CO₂-rich glass, round calcite crystals, and no liquid immiscibility in the system CaO-SiO₂-CO₂ at 2.5 GPa

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ABSTRACT

Following reports that the miscibility gap between silicate and carbonate liquids located experimentally on feldspar-calcite joins extended to the alkali-free side of the system CaO-Na₂O-Al₂O₃-SiO₂-CO₂, the melting of a mixture of calcite (70 wt%) and quartz was investigated at 2.5 GPa. The isobaric reaction, calcite (CC) + quartz (Qz) = liquid (L) + vapor (V), was reversed at 1350 °C. Quartz and rounded calcite crystals were concentrated at the bottom of the capsule, and CO₂ was distributed in large vapor bubbles in the glass layer and at the top of the capsule. The liquid quenched to transparent glass, which is unusual in carbonate-rich systems. In two-stage reversal experiments, a sample of L + V that was heated to the subsolidus temperature of 1300 °C produced a few rounded calcite grains organized in dendritic patterns; at 1200 °C, dendritic intergrowths of CC + Qz were produced with some coarser-grained areas. The glass was found to contain about 20 wt% CO₂ on the basis of the geometry of phase boundaries and EDS analysis. There was no evidence for immiscible liquids. The round calcite crystals are equilibrium mineral phases, not quenched CaCO₃ liquids, and surface tension effects control their shapes. Infrared spectroscopic studies indicated that (CO₃)²⁻ is the dominant CO₂ species in the glass, and the silicate structure is partially polymerized, probably as a result of interaction between Ca²⁺ and SiO₄ tetrahedra. The phase relationships in the CaCO₃-SiO₂ system, the simplest model for subducted oceanic crust with limestone (or for basalt altered by sea water), show that subducted crust potentially could transport calcite to great depths for long-term storage in the mantle and could also yield low-SiO₂ carbonate-rich magmas under some thermal conditions. Such carbonate-rich melts may be efficient agents for mantle metasomatism.

INTRODUCTION

Melting relations in silicate-carbonate systems involving carbonate-rich melts are relevant for problems related to mantle metasomatism (Green and Wallace, 1988; Baker and Wyllie, 1992; Yaxley et al., 1991; Hauri et al., 1993), the possible involvement of carbonate-rich melts above subducted oceanic slabs (Sweeney et al., 1992; McInnes and Wyllie, 1992), and the generation of carbonatites and associated volcanism (Bell, 1989). The simplest petrologically relevant system combining silicate and carbonate liquids is CaO-SiO₂-CO₂. Melting relationships in this system (with and without H₂O) have been studied at crustal and mantle pressures (Wyllie and Haas, 1965, 1966; Boettcher and Wyllie, 1969; Huang and Wyllie, 1974; Huang et al., 1980).

Liquid immiscibility occurs between silicates and carbonates, extending across a wide range of compositions (Koster van Groos and Wyllie, 1966, 1973; Koster van Groos, 1975; Freestone and Hamilton, 1980), but there are also continuous liquidus paths between some silicate and carbonate-rich melts (Wyllie, 1989; Otto and Wyllie, 1993). There is excellent petrological and experimental evidence that some carbonatites separated from silicate magmas by liquid immiscibility (Le Bas, 1977; Kjarsgaard and Peterson, 1991; Macdonald et al., 1993), but the limits for the intervention of liquid immiscibility have not been completely determined (Lee and Wyllie, 1992a).

Kjarsgaard and Hamilton (1988, 1989) and Brooker and Hamilton (1990) reported miscibility gaps in several feldspar-carbonate systems at 0.2 and 0.5 GPa between 1100 and 1250 °C, at 1.5 GPa and 1225 °C, mostly in the presence of CO₂ vapor. Kjarsgaard and Hamilton (1989) presented a generalized projection of miscibility gaps from all their systems into the triangle (Al₂O₃ + SiO₂)-CaO-Na₂O, as shown in Figure 1. Their miscibility gap at 0.5 GPa and 1250 °C extends into alkali-free silicate-carbonate systems, with nearly pure CaCO₃ as the end-member immiscible carbonate liquid. Brooker and Hamilton (1990) added a second immiscible carbonate-rich liquid to this arrangement. Remarkably rounded, almost-pure CaCO₃ phases were interpreted as quenched, immiscible, carbonate liquid. The occurrence of almost-pure CaCO₃ as an immiscible liquid at temperatures several hundred degrees below the fusion temperature of calcite (Irving and Wyllie, 1975) is puzzling. According to these results, a miscibility gap might be expected in the joins CaCO₃-SiO₂, CaCO₃-CaSiO₃, and CaCO₃-grossu-
larite previously investigated by Huang et al. (1980),
Huang and Wyllie (1974), and Maaloe and Wyllie (1975)
at higher pressures (up to 3 GPa). No immiscible liquids
were found in these earlier studies, although rounded cal-
cite grains were reported. Lee and Wyllie (1992a, 1992b,
and in preparation) studied the phase fields intersected
by the join NaAlSiO₃-CaCO₃ at high pressures. The phase
relationships required that the rounded CaCO₃ phase rep-
resent solid calcite under conditions of the experiments,
and Lee and Wyllie defined the miscibility gap at 2.5
GPa, as shown in Figure 1. Note that the conjugate Ca
carbonate liquid boundary is quite far removed from
pure CaCO₃.

Macdonald et al. (1993) compared the original 0.5-GPa
miscibility gap of Kjarsgaard and Hamilton (1989) with
a revised version (Kjarsgaard and Hamilton, in prepa-
ration), which is similar to our result at 2.5 GPa (Fig. 1).
According to Kjarsgaard (1994 personal communica-
tion), the revision involves textural reinterpretation; he
is now satisfied that the rounded, almost-pure CaCO₃
globules previously interpreted as immiscible liquids are
indeed solid calcite under experimental conditions.

In this contribution, we present experiments in the end-
member system of Figure 1 to check how the results of
Huang et al. (1980) compare with the results of Kjars-
ggaard and Hamilton (1988, 1989) and Brooker and Ham-
ilton (1990). The original studies of Huang and Wyllie
(1974) and Huang et al. (1980) in the system CaO-SiO₂-
CO₂ at pressures up to 3 GPa relied on optical exami-
nation of crushed fragments in immersion oils and pow-
der X-ray diffraction measurements for the identification
of phases. Physical evidence for immiscible liquids was
sought on the basis of criteria established by previous
experience (e.g., Koster van Groos and Wyllie, 1966,
1973). Glassy beads were present above the solidus, but
evidence overall was interpreted in terms of simple liqui-
dus surfaces, without miscibility gaps in the range of
compositions studied. Given the recent results cited above,
however, reexamination of this boundary system is nec-
essary, using the improvements in textural visualization
and interpretation offered by the scanning electron mi-
croscope (SEM). The previously reported liquids that
quenched to glasses with up to 20 wt% dissolved CO₂
have now been studied by infrared spectroscopic meth-
ods (FTIR). Our new results at 2.5 GPa are consistent
with the revised 0.5-GPa results in Figure 1 (Kjarsgaard,
1994 personal communication).

**PHASE RELATIONSHIPS IN CaO-SiO₂-CO₂:
PREVIOUS STUDIES**

Subsolidus reactions in this system have received much
attention (Harker and Tuttle, 1956; Tuttle and Harker,
1957; Haselton et al., 1978). Huang et al. (1980) pub-
dlished a detailed P-T projection for the system to 3 GPa,
combining their new melting results with all relevant
published data, including a tabulation of the experimen-
tal data for the phase diagram for CaSiO₃-CaCO₃ of Huang
and Wyllie (1974). Our experiments were conducted on
a quartz + calcite mixture. Selected univariant curves
emphasizing reactions involving calcite and quartz are

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**Fig. 1.** Silicate-carbonate two-liquid fields projected onto the
compositional triangle (Al₂O₃ + SiO₂)-CaO-Na₂O, CO₂-saturat-
ed, showing results (heavy curves) from (A) Kjarsgaard and
Hamilton (1988, 1989) at 0.5 GPa and 1250 °C, (B) Lee and
Wyllie (1992a, and in preparation) at 2.5 GPa, and (C) Kjars-
ggaard and Hamilton (in preparation) at 0.5 GPa and 1250 °C,
presented in Macdonald et al. (1993, Fig. 5b). The tie line C'
connects coexisting silicate-rich liquid (Ls) and carbonate-rich
liquid (Lc). Note that for A the field Ls + Lc extends all the way
to the Na₂O-free side.

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**Fig. 2.** Selected phase relationships in CaO-SiO₂-CO₂ show-
ing calcite-quartz reactions (Fig. 4 in Huang et al., 1980). Ab-
Breviations: CC = calcite; Wo = wollastonite; Qz = quartz; Sp = spurrite; L = liquid; V = vapor; I = invariant point; S =
singular point. The positions of new experiments at 2.5 GPa
(Table 1) are given by open circles. Reactions specified in
the text are denoted by numbers 1 and 2.
reproduced in Figure 2. Calcite and quartz dissociate together in the reaction

\[
\text{calcite + quartz = wollastonite + vapor } \quad (1)
\]

(Harker and Tuttle, 1956, to 0.3 GPa) at pressures up to the invariant point \( I_1 \) at 1.85 GPa and 1325 °C (Huang et al., 1980), where the assemblage is joined by liquid. Haselton et al. (1978) studied several subsolidus decarbonation reactions and derived \( f_{CO_2} \) data from the results. Their curve for Reaction 1 is somewhat higher than that in Figure 2 at lower pressures, but within error limits it passes through point \( I_1 \). Among the array of curves emanating from \( I_1 \), the incongruent melting reaction

\[
\text{calcite + quartz = liquid } e + \text{ vapor } v \quad (2)
\]
rises to higher pressure. The phase relationships and Reaction 2 are illustrated in Figure 3.

Figure 3A shows the phase compositions in the ternary reactions at 2.7 GPa, with liquid field-boundary compositions constrained by the several joins studied by Huang et al. (1980). The curve a-b-e-c-d is the vapor-saturated liquidus boundary, giving liquid compositions coexisting with minerals and almost-pure CO\(_2\) vapor at \( v \). The field boundaries divide the CO\(_2\)-undersaturated liquidus surface into areas for the primary precipitation of calcite, quartz, wollastonite, larnite, and lime. Reaction 2, like Reaction 1, is a dissociation reaction, and the liquid compositions formed from mixtures on the join CaCO\(_3\)-SiO\(_2\) between \( b \) and \( c \) have lost CO\(_2\). Figure 3B shows the phase fields intersected by the join CaCO\(_3\)-SiO\(_2\) at 2.7 GPa, according to Huang et al. (1980). The eutectic-like liquid \( e' \) in Figure 3B is a projection of the ternary liquid \( e \), as shown in Figure 3A.

Kjarsgaard and Hamilton (1989) did not claim that immiscible liquids exist between quartz or calcium silicates and calcite, but a literal interpretation of their original results in Figure 1, and those of Brooker and Hamilton (1990), implies a miscibility gap between about \( f' \) and \( g' \) on the line CC-Qz in Figure 3A and 3B, which corresponds to the interval \( f \) and \( g \) on the field boundary a-d in Figure 3A. [This is no longer required by Kjarsgaard’s reinterpretation (in preparation); Fig. 1.]

**Experimental Methods**

Primary standard-grade CaCO\(_3\) powders (Alfa Products) and reagent-grade SiO\(_2\) powders (Puratronic) were dried at 110 °C for at least 1 d. A mixture of 30 wt% quartz and 70 wt% calcite was ground under ethanol in an agate mortar to obtain a fine-grained, uniform mixture. About 5 mg of the mixture for each experiment was dried for >1 h in a vacuum drying oven at 110 °C before being loaded into Pt capsules about 4 mm long. This starting bulk composition is identified in Figure 3 by SM.

Experiments were done in a piston-cylinder apparatus 1.27 cm in diameter, with calcium fluoride as a pressure medium. Temperature was controlled and monitored by \( W_5\)-Rh\(_5\)-W\(_5\)-Rh\(_{30}\) thermocouple, with no correction for the effects of pressure on the emf. Experiment duration varied from 0.5 to 4 h, depending on the temperature. Experiments were terminated by turning off the electrical power. The quenching rate was about 100 °C/s for the first 600 °C, and it dropped to about 50 °C/s or less at
lower temperatures. Pressure was not held constant during quenching. The pressure accuracy is about \( \pm 0.05 \) GPa, and temperature accuracy is estimated to be \( \pm 10 \) °C, according to many previous calibrations with this furnace assembly.

Polished surfaces of experimental products were analyzed with a Camscan scanning electron microscope fitted with an energy-dispersive system (EDS). Quenched liquids and crystals were analyzed using a beam current of 0.1 nA on brass with 15 kV acceleration voltage. The quenched liquids were analyzed in raster modes, with a covered area of about 100 \( \mu \text{m}^2 \). The identification of phases was based on texture and morphology observed with a petrographic microscope (with both reflected and transmitted light) and SEM, and compositions were determined by EDS. A thin, plane-parallel plate of glass from the sample of experiment 151, prepared by polishing both sides, was used in a spectroscopic study to identify the \( \text{CO}_2 \) species in glass and to obtain quantitative analyses utilizing infrared absorption spectra from a Nicolet 60SX FTIR spectrometer. In addition, a polished surface of the sample in experiment 151 was prepared to obtain the infrared reflectance spectra with a Nicolet NicoPlan IR microscope.

**Experimental results**

The starting mixture was selected to be near the projected eutectic e' in Figure 3, just within the field for liquidus quartz. The criteria for identifying Reaction 2 are (1) the appearance of liquid, (2) the disappearance of calcite, (3) the appearance of a vapor phase, and (4) probably an increase in the size of quartz compared with the subsolidus assemblage. Few melting reactions in silicate-carbonate systems have been satisfactorily reversed because the liquid commonly crystallizes to a dendritic aggregate during normal rapid quench, and this texture is retained during subsolidus annealing. Therefore, it has been difficult to reach unambiguous conclusions about whether the texture in a reversal experiment was formed during the quench or during the experiment at temperature below the solidus. Because the liquid in our experiments quenched to a glass, we were able to study textures of minerals grown from liquid during annealing in two-stage reversal experiments.
Experiments are listed in Table 1, and their positions are shown in Figure 2. Experimental products are illustrated in Figures 4, 5, and 6, and the results for experiments of various duration (0.5-4 h) are plotted in Figure 7. For comparison, experiments by Huang et al. (1980) on this join at 2.7 GPa varied from 30 to 40 min.

Phase relationships

Figure 4 illustrates an experiment with all phases present (expt. 24, Fig. 7A). The top part of the sample consists of clear glass enclosing large vapor bubbles, with additional vapor cavities above the glass. The lower part consists of crystals of calcite and quartz, with interstitial glass. The corresponding experiment of longer duration (expt. 28) contains the same assemblages, with the sharp interface between the two parts shown in the SEM microphotographs (Fig. 5A). There are fine dendrites extending about 200 µm into glass from the interface between glass and the crystal layer (Fig. 5B), presumably resulting from nucleation during the quench on calcite crystals. The small percentage of glass between the crystals decreases with distance below the interface. The rounded calcite grains in contact with liquid reported by Huang et al. (1980) are clearly displayed. Quartz grains vary from subangular to rounded.

The short-duration experiment at 1400 °C (expt. 23) produced the same four-phase assemblage with similar phase distribution, but the 2-h experiment (no. 26, Fig. 7A) was essentially L + V, with traces of calcite and quartz remaining at the bottom of the capsule. The experiment at 1300 °C (no. 27) is similar to the lower portion of Figures 4 and 5, but with only a trace of interstitial liquid (estimated 1%), distributed sporadically, and small vapor pits along the boundaries among crystals and liquid. At constant pressure, the four-phase assemblage should be isothermal, so we conclude that Reaction 2 occurs at 1350 °C and attribute the trace of liquid at lower temperature to a trace of absorbed H₂O in the starting material. Calcite formed with polyhedral morphology in contact with other calcite grains, but in the presence of interstitial liquid it became subrounded.

The results of three reversal experiments are shown in Figure 7B. Experiments 34R and 41R were first held at 1500 °C for 0.5 h to produce liquid and vapor (Fig. 7A), and temperatures were then rapidly lowered into the subsolidus region at 1300 and 1200 °C, respectively, for 2 h (Fig. 7B). The reversal to 1300 °C consisted largely of glass, with vapor bubbles and some calcite (Fig. 6A), whereas the reversal to 1200 °C contained a much higher proportion of crystals (Fig. 6B). Figure 6A shows sparse,
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**Table 1.** Experiment results in the system CaO-SiO$_2$-CO$_2$ at 2.5 GPa for the mixture 70% calcite, 30% quartz

<table>
<thead>
<tr>
<th>Expt.</th>
<th>$T$ (°C)</th>
<th>$t$ (h)</th>
<th>Resulting assemblage</th>
</tr>
</thead>
<tbody>
<tr>
<td>21</td>
<td>1500</td>
<td>0.5</td>
<td>L + V</td>
</tr>
<tr>
<td>151</td>
<td>1500</td>
<td>0.67</td>
<td>L + V</td>
</tr>
<tr>
<td>22</td>
<td>1450</td>
<td>0.5</td>
<td>L + V</td>
</tr>
<tr>
<td>23</td>
<td>1400</td>
<td>0.5</td>
<td>L + V + CC + Oz</td>
</tr>
<tr>
<td>26</td>
<td>1400</td>
<td>2</td>
<td>L + V + CC + Oz (tr)</td>
</tr>
<tr>
<td>24</td>
<td>1350</td>
<td>2</td>
<td>L + V + CC + Oz</td>
</tr>
<tr>
<td>28</td>
<td>1350</td>
<td>4</td>
<td>L + V + CC + Oz</td>
</tr>
<tr>
<td>34R</td>
<td>1500 to 1300</td>
<td>0 to 2</td>
<td>L + V + CC + Oz (tr)</td>
</tr>
<tr>
<td>35R</td>
<td>1350 to 1300</td>
<td>2 to 2</td>
<td>L + V + CC + Oz</td>
</tr>
<tr>
<td>41R</td>
<td>1500 to 1200</td>
<td>0.5 to 2</td>
<td>L + V + CC + Oz</td>
</tr>
</tbody>
</table>

Note: abbreviations are defined as follows: CC = calcite, Oz = quartz, L = liquid, V = vapor, tr = trace amount. The expt. nos. marked by R indicate reversals. For example, expt. 34R was first held at 1500 °C for 0.5 h and then quenched to 1300 °C and held for 2 h.

Fig. 6. SEM photomicrographs of experiments reversed from the assemblage L + V at 1500 °C (Table 1). Phases in direction of lighter gray: quartz → calcite → liquid. (A) Expt. no. 34R, held at 1300 °C. Glass with calcite dendrites, composed of small rounded units arranged in linear structures. (B) Expt. no. 41R, held at 1200 °C. Calcite dendrites and very small quartz grains with interstitial quenched liquid, with glass increasing in amount locally in regions of coarser grain size.

rounded grains to elongated ovoids of calcite with organized structure in dendritic form. A few quartz crystals were found near the edge of the sample. The reversal to 100 °C lower consisted for the most part of glass containing fine-grained dendritic calcite-quartz intergrowths, enclosing some irregular areas with much larger, elongated calcite and a few larger prismatic quartz crystals. Reversal experiment 35R was conducted initially to generate the four-phase assemblage of Figure 4 and then lowered through 50 °C into the subsolidus region. The upper, originally liquid part produced sparse, rounded calcite in dendritic arrays (compare Fig. 6A), the interface between the two parts became less well defined, and a few hexagonal or rounded prismatic crystals of quartz grew in the glass near the interface.

Many more experiments would be necessary to work out the detailed nature of mineral nucleation and growth during subsolidus annealing, but these experiments confirm that Reaction 2 is in reversible equilibrium at temperatures near 1350 °C, with results consistent with the topology of the phase diagram in Figure 3B. This temperature is about 50 °C lower than the curve of Huang et al. (1980). Despite the observation that liquids in many carbonate-rich systems cannot be quenched to glasses, the liquid compositions near e (Fig. 3A) crystallize only slowly, even 150 °C below the solidus. For the equilibrium assemblage of calcite + quartz to grow, it is necessary for the free CO$_2$ to diffuse into the subsolidus glass and to combine with Ca$^{2+}$, which probably has to be extracted from strong bonding in the SiO$_2$ structural environment.

**Rounded calcite without liquid immiscibility**

Kjarsgaard and Hamilton (1988, 1989) and Brooker and Hamilton (1990) have interpreted rounded and dumbbell-shaped calcite grains as liquids with compositions equal to or very close to pure CaCO$_3$, immiscible with the silicate-rich liquid in which they were suspended, but Kjarsgaard (1994 personal communication) is now satisfied with the interpretation of Lee and Wyllie (1992b) that round CaCO$_3$ in the join albite-calcite represents a solid mineral. Although the round calcite in our experiments with liquid certainly has the physical appearance of an immiscible liquid (Fig. 5), we are satisfied that its distribution proves it was a mineral under experimental conditions and not a CaCO$_3$ liquid.

Evidence that the round calcite represents a mineral phase rather than quenched liquid includes the following: (1) the intermixed distribution of round calcite and quartz beneath a layer of silicate liquid (Fig. 5), (2) the restriction of the assemblage (rounded calcite and other phases) to a few tens of degrees between the solidus and a homo-
geneous silicate-CO$_2$ liquid, (3) the rounded calcite grains in the experiment reversed from the liquid field being parts of an organized dendritic feature (Fig. 6A), which represents the growth of crystals and not the exsolution of liquid (which subsequently solidified to pure CaCO$_3$), (4) the melting temperature of calcite at 2.7 GPa being 1615 °C (Fig. 3B), which is more than 250 °C higher than the disappearance of the round phase (labeled calcite) in Figure 7, and (5) the internally consistent phase relations by Huang et al. (1980) for the complete system CaO-SiO$_2$-CO$_2$ without the intervention of liquid immiscibility, which are now confirmed by our test of the phase fields intersected by the join SiO$_2$-CaCO$_3$ at 2.5 GPa (Figs. 2 and 3).

The results in Table 1 and Figure 7 and the samples illustrated in Figures 4, 5, and 6 are consistent with the previous interpretation of a simple eutectic reaction, and we cannot see how these results could be reconciled with a miscibility gap in Figure 3B. The same arguments apply to the round calcite grains in the join grossular-CaCO$_3$ at 3.0 GPa (Maaloe and Wyllie, 1975). The rounded shapes of the calcite crystals are most probably the result of surface tension effects between calcite and the carbonate-silicate liquids.

**Composition of glass**

The composition of the liquid formed from the starting mixture in our experiments was estimated from EDS analyses using SEM, complementing the chemographic solubilities represented by the phase diagram in Figure 3A of Huang et al. (1980). No attempt was made to measure CO$_2$ solubility by weight loss after ignition or by FTIR.

The ratio of SiO$_2$ to CaO in the glass from L + V experiments measured by EDS/SEM was coincident with that in the starting mixture SM. The presence of vapor confirmed that the glass composition contained less CO$_2$ than did SM and therefore lay on the line v-SM extended in Figure 3A. The determination of CO$_2$ in the quenched glasses using EDS analysis was based on the deficiency of detectable elements below 100%, using calcite as a reference phase and applying a correction based on quartz analyses. The results (probable error 1-2 wt%) showed decreasing CO$_2$ solubility with increasing temperature: 1350 °C = 17.0%, 1400 °C = 16.9%, 1450 °C = 15.6%, and 1500 °C = 14.2%. These values straddle m in Figure 3A.

Huang et al. (1980) estimated the position of the CO$_2$-saturated liquidus field boundary a-b-e-c-d (Fig. 3A) from the geometry of phase fields intersected by several joins through the system. Their field boundary gave a value of about 16 wt% for the CO$_2$ content of liquid m, which is the liquid composition obtained by projection through v-SM. Complete melting of the mixture SM should yield liquid m and vapor v (compare Fig. 3B).

**Structure of glass**

The infrared reflectance spectrum of the glass from experiment 151 is illustrated in Figure 8A. It shows a broad band between 1160 and 1570 cm$^{-1}$, resulting from the $v_1$ vibrational mode of the (CO$_3$)$_2^-$ unit, which is split into two peaks at 1477 and 1392 cm$^{-1}$. The peak positioned at 854 cm$^{-1}$ is assigned to the $v_2$ vibration of carbonate. Other bands in Figure 8A between 550 and 1160 cm$^{-1}$ are attributed to silicate vibrations. The broad feature between 550 and 800 cm$^{-1}$ is probably due to the Si-O-
Si asymmetric bending and the bridging O asymmetric stretching vibrations. The features between 800 and 1160 cm\(^{-1}\) result from the asymmetric Si-O stretching of SiO\(_4\) tetrahedra. These assignments are summarized in McMillan (1984). The two peaks at 995 and 917 cm\(^{-1}\) probably arise from the stretching of silica tetrahedra with two and three nonbridging O atoms, respectively. This gives an indication of partial polymerization in the glass. Figure 8B is an infrared absorption spectrum of the glass quenched from 1500 °C (expt. 151), showing a broad band between 3700 and 2300 cm\(^{-1}\), superimposed with two narrower bands centered at 2520 and 2920 cm\(^{-1}\). The broad band is caused by OH\(^-\) stretching (equivalent to about 0.5 wt% total H\(_2\)O in the glass, using the calibration of Newman et al., 1986), and the two bands around 2520 and 2920 cm\(^{-1}\) are assigned to the combination of the \(\nu_1\) + \(\nu_3\) vibrational modes and the first overtone of \(\nu_3\), for the (CO\(_3\))\(^2-\) complex, respectively. The split on the \(\nu_3\) mode in Figure 8A is mimicked in the 2520 cm\(^{-1}\) band but no longer is resolved in the 2920 cm\(^{-1}\) band. The intensity of the characteristic band around 2350 cm\(^{-1}\) for the \(\nu_3\) antisymmetric stretching mode of molecular CO\(_2\) is insignificant compared with the carbonate bands in the glass spectrum. Therefore, the CO\(_2\) species in the glass is mainly carbonate.

The bands at 2520 and 2920 cm\(^{-1}\) discussed above may be used for the quantitative analysis of CO\(_2\) concentrations in glasses, but such an analysis requires a detailed calibration for this purpose. Table 2 compares the CO\(_2\) molar absorption coefficients of a calcite crystal with the glass from experiment 151, on the basis of chemical data from EDS/SEM and the IR spectroscopy, with the technique described elsewhere (e.g., Fine and Stolper, 1985). The uncertainty is somewhat large if both the uncertainties on the base-line correction for the peaks (Table 2) and the CO\(_2\) concentration from EDS are considered. However, it indicates that the absorption coefficients of the carbonate bands in the glass are generally lower than those of calcite (CC\(_{\text{mean}}\), a calculated mean over all possible orientation of calcite crystals) and likely are due to the greater distortion on the carbonate ions in the glass. The transparent nature of the quenched liquids and the slow growth rate of calcite crystals during both quenching and reversal experiments are likely to be controlled by the liquid structure. However, the detailed effects of liquid structures on the growth mechanism of calcite are not established. Probably the sluggish crystallization is due to the strong interaction of Ca\(^{2+}\) with SiO\(_4\) tetrahedra.

**Discussion**

The main objective of this study was to test for immiscible liquids in the system CaO-SiO\(_2\)-CO\(_2\), given the
results of Kjarsgaard and Hamilton (1988, 1989) and Brooker and Hamilton (1990) in related systems summarized in Figure 1. Associated objectives were to examine their interpretation of the rounded and dumbbell-shaped CaCO\textsubscript{3} phase as a quenched liquid, with the resultant conclusion that the immiscible carbonate liquids could be extremely rich in CaCO\textsubscript{3}.

No miscibility gap was found between silicate and carbonate liquids at high pressures, which is consistent with the previous conclusion (Huang et al., 1980) and with Kjarsgaard’s revised interpretation in related systems at 0.5 GPa (in Macdonald et al., 1993). The rounded calcite crystals shown in Figure 5 illustrate the tendency of calcite to form in rounded habits in many systems, including those without silicates (e.g., Wyllie and Tuttle, 1960). We presented evidence supporting our conclusion (Lee and Wyllie, 1992a, 1992b) that the round CaCO\textsubscript{3} phase in quenched samples in parts of the system Na\textsubscript{2}O-CaO-Al\textsubscript{2}O\textsubscript{3}-SiO\textsubscript{2}-CO\textsubscript{2}, represented crystalline calcite under experimental conditions, which is now accepted by Kjarsgaard’s reinterpretation (1994 personal communication). This interpretation, along with our results from NaAl\textsubscript{2}Si\textsubscript{2}O\textsubscript{5}-CaCO\textsubscript{3}, from 1.0 to 2.5 GPa and Kjarsgaard’s revised miscibility gap at 0.5 GPa, indicates that, although immiscible carbonate liquids may be sówki, they appear to remain well separated from pure CaCO\textsubscript{3} in composition.

The occurrence of metasomatism by carbonate-rich melts in mantle lherzolites is now well established (e.g., Yaxley et al., 1991; Hauri et al., 1993; Ionov et al., 1993). These melts may be derived by the action of CO\textsubscript{2} on lherzolite (e.g., Wyllie and Huang, 1976; Egglger, 1978; Wallace and Green, 1988; Dalton and Wood, 1993) or by CO\textsubscript{2} or melts derived from subducted oceanic crust (Huang et al., 1980; Sweeney et al., 1992). The conditions under which partially carbonated oceanic crust might transport carbonate deep into the Earth’s interior for long-term storage, or might become reduced to generate diamonds (Luth, 1993), or might yield a carbonate-rich melt are of particular interest.

The two reactions in Figure 2 showing the stability limit of dry pelagic limestones, represented by the assemblage calcite + quartz, are situated well above expected temperatures in subducted oceanic crust, confirming that calcite may survive subduction for storage in the mantle (Huang et al., 1980). In the presence of H\textsubscript{2}O, however, reaction and partial melting would yield traces of carbonate-rich melt at much lower temperatures (Wyllie and Haas, 1965; Boettcher and Wyllie, 1969; Huang et al., 1980). The results in Figure 3 focus on the existence of a eutectic, vapor-saturated liquid, with composition near 45% CaO, 35% SiO\textsubscript{2}, and 20% CO\textsubscript{2}, coexisting with calcite and quartz at 2.5 GPa, and with calcite and wollastonite at pressures below about 1.8 GPa (Fig. 2, Huang et al., 1980). If pockets of pelagic limestones survive subduction, they can generate liquids of this composition at depths of 100 km or more where temperatures reach 1500 °C. Such liquids would change composition rapidly by reaction with lherzolite or eclogite, causing significant metasomatic changes in the host rocks as they changed into the carbonate-rich melts in equilibrium with their more mafic hosts.

The significance of the predominance of \(\text{(CO}_2\text{)}^{2-}\overline{\text{ }}\) over molecular CO\textsubscript{2} dissolved in the liquids at 2.5 GPa in the system CaO-SiO\textsubscript{2}-CO\textsubscript{2}, along with the structural information, indicates that up to 20% of carbonates can be incorporated into partially polymerized silicate liquid in the presence of a high content of network modifiers, such as Ca\textsuperscript{2+}. Hence, a CO\textsubscript{2}-rich silicate liquid at mantle conditions can evolve to be highly carbonated as silicates are fractionated during magma transportation. The liquid is slow to crystallize, however, compared with most carbonate-rich liquids, which cannot be quenched to glasses. Figure 6A shows the sluggish behavior of liquid carried 50 °C below its solidus temperature, where only a few calcite dendrites have crystallized.

Our partly melted experimental samples demonstrated the ease with which carbonate-rich magma, if formed, could be separated from its source rock (Hunter and McKenzie, 1989; White and Wyllie, 1992). Figure 4 shows a 2-h experiment in which all crystals were concentrated into a lower layer, with only little interstitial liquid remaining. It appears that it takes longer for the large CO\textsubscript{2} vapor bubbles to rise through the liquid than it does for minerals to set out from the liquid or for liquid to rise from the mineral matrix.

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