function of composition, ΔG°(MnₓCa₁₋ₓC₀₃). Moreover, the authors provide an empirical expression for the solubility product of the MnₓCa₁₋ₓC₀₃ solid solution:

\[ K_{\text{Mn},\text{Ca},\text{CO}_3} = \exp \left( 212.91 + 223.31(1 - x) + 92.01 x + 0.40339 \left( \Delta \tilde{G}^\theta_{\text{Mn},\text{Ca},\text{CO}_3}/k\text{Jmol}^{-1} \right) \right) \]  (1)

where x is the mole fraction of MnCO₃ in the solid solution. This equation can be used to evaluate the supersaturation states in the Mn-Ca-CO₃-H₂O SS-AS system at 25°C. Figure 1 shows the variation of the solubility product for MnₓCa₁₋ₓC₀₃ as a function of the composition. It is worth noting that for intermediate solid compositions (XₘnC₀₃ ≈ 0.5) the solubility exhibits a relative minimum.

2.1.2. Supersaturation evaluation

Supersaturation in SS-AS systems cannot be described by a single value but by a function of both solid and fluid composition. For the Mn-Ca-CO₃-H₂O SS-AS system, the function \( \beta(x) \) that expresses the supersaturation of aqueous solutions with respect to solid solutions has the following form (e.g., Prieto et al., 1993; Pina and Pumis, 1995):

\[ \beta(x) = \frac{[\text{Mn}^{2+}]^{a}\text{[Ca}^{2+}]^{b-1}\text{[CO}_3^{-2}]^{c}}{K_{\text{Mn},\text{Ca},\text{CO}_3}} \]  (2)

where [Mn²⁺], [Ca²⁺], and [CO₃⁻²] are the activities of the ions in the solution and \( K_{\text{Mn},\text{Ca},\text{CO}_3} \) is the solubility product as a function of the solid solution composition (Eq. (1)). Figure 2a shows, for example, the supersaturation distribution as a function of the solid composition for an aqueous solution with activities: \( [\text{Mn}^{2+}] = 0.01 \), \( [\text{Ca}^{2+}] = [\text{CO}_3^{-2}] = 0.1 \). As can be seen, the supersaturation maximum is reached for a solid composition with molar fraction \( X_{\text{MnC0}_3} \approx 0.5 \), even though the Ca activity in the aqueous solution is 10 times higher than the Mn activity. This is a consequence of the particular shape of the solubility product function (Eq. (1)). Other compositions of the aqueous solution will result in supersaturation functions with maxima located at different solid solution compositions.

2.1.3. Nucleation rates

Once the supersaturation function of an aqueous solution with respect to a solid solution is known, it is possible to evaluate the nucleation kinetics for each composition of the solid. This can be done by means of the nucleation rate function, \( J(x) \), which is obtained by considering some of the basic parameters of the classical nucleation rate equation as functions of the solid composition (see references in Pina et al., 2000):

\[ J(x) = \Gamma(x) \cdot \exp \left( \frac{-B \cdot \xi \cdot \sigma^3(x) \cdot \Omega^2(x)}{k^4 \cdot T^4 \cdot (\ln S(x))^2} \right) \]  (3)

where, \( S(x) = \sqrt{\beta(x)} \), o(x) is the interfacial free energy function, \( \Omega(x) \) is the molecular volume as a function of solid composition, and \( \Gamma(x) \) is the preexponential factor as a function of solid composition (which involves a volume diffusion step and is dependent on the interfacial free energy and solubility). The other parameters in Eq. (3) are: \( k \), the Boltzmann constant; \( T \), the absolute temperature; \( B \), a geometrical factor, which is 2πr for a spherical nucleus. Finally, \( \xi \) is a correction factor \((0 < \xi < 1)\) related to the formation of heterogeneous nuclei (i.e., \( J(x) \), expressed as nuclei per unit of area and unit of time). Such a parameter can be adjusted by considering experimental nucleation densities (e.g., obtained from AFM experiments). Although absolute values for nucleation rates given by Eq. (3) are not very reliable (the function is very sensitive to small variations of the parameters in the equation), the \( J(x) \) function is useful to determine the solid solution compositions that are kinetically favoured (Pina et al., 2000; Pina and Pumis, 2002). Figure 2b shows the nucleation rate function corresponding to...
the supersaturation distribution shown in Figure 2a. The endmember parameters required to calculate the \( J(\alpha) \) functions were taken from the literature. They are: \( \sigma_{\text{calcite}} = 97 \text{ mJ/m}^2 \) and \( \sigma_{\text{rhodochrosite}} = 81 \text{ mJ/m}^2 \) (Söhnle, 1982; Wu and Nancollas, 1999). The calculated molecular volumes are \( V_{\text{calcite}} = 6.1 \times 10^{-29} \text{m}^3 \) and \( V_{\text{rhodochrosite}} = 5.16 \times 10^{-29} \text{m}^3 \). Preexponential factors of the equation were estimated using the approximation: \( \Gamma(\alpha) = D/\Omega^{\frac{2}{3}}(\alpha) \), where \( D \approx 10^{-9} \text{m}^2\text{s}^{-1} \) is the mean diffusion coefficient for ions in water.

2.2. Structure of the Calcite \( \{1014\} \) Surface, Mn Uptake and Adsorption Processes

Since crystal growth takes place at the solid-aqueous interface, a comprehensive study of growth processes requires not only a macroscopic thermodynamic model but also a knowledge of crystal structure and surface microtopography of the growing phase. The interrelationships between such parameters and the anisotropic growth observed on calcite \( \{1014\} \) surfaces (Reeder, 1996; Astilleros et al., 2000) can be elegantly established by means of the periodic bond (PBC) chain model of Hartman and Perdok (1955). The PBCs are defined as continuous connections through the structure by strong bonds. Because such bonds have the highest attachment energies, they play an important role in determining surface microtopography and growth anisotropy. Calcite \( \{1014\} \) faces, which define the calcite cleavage rhombohedron, contain three non-equivalent PBCs parallel to the \( <441>, <221>, \) and \( <010> \) directions and, therefore, are flat (F) faces (Heijnen, 1985). Steps parallel to a given set of PBCs are structurally identical, but because of the orientation of such steps to the symmetry elements, opposite directions of advancement for a given step are not equivalent (\( [441]_+, [481]_+ \) and \( [441]_-, [481]_- \), according to the notation used by Staudt and Reeder, 1994). Figure 3 shows a representative example of etch pit development on a calcite \( \{1014\} \) surface in contact with water. Etch pits show a typical rhombohedral shape defined by \( [441]_+, [481]_+ \) or \( [441]_-, [481]_- \), according to the notation used by Staudt and Reeder, 1994) are symmetrically related by means of a c-glide, no symmetry operation relates the \( [441]_+, [481]_+ \) steps to either \( [441]_- \) or \( [481]_- \) steps, respectively. This non-equivalence of growth steps, which also determines growth and dissolution rates (positive steps migrate faster than negative steps; Hillner et al., 1992), is considered by Paquette and Reeder (1990, 1995) and Staudt and Reeder (1994) as the main reason for differential incorporation of trace elements into the calcite structure. In the model proposed by these investigators, two general types of kinks along \( <441> \) steps can be distinguished: the more constrained and smaller sites, distributed along \( [441]_+, [481]_+ \) steps, and the less constrained and larger ones, which are present in the \( [441]_-, [481]_- \) steps. Cations smaller than Ca (Mn, Mg, Co, Cd) will tend to be incorporated in nega-
tive steps, whereas cations larger than Ca (Ba, Sr) will be preferentially incorporated in positive steps.

The ionic radius of Mn$^{2+}$ is 0.80 Å, considerably smaller than Ca$^{2+}$ (0.99 Å). This difference will result in a preferential incorporation of Mn$^{2+}$ in the smaller sites located along negative steps and, probably, a lower detachment probability. Therefore, faster step advancement in comparison to pure calcite can be expected. However, the simultaneous incorporation of Mn and Ca into the calcite structure, particularly at high growth rates, will have an important effect on the distribution of defects.

3. MATERIAL AND METHODS

Crystal growth experiments were carried out at 25° C in situ in a fluid cell of a Digital Instruments Multimode Atomic Force Microscope working in contact mode. Care was taken to ensure that the tip did not affect the surface during the scan. Repeated scans of the tip over the same area did not cause any appreciable differences at this location relative to the rest of the image. Before each growth experiment, deionised water was passed over the crystal to clean the cleaved {10$ar{1}$4} calcite, as well as to adjust the AFM parameters. Calcite samples used as substrates were cleaved from a block of optically clear Iceland Spar. Ca$^{2+}$-Mn$^{2+}$-CO$_3^{2-}$ aqueous solutions were prepared by mixing Na$_2$CO$_3$, MnCl$_2$, and CaCl$_2$ aqueous solutions (SIGMA 99+%). The pH of the solutions was measured using a TESTO 230 pH meter. A value of 10.20 ± 0.05 was obtained in all cases. The pH meter electrode (type 05 pH) was calibrated using 7.00 and 10.00 buffers (Shott Geräte). Table 1 shows the composition of the solutions used in the growth experiments. Solutions were prepared immediately before injection in the fluid cell to avoid the precipitation of MnO$_{2(s)}$.

Supersaturation states of the solutions were evaluated using Eq. (2). For this purpose, it is necessary to calculate the

![Fig. 2. (a) Supersaturation function, $\beta(x)$, and (b) nucleation rate function, $J(x)$, expressed in nuclei/μm$^2$·s for a hypothetical aqueous solution with activities $a$(Mn$^{2+}$) = 0.01; $a$(Ca$^{2+}$) = $a$(CO$_3^{2-}$) = 0.1.](image)

- 64
- 56
- 48
- 40
- 32
- 24
- 16
- 8
- 0
activities of the ions in solution. The activity coefficients for Ca\(^{2+}\), Mn\(^{2+}\), and CO\(_3\)^{2-} were computed using the extended Debye-Hückel formula (Shinn and Morgan, 1981). The aqueous speciation program used considered the following species: H\(^{+}\), OH\(^{-}\), Ca\(^{2+}\), Mn\(^{2+}\), CO\(_3\)^{2-}, Cl\(^{-}\), Na\(^{+}\), HCO\(_3\)^{-}, H\(_2\)CO\(_3\), CaOH\(^{+}\), CaCO\(_3\), CaHCO\(_3\), MnOH\(^{-}\), MnCO\(_3\), MnHCO\(_3\), NaCl\(^{-}\), NaOH\(^{-}\), NaCO\(_3\), Na\(_2\)CO\(_3\), and NaHCO\(_3\). Ion-size parameters required to apply the extended Debye-Hückel formula were taken from Truesdel and Jones (1974). Table 1 (last column) shows the calculated ionic strength for the solutions. The ionic strength values (I < 0.0134) are within the range of applicability of the extended Debye-Hückel formula. As an example of calculated solution compositions, Table 2 shows free ion and ion pair concentrations in experiment number three. Note that the calculated pH value is very similar to our electrometric measurements. This is a test of the applicability of the speciation model used.

For all solutions used in the experiments, the supersaturation with respect to the endmember calcite was constant. The supersaturation with respect to rhodochrosite was increased in each experiment by increasing the concentration of Mn. To maintain a constant supersaturation value with respect to calcite, the Ca concentrations in the solutions were adjusted. In all the experiments, the dissolved inorganic carbon concentration was maintained constant.

As mentioned in the previous section, before each growth experiment, deionized water was injected in the fluid cell of the AFM. The calcite (10 14) surface in contact with water rapidly begins to dissolve by both step retreat and formation of etch pits. Etch pits on calcite usually show a typical rhombohedral shape defined by the [481] and [441] steps, which dissolve at different velocities. Therefore, by direct observation of the development of such etch pits, it is possible to determine unequivocally the crystallographic directions on calcite (10 14) surfaces before each growth experiment.

Once the orientation of the calcite substrate was established, the Ca\(^{2+}\)-Mn\(^{2+}\)-CO\(_3\)^{2-} solutions described in Table 1 were injected into the fluid cell of the AFM. In all the experiments, growth on the surface was observed. Figure 4 shows a growth sequence on a calcite (10 14) surface using solution 1. The first image (a) was taken immediately after flowing deionised water and a few seconds before injection of the growth solution. In successive images, step advancement occurs. At certain places, it is possible to observe a temporary stoppage of the step advancement. As a result, the initially straight step edges are distorted and become rounded, developing lobes during growth. Arrows 1 and 2 in Figure 4 indicate sites along the step edges where the growth slows down. A detailed observation of the sequence shows that such sites correspond to the previous corners of the etch pits. From these observations, a relationship between the original topography (arrows 1 and 2) and the lobed areas can be established.

When the concentration of Mn in the solution is slightly higher (solution 2), the growth behaviour is considerably modified. Figure 5 shows the evolution of the calcite (10 14) surface in contact with this solution. Although lateral step advancement is again the operating growth mechanism, only the first elementary growth layer advancing on the preexisting surface grows normally. This first newly formed substrate exerts a strong control on the subsequent surface evolution: when a second growth layer reaches previously grown areas, it abruptly reduces its velocity and growth stops. The growth is, however, normal on the original pure calcite surface. This inhomogeneous growth leads to an almost perfect reproduction of the topography of the original surface (cf. Fig. 5a with Fig. 5c). After some minutes, the stopped steps begin again to spread and edges become wavy (Fig. 5d). The velocity of step advancement is now considerably lower that in the former case.

Table 1. Concentrations of the solutions used in the AFM growth experiments.

<table>
<thead>
<tr>
<th>Solution n°</th>
<th>[CaCl(_2)] (mmol/L)</th>
<th>[MnCl(_2)] (mmol/L)</th>
<th>[Na(_2)CO(_3)] (mmol/L)</th>
<th>X(_{\text{aq}}^{\text{M}^{2+}})</th>
<th>Ionic Strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.215</td>
<td>0.01</td>
<td>0.3</td>
<td>0.021</td>
<td>1.31 x 10(^{-3})</td>
</tr>
<tr>
<td>2</td>
<td>0.215</td>
<td>0.025</td>
<td>0.3</td>
<td>0.051</td>
<td>1.32 x 10(^{-3})</td>
</tr>
<tr>
<td>3</td>
<td>0.20</td>
<td>0.05</td>
<td>0.3</td>
<td>0.046</td>
<td>1.34 x 10(^{-3})</td>
</tr>
</tbody>
</table>
are related to the Mn concentration in solution. For solution 3, step growth is completely inhibited during the observation time. However, the higher Mn concentration in solution 3 results in an important modification of growth behaviour: the transition to a two-dimensional growth mechanism. This can be observed in Figure 7 where the formation of a number of two-dimensional nuclei on the surface occurs simultaneously with the step advancement. As growth proceeds, neighbouring nuclei come into contact and coalesce, forming a whole new layer after a short time. The growth sequence shows again the structural control of the calcite surface on the growth process. In this case, the effect is even more pronounced. The newly formed surface not only avoids the step advancement, but also prevents the formation of these two-dimensional nuclei on it. The dotted line in Figure 7b indicates the original position of the step edge, and it separates a region where nucleation density is very high (left) from another region, generated during the growth experiment, where nuclei formation is inhibited. Such a phenomenon, together with the coalescence of nuclei, results in the reproduction of the preexisting microtopography. Once the coating of the preexisting surface has taken place, the growth process seems to freeze. However, at higher magnification, the formation of a second generation of tiny nuclei, different from the first ones, can be detected (Fig. 7d). Such nuclei are characteristic of growth on the newly formed surface, and form preferentially close to step edges. Figure 8 shows the two different types of small nuclei. The first type includes very small nuclei exhibiting a high nucleation density. They are invariably formed on the surface created by the advancement of slow steps (arrow 1). In contrast, the second type consists of nuclei that are comparatively bigger and scarcer. Their formation preferentially occurs on the surface generated by the advancement of faster steps (arrow 2).

The growth behaviour described above exhibits a variety of phenomena that occur almost simultaneously: (1) a primary two-dimensional nucleation event, (2) stoppage of growth when they reach previously coated surfaces, and (3) a second-
5. DISCUSSION

The AFM observations described above indicate that the growth behaviour of calcite (1014) faces in the presence of Mn is a consequence of the interplay of thermodynamic, kinetic, structural, and microtopographic factors. Therefore, we will discuss all the phenomena observed on the basis of the ideas presented in section 2 (above).

The most obvious phenomenon revealed by our experiments is the transition from simple step advancement (solutions 1 and 2) to a two-dimensional nucleation growth mechanism (solution 3). This can be explained by comparing the supersaturation distributions for the growth solutions. Figure 9a shows the supersaturation functions, $\beta(x)$, calculated for solutions 1, 2, and 3. As can be seen, these three solutions are supersaturated with respect to the whole range of $\text{Mn}_x\text{Ca}_{1-x}\text{CO}_3$ solid solution compositions. However, while the supersaturation for pure calcite remains constant, the increase of Mn from solution 1 to
Fig. 6. (a) Step advancement for the [441]+ and [481]+ steps vs. time from solutions of different concentrations of Mn. (b) Comparison of the step velocities on the original and the new substrate for the three solutions used.
solution 3 leads to an increase in the supersaturation with respect to the rest of solid solution compositions. Since growth mechanisms operating on a surface depend on supersaturation (Sunagawa, 1987), by increasing the supersaturation, the transitional supersaturation value, $\beta^*$, which separates step advancement and two-dimensional nucleation growth mechanisms, will be exceeded. For the three solutions used, the maximum supersaturation corresponds to a solid composition with a mol fraction $X_{\text{MnCO}_3} \approx 0.5$. Therefore, the transitional supersaturation between growth mechanisms for a solid solution $X_{\text{MnCO}_3} = 0.5$ has to be placed somewhere between the maxima for curves 2 and 3 (point $\beta^*_{\text{MnCO}_3=0.5}$ in Figure 9). In addition, AFM experiments conducted by Teng et al. (1997) have shown that the transitional supersaturation for pure calcite...
Fig. 9. (a) Supersaturation functions, $\beta(x)$, for solutions 1, 2, and 3 calculated using Eq. 2. The $\beta^*$ line represents the variation of the transitional supersaturation between step advancement and two-dimensional nucleation as a function of the solid composition. The dotted lines show the solid compositions for which the two-dimensional nucleation is the only possible growth mechanism (solution 3). (b) Nucleation rate function, $J(x)$, expressed in nuclei/$\mu$m$^2$·s for solution 3. The basic parameters introduced in Eqn. 3 were taken from the literature (see text), and the correction factor for heterogeneous nucleation, $\xi$, was adjusted considering the nucleation densities observed in the AFM experiments. The maximum of such a function gives the most kinetically favoured solid solution composition.
differences in chemistry and surface properties of pure calcite
kinetics. The calculation of the nucleation rate function, \( J(x) \), 
Thus, an explanation of such an effect must be found in the 
tween the calcite {1014} surface structure and the supersatu­
tion function using 
the f3(x) function does not provide information about the nucleation 
ation rate as a fimction of the solid composition for solution 
saturations. These authors obtained crystals with molar frac­
tions similar to our solution 
Minerals with molar fractions 0.05 < 0.2. 
Note that we do not assume equilibrium compositions for the 
precipitates. On the contrary, the composition will be deter­
determined by the nucleation kinetics, i.e., the value of the nucle­
aton rate for the different solid solution compositions. The 
most probable composition will be that for which nucleation 
rate is maximum. Moreover, our considerations are indepen­
dent of the KmcCO3 chosen. Note that we calculated the super­
saturation function using KmcCO3 = 10^{-9.43} (McKeeth et al., 
1998). This is the highest value proposed for KmcCO3 by 
different authors (see section 2.1.1 above). Calculation of super­
saturation using any other value for KmcCO3 would lead to 
higher f3(x) distributions. In such a case, the range of possible compositions for the precipitate would be wider. However, our main arguments would still be valid.

Although the characteristics of supersaturation and nucleation 
rate functions provide a general view of the growth process, our AFM observations clearly reveal that the actual growth behaviour is strongly controlled by the interaction between the calcite (1014) surface structure and the supersatu­
ated solutions. Such a control has as the most obvious effect the inhibition by first-grown layers on subsequent step advancement, which results in the reproduction of the preexisting topography. This is directly related to the Mn concentration in the aqueous solution. The higher Mn concentration in solution 
and, therefore, the higher incorporation in growing steps), the more exact is the reproduction of the preexisting topography. Thus, an explanation of such an effect must be found in the differences in chemistry and surface properties of pure calcite and the newly formed Mn,Ca_{1-x}CO_{3} layers. Because of the differences in ionic radii between Ca^{2+} (0.99 Å) and Mn^{2+} (0.80 Å), the incorporation of a significant amount of Mn in the growing calcite steps can produce distortions around the Mn and Ca atoms. As a result, the terrace surface becomes crum­
pred and irregular and, on average, the lattice parameters are reduced, causing a decrease of step height. Both (the difference in height and the formation of irregular terraces, which will be more important for larger substitutions of Ca by Mn), are the first barriers to the advancement of the growing step on the new-formed surface. In addition, as we mentioned above, the increase in Mn concentration of the aqueous solutions implies an increase in supersaturation and, consequently, an increase in both growth rates and Mn incorporation. The formation of Mn-rich calcites at high supersaturations seems to be associated with poorly organised and defective surfaces. This has been demonstrated by the diffuseness of X-ray patterns (Goldsmith and Graf, 1957) that can be related to random inclusions of water molecules in the structure (Lippmann, 1973) and by the observation of crystals with rounded and rough faces (Fernández-Gonzalez and Prieto, 1999). Evidence of the accumulation of defects on calcite (1014) surfaces containing Mn was obtained by passing deionised water over a surface after growth 
from an Mn^{2+}-Ca^{2+}-CO_{3}^{2-} solution (solution 1). As can be observed in Figure 10, dissolution proceeds quickly by develop­
ing a high density of small etch pits on those areas where Mn,Ca_{1-x}CO_{3} layers were previously formed. This is clearly indicative of the presence of a high number of point defects (Sangwal, 1987). Finally, Mn and Ca have a different physico­
chemical behaviour (e.g., hydration and absorption energies), and this will be reflected in differences in surface speciation (Van Cappellen et al., 1993). All these factors can be sufficient to prevent, at least momentarily, the advancement of the growth step on the newly formed Mn-bearing substrate, resulting in the reproduction of the microtopographic features of the original surface.

In situ AFM observations have also shown that the new substrate affects not only the step advancement, but also the formation of two-dimensional nuclei (solution 3). On the newly grown surfaces, nuclei are scarcer and smaller than those previously grown on pure calcite (1014) surfaces. In addition, as we described in section 4 (above), the second generation of nuclei are of two types: (1) small and with high nucleation density on the slow-growth steps and (2) bigger and with low nucleation density on the fast-growth steps. This heterogeneous distribution of nuclei is consistent with the specific incorporation behaviour expected for the Mn in the calcite model by Reeder (1996). The steps with lower velocities (denoted by −) correspond to those steps presenting an acute angle (close kink site), where the incorporation of Mn is preferred. A higher Mn content in these areas promotes the formation of a high number of small nuclei. In contrast, on the surfaces generated by the + steps, where the Mn content is presumably lower, nuclei have a low nucleation density and they reach a larger size.

The phenomena described and discussed in this paper demon­
strate the complexity of the surface processes, which can be observed when the Mn,Ca_{1-x}CO_{3} solid solution grows on cal­
cite (1014) surfaces. The main reason for such complexity is the fact that growing surfaces are not only the simple result of the crystallization of a solid from a supersaturated solution, but they also play an important role on the whole growth processes.
Although a complete elucidation of the relations between composition and microtopographic features of the surfaces in the Mn-Ca-CO$_3$H$_2$O system was beyond the scope of this work, it seems clear that the combination of in situ AFM observations, calculations based on the SS-AS theory, and surface structure models provide an adequate approach to the study of the formation of solid solutions on preexisting surfaces.

Acknowledgments—We thank the Deutsche Forschungsgemeinschaft (DFG) for financial support of this work. J. M. Astilleros thanks the Universidad Complutense de Madrid for a research grant (2000) and the European Commission for the receipt of a Marie Curie Fellowship (from July 2001). The authors appreciate the careful reviews by Dr. H. Gamsjäger, Dr. R. Reeder, an anonymous reviewer, and comments by Dr. A. Mucci.

Associate editor: A. Mucci

REFERENCES


