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Reaction kinetics of dolomite rim growth

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Abstract

Reaction rims of dolomite (CaMg(CO3)2) were produced by solid-state reactions at the contacts of oriented calcite (CaCO3) and magnesite (MgCO3) single crystals at 400 MPa pressure, 750-850 °C temperature and 3-146 h annealing time to determine the reaction kinetics. The dolomite reaction rims show two different microstructural domains. Elongated palisades of dolomite grew perpendicular into the magnesite interface with length ranging from about 6 µm to 41 µm. At the same time, a 5-71 µm wide rim of equiaxed granular dolomite grew at the contact with calcite. Platinum markers showed that the original interface is located at the boundary between the granular and palisade-forming dolomite. In addition to dolomite, a 12 to 80 µm thick magnesio-calcite layer formed between the dolomite reaction rims and the calcite single crystals. All reaction products show at least an axiotactic crystallographic relationship with respect to calcite reactant, while full topotaxy to calcite prevails within the granular dolomite and magnesio-calcite. Dolomite grains frequently exhibit growth twins characterized by a rotation of 180° around one of the [1120] equivalent axis. From mass balance considerations it is inferred that the reaction rim of dolomite grew by counter diffusion of MgO and CaO. Assuming an Arrhenius type temperature dependence, activation energies for diffusion of CaO and MgO are Ea (CaO) = 192 ± 54 kJ/mol and Ea (MgO) = 198 ± 44 kJ/mol, respectively.

Introduction

Mineral reactions in the solid-state are ubiquitous in the Earth’s crust and mantle. Unless deformed or overprinted, the product phases of such solid-state reactions are commonly located along former phase boundaries between reactants, forming polycrystalline or polyphase reaction rims or coronas (ASHWORTH ET AL. 1998; KELLER ET AL. 2006; KELLER, WIRTH, ET AL. 2008; KELLER, WUNDER, ET AL. 2008). They may contain information about the pressure, temperature and time conditions of their formation and therefore provide a useful indication on the geological history (KELLER, WIRTH, ET AL. 2008; JOACHIM ET AL. 2010).

For a solid-state reaction under “dry” conditions, the necessary transfer of matter between the reactants occurs by diffusion leading to reaction rim growth with kinetics that depends on P-T-t conditions (LASAGA 1983). To constrain the diffusion kinetics using microstructure analysis, knowledge of the component mobility and the transport mechanisms are required. In a polyphase aggregate, the transfer of chemical components may occur via a
combination of grain boundary- and volume diffusion, with grain boundary (GB) diffusion
typically being several orders of magnitude faster than diffusion through the volume of the
grains. GB diffusion thus is effective over larger spatial scales (DOHMEN AND MILKE 2010).
However, at elevated temperatures the contribution of volume diffusion to the bulk mass transfer
may increase considerably owing to higher activation energy than for grain boundary diffusion
(MROWEĆ 1980). In general, the growth behaviour of solid-solid reaction rims can be described
by the relation: \(\Delta x^n = kt\), where \(\Delta x\) is the thickness of the reaction layer, \(k\) is a rate constant, \(t\) is
time and \(n\) is a parameter depending on the reaction controlling mechanism (SCHMALZRIED 1978;
KELLER ET AL. 2010). For interface-reaction controlled growth \(n = 1\) and for diffusion controlled
growth \(n = 2\). It has been demonstrated by ABART AND PETRISHCHEVA (2011) that, after an
incipient nucleation phase, reaction rim growth is initially interface reaction-controlled, replaced
by diffusion-controlled growth as the rim thickness increases. From the rate of reaction rim
growth in the diffusion controlled kinetic regime the mobility of the respective components can
be inferred. This motivated a series of rim growth experiments that aimed to determine
component mobility. Dedicated studies exist for several systems such as CaCO\(_3\)-SiO\(_2\), where
wollastonite is formed between calcite and quartz (MILKE AND HEINRICH 2002), MgO-SiO\(_2\),
where enstatite is formed between forsterite and quartz, or enstatite-forsterite double layers are
formed between periclase and quartz (e.g. GARDÉS & HEINRICH, 2011; GÖTZE ET AL., 2009;
MILKE, WIEDENBECK, & HEINRICH, 2001), and MgO-Al\(_2\)O\(_3\) where spinel is formed between
periclase and corundum (e.g. GÖTZE ET AL., 2009; KELLER ET AL., 2010; WATSON & PRICE,
2002). For grain boundary diffusion-controlled growth, the reaction kinetics may be influenced
by grain size (GARDÉS ET AL. 2011; GARDÉS AND HEINRICH 2011), water content (GARDÉS ET
AL. 2012; MILKE ET AL. 2009; MILKE ET AL. 2012), and non-hydrostatic stress (GÖTZE ET AL.
2009; KELLER ET AL. 2010). In contrast, little is known about diffusion-controlled reactions in the
system CaCO\(_3\)-MgCO\(_3\), i.e. the formation of dolomite [(CaMg(CO\(_3\)))\(_2\)] between calcite (CaCO\(_3\))
and magnesite (MgCO\(_3\)). So far, few cation diffusion data are available for diffusion in dolomite.
ANDERSON (1972) determined self-diffusion of C and O in dolomite at \(T = 645\) to 785°C and \(P = 12\) to 93.5 MPa and MÜLLER, CHERNIAK, & WATSON (2012) investigated cation exchange
between dolomite and siderite or rhodochrosite to determine volume diffusion of (Mn, Fe) – Mg – Ca at \(T = 400-625°C\), \(P = 0.1\) MPa.

In this study reaction rims of dolomite were produced at contacts between oriented
natural calcite and magnesite single crystals following the reaction:

\[
\text{MgCO}_3\text{solid} + \text{CaCO}_3\text{solid} = \text{CaMg(CO}_3\text{)}_2\text{solid}
\]  

(1)

The experiments were performed at \(T = 750-850°C\) and \(P_c = 400\) MPa to determine the reaction
kinetics of dolomite rim growth in the carbonate system under static annealing conditions.
Experimental and analytical details

**Starting material**

The starting materials were prepared from natural calcite single crystals from Mina Prieta Linda, Naica (Chihuahua, Mexico) and Minas Gerais (Brasil) and from magnesite single crystals from Bahia (Brumado, Brasil). The chemical composition was analyzed by field emission gun electron microprobe (JEOL JXA-8500 F HYPERPROBE), using 15 keV accelerating voltage, 5 nA beam current and 1 µm beam diameter. The results are given in Table 1, revealing trace amounts of Fe and Mg in calcite. Magnesite contained impurities of Ca, Fe and traces of Mn, Ba, and Sr. The water content estimated from Vario EL III element analyser (Elementar Analysysysteme GmbH) is about 0.15 wt.% and 0.25 wt.% in calcite and magnesite, respectively.

<table>
<thead>
<tr>
<th></th>
<th>Calcite ± 1σ</th>
<th>Magnesite ± 1σ</th>
</tr>
</thead>
<tbody>
<tr>
<td>MgO</td>
<td>0.03 ± 0.04</td>
<td>46.08 ± 1.65</td>
</tr>
<tr>
<td>FeO</td>
<td>0.04 ± 0.05</td>
<td>0.18 ± 0.05</td>
</tr>
<tr>
<td>CaO</td>
<td>53.37 ± 0.76</td>
<td>0.26 ± 0.04</td>
</tr>
<tr>
<td>SrO</td>
<td>0.02 ± 0.03</td>
<td>0.02 ± 0.01</td>
</tr>
<tr>
<td>MnO</td>
<td>0.01 ± 0.03</td>
<td>0.04 ± 0.03</td>
</tr>
<tr>
<td>TiO₂</td>
<td>0.01 ± 0.02</td>
<td>0.01 ± 0.01</td>
</tr>
<tr>
<td>SiO₂</td>
<td>0.00 ± 0.01</td>
<td>0.01 ± 0.02</td>
</tr>
<tr>
<td>BaO</td>
<td>0.00 ± 0.01</td>
<td>0.04 ± 0.03</td>
</tr>
<tr>
<td>CO₂</td>
<td>46.51 ± 0.74</td>
<td>53.36 ± 1.66</td>
</tr>
</tbody>
</table>

*Table 1* Composition of starting materials in wt.%. Values represent mean data based on 5 point analyses. CO2 contents were calculated assuming an oxide total of 100%

**Annealing experiments**

Cylindrical samples of 3-5 mm length and 7 mm diameter were drilled out of the single crystals, oriented with the cylinder axis perpendicular to the cleavage planes of the rhombohedra \(((10\overline{4}), (\overline{1}104), (0\overline{1}14))\]. Subsequently samples were polished with diamond powder to a surface roughness of 1 µm. In three experiments (CaMg-13, CaMg-14, CaMg-15, Table 2), both contact surfaces of calcite and magnesite were polished to 0.25 µm finish and sputtered with a thin platinum layer (< 2nm), which served as a marker to locate the initial calcite-magnesite interface after formation of the rims. The assembly for each run consisted of a calcite-magnesite sandwich, located between alumina spacers and adjacent alumina pistons Figure 1.
Figure 1 Experimental setup. The sample assembly consists of polished cylinders of calcite and magnesite single crystals. The dolomite reaction rim grows from the initial calcite-magnesite contact into both directions. The initial calcite-magnesite contact is marked by platinum markers (white dots) in some experiments. Adjacent magnesio-calcite evolved next to calcite reactant.

The whole assembly was jacketed by a copper-sleeve to prevent intrusion of the confining pressure medium (argon-gas). Temperature was measured using a Pt/Pt-13%Rh (R-type) thermocouple 3 mm from the sample assembly. All experiments were performed in an internally heated pressure vessel using a Paterson type gas deformation apparatus (PATerson 1970) at a confining pressure of $P_c = 400$ MPa. Temperatures were kept constant at 750°C, 800°C, 825°C or 850°C with a heating rate of 20°C/min and a cooling rate of 2°C/min (Table 2). After the experiments, the jacketed sample assembly was cut along the cylinder axis using a slow-speed diamond saw, embedded in epoxy and the surfaces were polished down to 1 µm roughness for analysis of the reaction rim.
Table 2 Average thickness of entire dolomite ($\Delta x_{\text{Dol}}$), dolomite palisades ($\Delta x_{\text{Pal}}$), granular dolomite layers ($\Delta x_{\text{gran}}$), and magnesio-calcite ($\Delta x_{\text{Mg-Cal}}$). $a_{\text{Pal}}$, $a_{\text{gran}}$ and $a_{\text{Mg-Cal}}$ are the mean grain diameter of dolomite palisades, granular dolomite and magnesio-calcite, respectively. $D_{\text{CaO}}$ and $D_{\text{MgO}}$ are the derived diffusion coefficients of CaO and MgO. Uncertainties are given as 1σ value.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>$T$ (K)</th>
<th>$t$ (h)</th>
<th>$\Delta x_{\text{Dol}}$ ($\mu$m)</th>
<th>$\Delta x_{\text{Pal}}$ ($\mu$m)</th>
<th>$\Delta x_{\text{gran}}$ ($\mu$m)</th>
<th>$\Delta x_{\text{Mg-Cal}}$ ($\mu$m)</th>
<th>$a_{\text{Pal}}$ ($\mu$m)</th>
<th>$a_{\text{gran}}$ ($\mu$m)</th>
<th>$a_{\text{Mg-Cal}}$ ($\mu$m)</th>
<th>$\log D_{\text{CaO}}$ (m²/s)</th>
<th>$\log D_{\text{MgO}}$ (m²/s)</th>
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<tr>
<td>CaMg-02</td>
<td>1073</td>
<td>74.03</td>
<td>36.3 ± 5.3</td>
<td>16.5 ± 3.1</td>
<td>55.0 ± 10.7</td>
<td>3.6 ± 0.9</td>
<td>5.4 ± 1.9</td>
<td>39.0 ± 20.1</td>
<td>-14.4 ± 0.3</td>
<td>-14.2 ± 0.3</td>
<td></td>
</tr>
<tr>
<td>CaMg-03</td>
<td>1023</td>
<td>73.95</td>
<td>14.9 ± 5.3</td>
<td>10.5 ± 4.0</td>
<td>40.9 ± 10.4</td>
<td>3.1 ± 1.1</td>
<td>2.7 ± 1.1</td>
<td>23.3 ± 12.9</td>
<td>-14.6 ± 0.4</td>
<td>-14.6 ± 0.4</td>
<td></td>
</tr>
<tr>
<td>CaMg-04</td>
<td>1023</td>
<td>50.07</td>
<td>17.5 ± 5.7</td>
<td>9.0 ± 2.8</td>
<td>27.3 ± 8.3</td>
<td>2.9 ± 0.9</td>
<td>3.2 ± 1.3</td>
<td>22.2 ± 16.0</td>
<td>-14.6 ± 0.5</td>
<td>-14.5 ± 0.4</td>
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<tr>
<td>CaMg-05</td>
<td>1023</td>
<td>145.77</td>
<td>26.0 ± 6.0</td>
<td>12.3 ± 3.0</td>
<td>37.2 ± 6.1</td>
<td>3.9 ± 1.0</td>
<td>4.0 ± 1.3</td>
<td>37.3 ± 17.3</td>
<td>-14.8 ± 0.4</td>
<td>-14.7 ± 0.4</td>
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<td>CaMg-06</td>
<td>1073</td>
<td>144.71</td>
<td>40.9 ± 7.1</td>
<td>16.2 ± 4.2</td>
<td>74.2 ± 11.6</td>
<td>4.3 ± 1.4</td>
<td>4.2 ± 1.4</td>
<td>44.3 ± 23.9</td>
<td>-14.7 ± 0.3</td>
<td>-14.4 ± 0.3</td>
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<tr>
<td>CaMg-07</td>
<td>1073</td>
<td>84.82</td>
<td>30.0 ± 7.2</td>
<td>14.7 ± 3.3</td>
<td>37.0 ± 7.3</td>
<td>4.1 ± 1.2</td>
<td>4.6 ± 1.4</td>
<td>35.7 ± 15.4</td>
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<td>CaMg-08</td>
<td>1073</td>
<td>48.1</td>
<td>25.4 ± 7.1</td>
<td>12.1 ± 3.4</td>
<td>48.5 ± 8.2</td>
<td>3.3 ± 0.8</td>
<td>3.9 ± 1.2</td>
<td>38.2 ± 23.4</td>
<td>-14.4 ± 0.5</td>
<td>-14.2 ± 0.5</td>
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<td>CaMg-09</td>
<td>1023</td>
<td>2.44</td>
<td>2.9 ± 1.2</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>12.7 ± 4.7</td>
<td>NA</td>
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<tr>
<td>CaMg-10</td>
<td>1123</td>
<td>74.15</td>
<td>103.7 ± 33.5</td>
<td>41.3 ± 10.2</td>
<td>80.3 ± 21.2</td>
<td>5.0 ± 1.2</td>
<td>4.3 ± 1.3</td>
<td>45.8 ± 32.9</td>
<td>-13.4 ± 0.7</td>
<td>-13.4 ± 0.7</td>
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<td>CaMg-12</td>
<td>1073</td>
<td>4.03</td>
<td>11.3 ± 4.2</td>
<td>6.0 ± 2.4</td>
<td>20.0 ± 5.5</td>
<td>1.7 ± 0.4</td>
<td>2.6 ± 0.9</td>
<td>18.5 ± 9.7</td>
<td>-13.9 ± 0.6</td>
<td>-13.8 ± 0.6</td>
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<tr>
<td>CaMg-13*</td>
<td>1098</td>
<td>74.92</td>
<td>40.1 ± 9.5</td>
<td>17.3 ± 5.1</td>
<td>52.3 ± 13.3</td>
<td>4.3 ± 1.1</td>
<td>1.4 ± 0.8</td>
<td>42.0 ± 27.7</td>
<td>-14.2 ± 0.5</td>
<td>-14.2 ± 0.5</td>
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<tr>
<td>CaMg-14*</td>
<td>1073</td>
<td>28.94</td>
<td>14.2 ± 4.9</td>
<td>8.3 ± 3.1</td>
<td>31.2 ± 8.3</td>
<td>2.4 ± 0.8</td>
<td>2.9 ± 1.2</td>
<td>26.1 ± 15.4</td>
<td>-14.4 ± 0.5</td>
<td>-14.4 ± 0.5</td>
<td></td>
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<tr>
<td>CaMg-15*</td>
<td>1023</td>
<td>29.13</td>
<td>13.0 ± 1.6</td>
<td>6.8 ± 2.4</td>
<td>31.5 ± 10.2</td>
<td>2.6 ± 0.6</td>
<td>2.5 ± 0.8</td>
<td>26.6 ± 14.0</td>
<td>-14.6 ± 0.2</td>
<td>-14.6 ± 0.2</td>
<td></td>
</tr>
</tbody>
</table>

*Platinum marker experiment
The run conditions and the results of 13 annealing experiments are presented in Table 2. Dolomite reaction rims were produced during static annealing tests of 13 samples with annealing times between 3 and 146 hours (Table 2).

Figure 2 shows the polybaric (200-850 MPa) phase diagram for CaCO$_3$-MgCO$_3$ adopted from Goldsmith and Heard (1961), revealing that both dolomite and magnesio-calcite are stable at the applied experimental conditions. Compared to the solvi shown in Figure 2, the experimental pressure of 400 MPa slightly increases the solubility of MgCO$_3$ in calcite by about 1 mol % at 750°C and 2% mol % at 850°C, respectively (Gottschalk and Metz 1992).

Analytical methods

For determination of the dolomite and magnesio-calcite reaction layer thicknesses we used an optical microscope (Leica DM RX) to obtain a complete set of reflected-light micrographs across the whole sample diameter. The width of reaction rims was subsequently analyzed by digitizing the location of phase boundaries and calculation of average values owing to the relatively large thickness variation up to 41 % along a single profile. The associated run durations were corrected for additional growth during heating and cooling ramps (Lasaga 1983), but the effect is negligible.

Microprobe analysis of the composition profiles across selected reaction rims was performed by quantitative line scans using wavelength dispersive X-ray (WDS) measurements, counting only the major elements Ca (K$_\alpha$, PETJ) and Mg (K$_\alpha$, TAP). For the analysis, we applied an accelerating voltage of 15 kV and a beam current of 5 nA. Counting time was 5 s on the peak, step size 2 µm and the beam diameter was 1 µm.

The crystallographic orientations of the reactant and product phases were measured using a dual-beam scanning electron microscope (FEI Quanta 3d FEG SEM-FIB), equipped with an electron backscatter diffraction detector (EBSD, TSL DigiView) and a semi-quantitative energy dispersive spectroscopy (EDS) mapping device. Since calcite, dolomite and magnesite have very close unit cell parameter a, b, c the EBSD patterns of these three phases are very similar. Therefore, we used EBSD and EDS mapping to discriminate the three minerals by measuring the
Mg and Ca content. First, we mapped the reactants and the reaction rim using only the calcite reflector list, together with EDS analysis. Subsequently, we identified the three-phases based on the Mg/Ca ratios (e.g., Mg-rich – magnesite; Mg/Ca – dolomite; Ca-rich – calcite). Finally, the inspected area was remapped using the reflectors of all three phases. The magnesio-calcite was discriminated separately using microprobe analysis. For EBSD analysis, the samples were first mechanically polished to 0.25 µm roughness using a diamond paste and afterwards chemically-mechanically polished for one hour using an alkaline solution of colloidal silica. To avoid carbon coating, measurements were conducted in low-vacuum with a chamber pressure of 10 Pa of H₂O. Automatic mappings of selected areas were performed using an accelerating voltage of 15 keV, a step size between 0.5 and 2 µm, a working distance of 15 mm and a beam current of 8 nA. The TSL-OIM software (ADAMS, WRIGHT, AND KUNZE 1993) was used to index and analyze the EBSD patterns. Post-acquisition treatment of the raw EBSD data included the standardization of the confidence index (CI) and the CI correlation between neighbour points, assuming a value of 0.2. After processing, the points with CI < 0.2 were removed from the datasets to avoid incorrect measurements. In general, the CI ranges from 0 to 1 and quantifies the relationship between the number of votes that each phase receives during the indexation of patterns in EBSD mapping. A CI value of 0.2 indicates that the patterns are correctly indexed by about ~99 %. Pole figures were plotted as one point per grain in a reference frame where the E-W plane is parallel to the reaction interface, and the pole of this plane indicates the growing direction (GD) of the reaction rim.

Results

Microstructure of reaction rims

All experiments resulted in polycrystalline dolomite reaction rims with two spatially separated regions of 1) elongated palisades oriented perpendicular to the interface with magnesite and 2) granular equiaxed dolomite grains in contact with calcite. In addition, the pure calcite reactant close to the dolomite rim transformed to polycrystalline magnesio-calcite. The layer width was defined petrographically by the occurrence of new formed grains using an optical microscope. Microprobe analyses confirm these new formed grains to be magnesio-calcite with decreasing magnesium content towards pure calcite (see Figure 7). The thickness of the magnesio-calcite layer is about 4 times that of the entire dolomite rim (Table 2). The size of the magnesio-calcite grains increases from the dolomite reaction rim towards the pure calcite starting material (
The grains often show curved grain boundaries with equilibrium angles at triple junctions. The platinum marker experiments indicate that the original interface between the reactants is located between dolomite palisades and granular dolomite. Open cracks between interfaces are expected to evolve during cooling and sample preparation.

**Figure 3** Pt-marker experiment CaMg-14, annealed at 800°C and 29 h run duration. **a** Optical micrograph of polycrystalline dolomite and magnesio-calcite layers between magnesite and calcite single crystals. The dolomite rim consists of palisades oriented perpendicular to the interface with magnesite and of granular dolomite in contact to magnesio-calcite. **b** BSE image showing platinum markers (white spots), which indicate that the original calcite-magnesite interface is located at the boundary between the domains of the palisade-like and granular dolomite. Outliers are attributed to dolomite grain growth during annealing. Open cracks between interfaces are expected to evolve during cooling and sample preparation.
Figure 3 Pt-marker experiment CaMg-14, annealed at 800°C and 29 h run duration. 

**a** Optical micrograph of polycrystalline dolomite and magnesio-calcite layers between magnesite and calcite single crystals. The dolomite rim consists of palisades oriented perpendicular to the interface with magnesite and of granular dolomite in contact to magnesio-calcite. 

**b** BSE image showing platinum markers (white spots), which indicate that the original calcite-magnesite interface is located at the boundary between the domains of the palisade-like and granular dolomite. Outliers are attributed to dolomite grain growth during annealing. Open cracks between interfaces are expected to evolve during cooling and sample preparation.

b)
Kinetics of reaction rim growth

Time series experiments were done at temperatures of 750 and 800°C by varying the annealing time between 3 and 146 hours. For determining the temperature dependence of the reaction kinetics, the temperature was varied between 750 and 850°C at a fixed time of 74 hours. Generally, we find increasing rim thicknesses and grain sizes with increasing annealing time and temperature (Table 2). Figure 4 shows rim thickness as a function of square root of time for all rim growth experiments. Both time series at temperatures of 750°C and 800°C reveal a linear increase of dolomite rim width with increasing square root of time, following a power law form of $\Delta x^2 = kt$ and consequently indicating diffusion-controlled rim growth (JOESTEN 1977; JOACHIM ET AL. 2010; ABART ET AL. 2009). In comparison, at similar temperature and time the magnesio-calcite layer is thicker than the entire dolomite layer, whereas granular and elongated dolomite rims are approximately equal in width.
Figure 4 Reaction product layer width versus square root of time, linear relations are indicated for $T = 750^\circ C$ (dashed lines) and $800^\circ C$ (solid lines). a Total dolomite reaction rim width, b magnesio-calcite width, c dolomite palisades width, and d granular dolomite thickness. Error bars reflect the relatively large thickness variation (12-41%) along each profile of 7 mm length. Note different scales.

For example, for the time series performed at $T = 750^\circ C$, the average thickness of dolomite reaction rims varies between 2.9 and 26 $\mu$m and that of magnesio-calcite between 12.1 and 40.9 $\mu$m, respectively. At $800^\circ C$ the values range from 11.3 to 40.9 $\mu$m for the entire dolomite and 31.2 to 74.2 $\mu$m for the magnesio-calcite layer. The thicknesses of magnesio-calcite and dolomite palisades show a fast non-linear incipient stage (Figure 4 b, c), while granular dolomite growth is retarded (Figure 4 d). Afterwards the layer thicknesses increase linearly with the square root of time following a parabolic growth behavior. Grain coarsening of dolomite palisades, granular dolomite and magnesio-calcite takes place during the experiments. Palisades and granular
dolomite grain sizes increase by a factor of 1.5-1.8 at 750°C and 1.5-1.6 at 800°C during run durations between 29 and 145 hours (Table 2). In the same time span, the average magnesio-calcite grains coarsen about a factor of 1.4 and 1.7 (Table 2).

The different slopes of the regression lines for $T = 750°C$ and $800°C$ shown in Figure 4 indicate that the temperature sensitivity of rim growth is lowest for dolomite palisades and highest for magnesio-calcite. Extending the temperature range to $850°C$ suggests that at a constant annealing time of 74 h the ratio of $\Delta x_{Dol}/\Delta x_{Mg-Cal}$ increases with $T$ and exceeds 1 at about $840°C$ (Figure 5). In contrast, the ratio of $\Delta x_{Pal}/\Delta x_{gran}$ decreases with temperature from 1.4 to 0.58 (Figure 5 b), suggesting a lower temperature sensitivity for growth of dolomite palisades than of granular dolomite.

![Figure 5](image)

**Figure 5** a Dolomite rim/magnesio-calcite thickness ratio versus temperature. The thickness ratio increases with increasing temperature. b Dolomite palisades/granular dolomite thickness ratio versus temperature. The thickness ratio tends to decrease with increasing temperature.

**Texture analyses**

Detailed crystallographic orientation maps were measured in 9 selected samples including the orientation of calcite and magnesite single crystals, polycrystalline dolomite (palisade and granular shapes) and polycrystalline magnesio-calcite. A typical orientation map and pole figures are presented in Figure 6, using the inverse pole figure color coding (see inset for details). The orientations of calcite and magnesite starting material single crystals are constrained by the orientation of one of the three $\{1014\}$ poles parallel to the growing direction (GD), since one of the three symmetric rhomb planes was parallel to the initial reaction interface (Figure 6 a). The resulting bulk texture for dolomite shows $[0001]$ axes for both palisades and granular dolomite concentrated in a position normal to the GD, with secondary concentrations (principally for the palisades) spreading all over the pole figures (Figure 6 a). The poles of $\{2110\}$ prismatic planes are distributed in a more complex form with no clear relation between maximum concentrations and external reference frame. The poles of $\{10\overline{1}0\}$ prismatic planes on the other hand form broad girdles normal to the reaction interface with weak maxima parallel to GD. The $\{10\overline{1}4\}$
rhombs are also distributed in a complex way, and the maxima of these poles tend to lie within the reaction interface plane (Figure 6a). In contrast, magnesio-calcite forms [0001] axes subparallel or oblique to GD, poles of \{2\overline{1}0\} and \{10\overline{1}0\} form broad girdles subparallel to the reaction interface, and poles to \{1\overline{0}1\} form small circles around the growing direction axis (Figure 6a). Although the dolomite orientations show considerable scattering, the bulk texture still indicates some crystallographic orientation relationship to magnesite but primarily to calcite single crystals (see labels 1-6 in Figure 6a). Detailed analysis of EBSD data revealed the presence of dolomite growth twins characterized by a rotation of 180° around one of the three equivalent a-axes (\{1\overline{1}20\} twin axis). The twin grain boundaries represent between 25 and 40% of dolomite-dolomite boundaries in all the analyzed samples (see black lines decorating the twin boundaries in the EBSD map and histogram of misorientation in Figure 6a). Although the twin grains occur within granular and palisades parts of the dolomite rim, the proportion of twin grains prevails in granular dolomite. Furthermore a topotactic relationship, where all crystallographic directions of dolomite are fixed with respect to the calcite single crystal, could be observed in all samples (e.g. Figure 6b). These topotactic grains often appear equiaxed in shape and are characterized by the above-described twin relationship. Even within the magnesio-calcite layer, grains with the topotactic relationship to the calcite single crystal are present in most analyzed samples (e.g. Figure 6b). The dolomite bulk texture diagrams (Figure 6a) indicate axiotactic (only one crystallographic direction parallel) relationships of dolomite \{2\overline{1}10\} and \{10\overline{1}0\} axes to both calcite and magnesite single crystals (labeled 2, 5 and 6 in Figure 6a). Furthermore, weak axiotactic relationship of rhombohedral crystallographic directions in dolomite to single crystal calcite may be suspected from the dolomite bulk texture diagrams (labeled 3 and 4 in Figure 6a).
Figure 6 a EBSD inverse pole figure (IPF) color-coded map of sample CaMg-05 (146 h, 750 °C), showing the orientation of magnesite and calcite single crystals, polycrystalline dolomite (palisades and granular shapes) and magnesio-calcite. IPF color-code indicates the orientation of reference axes parallel to the growing direction (GD). For example, red or light blue grains have the $[\bar{1}2\bar{1}0]$ / [0001] or $[\bar{1}2\bar{1}0]$ axis pointing to the reader (note different colors for different phases). Black lines within dolomite rim highlight the twin relationship between neighboring dolomite grains. The histogram below shows the correlated misorientation angles of dolomite indicating that 26 % of all dolomite-dolomite boundaries are represented by twin boundaries. Pole figures for [0001] axes, poles to prismatic $\{2\bar{1}0\}$ and rhomb $\{10\bar{1}4\}$ planes are plotted with respect to the external reference frame. The reference frame is defined by E-W plane, which represents the reaction rim interface and GD of the rim which is oriented N-S. The numbers on the pole figures indicate individual orientation relationships between calcite single crystal and dolomite (1-4) and magnesite and dolomite (5-6). All pole figures are equal-area projections (lower hemisphere) with a Gaussian half-width of 10° and a confidence index (CI) >0.2. Contour color densities are drawn based on multiples of random distribution with a factor of 9 from blue to red. b EBSD inverse pole figure color-coded map of sample CaMg-08 (48 h, 800 °C), which best documents the totopastic relations to single crystal calcite reactant. The map on the right shows grains of dolomite (dark blue) and magnesio-calcite (red) with totopastic relations to calcite (light blue) demonstrated in the pole figures. The light green grains in the totopaxy map represent the twinned dolomite grains in relation to the dark blue ones that show the totopastic relationships described before.
Microprobe analyses

Compositional line scans were performed from one reactant to the other across the dolomite rim and magnesio-calcite layer to measure chemical profiles (Figure 7). While the composition of the reactant phases is homogenous and flat also adjacent to the reaction rims, composition gradients occur within the newly evolved magnesio-calcite and dolomite rims. The dolomite is close to stoichiometric at the dolomite-magnesite interface, and it becomes successively more non-stoichiometric with increasing Mg deficiency towards the dolomite-calcite interface. According to the phase diagram in Figure 2, dolomite in equilibrium with magnesio-calcite at 800°C should have a composition with about 47 mol % MgCO₃ component and a dolomite in equilibrium with magnesite should have about 51 mol % MgCO₃ component. The composition of dolomite at the dolomite-magnesite interface thus closely corresponds to local equilibrium. In contrast, the dolomite is more calcium-rich and the magnesio-calcite is somewhat more Mg-rich than what is expected for local equilibrium at the calcite-dolomite interface. The processes underlying this deviation from local equilibrium are not known. A possible explanation could be the three dimensional character of the microstructures, which would especially influence analyses at phase boundaries, or the effect of nucleation and interface-reaction at the initial stage of rim growth, which appears to be fast for dolomite palisades (Figure 4 c, d). Alternatively, the depletion of Mg in granular dolomite due to the preceding magnesio-calcite formation may induce local disequilibrium in granular dolomite. The effect of deviations from local equilibrium element partitioning on the overall reaction kinetics are considered as minor and this complication has not been accounted for in our thermodynamic model. Magnesio-calcite always shows a curved profile with progressively decreasing Mg content from granular dolomite towards calcite. Measurements on 10 samples at different run durations and temperatures reveal two different CaO and MgO distributions in dolomite, illustrated in Figure 7. At T = 850°C, the dolomite palisades are nearly stoichiometric. The granular dolomite is substantially more Ca-rich and is clearly non stoichiometric with respect to its Ca/Mg ratio towards the contact with magnesio-calcite (Figure 7 a). At T = 825°C, the molar concentrations of CaO and MgO are similar at the initial interface (dashed line in Figure 7 b). Chemical profiles of the other samples show the same two trends, but no systematic variation with annealing time or temperature.
Figure 7 Chemical profiles in a sample CaMg-10 \(t = 74.28\) h, \(T = 850\) °C and b sample CaMg-13 \(t = 75.05\) h, \(T = 825\) °C. The mole fractions of calcium \(X_{CaO}\) and magnesium \(X_{MgO}\) are shown in red and black, respectively. Solid vertical lines indicate locations of phase boundaries while the dashed line indicates the original interface.

**Discussion**

**Microstructure and texture evolution**

Initial reaction phase during experiments is associated with the formation of magnesio-calcite (Figure 2), which grew towards the calcite single crystal. This initial re-equilibration is assumed to affect the reaction interface, so that dolomite formed in between single crystal magnesite and polycrystalline magnesio-calcite. Such setup is probably responsible for the development of the two distinct dolomite regions. Granular dolomite encountered boundaries of already existing magnesio-calcite grains, while palisades could grow into the magnesite single crystal. The elongated shape of dolomite in contact with magnesite is suggested to be a stress-induced phenomenon resulting from a positive volume change at the dolomite-magnesite reaction interface (Milke and Wirth 2003). Although, the overall reaction is more or less balanced regarding the volume change, the partial reaction at the magnesite-dolomite interface implicates a positive volume change, while the incorporation of magnesium in calcite results in a negative volume change.

Since magnesio-calcite results from magnesium incorporation into calcite, it is likely that magnesio-calcite grains may inherit the crystallographic orientation of pure calcite. Indeed, some of the magnesio-calcite grains show a full crystallographic relation to calcite, which is then transferred also into dolomite (Figure 6 b). The relatively large size of the magnesio-calcite grains implies few nucleation sites, possibly because of the small composition contrast across the magnesio-calcite / calcite interface (Figure 7). The dolomite grains with topotaxy to calcite reactant occur mainly in the granular part of the rim or near the original reaction interface. They are characterized by a slightly larger grain size than the average and their spatial occurrence usually coincide with slightly thinner rim portions (Figure 6 b). This correlation may be
explained by a slightly hindered grain boundary diffusion and nucleation during the initial phases of the rim growth in topotactic domains.

1.1.1 Diffusion components

The parabolic growth behavior of the dolomite reaction rim suggests that diffusion of Mg and/or Ca components was rate limiting (Figure 4a, Fisher 1978). The term component is used here as a mere chemical entity that suffices to describe the chemical variation in the system of interest. It does not necessarily correspond to the composition of a phase or species. Within a single crystal of dolomite mass transfer may occur via interdiffusion of Mg$^{2+}$ and Ca$^{2+}$ ions. In this case, the cation fluxes are forced to be equal and directed in opposite directions due to the charge-balance constraint. Alternatively, Mg$^{2+}$ and Ca$^{2+}$ may diffuse coupled with O$^{2-}$, which would make Ca- and Mg-fluxes independent. Such behavior is probably enhanced through the presence of lattice imperfections such as edge dislocations and, more importantly, of grain boundaries, which could act as sources and sinks of vacancies. In case of a fine-grained polycrystalline reaction rim the coupled diffusion of divalent cations and oxygen is thus conceivable. Based on experiments in the MgO-SiO$_2$ system (Gardès and Heinrich 2011) and in the CaO-MgO-SiO$_2$ system (Joachim et al. 2010), which were both conducted under "dry" conditions, coupled diffusion of Mg$^{2+}$ and O$^{2-}$ could indeed be demonstrated. During growth of a polycrystalline dolomite reaction rim in the two component system MgCO$_3$-CaCO$_3$ the diffusion fluxes of the two components may be independent. In our study, we do not consider transfer of carbonate. It is thus sufficient to consider MgO and CaO as the two mobile components. Although the speciation of the diffusing matter is not known, the bulk effect of its transfer can be described in terms of the MgO and CaO components. If the Mg and the Ca–bearing species indeed diffuse independently, the rim growth rate depends on the mobilities of both components. The relative fluxes of the two components can be inferred from the position of the original calcite-magnesite interface (Abart et al. 2004). The off-center position of the original calcite-magnesite interface as indicated by the platinum markers (}
suggests that the diffusive fluxes of the two components were different.

1.1.2 Dolomite rim growth model

The overall reaction (1) can be split into two partial reactions, taking place at the dolomite-calcite (1) magnesite-dolomite (2) and the interface:

Interface I: \[ \text{CaCO}_3\text{solid} + \nu \text{MgCO}_3\text{mobile} = \nu \text{CaMg(CO}_3)_2\text{solid} + (1-\nu) \text{CaCO}_3\text{mobile} \] (1)

Interface II: \[ \text{MgCO}_3\text{solid} + (1-\nu) \text{CaCO}_3\text{mobile} = (1-\nu) \text{CaMg(CO}_3)_2\text{solid} + \nu \text{MgCO}_3\text{mobile} \] (2)

where \(\nu\) is the molar amount of MgO forming granular dolomite at the calcite-dolomite interface and \((1-\nu)\) is the molar amount of CaO producing dolomite palisades from magnesite at the dolomite-magnesite interface. In

Figure 8 the geometry of the rim growth setting and the associated component fluxes are illustrated schematically.
Within the dolomite rim the domains with the palisade and the granular microstructures are discerned. Note that the flux of CaO is restricted to the rim of newly formed dolomite; in contrast a fraction of the MgO that is derived from the consumption of the magnesite at the dolomite-magnesite interface diffuses across the dolomite-calcite interface and into the calcite forming magnesio-calcite. The fraction \( \nu \) is related to the molar amount \( (n) \) of components that are transferred during reaction rim growth

\[
\nu = \frac{n_{\text{MgO}}}{n_{\text{MgO}} + n_{\text{CaO}}}
\]

where \( n \) can be derived from the thickness of dolomite layers \( \Delta x_{\text{pal}} \) and \( \Delta x_{\text{gran}} \), the molar volume \( \bar{V}_y \) of phase \( y \) and the unit cross section \( A \).

\[
n_{\text{CaO}} = \frac{\Delta x_{\text{pal}} A}{\bar{V}_{\text{dot}}} \quad \text{(4)}
\]

\[
n_{\text{MgO}} = \frac{\Delta x_{\text{gran}} A}{\bar{V}_{\text{dot}}} + \frac{\Delta x_{\text{Mg-Cal}} c_T A}{\bar{V}_{\text{cal}}} \quad \text{(5)}
\]

The second term in equation (5) accounts for the formation of magnesio-calcite. The correction factor \( c_T \) represents the molar Mg concentration, which was determined from measured chemical profiles leading to average values of \( c_{750,850} = 0.03 \), \( c_{800} = 0.04 \) and \( c_{825} = 0.06 \) at \( T = 750^\circ \text{C} \) and \( 850^\circ \text{C} \), \( 800^\circ \text{C} \) and \( 825^\circ \text{C} \), respectively.

To determine the diffusion coefficients of CaO and MgO the thermodynamic model for diffusion controlled reaction rim growth in a binary system of ABART ET AL. (2009) with extension of GÖTZE ET AL. (2009) was used. This model is based on the assumption that the diffusion of chemical components across the dolomite reaction rim is the only dissipative process and that the potential contribution of dolomite nucleation to the total energy budget can be neglected. Rearrangement of equations 26, 34 and 36 from ABART ET AL. (2009) leads to the combined diffusion coefficient \( (D_{\text{com}}) \) which accounts for the simultaneous diffusion of both mobile components MgO and CaO.
\[ D_{\text{com}} = R_g T \frac{\Delta x_{\text{pol}}^2 (1 - u)}{2t \Delta G_{\text{rim}}} \left( X_{\text{Mgo}}^{\text{Mgs}} X_{\text{CaO}}^{\text{Dol}} - X_{\text{Mgo}}^{\text{Dol}} X_{\text{CaO}}^{\text{Mgs}} \right)^2 V_{\text{Dol}} \frac{1}{(v - 1)X_{\text{Mgo}}^{\text{Dol}} - vX_{\text{CaO}}^{\text{Dol}} (V_{\text{Mgs}})^2 X_{\text{CaO}}^{\text{Dol}} X_{\text{Mgo}}^{\text{Dol}}} \]

where \( R_g \) is the gas constant, \( V \) is the specific molar volume of phase \( \gamma \), and \( X_{\gamma}^i \) is the mole fraction of component \( i \) in phase \( \gamma \). The parameter \( u \) incorporates the mass balance at the reaction fronts and information of interface motions (A\textsc{bart} et al. 2009). Since the original model assumes stoichiometric composition of the reaction rim, we account for the measured chemical gradient in dolomite by using a modified expression for \( u \) using equations 14, 15, 17, 18 and 21 of Abart et al. (2009), which leads to (G\textsc{ö}t\textsc{ze} et al. 2009):

\[ u = 1 - \frac{V_{\text{Dol}}}{V_{\text{Mag}}} \left[ \frac{v (1 - 2X_{\text{CaO}}^{\text{Mgs}}) - (1 - X_{\text{CaO}}^{\text{Dol}})}{v (1 - 2X_{\text{CaO}}^{\text{Pol}}) - (1 - X_{\text{CaO}}^{\text{Pol}})} \right] - \frac{-v (1 - 2X_{\text{CaO}}^{\text{Cal}}) + (1 - X_{\text{CaO}}^{\text{Cal}})}{X_{\text{CaO}}^{\text{Cal}} X_{\text{Mgo}}^{\text{Gran}} - X_{\text{CaO}}^{\text{Cal}} X_{\text{Mgo}}^{\text{Dol}}} \]

\[ \Delta G_{\text{rim}} = \frac{1}{V_{\text{Mgs}}} \Delta G \]

The thermodynamic data of the phases in our experiments used to calculate \( \Delta G_{\text{rim}} \) are given in Table 3. The combined diffusion coefficient \( D_{\text{com}} \) can also be expressed by (A\textsc{bart} et al. 2009):

\[ D_{\text{com}} = k D_{\text{Mgo}} + (1 - k) D_{\text{CaO}}, \]

where \( k \) gives the proportion of the component fluxes within the dolomite reaction rim defined as

\[ k = \frac{D_{\text{Mgo}}}{D_{\text{Mgo}} + D_{\text{CaO}}} \]

Combining equations (9) and (10) allows determining the individual component diffusivities:

\[ D_{\text{CaO}} = D_{\text{com}} \frac{(1-k)}{(2k^2-2k+1)} \quad \text{and} \quad D_{\text{Mgo}} = \frac{D_{\text{com}}-D_{\text{CaO}}(1-k)}{k} \]

The diffusion coefficients obtained from our rim growth experiments are shown in an Arrhenius diagram in Figure 9, following the relation:
with $k_0(\text{CaO}) = 10^{-4.9 \pm 2.7} \text{ m}^2/\text{s}$, $E_a(\text{CaO}) = 192 \pm 44 \text{ kJ/mol}$ and $k_0(\text{MgO}) = 10^{-4.5 \pm 2.2} \text{ m}^2/\text{s}$, $E_a(\text{MgO}) = 198 \pm 54 \text{ kJ/mol}$ respectively.

Figure 9 Arrhenius plot of the effective diffusion coefficients of a CaO and b MgO in dolomite in the range of 750-850°C. Values are from Table 2

Both components show almost identical temperature dependence of their diffusivities, which yields quite similar activation energies in the order of 200 kJ/mol for both components. Out of Figure 5, a huge difference in activation energies may have been expected due to the increasing or decreasing ratio of layer thicknesses with temperature. Indeed, if plotting the ratio of $\Delta x_{\text{Pal}}/ (\Delta x_{\text{gran}} + \Delta x_{\text{Mg-Cal}})$ against time the ratio remains nearly constant between 0.22 – 0.27, implying similar activation energies for diffusing components.

It must be noted, that the diffusion coefficients refer to the self-diffusion of CaO and MgO in polycrystalline dolomite, where diffusion may occur by a combination of volume- and grain-boundary diffusion. Diffusion coefficients obtained in this study should be regarded as effective diffusion coefficients. The relatively large size of magnesio-calcite grains compared to dolomite grains with magnesio-calcite grain boundaries oriented mainly perpendicular to the growth direction, and the measured chemical profile within magnesio-calcite may indicate dominantly volume diffusion. In contrast, abundant grain boundaries within the small granular dolomite with abundant twins and relatively straight grain boundaries of the dolomite palisades may suggest preferred grain boundary diffusion.

<table>
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<tr>
<th>Temperature</th>
<th>Calcite</th>
<th>Magneise</th>
<th>Dolomite</th>
</tr>
</thead>
<tbody>
<tr>
<td>(750°C)</td>
<td>37.56</td>
<td>28.72</td>
<td>65.84</td>
</tr>
<tr>
<td>(800°C)</td>
<td>37.63</td>
<td>28.79</td>
<td>65.98</td>
</tr>
<tr>
<td>(825°C)</td>
<td>37.67</td>
<td>28.82</td>
<td>66.05</td>
</tr>
<tr>
<td>(850°C)</td>
<td>37.79</td>
<td>28.85</td>
<td>66.13</td>
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<td>-1,114,861</td>
<td>-2,354,678</td>
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<tr>
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<td>-1,124,567</td>
<td>-2,375,712</td>
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<tr>
<td>$\Delta f G$ (825°C)</td>
<td>-1,252,291</td>
<td>-1,129,530</td>
<td>-2,386,457</td>
</tr>
</tbody>
</table>
Table 3 Molar volumes $V$ and Gibbs free energies $\Delta_f G$ of formation of calcite, magnesite and dolomite at $T = 750^\circ$C - 850$^\circ$C and $P = 400$ MPa (from Holland and Powell 1998)

<table>
<thead>
<tr>
<th>Temperature</th>
<th>$\Delta_f G$ [J/mol]</th>
</tr>
</thead>
<tbody>
<tr>
<td>750$^\circ$C</td>
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<tr>
<td>850$^\circ$C</td>
<td>-1,134,565</td>
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<td>-2,397,349</td>
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</tbody>
</table>

Comparison with other diffusion coefficients in carbonates

To the best of our knowledge only two studies determined component diffusion in dolomite so far. Anderson (1972) determined the self-diffusion coefficients of C and O in crushed dolomite using isotope exchange with CO$_2$ at $T = 645$-785$^\circ$C and $P = 12$-93.5 MPa. Diffusion rates of C and O in dolomite are nearly identical with activation energies of $E_a$ (C) = 468 kJ/mol and $E_a$ (O) = 485 kJ/mol (Figure 10).

Müller, Cherniak, and Watson (2012) performed thin film experiments between dolomite and siderite or rhodochrosite to determine interdiffusion coefficients of (Mn, Fe) – Mg – Ca in the temperature range of 400-625$^\circ$C at 1 atm pressure. The authors determined coupled diffusion of Mn-(Ca+Mg) and Fe-(Ca+Mg) with activation energies of $E_a$ (Mn-(Ca+Mg)) = 63 ± 5 kJ/mol and $E_a$ (Fe-(Ca+Mg)) = 123 ± 10 kJ/mol respectively. The temperature dependence of the quasi-binary exchange of Mn-Mg and Fe-Mg shows a kink at $T \sim 525^\circ$C with an increase of activation energies from $E_a$ (Mn-Mg) = 23 ± 4 kJ/mol to 168 ± 15 kJ/mol and from $E_a$ (Fe-Mg) = 34 ± 10 kJ/mol to 183 ± 14 kJ/mol (Figure 10) in a low-temperature regime. A proposed explanation is an increased disorder of dolomite at high temperatures, rather than a transition from an extrinsic to an intrinsic diffusion regime (Müller, Cherniak, and Watson 2012). When extrapolated to the temperatures of our experiments, the interdiffusion coefficients in the Ca, Mg and Ca, Mg, Mn, Fe-couples determined by Müller, Cherniak, and Watson (2012) are several orders of magnitude lower than the diffusion coefficients we determined for the CaO and MgO components.
Figure 10 Arrhenius diagram for diffusion coefficients of various elements in calcite, magnesite and dolomite. Published data on diffusion in dolomite and calcite are plotted as solid lines and dashed lines, respectively. Dots and triangles represent chemical diffusion of Ca and Mg in magnesite, respectively. 1 = FISLER AND CYGAN (1999), 2 = FARVER AND YUND (1996), 3 = MÜLLER, CHERNIAK, AND WATSON (2012), 4 = KENT ET AL. (2001), 5 = ANDERSON (1972). Indices at the lower left of the chemical symbols represent activation energies in kJ/mol. All data are determined for volume diffusion, except those marked by GB, which represents grain boundary diffusion.

For comparison Figure 10 also shows diffusion coefficients of Ca and Mg in calcite and magnesite. FARVER AND YUND (1996) determined activation energies for Ca volume and grain boundary diffusion in natural samples at P = 0.1 MPa and T = 650-900°C of 382 ± 37 kJ/mol and 267 ± 47 kJ/mol, respectively. Assuming an effective grain boundary width of 3 nm, diffusion rates would be 6 orders of magnitude higher for diffusion along the grain boundaries than for volume diffusion. Besides, they conducted calcium self-diffusion experiments parallel to the c-axis and perpendicular to the natural cleavage planes of natural calcite single crystals to investigate the anisotropy of diffusion coefficients. Despite the trigonal crystal system there was no anisotropy measurable regarding Ca diffusion in calcite.

FISLER AND CYGAN (1999) measured Ca and Mg self-diffusion coefficients perpendicular to the rhomb plane in natural calcite single crystals. Experiments were performed at T = 550-800°C and P = 0.1 MPa, for which activation energies were quite similar with $E_a$(Ca) = 271 ± 80 kJ/mol and $E_a$(Mg) = 284 ± 74 kJ/mol, although Mg self-diffusion appears to be one order of magnitude faster than Ca self-diffusion. Interestingly, the Ca-diffusivity measured by Fisler and Cygan (1999) is one order of magnitude faster than measured by FARVER AND YUND (1996), which may be related to different amounts of Mg and Mn present in the used starting materials (FISLER AND CYGAN, 1999). KENT ET AL. (2001) investigated Mg chemical diffusion in calcite as well as Ca chemical diffusion and Mg self-diffusion in magnesite. Experiments were performed...
using fragments of natural cleaved material, calcite or magnesite, which were placed together with dried powder of MgO and MgCO₃ for calcite or CaCO₃ for magnesite into Pt-capsules. The material was pre-dried and subsequently sealed inside evacuated silica glass tubes for annealing experiments using a muffle furnace. Annealing conditions were between 400 and 600°C at P = 0.1 MPa. For Mg chemical diffusion in calcite an activation energy of only Eₐ(Mg) = 76 ± 16 kJ/mol was determined, which may indicate a switch from intrinsic to extrinsic diffusion at low T. Chemical diffusion of Ca in magnesite yielded an activation energy of Eₐ(Ca) = 214 ± 60 kJ/mol. The measured self-diffusion coefficient of Mg in magnesite at 450°C is similar to Mg chemical diffusion in calcite.

These values are higher than those obtained for diffusion of CaO and MgO in our study (Eₐ(CaO) =192 ± 54 kJ/mol, Eₐ(MgO) = 198 ± 44 kJ/mol), but overlap within error bars at least for MgO. The associated diffusivities for calcium, magnesium and oxygen volume diffusion in calcite are 3 to 5 orders of magnitude lower than those estimated for dolomite (Figure 10).

Farver and Yund (1996) measured also the grain boundary diffusion of calcium in natural and hot-pressed calcite aggregates, yielding an activation energy of Eₐ,GB(Ca) = 267 ± 47 kJ/mol. Assuming a grain boundary width of 1 nm, Farver and Yund (1996) obtained a Ca grain-boundary diffusivity in calcite that is about two orders of magnitude faster than the component diffusivities in dolomite measured in our study (Figure 10). Although differences in the calculated diffusivities may arise from the applied experimental techniques, comparison of all of these diffusion data does not clearly demonstrate whether volume or grain boundary predominates in our dolomite rim growth experiments.

Conclusions

Annealing experiments on calcite and magnesite single crystals produced a magnesio-calcite layer and a polycrystalline dolomite reaction rim at temperatures between 750 and 850°C and 400 MPa confining pressure. Within dolomite, two different microstructural domains formed. Stress-induced palisades grew perpendicular to the interface on magnesite and granular dolomite formed on magnesio-calcite. Platinum markers showed that this microstructural boundary represents the original calcite-magnesite interface. Chemical composition of palisades is nearly stoichiometric, while the granular portions show a slight gradient diverging toward magnesio-calcite. All reaction domains, magnesio-calcite, palisades and granular dolomite show a crystallographic orientation relationship to the calcite reactant. Full crystallographic relationships with respect to calcite are restricted to magnesio-calcite and granular dolomite. Axiotactic dolomite grains also appear in the palisade domain, associated with randomly distributed growth twins. The entire dolomite rim thickness increases linearly with the square root of time, indicating a diffusion-controlled mass transport. Thermodynamic considerations lead to activation energies of Eₐ(CaO) =192 ± 54 kJ/mol and Eₐ(MgO) = 198 ± 44 kJ/mol.

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