CRYSTAL CHEMISTRY AND DIFFERENTIAL THERMAL EFFECTS OF DOLOMITE*†
W. F. BRADLEY,1 J. F. BURST,2 AND D. L. GRAF.3

ABSTRACT

It is observed that the lower temperature endothermal effect of dolomite takes place at a considerably lowered temperature after the dolomite has been subjected to prolonged vigorous grinding.

The mechanisms and consequences of plastic deformation in the dolomite crystallization are discussed, and are contrasted with the analogous deformations in the simple carbonates. Multiply deformed dolomite crystals actually contain island nuclei locally enriched in one or the other of the positive ions. Powder diffraction diagrams of such deformed crystallites exhibit greatly attenuated intensities for those reflections for which the form factor includes the term Ca-Mg.

An approximate evaluation of the four variable parameters of dolomite (in R 3) has been made, based on the powder diffraction. 2 C are in 2(c) with x0=0.243 and 6 O are in 6(f) with x0=0.500, y0=−0.042 and z0=0.271.

Among the abnormal differential thermal features exhibited by many dolomites, it has become apparent that some, at least, may be consequences of structural factors. An attempt has been made through extended grinding to alter mechanically the structural arrangement of a dolomite lattice and to record this alteration through variations of the differential thermal pattern and the x-ray diffraction effects.

The nature of the material resulting from this type of mechanical stressing precludes single crystal work, making the use of powders necessary for both x-ray diffraction and thermal analysis. It is thus desirable to establish with moderate accuracy the variable carbon and oxygen parameters. By drawing heavily on the crystal chemistry data available for the various simple rhombohedral carbonates, a suitable model is readily established from which refinements can be made.

The dolomite structure consists of hexagonally packed layers of carbonate groups between which alternate layers of octahedral interstices are occupied respectively by calcium and magnesium. A trial model is obtained by combining one layer of CaCO3 from calcite with one layer of MgCO3 from magnesite in such a manner that each layer assumes the dolomite translation, and the carbonate groups of each become coincident. This operation moves the carbonate groups toward the magnesium

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2 Exploration and Production Research Division, Shell Oil Company, Houston, Texas.
layers and maintains one of the carbonate oxygens in each of the three cleavage planes.

The simple rhombohedral carbonates crystallize in $R\overline{3}c$ and dolomite in $R\overline{3}$; the lattice translations are all accurately known (Bragg) (1). The best established C—O bond length is the 1.31 Å value deduced by Elliott (3). Using this value, the Ca—O bond in calcite is 2.34 Å, and the Mg—O bond in magnesite is 2.06 Å.

When set up so that planar CO$_3$ groups of C—O bond 1.31 Å are oriented with one oxygen in each cleavage plane, the structure is effectively reduced to the one variable parameter $x_0$. Best agreement between observed and calculated powder diffraction intensities is near $x_0 = 0.243$. Agreement deteriorates noticeably at $x_0$ deviations of ±.003. At $x_0 = 0.243$ the Ca—O and Mg—O bonds are respectively 2.34 and 2.08 Å. In Table 1, spectrometer counts from a G.E. recording spectrometer reduced according to the formula:

$$q^u \frac{1 + \cos^2 2\theta}{\sin^2 \theta \cos \theta} \frac{pF^2}{\sin \theta}$$

and, approximately scaled to absolute F’s, are compared with calculated form factors for Ca at 0, 0, 0; Mg at $\frac{1}{2}, \frac{1}{2}, \frac{1}{2}$; 2C at $\pm x_0 = 0.243$; and 6O in 6(f) at $x_0 = 0.500$, $y_0 = -0.042$, $z_0 = 0.271$, etc. The factor

$$\frac{1}{\sin \theta}$$

is introduced as an arbitrary correction for focusing conditions.

The atomic scattering factors employed are those of Pauling and Sherman (8). Some improvement in agreement for the first few lines would be realized by allowance for the partial ionic character of the Me—O bonds. Further refinement of parameters would require recourse to higher angle diffraction and single crystal methods, which would differentiate reflections to which the oxygen contributions are of two kinds.

The quality of agreement of the data in Table 1 establishes that the above-described model is an adequate basis on which to discuss the mechanical processes described below.

The effects of grinding field-samples of dolomite were investigated as a function of time by placing them in an electric-powered mortar and pestle from which samples of the material being ground were withdrawn at various intervals for differential thermal analysis and x-ray diffraction analysis. It was soon noted that prolonged grinding produced definite abnormalities in the physical characteristics of the materials. Therefore, the experiments were duplicated with U. S. B. S. Dolomite No. 88 in an attempt to assign the abnormalities directly to the dolomite structure rather than to some unknown condition within the field sample. The
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Table 1. Comparison of Calculated and Observed Relative F Values for Powder Diffraction Lines up to 2θ = 70°

<table>
<thead>
<tr>
<th>Index</th>
<th>sin θ</th>
<th>Form Factor</th>
<th>F Calc.</th>
<th>Reduced from Spectrometer Count</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>.1243</td>
<td>Ca−Mg</td>
<td>+0.09 C</td>
<td>+4.3</td>
</tr>
<tr>
<td>110</td>
<td>.1357</td>
<td>Ca+Mg</td>
<td>−1.99 C</td>
<td>+7.4</td>
</tr>
<tr>
<td>122</td>
<td>.1736</td>
<td>Ca+Mg</td>
<td>+1.97 C</td>
<td>+41.7</td>
</tr>
<tr>
<td>222</td>
<td>.1878</td>
<td>Ca+Mg</td>
<td>−1.93 C</td>
<td>−18.4</td>
</tr>
<tr>
<td>221</td>
<td>.1974</td>
<td>Ca−Mg</td>
<td>+0.43 C</td>
<td>+10.7</td>
</tr>
<tr>
<td>110(1)</td>
<td>.2084</td>
<td>Ca+Mg</td>
<td>+2 C</td>
<td>+15.3</td>
</tr>
<tr>
<td>210</td>
<td>.2286</td>
<td>Ca−Mg</td>
<td>−0.26 C</td>
<td>+28.0</td>
</tr>
<tr>
<td>120</td>
<td>.2427</td>
<td>Ca−Mg</td>
<td>+0.09 C</td>
<td>+11.1</td>
</tr>
<tr>
<td>322</td>
<td>.2499</td>
<td>Ca−Mg</td>
<td>−0.60 C</td>
<td>+2.2</td>
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<tr>
<td>220</td>
<td>.2713</td>
<td>Ca+Mg</td>
<td>+1.97 C</td>
<td>+15.5</td>
</tr>
<tr>
<td>332</td>
<td>.2778</td>
<td>Ca+Mg</td>
<td>+1.86 C</td>
<td>+29.7</td>
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<tr>
<td>123</td>
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<td>−1.93 C</td>
<td>+29.6</td>
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<tr>
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<td>Ca−Mg</td>
<td>+0.80 C</td>
<td>+16.4</td>
</tr>
<tr>
<td>311</td>
<td>.2870</td>
<td>Ca−Mg</td>
<td>+0.43 C</td>
<td>−2.5</td>
</tr>
<tr>
<td>210(1)</td>
<td>.3199</td>
<td>Ca−Mg</td>
<td>+0.09 C</td>
<td>+15.4</td>
</tr>
<tr>
<td>120</td>
<td>.3245</td>
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<td>−1.99 C</td>
<td>+10.5</td>
</tr>
<tr>
<td>112</td>
<td>.3353</td>
<td>Ca+Mg</td>
<td>−1.80 C</td>
<td>+22.6</td>
</tr>
<tr>
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<td>+1.97 C</td>
<td>+8.3</td>
</tr>
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<td>+15.9</td>
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<tr>
<td>234</td>
<td>.3504</td>
<td>Ca−Mg</td>
<td>+3.08 C</td>
<td>+18.0</td>
</tr>
<tr>
<td>324(1)</td>
<td>.3546</td>
<td>Ca−Mg</td>
<td>−4.95 C</td>
<td>+12.5</td>
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<tr>
<td>230</td>
<td>.3610</td>
<td>Ca+Mg</td>
<td>−2.31 C</td>
<td>+14.7</td>
</tr>
<tr>
<td>112</td>
<td>.3610</td>
<td>Ca+Mg</td>
<td>+2 C</td>
<td>+31.5</td>
</tr>
<tr>
<td>300</td>
<td>.3730</td>
<td>Ca−Mg</td>
<td>+0.26 C</td>
<td>+3.1</td>
</tr>
<tr>
<td>221</td>
<td>.3731</td>
<td>Ca−Mg</td>
<td>−0.26 C</td>
<td>−0.8</td>
</tr>
<tr>
<td>444</td>
<td>.3754</td>
<td>Ca+Mg</td>
<td>+1.73 C</td>
<td>+37.8</td>
</tr>
</tbody>
</table>

Entries marked (*) include some spurious count from incompletely resolved adjacent strong lines.

same striking results, obtained with the field sample, were duplicated precisely through the grinding of the standard material.

After continuous grinding for a period of eight hours, dolomite was noted by differential thermal analysis to undergo an additional, small
endothermic reaction between $500^\circ$ and $600^\circ$ C. This endothermic reaction is shown in curve 8, figure 1a. Curves on this plate show changes resulting from grinding, as recorded by differential thermal analysis. By employing Rowland and Lewis' (9) technique of furnace-atmosphere control, the carbonate peaks are separated and the area of each loop easily discernible. Curves in figure 1a show the thermal expressions from $500^\circ$ to $1000^\circ$ C. for material ground 3, 8, 18, 25, and 37 hours, respectively. Curves in figure 1b show thermal expressions for the entire range,
or room temperature to 1000° C., for material ground 50, 66, 125, 228, and 400 hours, respectively. The chart speed was decreased after the sample ground for 37 hours was analyzed because it had become apparent that a new endothermic loop was being formed which would be more suitably represented in a recording made at a slower chart speed.

The continued grinding increases the magnitude of the 500°–600° C. endothermic reaction at the expense of the 700°–800° C. endothermic reaction, commonly spoken of as the evolution of CO$_2$ from the MgCO$_3$ portion of the dolomite composition. This reciprocal variation is illustrated by the nested curves in figure 1. Also, it was found that the ratio of the sum of the areas of the two low-temperature peaks to the area of the high-temperature peak was virtually identical in each record. Several interesting relationships between the areas of the various loops are presented in figure 2.

![Figure 2](attachment:image.png)

**Fig. 2.** Areal relationships of the three separate endothermal DTA features of various ground dolomite specimens.

It should be noted that the newly formed loop (figure 1) is considerably removed along the temperature axis from its progenitor and is, in this respect, unlike thermal anomalies which arise from particle-size diminution alone. When particle size is the controlling factor, a gradual lowering of the decomposition temperature usually is noted with decreas-
ing particle size. Interest is also aroused by the fact that after the 500°-
600° C. loop is fully developed and the 700°-800° C. loop completely
eliminated (228-hour grind), no further significant change is recorded
even after prolonged grinding (400 hours).

X-ray powder diffraction diagrams of three ground samples, irradiated
and recorded simultaneously under the strictest comparable conditions,
show at least three lines which are selectively reduced in intensity upon
continued grinding. These lines are recorded at \( d = 4.02 \, \text{Å}, \)
\( d = 2.53 \, \text{Å}, \) and \( d = 2.06 \, \text{Å}, \) and represent, respectively, the crystal planes (100),
(221), and (111).

From Table 1 it can be noted that each of these planes \((h+k+l=\text{odd})\)
is characterized by a form factor of the type \( \text{Ca} - \text{Mg} \), modified by
small to moderate carbon and oxygen contributions. The line-intensity
diminutions are thought therefore to result from some specific structural
alteration such as a glide mechanism whereby \( \text{Ca}^{++} \) and \( \text{Mg}^{++} \) ions would
be migrated along certain directions within the lattice. When sufficient
\( \text{Ca}^{++} \) ions have been translated to positions formerly occupied by \( \text{Mg}^{++} \)
ions, so that a random distribution of these two ions is produced in the
projection of the whole structure onto the poles to the planes in question,
the lines representing those planes should almost disappear.

Examination of figure 3 indicates that reflections from planes (100),
(221), and (111), which were relatively intense after three hours of grind-
ing, became selectively less intense after 50 hours of grinding and virt-
ually disappeared after 228 hours of grinding. It was at this point that
the DTA curves also became stabilized (figure 1). The fact that these
two circumstances prevail concurrently indicates the attainment of a
new equilibrium established through the continued application of the
nondirectional stress load. Crystallographic studies of plastic deforma-
tion in ionic crystals have been concerned primarily with directed loads
as exemplified by Buerger's (2) analyses of the NaCl type face-centered
cubes. There is no reason to assume that the translations so induced are
the only ones which occur, and indeed, Buerger infers that deformations
along octahedral planes are probably induced by random loading. De-
formation along octahedral planes is also a frequent inference in diffrac-
tion analyses of orientation effects produced in the working of face-cen-
tered cubic metals.

The close structural analogy between the face-centered cubic alkali
halides and the rhombohedral carbonates suggests that such structural
consequences of deformations as could have resulted from the above
grinding experiments could profitably be examined.

In order to avoid the confusion which arises from the various sets of
reference axes to which rhombohedral carbonates are at times referred,
Miller indices will be used in the following discussion to refer only to the true rhombohedral cell. Analogous cubic directions can then be referred to by name. Table 2 presents the various index correlations for the common dolomite forms.

Translation gliding of dolomite with \( T\{111\}, \{10\overline{1}\} \) was produced experimentally by Johnsen (7), and twin gliding on \{110\}, probably along the short diagonals of the cleavage rhomb, is discussed by Fairbairn and Hawkes (4). Both of these are octahedral glides along dodecahedral poles in the cubic system. The familiar pressure twins of calcite across (332) are not known in dolomite. This is the “forbidden” dodecahedral translation along a cube pole. It occurs in calcite only as a twin glide, not as a full translation.

Figures 4, 5, and 6 are projections of the dolomite structure onto (110) arranged as discussed by Fairbairn and Hawkes. Figure 4 is a projection of the aspect a calcite pressure twin would assume in dolomite if it did occur.
When calcite is pressure-twinned the ions of each successive (332) level are displaced along a "cube" pole to the alternative equivalent position, which represents the exchange of the long for the short diagonal of the cleavage rhomb, and CO₃ ions rotate according to the steric requirements. Each successive level makes this exchange as the twin boundary.
progresses through successive 332 levels, but no level glides beyond the like-ion barrier of the "cube" pole glide. The result is that the (222) and one of the (110) directions interchange, and inasmuch as the projection of the structure upon each is a simple alternation of Ca and CO₃ ions, no drastic rearrangement has been required.

As is seen in figure 4, the application of the identical mechanism to dolomite fails as a physical possibility because of the unequal Ca and Mg ion sizes and dispositions. Projected ion sequences along [111] and [110] are, respectively, Ca, CO₃, Mg, CO₃, etc., and Ca+Mg, 2CO₃, Ca+Mg, 2CO₃, etc., which are not interchangeable. A pressure twin of composition face (332) would require a complete redistribution of the positive ions.
Figure 5 illustrates the apparent mechanism of the formation of the lamellar dolomite twins across (110). Not apparent in this projection, but clear in three dimensions and in the supplemental sketch, is the fact that alternating CO₃ levels as projected onto [111] have oxygen atoms pointed alternately toward and away from the possible short diagonal glide lines. Thus in gliding, a protruding oxygen atom can pass a Mg position with less friction than it can a Ca position, and a mechanism is set up to favor translations of one level only. It is seen that such unit glides (modified only by a small twin glide) along succeeding short diagonals for each successive (110) level redistribute the positive ions into dolomite successions in the synthesized twin, and simple rotation of the affected carbonate groups builds a twinned lamella. It seems reasonable that the natural twinned lamellae are so produced.

The probable mechanism of the deformation in the grinding experiments described earlier is illustrated in figure 6. The section depicts a detail of the crossing of a unit glide on (110) by a longer glide on (111). Under the varying directions and magnitudes of the loads impressed in grinding, this detail must be reproduced an enormous number of times in the reduction of natural texture dolomite rocks to impalpable powders. Each such intersection of glides produces a nucleus within the dolomite structure in which all neighbors of some carbonate groups are either all Ca or all Mg, and therefore repeated crosses can build up substantial Ca-rich and Mg-rich islands. In the same mechanism the phase relations of positive ions within coherently scattering domains are disrupted. Consequently, the diffraction lines which are peculiar to dolomite, and which
derive their intensity mainly from the difference in scattering power between Ca and Mg, are attenuated. Such lines are thus distinguished from those to which the Ca and Mg make contributions of equal sign, and from those to which the major contributor is oxygen.

Haul and others (5, 6, 10) have recently discussed the thermal decomposition of dolomite and have noted the evidence for an activation energy. In consideration of the observation that calcite and periclase are produced essentially simultaneously, the calcite, at least, being synthesized at orientations controlled by the original dolomite structure, it seems reasonable to assume that the activation energy is associated with interdiffusion of positive ions to produce Ca-rich and Mg-rich localities. This kind of sorting having already been accomplished mechanically by grinding, in the case of the vigorously ground material above, thermal decomposition of this powder proceeds at a notably reduced temperature, more comparable with that at which magnesite itself decomposes.

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References

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