1 INTRODUCTION

1.1 Recent progress in carbonate mineral dissolution kinetics

Significant progress has been made in understanding the mechanistic details of dissolution dynamics on the (104) cleavage surface of carbonate minerals. Much of this recent work has focused on AFM observations of the rate of step and kink movement (e.g. Liang et al. 1996a, b, Jordan & Rammensee 1998, Lea et al. 2001, Higgins et al. 2002). Observations of dissolution rates over larger lateral scales have also provided basic insight into the relationships of etch pit development and defect distribution (MacInnis & Brantley 1992, 1993, Luttge et al. 2003), as well as comparison with the overall “bulk” rate measured in mineral powders (Arvidson et al. 2003). In addition, considerable insight has been generated by integrating surface complexation models and observations (Van Cappellen et al. 1993, Stipp et al. 1994, Pokrovsky et al. 1999, Fenter et al. 2000, Pokrovsky & Schott 2002) with an atomistic description of surface topography and measurement of step velocities (Duckworth & Martin 2003). This general approach also has the potential to yield substantial detail regarding inhibition mechanisms in dissolution reactions as well. In this brief paper, we describe preliminary experimental observations of magnesium inhibition of calcite dissolution, and suggest how this inhibition may be related mechanistically to site-specific surface properties.

1.2 Previous work on Mg inhibition of calcite dissolution

Previous data on magnesium inhibition of calcite dissolution kinetics have been derived almost entirely through conventional experiments using mineral powders. These early data in general confirm inhibition, but differ in terms of the concentration required to impose a given reduction in rate. These variations most likely reflect the inherent diversity of experimental conditions: pH, $\Delta G^r$, ionic strength, composition of supporting electrolytes, solid phase characteristics, and other factors (e.g. Berner 1967). Subsequent to work on trace metal inhibition by Terjesen et al. (1961) and Nestaa & Terjesen (1969), Sabbides & Koutsoukos (1995) observed dissolution to be inhibited at Mg$^{2+}$ concentrations between 40 and 460 $\mu$M (rates reduced by 0.3 and 0.5 log$_{10}$ units, respectively) in experiments performed close to calcite saturation. In contrast, although Gutjahr et al. (1996) found a reduction in calcite growth rates, they found Mg to have no effect on dissolution at concentrations of 300 $\mu$M Mg(NO$_3$)$_2$. In dilute, mixed Mg$^{2+}$–Ca$^{2+}$ solutions, Sjöberg (1978) concluded inhibition of calcite dissolution to be sufficiently complex as to preclude incorporation of both these components in a general rate equation. Compton & Brown (1994) found a significant reduction in rate in the presence of 10 – 80 mM Mg$^{2+}$, (~50% at 40 mM, pH 8 – 9), which they found to be consistent with a Langmuir-Volmer adsorption isotherm. Alkattan et al. (2002) studied dissolution of calcite single crystals in rotating disk experiments and found no inhibition at MgCl$_2$ concentrations up to 0.1 molal (pH 1 – 3).
2 METHODS

We measured Mg inhibition of calcite dissolution on single crystal (104) surfaces by two approaches. Absolute surface-normal retreat rates were derived from time-lapse changes in surface topography by \textit{ex situ} vertical scanning interferometry (VSI), after flowing reactant solution over this surface in a fluid microcell (for details, see Luttge et al. 1999, 2003, Arvidson et al. 2003). This method allowed quantitative mapping of relatively large areas of the mineral surface at subnanometer vertical resolution (Fig. 1). The second approach involved monitoring similarly prepared calcite surfaces with reactant solutions in an AFM flow cell (Digital Instruments Nanoscope III Multimode SPM, operated in contact mode). All experiments used cleavage rhombs derived from high purity synthetic calcite. Carbonated reactant solutions were prepared from Na$_2$CO$_3$ or NaHCO$_3$ (4.4 meq/L) continuously equilibrated with atmospheric CO$_2$ (pH 8.8 – 9.0, 25°C; cf. Liang et al. 1996b, Lea et al. 2001). Mg was added (as MgCl$_2$) in concentrations of 0, 10, 50, 300, and 800 µM. Additional experiments were also performed as a function of pH in N$_2$-sparged (CO$_2$-free) solutions at the same ionic strength and MgCl$_2$ concentrations.

![100 µm scale bar](image)

**Figure 1.** VSI data (10× magnification, white light) from coalescing etch pits on calcite (104) surface, reacted in 4.4 mM NaHCO$_3$, 800 µM MgCl$_2$, maintained at pH ~ 9, PCO$_2$ ~10$^{-3.5}$, 25°C. Positive step directions [−441] and [48–1] are oriented to the right and down, respectively.

3 RESULTS AND DISCUSSION

In our VSI runs, we observed that inhibition occurs in carbonated solutions (pH ~9) at relatively low concentrations: relative to the rate in Mg-free NaHCO$_3$ solution, the addition of 10 µM MgCl$_2$ reduces the dissolution rate by ~80%, and 300 µM reduces the rate by ~90%. At higher concentrations, under conditions in which the solution is supersaturated with respect to pure magnesite, we observed that etch pit morphology departed significantly from the symmetrical rhombic shape observed in Mg- and CO$_2$-free solutions. Etch pit morphologies in carbonated Mg-free solutions (pH ~ 9) showed rounding at the intersection of the obtuse (+) step edges, such that the radius of the (+) steps decrease toward their common terminus. Lea et al. (2001) showed this morphology to be associated with a reduction in (+) step velocity, and attributed this inhibition to probable adsorption of carbonate ion at (+,+ kink sites. In our experiments, carbonated solutions having the highest (~800 µM) Mg concentrations produced (+,+ step intersections that remained relatively sharp but instead curved toward the (+,−) step intersections (Fig. 1). These changes implied changes in kink density along step edges, and variations in the kink nucleation rate.

![Figure 2](image)

**Figure 2.** Evolution of step edges showing relationships between double kink nucleation rate ($R_{KK}$), kink migration rate ($R_{K}$), kink density, and step movement (adapted from Liang et al. 1996a). Time $t$ increases from top to bottom. (A) In order for steps to remain straight, kink density must remain low: either no new double kinks can nucleate prior to the exit of single kinks at the step terminus (self annihilation), or single kinks formed from adjacent double kinks must annihilate each other (kink-kink annihilation). In both cases, $R_K$ must remain high relative to $R_{KK}$. (B) Rough, irregular step profiles should thus result from inhibition of $R_K$ relative to $R_{KK}$. “Curved” step profiles can be produced by a coupled reduction in $R_K$ relative to $R_{KK}$ and a pronounced anisotropy in single kink migration rates, i.e. $R_{K+} \neq R_{K−}$.

Step curvature and roughness can be understood in the context of the rate of two related processes: double kink nucleation ($R_{KK}$) and single kink propagation of either positive ($R_{K+}$) or negative ($R_{K−}$) kink sites. As shown in Figure 2, if $R_{KK}$ is slow relative to $R_K$, then kinks will tend to annihilate one another before new kinks can form. Conversely, if $R_K$ is reduced to the point where $R_{KK}$ is competitive, then existing kinks may be preserved prior to the next cycle of kink nucleation. Liang et al. (1996a) elegantly demonstrated that simple kink annihilation at the step terminus was insufficient to explain both straight step morphology and constant step velocity, and postulated that the persistence of straight steps must reflect a balance between $R_{KK}$ and the annihilation of adjacent kink pairs.
Figure 3. Calcite 104 cleavage surface. (A) Orientation of cleavage planes, hexagonal unit cell, obtuse (+) step faces, Ca atoms (large gray), and trigonal CO3 groups. Out-of-plane, in-plane, and below-plane oxygens coordinate each carbon atom. (B) Ca-O coordination at kink sites, showing monatomic etch pit (black outline), trigonal CO3 groups (gray), and Ca atoms (shown either as large black atoms, stars, or omitted for clarity). The length of Ca-O bonds between surface calciums (large black) and coordinating oxygens (small gray) at different kink sites have the same length. However, oxygen-oxygen distances within these truncated coordination octahedra are not equivalent: oxygens coordinating Ca at the (+,+), (+,–) and (–,–) metal sites (Fig. 3).

Morphologic changes occur at Mg concentrations substantially greater than those required to induce significant overall inhibition. Because movement of the “fast” (+) steps is responsible for much of the bulk dissolution (Liang et al. 1996b), inhibition at these sites may occur with relatively little inhibitor addition, similar to the effect observed for dissolved carbon and Mn2+ (Lea et al. 2001, Arvidson et al. 2003). It is unlikely that these morphologies reflect simple adsorption of dehydrated Mg2+ ions at specific kink sites, as they were not observed in CO2-free solutions (pH 8 – 11+). Formation of ion pairs MgCO3 or MgHCO3+ scavenge free carbonate ion from solution, and thus may reduce the carbonate ion concentration at kink sites. These components (as well as Mg2+) may also form significant surface complexes as well, and their interactions result in overall inhibition (Arvidson et al., in prep.).

Figure 4. AFM reaction sequence. (A) 10×10 µm image of large etch pit previously formed in Mg-free solution, showing “CO3” inhibition of obtuse (+) step directions by rounding towards the (+,+) intersection. Etch pit at lower right formed immediately after introduction of 800 µM MgCl2. (B) 1×1 µm image detail of (+,+) corner (left of arrows in A), contrasting rough step edges of “CO3”-inhibited etch pit with those of newly formed pit (appearing after MgCl2 introduction) having sharp (+,+) corners. (C, D) Growth of etch pit via advance of acute (–) steps in 800 µM MgCl2 solution, showing coalescence with static (+,+) intersection of older etch pit. Scan field is 3×3 µm.

The basic anisotropy of obtuse versus acute step velocities on the calcite surface may reflect a complex interaction between the energetics of removal of cationic and anionic components broadly mediated by dominant surface complex populations (>MOH2+ versus >CO3H; Duckworth & Martin, 2003). Because of the oblique orientation of carbonate groups with respect to the (104) surface, these sites differ in terms of coordination. For example, although removal of surface calcium demands cleavage of Ca-O bonds, the O-O distances within the surface-truncated Ca-octahedra vary according to their crystallographic orientation at (+,+), (+,–) and (–,–) metal sites (Fig. 3).

The relationship of step movements and inhibition can also be clarified by study of time-lapse AFM data. Figure 4 shows a sequence in which a previously developed etch pit, grown in carbonated Mg-free solution, is the site of new etch pit formation upon introduction of 800 µM carbonated MgCl2 into the flow cell. This sequence definitively shows that (1) the (+) steps are severely inhibited after Mg introduction, with little measurable advance, and (2) growth of new etch pits occurs by advance of (–) steps. Although dissolved carbon (“CO3”, Fig. 4) reduces (+) step velocities (Lea et al. 2001), the fact that step curvature towards the (+,+) corner, previ-
ously developed under Mg-free conditions, is retained, while newly formed pits at 800 μM MgCl₂ show relatively sharp (+,+) corners, suggests that the (+) steps are fully passivated, and that the rates of both double kink nucleation and single kink migration along either [48–1], or [–441], are low. Curvature of the (+) steps may instead reflect ineffective kink removal at the (–,+) corner, implying that the rate of single kink migration ($R_{K+}$) along the (+) steps is inhibited in the (–) direction as well. However, for the (–) steps to remain straight (Fig. 4) also implies that $R_K >> R_{KK}$.

4 CONCLUSIONS

We have documented magnesium inhibition of calcite dissolution in alkaline solutions far from equilibrium. As is the case for dissolved carbon addition, inhibition from increasing Mg concentration is associated with unique etch pit morphologies, reflecting possible changes in the relative rates of double kink nucleation and single kink migration. These changes are also linked to dissolved carbon concentration, and may reflect interaction of Mg ion pairs with the surface. These relationships will be explored in future modeling work.

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REFERENCES


