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Four-Phase Curve in the System CaAl₂Si₂O₈-SiO₂-H₂O between 1 and 10 Kilobars*)

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With 8 figures in the text and 3 tables

Abstract

The $CaAl_2Si_2O_8(An)$ -SiO₂(Qz) system has a eutectic at SiO₂ 49.5 weight per cent (SCHAIRER and BOWEN, 1947); the ternary system An-Qz-H₂O projected from H₂O is also of eutectic type. Anorthite, quartz, liquid, and gas coexist under the following conditions:

		$CaAl_2Si_2O_8$: SiO_2	
$P_{H,O} = P_{total}$	Temperature	in liquid	H_2O in liquid
(bars)	$^{\circ}\mathrm{C}$	(weight per cent)	(weight per cent)
1000	1040 ± 10	32:68	?
2000	922 ± 3	37:63	~ 6
5000	815 ± 5	42:58	~ 10
10000	757 ± 7	$\mathbf{48:52}$	> 12

The H₂O-saturated Qz-H₂O liquidus lies at $1130 \pm 5^{\circ}$ C at 2000 bars, $1065 \pm 5^{\circ}$ C at 5000 bars, and $1055 \pm 10^{\circ}$ C at 10,000 bars. The H₂O-saturated An-H₂O liquidus determined by YODER (1954) was confirmed to 5000 bars and extended to 10,000 bars at $1110 \pm 10^{\circ}$ C. The unit-cell parameters are given for four samples of anorthite synthesized from various compositions, temperatures, and pressures but only small differences were found, the most significant being for the cell volume and inter-axial angle α . An indexed powder-diffraction pattern is given for primitive anorthite.

Petrologic applications suggested include explanation of differentiation paths of magmas containing quartz and feldspar, a geobarometer for H_2O pressure, and explanation for resorbed quartz crystals in volcanic rocks.

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INTRODUCTION

Rocks consisting largely of quartz, plagioclase, and alkali feldspar are abundant, and the phase relations between silica and feldspar components have been the subject of intensive experimental study (TUTTLE and Bowen, 1958; STEWART, 1957, 1958; YODER, STEWART, and SMITH, 1957; SHAW, 1963; LUTH, JAHNS, and TUTTLE, 1964; and by WINKLER and VON PLATEN, summarized by WINKLER, 1965, p. 178-199). This study of the system CaAl₂Si₂O₈ (hereafter called An)-SiO₂ (called Qz)-H₂O supplements these investigations, and is of interest because discussion of the phase relations of quartz and feldspars over a wide range of temperatures and pressures requires this information. The data are applicable strictly to few rocks because anorthite-quartz assemblages are very rare (BIRKELAND, 1958, p. 351-352), but they do offer further evidence of the nature of silicate-water systems over a wide range of pressures that can be combined with similar results by others (DOLIVO-DOBROVOL'SKIY, 1961; ORLOVA, 1964; KADIK and KHITAROV, 1963; KHITAROV, KADIK and LEBEDEV, 1963; SHAW, 1964).

Professor LAVES has contributed much to our knowledge of many of the complexities of the structures of plagioclase feldspars. His discussions of the polymorphism of NaAlSi₃O₈ and CaAl₂Si₂O₈ have stimulated the author to investigate these compounds further, and his contributions to the study of unmixing in peristerites and labradorites suggest many experiments. In the present paper the unit-cell parameters of anorthites synthesized under various conditions will be presented, as will an indexed powder diffraction pattern of anorthite supplementing the partial indexing of GOLDSMITH and LAVES (1956, p. 399) and BROWN (1960) p. 315). Reviews of the crystallographic complexities of anorthite are given by GAY (1962, p. 42—45) and MEGAW (1962, p. 121—124) for those unfamiliar with past research.

EXPERIMENTAL METHODS

Starting materials

Glasses prepared by SCHAIRER and BOWEN (1947, p. 70-72) were used for starting materials. For some experiments the crystalline equivalents of these glasses were prepared either by anhydrous crystallization



Fig. 1. Index of refraction of anhydrous glasses in the system An-Qz at room temperature, and of hydrous glasses quenched to room temperature at the pressures shown by the numbers. Data for glasses containing less than 5% crystals are plotted, but most samples contained less than 2% crystals and the small correction involved was not applied.

Hydrous An-rich glass has a lower index of refraction than the anhydrous composition, and Qz-rich hydrous glass has a higher index of refraction than its anhydrous equivalent. The effect is enhanced the higher the pressure during quenching. Other factors than the H_2O content of the glass may be responsible in part for the effect, such as pressure densification of the glass at constant H_2O content, or the presence of residual stresses in the initial anhydrous glass that affected its original index of refraction. to anorthite and cristobalite by SCHAIRER and BOWEN or by hydrothermal crystallization to anorthite and quartz at 750—800°C at 2 kb for several hundred hours. Where necessary, new batches and additional compositions were prepared under the direction of J. F. Schairer. The index of refraction of each glass was measured with freshly calibrated oils in sodium light, and the results (Figure 1) plot as a curve at slightly lower values than previously reported by SCHAIRER and BOWEN (1947, Table 1).

Glass of $CaAl_2Si_2O_8$ composition was very carefully prepared from $CaCO_3$ (BAKER's lot 51735), Al_2O_3 (T 61 grade), and SiO_2 (Lisbon, Maryland, quartz inverted to cristobalite at 1500°C), and was crystallized at 1200°C for 12 hours. The index of refraction of the glass in sodium light was 1.574 ± 0.0005 at $25^{\circ}C$. The melting point of this crystalline anorthite was determined with great care under J. F. Schairer's direction to be $1551.0^{\circ}C \pm 1.5^{\circ}C$; the melting interval (first sintering to all glass) was $4^{\circ}C$. A Pt-Pt90Rh10 thermocouple calibrated against the melting point of diopside, $1391.5^{\circ}C$, and pseudowollastonite, $1544^{\circ}C$, was used, and a linear extrapolation was assumed.

Very pure quartz (0.03% residue on evaporation with HF and H₂SO₄) from Lisbon, Maryland, and a silica glass prepared by C. N. Fenner were used as sources of silica.

In a few experiments finely ground mixtures of crystalline synthetic anorthite plus silica glass were used as starting materials.

A sample of lawsonite from Panoche Pass, California, supplied by W. G. Ernst was used for a few experiments at 10 kb.

Procedure

The experiments were performed at the Geophysical Laboratory, Carnegie Institution of Washington, in internally heated high pressure apparatus designed by H. S. YODER, Jr. (1950). Samples were sealed in platinum tubes with H₂O present in excess of saturation requirements. The temperature given is the highest temperature measured during an experiment; the lowest temperature during an experiment was less than 15° C below the specified temperature of the experiment. Pressure oscillations resulting from temperature regulation normally were about 2%of the total pressure of the experiment. The total pressure was measured with calibrated gauges, and is accurate to within 1% of the stated pressure. Quenching to 500° C took place in 2 to 3 minutes following the shutting off of power to the furnace, and the samples were quenched isobarically by pumping during the quench.

CRYSTALLINE PHASES

Anorthite

Anorthite crystallized readily as euhedral crystals tabular on (010) (Figure 2) or as sub-parallel aggregates in experiments made in the presence of silicate liquid. Albite-twinned crystals were common, and occasionally twinning on other laws was observed. Experiments as short as one hour yielded crystals as large as those produced in the longest experiments, 45 hours. Liquids quenched from as much as 75°C above the anorthite plus liquid field yielded distinctive radiating fibrous aggregates of anorthite (Figure 3) formed during the quenching.

LAVES and GOLDSMITH (1955) and GOLDSMITH and LAVES (1956) have shown how strongly the polymorphism of anorthite as observed at room temperature is affected by the thermal history of the sample, and that the presence of lines with h + k even, l odd ("c-reflections") in the powder diffraction pattern indicates that the lattice is primitive with a c axis of \sim 14 Å. As observed at room temperature on x-ray photographs a weak 111 diffraction line appeared from anorthite synthesized from glass after as little as one hour under the temperature and pressure of the experiment, and this line was strong after as little as two hours under many experimental conditions. Geiger-counter diffractometer traces were only slightly less sensitive for detecting the $\overline{11}1$ line. The qualitative data at hand are inadequate to ascertain any temperature dependence on the rate of development of the phenomenon responsible for the change. BROWN, HOFFMAN, and LAVES (1963) have shown that the "c-reflections" disappear continuously on heating between 25°C and 350°C in a sample of natural anorthite from Vesuvius and reappear on cooling, and BLOSS (1964) has indicated other rapidly reversible optical changes at higher temperatures. The presence or absence of c-reflections at 25°C apparently indicates the structural state the crystal attained both under the experimental conditions and during the cooling from these temperatures to room temperature. The presence of "c-reflections" in either synthetic or natural crystals therefore lacks significance as either a geothermometer or geobarometer.

LAVES and GOLDSMITH (1955, p. 233) noted "... no major change in lattice constants can be correlated with the various thermal states. Some small changes are observable, however, that can be correlated with the degree of diffuseness of the type (c) reflections ..." and on p. 234 they said "The size of the 'out-of-step' domains thus has a small but definite influence on the lattice constants". To investigate this problem further,

the unit-cell dimensions were determined for four anorthites synthesized under a variety of experimental conditions yielding $\overline{111}$ reflections varying in intensity from very weak to very distinct. (Sample ANS-26 has the weakest $\overline{11}1$ reflection, and sample ANS-305 the most distinct.) The results are given in Table 1, together with other reported measurements for anorthite. Comparisons between data collected in different ways mean little, but comparisons of the changes shown by the natural and heated samples from Salem, India (with sharp and very diffuse "c-reflections", respectively), with the new data and with the sample from Monte Somma which has strong "c-reflections" all indicate that the primitive cell has a smaller volume, a distinctly smaller value of α , possibly smaller values for a and c, and possibly larger value of γ . Only a few differences are apparent by inspection of the powder-diffraction patterns, the most notable being the larger separation of the $\overline{2}42$ and $\overline{24}2$ reflections $(0.37^{\circ} 2 \theta \operatorname{CuK} \alpha_1$ in ANS 305 with distinct "c-reflections", and 0.30° in ANS-26 with very weak "c-reflections"), and smaller separation of $\overline{13}4$ and $\overline{13}4$ in ANS 305 (1.81° vs. 1.86°). Anorthite produced in the presence of excess silica was indistinguishable from that formed from pure An glass. An indexed x-ray powder-diffraction pattern of primitive anorthite obtained by least-squares methods using the latest version of the Evans, Appleman, and Handwerker (1963) computer program (D. E. APPLEMAN, written communication, 1966) is given in Table 2. Nine distinctive "c-reflections" were observed in this pattern, which was

Fig. 4. Hexagonal dipyramidal crystals of quartz with six-fold layers of fluid inclusions, interpreted to have been β -quartz under experimental conditions. Starting material An₃₀Qz₇₀ glass, 2 kb, 975°C, 22 hours, run ANS-37.

Fig. 5. Crystals of quartz showing the rapid axial growth that results in the entrapment of fluid inclusions. Sample interpreted to have been hexagonal under experimental conditions because of six-fold symmetry. Starting material $An_{30}Qz_{70}$ glass, 2 kb, 925°C, 2 hours, run ANS-9.

Fig. 6. Composite photomicrograph of quartz crystals with α -quartz morphology and three-fold layers of fluid inclusions. Starting material An₄₀Qz₆₀ glass, 10 kb, 765°C, 4 hours, run ANS-351.

Fig. 2. Anorthite crystals with the tabular habit commonly found in experiments near the liquidus. Starting material $An_{50}Qz_{50}$ glass, 10 kb, 765°C, 4 hours, run ANS-348. Glass nearly matching the index of refraction of the oil is present in this photomicrograph and in Figures 3 to 6.

Fig. 3. Radiating, fibrous aggregates of anorthite, formed on quenching, and plates of " β -alumina", believed to have been present during the experiment. Starting material An₉₀Qz₁₀ glass, 10 kb, 1100°C, 2 hours, run ANS-315.



obtained with a diffractometer equipped with a pulse height analyser for $CuK\alpha$ radiation. The lines at $d \sim 3.74$ and ~ 2.49 Å reported by LAVES and GOLDSMITH (1955, p. 233) to show spacing changes with variation in intensity of "c-reflections" seem to involve splitting of overlapping "a-reflections".

Silica polymorphs

Quartz most commonly formed sharp hexagonal dipyramids typical of β -quartz (Figure 4), though it all transformed to α -quartz on cooling. Many of these crystals contained inclusions arranged in two layers of six inclusions, one layer located above the girdle of the crystal and the other below it, with the inclusions in crystallographic register with each other. Such inclusions are entrapped by rapid initial growth along the axial directions (Figure 5). In some experiments at 10 kb the quartz plus liquid field occurs at temperatures below 815° C so that α -quartz is stable on the liquidus (YODER, 1950), and rhombohedral crystals of quartz grew with two layers of inclusions with the three inclusions in each layer displaced 60° relative to the other layer (Figure 6).

No evidence was obtained for solid solution in the quartz.

The products of some experiments at 1 kb contained tridymite and cristobalite. Most cristobalite originated from crystalline starting materials and all presumably was metastable. Tridymite occurred at 1040° C and quartz was obtained abundantly at 1025° C, thus confirming the temperature of 1040° C of the quartz-tridymite transition reported for 1 kb by TUTTLE and BOWEN (1958, p. 30).

Other phases

An-rich compositions held near the liquidus at high pressures contained from traces to 10% of thin hexagonal plates of " β -alumina" of unknown composition (Figure 3). The " β -alumina" arose from the incongruent solubility of anorthite in the gas, which was enriched in silica. The effect was pressure sensitive, being extensive at 10 kb, much less so at 5 kb, and not observable at lower pressures, in parallel with the decreased solubility in the gas at low pressures. On quenching, the gas condensed to liquid water and hydrous silicate glass balls, which represented the dissolved solids.

The surfaces of charges of silica-rich compositions $(An_{45}Qz_{55}-An_{30}Qz_{70})$ in many cases showed isolated clots of small euhedral bladed crystals deposited from the gas during the quench interval. The crystals were biaxial (+), with moderate 2V, $\alpha \sim 1.493$, $\gamma \sim 1.51$, and an extinction angle of 38°. The amount was always insufficient to be detected by x-ray examination by routine methods. The mineral was not identified, but presumably was a zeolite.

HYDROUS GLASSES

In many experiments the hydrous silicate liquid quenched to bubblefree hydrous glass on which the index of refraction, specific gravity, and H_2O content were measured. Such data are potentially useful in evaluating the structure of the melts and the role of H_2O in them, and will be discussed elsewhere. The trends of the variation of index of refraction with H_2O content are shown in Figure 1. Some of these data are given in Table 3, along with the H_2O content determined by the weight loss of glass slugs after drying to constant weight at $110^{\circ}C$ and then slowly heating to $1200^{\circ}C$.

Hydrous liquids produced at 5 kb and below can be quenched without loss of H₂O, and their H₂O content can be found by the weight loss technique, but the hydrous liquids produced at 10 kb cannot be quenched without loss of H₂O from the liquid. The system SiO₂-H₂O at 10 kb as studied by the author by a static equilibrium method yields hydrous silicate liquid containing 22% H₂O by weight at 1055°C. The following observations by my colleague EDWIN ROEDDER (written communication, 1963) on fluid inclusions in α -quartz synthesized at 10 kb (Figure 6) (sample ANS 351, $An_{40}Qz_{60}$ glass = 78.63%, H_2O = 21.37%, 765°C, 4 hours) indicate that the hydrous silicate liquid near the four-phase point at 10 kb contains more $\rm H_2O$ than the $\sim 12\%$ found in the quenched glasses. During growth the quartz crystal engulfed a portion of hydrous silicate liquid, and subsequently the quartz served as an impermeable container. The inclusions now consist of rounded bodies of hydrous silicate glass and a spherical drop of water with a bubble of gas. Under the conditions of formation the liquid that was trapped was homogeneous and in equilibrium with quartz and gas, but during quenching has become heterogeneous principally by exsolution of H_2O , probably initially in the form of gas. The gas has later condensed to hydrous liquid plus water vapor. Measured at 1250 X with a micrometer ocular, two such inclusions were found to contain 7.9, 6.2% water by volume, and 0.5, 0.3% vapor. Assuming that the hydrous silicate glass has properties similar to those produced by quenching other samples of the same composition, the volumetric relations observed in the inclusions indicate that the initial homogeneous liquid contained 14.6, 15.3% H₂O by weight. The accuracy

of the measurements is less than the precision of the two measurements reported. Optical measurements on such small objects with spherical shapes are very difficult, and the errors caused by the negative lenses of the spherical objects normally tend toward significant underestimation of the size of the water drop in the spherical glass inclusion. It is uncertain if the H_2O content of the original liquid in the sample studied ever was as high as the H_2O content of the charge, but it was higher than is now found in liquids free to exsolve H_2O in the capsule during quenching. The figure obtained by weight loss methods thus sets a lower limit on the H_2O content.

Under isobaric conditions the H_2O content of a H_2O -saturated liquid of fixed An : Qz ratio decreases as the temperature is raised. The H_2O content may vary isothermally with variation of the An : Qz ratio, decreasing as the content of An increases. In such a case the effects are probably not greater than the uncertainty of the individual measurements, about 10% of the amount of H_2O reported at that pressure.

The hydrous An-rich melts have such low viscosity that anorthite crystals settled 2 to 3 mm in times averaging a few hours; anorthite settling was observed in compositions richer in Qz as the H_2O pressure increased. No evidence was seen of settling of quartz.

PHASE RELATIONS

System An-H₂O

The H₂O-saturated An-H₂O liquidus was reported by YODER (1954, p. 107; oral communication, 1957) to be $1343 \pm 5^{\circ}$ C at 2 kb, and $1235 \pm \pm 5^{\circ}$ C at 5 kb. These points are consistent with the data given in Table 3, and the liquidus at 10 kb is shown to lie at $1110 \pm 10^{\circ}$ C. This point has also been determined as $1115 \pm 5^{\circ}$ C by YODER (1965, p. 85—88). The presence of β -alumina has been ignored in interpreting the liquidus relations, and the solubility in the gas has not been determined.

Under some conditions of temperature and pressure anorthite reacts with gas to form hydrous phases (zeolites), assemblages including one or more phases with aluminum in six-fold coordination (grossularite, corundum, aluminosilicates) and free silica, or to hydrous phases that contain aluminum in six-fold coordination (lawsonite, zoisite, pumpellyite, prehnite) (P. LE COMTE, written communication, 1959; D. H. LINDSLEY, oral communication, 1963; NEWTON and KENNEDY, 1963; CRAWFORD and FYFE, 1965; and NEWTON, 1966). None of these phases or assemblages were synthesized stably in the present research, and none would be expected in the pressure-temperature range studied on the basis of all information available to the author to date. The natural lawsonite held at 10 kb decomposed at 940°C and 750°C to clinozoisite, epidote, and gas, and the clinozoisite was in turn decomposing to anorthite when the experiments were terminated. It was concluded that anorthite was the stable calcium-bearing silicate at the H₂O-saturated liquidus throughout this investigation.

System Qz-H₂O

The H₂O-saturated Qz-H₂O liquidus has been studied by TUTTLE and ENGLAND (1955), OSTROVSKIY, MISHINA, and POVILAITIS (1959), KENNEDY, WASSERBURG, HEARD, and NEWTON (1962) and STEWART (unpublished data). There is general agreement that the H₂O-saturated liquidus at 2 kb is $1130 \pm 5^{\circ}$ C. This liquidus at 5 kb is $1065 \pm 5^{\circ}$ C and at 10 kb is $1055 \pm 5^{\circ}$ C according to the present work; OSTROVSKIY and others (1959) and KENNEDY and others (1962) report temperatures 15° to 25°C higher at 5 kb, which seem incompatible with data of Table 3 for the H₂O-saturated liquidus for quartz in the system Qz-An-H₂O. KENNEDY and others (1962) claim the upper critical end point lies at 1080°C and 9.7 kb; the results of the author based on the distribution of stable phase assemblages on an isobaric T-X section indicate that the liquid on the three-phase curve at 10 kb contains $22 \pm 2\%$ H₂O, and the gas contains $38 \pm 2.0\%$ Qz. These results indicate that the upper critical end point lies at some pressure higher than 10 kb, and probably at a temperature lower than 1055° C, although it is possible for the temperature to be either lower or higher with increasing pressure (KADIK and KHITAROV, 1963).

The steep slope of the univariant curve representing stable assemblages of β -quartz + liquid + gas in the H₂O-saturated Qz-H₂O system (see Figure 7) is probably caused by a change in the structure of the hydrous melt. On a plot of density against index of refraction such glasses plot on the line between quartz and water. A similarity of the structures of quartz and the melt could account for the observed slope.

System An-Qz-H₂O

The data of Table 3 locate the saturated liquidus at several pressures. The ternary system projected from the H_2O apex to the An-Qz sideline yields a diagram of eutectic type (Figure 8). Gas coexists with the phases

Fig. 7. P-T projection of the H₂O-saturated liquidus in the systems Qz-H₂O, An-H₂O, and An-Qz-H₂O. The curve for the α - β quartz inversion is from YODER (1950) and for the quartz-tridymite transition from TUTTLE and BOWEN (1958); An-H₂O liquidus at 2 and 5 kb is from YODER (1954; oral communication, 1957).

Dots represent liquid plus gas, and crosses represent crystals plus gas.

shown on the diagram, and only the An : Qz ratios of the liquids are shown. The amount of H_2O contained in the glasses (liquids) are shown in Table 3 where known. Four phases coexist at the "eutectic": anorthite, a silica polymorph, hydrous silicate liquid, and gas. The composition of the liquid at the four phase point varies with pressure, as does the temperature of the point:

		$CaAl_2Si_2O_8:SiO_2$	
$P_{H_{3}O} = P_{total}$	Temperature	in liquid	H_2O in liquid
(bars)	$^{\circ}\mathrm{C}$	(weight per cent)	(weight per cent)
1000	1040 ± 10	32:68	?
2000	922 ± 3	37:63	~ 6
5000	815 ± 5	$\mathbf{42:58}$	~10
10000	$575\pm~7$	48:52	> 12

The relations are also summarized graphically in Figure 8. The lowering of liquidus temperatures with increasing H_2O pressure is common to all

Fig. 8. Projections of the liquidus for H_2O -saturated melts at 1, 2, 5, and 10 kb in the system An-Qz-H₂O to the join An-Qz. A = anorthite, C = cristobalite, L = liquid, T = tridymite. Data for the anhydrous join were taken SCHAIRER and BOWEN (1947). The silica polymorph stable on the liquidus at 1 kb is tridymite; the four-phase point on this projection is experimentally indistinguishable from the invariant point quartz-tridymite-anorthite-liquid-gas, and quartz is the stable phase of silica at temperatures below this four-phase point. At higher pressures, quartz is the stable silica phase at all temperatures studied.

feldspar-silica systems, as is the variation with rising H_2O pressure of the feldspar-silica ratio in the liquid toward lower feldspar-silica ratios while cristobalite and tridymite are stable on the liquidus, and toward higher feldspar-silica ratios when quartz is stable at liquidus temperatures. The effect is more pronounced in the An-Qz-H₂O system than in the other feldspar-silica systems because the temperatures in the anhydrous binary system are high and cristobalite and tridymite occur over a long temperature interval.

The temperature of the invariant point quartz + tridymite + anorthite + liquid + gas can be estimated from a P-T projection (Figure 7) of the four-phase univariant curve and the quartz-tridymite univariant curve (TUTTLE and BOWEN, 1958) to be 1040° C, at a pressure of 1 kb, indistinguishable from the values of the four-phase point tridymite + anorthite + liquid + gas.

PETROLOGIC APPLICATIONS

The boundary surface in the quinary system $\text{Or-Ab-An-Qz-H}_2\text{O}$ at constant pressure along which liquids crystallize to quartz and feldspar (or feldspars) is of great petrologic interest, for the composition of the liquid once on this surface cannot leave it under equilibrium conditions. It follows that the differentiation paths of many magmas proceed away from the primary phase (commonly a plagioclase) toward the boundary surface, and on reaching it, change course to stay on the surface. One example of an application of these relationships is given by BATEMAN and others (1963, p. D33-D38). A reverse course would be obtained on partial melting of pre-existing rocks.

The ratio of feldspar to quartz forming from the liquid when it reaches the boundary surface has been shown by TUTTLE and BOWEN (1958), SHAW (1963), and LUTH, JAHNS, and TUTTLE (1964), and this study to be sensitive to H_2O pressure. STEWART (1957) suggested this variation as the basis for a geologic barometer, and attempted an application to zones in granitic pegmatites. Another application to the problem of the depth of differentiation of rhyolitic magma was made by WHITE, THOMP-SON, and SANDBERG (1964, p. B38-B39).

Another application of these relationships concerns an explanation for the resorption of phenocrysts in lavas, most commonly quartz phenocrysts, but sometimes feldspar phenocrysts instead. Rarely are both kinds of phenocrysts attacked. This problem is not at all as enigmatic as described by FOSTER (1960).

The basic premise of the argument is that the phenocrysts of most lavas form by growth deep within a magma chamber, and not during or following extrusion. Once feldspar and a silica polymorph have begun to crystallize (the liquid has reached the boundary surface), any process that results in movement of the boundary surface necessitates a change in the composition of the liquid in equilibrium with the quartz and feldspar phenocrysts. The liquid changes composition by dissolving one of the phenocrysts. The phase resorbed during small deviations from equilibrium will be the phase toward which the boundary surface moves. The explanation sought thus involves consideration of what processes can move the boundary surface toward quartz. It is a characteristic of the feldspar-silica system that the boundary surface moves toward Qz as the H₂O pressure is increased from below one bar to a maximum of 1 kb. However, the stable silica phase crystallizing under such conditions is tridymite, rarely found as phenocrysts. At higher pressures where quartz is stable on the liquidus the boundary surface moves toward feldspar with increasing pressure. A sudden drastic reduction in the H₂O pressure such as might accompany the early stage of an eruption would normally result in a change in the position of the boundary surface toward Qz and in disequilibrium in the composition of the liquid. If the silica polymorph crystallizing with feldspar was tridymite, decrease in H₂O pressure would result in resorption of feldspar in all compositions except those on the feldspar side of the new four-phase point at the reduced pressure. In the usual geologic case where quartz is the stable silica polymorph, partial decrease in pressure would enlarge the feldspar plus liquid field and result in the resorption of quartz crystals in compositions except those on the silica side of the new four-phase point. When equilibrium was reestablished normal growth of all phases would continue, and the appearance of resorptions of inner zones followed by normal growth would be observed.

If the phenocrysts have always coexisted, the ratio of phenocrysts of feldspars to those of silica minerals can vary only from 2.0 to 0.5 under geologically reasonable variations of H_2O pressure (<10 kb). A ratio outside this range indicates that the more abundant mineral crystallized first. Judging from geologic evidence of the ratio of feldspar and silica phenocrysts, it is unusual when silica minerals crystallize first.

The system Ab (NaAlSi₃O₈)-An-Di (CaMgSi₂O₆) is commonly used for teaching purposes as an example of a ternary system with one solid solution. However, there are complications introduced by solid solution in the pyroxene, and a more useful example is the tetrahedron with Ab, An, Qz, and H₂O as apices as projected from the H₂O apex onto the base of silicate components. The boundary curve between the plagioclase and silica polymorph fields can be readily sketched from the data for An-Qz-H₂O given above and from data for Ab-Qz-H₂O given in TUTTLE and BOWEN (1958, p. 50—53). The boundary curve between these points can

be readily estimated by analogy to the anhydrous Ab-An-Qz system given by SCHAIRER (1957, Figure 35). The resulting diagram is applicable to many rocks, and its interpretation involves the same principles as are involved in the system Ab-An-Di.

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Table 1. Cell dimensions of anorthite

a, Å	b, Å	c, Å	α	β	γ	Volume Å ³	
$8.1768 \\ 0.0005$	$\begin{array}{c} 12.8768\\ 0.0005\end{array}$	$\begin{array}{c} 14.1690\\ 0.0005 \end{array}$	93° 10.0′ 1.0′	115° 50.8′ 1.0′	91° 13.3′ 1.0′	1338.84	Monte Somma, Italy. Cole, Sorum, Taylor (1951)
8.1615	12.8779	14.1670	93° 6.9′	$115^\circ~48'$	$91^\circ~10.2'$	1336.96	Salem, India. LAVES and GOLDSMITH (1955)
8.1752	12.8719	14.1707	$93^\circ14.8'$	115° 46'	91° 14.1′	1339.06	Salem, India heated 1 hour at 1425°C. LAVES and GOLDSMITH (1955)
8.17	12.87	14.14	$93^\circ16'$	$115^\circ48^\prime$	91° 16'	1334.6	"Synthetic". GOLDSMITH and LAVES (1955, p. 215)
8.18_{15} 0.0026	12.87 ₃₃ 0.0024	14.17 ₇₆ 0.0027	93° 12.4′ 1.1′	115° 50.1′ 1.4′	91° 8.2′ 1.2′	$\begin{array}{c} 1340.25\\ 0.39\end{array}$	Hydrothermal synthesis from An 90 Qz 10 glass, 1300°C, 2 kb, 1 hour, ANS-26, 32 lines, $\sigma =$ 0.0166° 2 θ
8.18 ₂₅ 0.0022	12.87 ₁₈ 0.0022	14.18 ₁₃ 0.0025	93° 7.3′ 1.2′	115° 53.3′ 1.1′	91° 15.0′ 1.1′	1340.02 0.33	From An 100 glass crys- tallized at 1200°C, 12 hours, then hydrother- mally at 1060°C, 10 kb, 2 hours, ANS-304, 33 lines, $\sigma = 0.0151^{\circ} 2 \theta$
8.17 ₃₇ 0.0024	12.8753 0.0025	14.1674 0.0027	93° 6.2′ 1.2′	115° 50.4′ 1.1′	91° 18.3′ 1.2′	1338.13 0.38	Hydrothermal synthesis from An 95 Qz 5 glass, 750° C, 10 kb, 3 hours, ANS-324, 28 lines, $\sigma =$ $0.0151^{\circ} 2 \theta$
8.17 ₉₃ 0.0019	12.86 ₈₇ 0.0020	14.17 ₃₉ 0.0021	93° 6.9′ 1.0′	115° 53.2′ 0.9′	91° 14.1′ 0.9′	1338.52 0.28	Hydrothermal synthesis from An 100 glass, 1060°C, 10 kb, 2 hours, ANS-305. See Table 2 for indexed pattern. 33 lines, $\sigma =$ 0.0130° 2 θ

Table 2. Indexed powder-diffraction pattern for primitive anorthite synthesized at 10 kb, 1060°C, 2 hours from CaAl₂Si₂O₈ glass in the presence of steam

Triclinic CaAl₂Si₂O.; P₁, $a = 8.179 \pm 0.002$ Å, $b = 12.869 \pm 0.002$, $c = 14.174 \pm 0.002$, $\alpha = 93^{\circ}$ 6.9 $\pm 1.0'$, $\beta = 115^{\circ}$ 53.2 $\pm 0.9'$, $\gamma = 91^{\circ}$ 14.1 $\pm 0.9'$, cell volume 1338.5 ± 0.3 Å³ as determined by least squares refinement of measured data italicized. No unique reflections with h+k odd and l even were observed, and all calculated reflections of this type have been omitted from the tabulated data.

hkl	Reflection	Calcu	ilated ²)	Measured ³)		
	type ¹)	d _{hkl} (Å)	2 θ Cu K α_1	$2 \theta K \alpha_1$	$\mathbf{I_0}/\mathbf{I_{100}}$	
001	е	12.720	6.943			
011	b	9.372	9.428			
011	Ъ	8.730	10.124			
$\overline{1}01$	b	8.081	10.939			
111	с	6.870	12.874	12.850	2	
111	\mathbf{c}	6.806	12.996	12.960	4	
11 0	a	6.521	13.567	13.542	10	
020	\mathbf{a}	6.417	13.789	13.770	5	
002	a	6.360	13.912			
110	a	6.244	14.173			
021	с	5.899	15.005	15.000	3	
112	\mathbf{a}	5.785	15.302			
112	a	5.660	15.643			
021	с	5.572	15.890	15.870	1	
101	b	5.417	16.349			
111	с	5.116	17.320	17.310	3	
121	b	5.051	17.544			
$\overline{121}$	b	5.000	17.724	17.700	1	
111	с	4.875	18.183			
$0\overline{2}2$	a	4.686	18.921	18.914	25	
103	b	4.663	19.017			
113	с	4.448	19.942			
022	a	4.365	20.327			
113	е	4.319	20.546			
$1\overline{2}1$	b	4.284	20.716			
003	\mathbf{c}	4.240	20.934	20.950	3	
031	b	4.144	21.422			
$0\overline{1}3$	b	4.114	21.584			
$\overline{202}$	a	4.040	21.980	21.282	50	
201	е	4.033	22.020			
121	b	4.008	22.158			
031	b	3.971	22.372			
013	Ъ	3.943	22.527			
$1\overline{1}2$	a	3.914	22.698	22.689	15	
$\overline{211}$	b	3.884	22.875			
$\overline{123}$	b	3.857	23.038			
$\overline{211}$	b	3.812	23.315			
$\overline{1}31$	С	3.797	23.408			

¹) GOLDSMITH and LAVES (1956, p. 397).

²) Calculated spacings are shown for $d \ge 2.500$ Å — calculated spacings less than 2.100 Å are listed only when they correspond to an indexed observed reflection.

³) Average of three observations with annealed CaF₂ as internal standard, a = 5.4622 Å at 25°C. Ni-filtered CuK α_1 radiation ($\lambda = 1.54050$ Å). Lower limit of 2 θ measured = 6° (14.72 Å). Pattern obtained at 25°C.

hkl	Reflection	Calculated ²)		Measure	ed 3)
	type ¹)	d _{<i>hkl</i>} (Å)	2θ Cu K α ₁	2 θ K α1	$\mathbf{I_0}/\mathbf{I_{100}}$
ī 30	a	3.781	23.510	23.512	45
131	е	3.765	23.613		
112	a	3.756	23.666	23.650	25
$\overline{1}23$	b	3.692	24.083		
$\overline{2}03$	с	3.691	24.089		
200	a	3.675	24.199		
$0\overline{2}3$	\mathbf{c}	3.659	24.307		
130	a	3.619	24.579	24.591	50
$\overline{13}2$	a,	3.601	24.702		
$\overline{213}$	b	3.558	25.006		
$\overline{2}13$	b	3.537	25.156		
132	a	3.511	25.346	25.290	5
$1\overline{3}1$	е	3.473	25.628		
$\overline{2}21$	с	3.466	25.677		
$\overline{1}\overline{1}4$	a	3.460	25.728	25.723	25
222	a	3.435	25.914	25.890	3
023	с	3.427	25.974		
$\overline{222}$	a	3.403	26.162	26.200	5
$\overline{221}$	с	3.365	26.463		
114	a	3.364	26.472	26.475	45
$\frac{1}{2}20$	a	3.260	27.330	27.336	70
131	с	3.252	27.398		
133	с	3.227	27.619		
223	с	3.215	27.724		
040	a	3.208	27.782		
$\frac{1}{2}04$	a	3.196	27.891	27.940	100
$\frac{-01}{223}$	с	3.184	27.996		
004	a	3.180	28.036		
201	с	3.178	28.051		
041	е	3.165	28.175		
$2\overline{1}1$	b	3.130	28.491		
103	b	3.127	28.523		
033	Ъ	3.124	28.547		
220	a	3.122	28.570	28.581	50
$1\overline{1}3$	с	3.091	28.862		
$\bar{1}33$	е	3.082	28.942		
041	е	3.060	29.159		
211	b	3.042	29.337		
$1\overline{3}2$	a	3.040	29.358	29.347	30
141	е	2.993	29.830		
113	С	2.988	29.881		
$\overline{2}31$	b	2.984	29.919		
141	b	2.971	30.048		
$0\bar{4}2$	a	2.950	30.273	30.267	50
$0\overline{2}4$	a	2.933	30.449	30.437	35
$2\overline{2}1$	С	2.920	30.589	30.550	10
033	Ъ	2.910	30.697		
$1\overline{2}3$	b	2.896	30.848		
$\overline{224}$	a	2.893	30.886	30.897	10
$\overline{231}$	b	2.888	30.941		
$1\overline{4}1$	b	2.846	31.407		
$\overline{2}24$	a	2.830	31.587)	31 616	35
132	a	2.828	31.614 ∫	01.010	00
$\overline{1}05$	b	2.816	31.751		
233	b	2.810	31.817		
$\overline{134}$	a	2.803	31.904	31.910	10
042	a	2.786	32.097		

Four-Phase Curve in the System ${\rm CaAl_2Si_2O_8-SiO_2-H_2O}$

hkl	Reflection	Cale	ulated ²)	Measu	red ³)
	type ¹)	\mathbf{d}_{hkl} (Å)	$2 \theta Cu K \alpha_1$	$2 \theta \mathbf{K} \alpha_1$	I_0/I_{100}
$\overline{115}$	е	2.785	32 111		
221	ē	2.781	32.159		
$\overline{2}33$	b	2.779	32.177		
024	a	2.772	32.264		
123	b	2.733	32.744		
$\overline{2}05$	е	2.724	32.850	32.850	2
$\overline{1}15$	е	2.717	32.939	01.000	-
202	a	2.708	33.043		
$\overline{1}\overline{4}3$	b	2.702	33,130		
303	Ъ	2.694	33.231		
215	\mathbf{b}	2.686	33.332		
141	b	2.682	33.374		
312	a	2.676	33.459	33.460	2
134	a	2.656	33.715	33.722	20
043	b	2.650	33.790		
312	a	2.648	33.820		
$\frac{215}{215}$	b	2.644	33.873		
$\frac{313}{313}$	с	2.642	33.904		
$\frac{125}{212}$	b	2.636	33.974		
$\frac{313}{2}$	e	2.631	34.050		
231	b	2.629	34.067		
301	d	2.629	34.071	64 66	
100	C b	2.618	34.222	34.20	3
011 149	U b	2.096	34.521		
113	u a	2.988	34.028 25.052)		
311	a	2.556	25.078	35.079	5
	c	2.550	35.070)		
0.51	b b	2.552	35 145		
005	c	2.544	35 949		
$\overline{2}\overline{2}5$	c	2.543	35 264		
$\overline{015}$	b	2.530	35.453		
$1\overline{1}4$	a	2.528	35.478		
$\overline{2}42$	a	2.525	35.517	35.509	25
$\overline{1}25$	b	2.524	35.536		
314	a	2.510	35.744		
$\overline{3}14$	a	2.506	35.806		
$\overline{24}2$	a	2.500	35.890	35.902	30
222	a	2.437	36.846	36.850	5
310	a	2.429	36.977	36.950	3
152	a	2.405	37.363	37.360	3
321	\mathbf{b}	2.399	37.441	37.450	2
150	a	2.386	37.667	37 679	9
$\frac{310}{10}$	a	2.384	37.695 J	01.010	0
152	a	2.360	38.105	38.095	4
$\frac{240}{200}$	a	2.359	38.121	00.000	т
332	a	2.323	38.725	38.725	5
244	a	2.296	39.204	39.210	3
110	a	2.265	39.765	39.755	10
102 911	a	2.237	40.274	40.296	5
<u>444</u> 291	it.	2.234	40.339	40.340	3
	iti Q	2.191 9 100	41.173	40.150	2
240	a	4.100 9.149	41.792 19 151)	41.821	3
060	a 9.	2.142	42.101	42.153	20
006	8	2.139	44.410 J 49.611	19 850	9
152	a	2.095	43 141	42.000	о 15
	7.		101111	20.200	10

53

N.

hkl	Reflection	Calc	ulated ²)	Measured ³)		
	type ¹)	d _{hkl} (Å)	$2 \theta Cu K \alpha_1$	$2 \theta \mathbf{K} \alpha_1$	I_0/I_{100}	
$\frac{1}{4}04$	a	2.020	44.825	44.870	5	
$\begin{array}{c} 402 \\ 062 \end{array}$	a a	$2.017 \\ 1.985$	44.910 J 45.659	45.610	3	
224	a	1.878	48.426	48.419	5	
$\overline{\overline{2}}$ 46 $\overline{4}$ 06	a a	1.846 1.846	49.321 49.334	49.356	5	
064	a	1.836	49.608	49.596	15	
260	a	1.809	50.391	50.401	5	
116	a	1.797	50.761	50.772	10	
$\overline{2}08$	a	1.769	51.621	51.623	15	

Table 3. Experiments in the system An-Qz- H_2O used to define the liquidus and solidus at several pressures, and H_2O content of quenched glasses

Run	Tem-				D. 1 (1) activated and the property
number	perature	Time	% H ₂ O in	An: Qz	Products ¹), estimated weight percent
ANS-	°C	hours	capsuie		
			Ea	cperiments a	at 1 kb H_2O pressure
404	1040	2	23.8	40:60*	A = 30, (C) = 40, G = 30 (1.494)
400	1025	3	16.7	40:60*	A = 40, (C) = 10, Q = 50
405	1040	2	20.9	35:65	A = 20, (C) = 5, G = 75 (1.494)
401	1025	3	17.9	35:65	A = 25, (C) = 35, (G) = 35 (1.494), Q = 5
406	1040	2	34.9	30:70	(C) = 12, G = 85 (1.492), Q = 3
402	1025	3	16.9	30:70	A = 20, (C) = 45, (G) = 30 (1.494), Q = 5
407	1040	2	44.4	20:80*	A = 19, (G) = 1, Q = 70, T = 10
403	1025	3	24.0	20:80*	A = 20, Q = 80
			E_{2}	cperiments e	u 2 ko H ₂ O pressure
HSY	1350	1	?	100:0*	(A), (B), G
HSY	1320	1	ş	100:0*	A = 100, (B)
53	1330	1	15.7	95:5	$(A) = 1; G = 99 (1.534), 5.9 \% H_2O$
62	920	36	16.8	95:5*	A = 95, Q = 5
26	1300	1	16.8	90:1 0	A = 90, G = 10 (1.531)
27	1300	1	15.1	80:20	(A) = 1, G = 99 (1.540)
21	1250	1	13.0	80:20	A = 85, G = 15 (1.513)
22	1250	1	10.9	70:30	(A) =1; G =99 (1.524), 5.1 % H_2O
30	1200	1	14.8	65:30	A = 20, G = 80 (1.524)
67	1320	0.5	18.1	60:40	$G = 100 (1.520), 4.8 \% H_2O$
23	1250	1	12.2	60:40	$G = 100 (1.523), 4.4 \% H_2O$
31	1200	1	17.4	60:40	G = 100 (1.517)
32	1200	1	14.0	51:49	$(A) = <1; G = 100 (1.510), 6.2 \% H_2O$
28	1300	1	15.3	50:50	$G = 100 (1.509), 5.5 \% H_2O$
15	1100	4	19.8	50:50	A = 20, G = 80 (1.501)
46	1100	6	19.1	45:55	$(A) = <1; G = 100 (1.504), 5.1 \% H_2O$
43	1050	7	16.9	45:55	$A = 1; G = 99 (1.503), 6.1 \% H_2O$
34	975	22	17.2	45:55	$A = 5$; $G = 95$ (1.494), 5.7 % H_2O
38	915	45	14.6	45:55	$\mathbf{A}\!=\!45,\ \mathbf{Q}\!=\!55$
63	920	36	15.3	45:55*	A = 45, (G) = <1, Q = 55
47	1100	6	18.7	40:60*	$G = 100 (1.499), 4.8 \% H_2O$
44	1050	7	23.1	40:60*	(A) = < 1 ; G = 100 (1.499), 5.3 % H ₂ O
60	1000	8	18.5	40:60	$A = 10; G = 90 (1.499), 5.9 \% H_2O$
35	975	22	19.3	40:60*	$A = 10; G = 90 (1.494), 6.2\% H_2O$
11	925	2	45.3	40:60	A = 25, G = 70, (Q) = 5

Run	Tem-								
number	perature	Time	% H ₂ O in	An : Qz	Products ¹), estimated weight percent				
ANS-	°C	hours	capsule	2					
	Experiments at 2 kb H_2O pressure — continued								
39	915	45	17.4	40:60*	A = 40, Q = 60				
1	1150	1	18.0	35:65	$G = 100 (1.494), 6.0 \% H_2O$				
61	1000	8	18.5	35:65	$G = 100 (1.494), 5.3 \% H_2O; (Q) = <1$				
36	975	22	17.9	35:65	$(A) = <1; G = 100 (1.493), 5.9\% H_2O; (Q) = <1$				
10	925	2	22.1	35:65	G = 80 (1.495), $Q = 20$				
64	920	36	19.2	35:65*	A = 35, (G) = <1, Q = 65				
66	1320	0.5	19.9	30:70	$G = 100 (1.492), 4.7 \% H_2O$				
2	1100	1	16.0	30:70	$G = 100 (1.493), 6.5 \% H_2O$				
3	1050	1	19.0	30:70	$G = 100 (1.493), 5.5 \% H_2O; (Q) = < 1$				
19	1000	6	24.1	30:70	$G = 95$ (1.493), 6.2 % H_2O ; $Q = 5$				
8	925	2	16.9	30:70*	(A) = 25, G = 10 (1.495), Q = 65				
9	925	2	25.4	30:70	(A) = <1, G = 70 (1.495), Q = 30				
41	915	45	18.3	30:70	A = 30, Q = 70 ,				
29	1300	1	22.4	20:80*	$G = 100 (1.481), 4.3 \% H_2O$				
45	1050	7	28.1	20:80*	$G = 95$ (1.485), 4.9 % H_2O ; $Q = 5$				
70	1125	5	20.9	10:90*	$G = 100 (1.486), 3.6 \% H_2O$				
48	1100	6	17.6	10:90*	$G = 85 (1.475), 4.3 \% H_2O; Q = 15$				
71	1125	5	27.9	5:95*	$G = 100$ (1.468), 4.1 % H_2O				
79	1115	4	22.8	5:95*	$G = 90$ (1.473), 4.0 % H_2O ; $Q = 10$				
65	920	36	24.0	5:95*	A = 5, Q = 95				
54	1330	1	20.6	0:100	$G = 100 (1.465), 3.6 \% H_2O$				
57	1135	7	19.6	0:100	$G = 100 (1.472), 4.5 \% H_2O$				
58	1135	7	14.4	0:100*	G = 45 (1.468), (Q) = 55				
75	1130	3	17.5	0:100	$G = 96$ (1.466), 3.5 % H_2O ; $Q = 4$				
72	1125	5	18.4	0:100	G = 5 (1.468), $Q = 95$				
73	1125	5	14.2	0:100*	G = 8 (1.468), $Q = 92$				
81	1115	4	28.0	0:100	$\mathbf{Q} = 100$				
			Ĺ	Experiments	at 5 kb pressure				
145	810	41	16.3	95:5	A = 95, Q = 5				
131	1190	1	16.3	80:20	A = 25, G = 75 (1.535)				
132	1190	1	14.9	70:30	(A) < 1; G = 99 (1.529), 7.7 % H ₂ O				
134	1140	2	14.3	70:30	$A = 2; G = 98 (1.527), 8.6 \% H_{\circ}O$				
135	1140	2	16.8	65:35	$G = 100 (1.523), 8.4 \% H_{2}O$				
136	1140	2	16.5	60:40	$(A) < 1; G = 100 (1.518), 9.8 \% H_{\circ}O$				
118	1050	4	19.6	60:40	$A = 5, G = 95 (1.519), 8.9 \% H_2O$				
117	1050	4	15.5	55:45	$G = 100 (1.515), 9.5 \% H_2O$				
160	1010	4	14.6	55:45	$A = 1; G = 99 (1.514), 9.9 \% H_2O, S.G. = 2.324 \pm 4$				
146	1000	5	15.2	55:45	$A = 5; G = 95 (1.514), 8.2 \% H_2O$				
147	1000	5	14.7	50:50	$G = 100 (1.511), 8.8 \% H_2O$				
128	950	5	18.8	50:50	$A = 5; G = 95 (1.508), 9.2 \% H_2O$				
124	900	17	15.7	50:50	$A = 10; G = 90 (1.504), 10.0 \% H_2O$				
161	1010	4	15.0	45:55	$G = 100$ (1.506), 8.8 % H_2O , S.G. = 2.294 ± 4				
166	950	4	16.2	45:55	$G = 100$ (1.504), 9.6 % H_2O , S.G. = 2.286				
141	920	6	17.1	45:55	(A) = 1; G = 99 (1.507), 10.0 % H_2O				
159	900	14	13.8	45:55	$A = 2; G = 98 (1.504), 8.9 \% H_2O$				
138	850	17	12.9	45:55	$A = 5$, $(A) = 2$; $G = 93$ (1.504), 9.5 % H_2O				
104	820	25	17.0	45:55	A = 45, G = 5, Q = 50				
144	810	41	16.5	45:55*	A = 45, Q = 55				
123	900	17	22.2	40:60	$G = 100$ (1.501), 10.2 % H_2O				
156	875	5	16.2	40:60	(A) =1; G =99 (1.503), S.G. = 2.285 ± 4 ; (Q) < 1				
139	850	17	17.4	40:60	$G = 99 (1.502), 9.3 \% H_2O; (Q) = 1$				
164	835	16	16.4	40:60	$G = 90$ (1.503), 9.8 % H_2O ; $Q = 10$				
101	820	25	18.0	40:60	(A) = 2, G = 60 (1.499), Q = 38				

Run	\mathbf{Tem} -				
number	perature	Time	% H ₂ O in	$\mathbf{An}: \mathbf{Qz}$	Products ¹), estimated weight percent
ANS-	°C	hours	capsule		
			Experi	ments at 5 kb	pressure — continued
10. 1259.11		1000			
143	810	41	21.1	40:60*	A = 40, Q = 60
167	950	4	16.1	35:65	$G = 100 (1.499), 10.6 \% H_2O; (Q) < 1$
151	920	8	17.5	35:65	$G = 99$ (1.497), 9.2 % H_2O ; $Q = 1$
158	900	14	17.0	35:65	$G = 95 (1.498), 9.2 \% H_2O; Q = 5$
162	1010	4	15.2	30:70	$G = 100 (1.495), 9.9 \% H_2O, S.G. = 2.250$
168	950	4	16.4	30:70	$G = 99$ (1.494), 8.5 % H_2O , S.G. = 2.23; $Q = 1$
150	920	8	11.9	30:70	$G = 90 (1.493), 7.43 \% H_2O; Q = 10$
163	1010	4	17.0	20:80*	G = 97 (1.486), 7.3 % H ₂ O; (Q) = 3
148	1000	5	18.6	20:80*	G = 93 (1.485), 7.8% H ₀ O: $O = 7$
109	1070	4	18.5	10:90*	G = 100 (1.473) with liquid inclusions, 10.9% H _• O
116	1055	4	18.3	10.00*	G = 100 (1.475) 11.0% H ₂ O (bubbles in glass)
105	1040	т 5	18.1	10.00*	$G = 98 (1.477) 11.9\% H_O (hubbles): O = 9$
100	1040	3	10.1	5.05*	G = 100 (1.417), 11.0 / 0 1120 (0000103), Q = 2 G = 100 (1.472), 0.50 / U O
110	1070	4	20.0	0:90*	$G = 100 (1.473), 9.5\% H_2 U$
115	1055	4	22.7	5:95*	G = 10 (1.413), $Q = 30$
142	810	41	26.0	5:95*	A = 5, Q = 95
111	1070	5	19.6	0:100	$G = 100 (1.468), 11.2 \% H_2O (bubbles)$
112	1070	5	12.9	0:100*	G = 75 (1.468), (Q) = 25
113	1055	4	20.1	0:100	$\mathbf{Q} = 100$
114	1055	4	16.1	0:100*	$\mathbf{Q} = 100$
			Ex	periments at .	10 kb H ₂ O pressure
000	1100	-	10.0	100 0	
333	1120	1	16.8	100:0	(A) = 25, (B) = 25, G = 50 (1.553)
332	1120	1	26.1	100:0*	(A) = 40, (B) = 10, G = 50 (1.551)
312	1100	2	22.3	100:0*	A = 95, (B) = 5, G < 1 (1.544)
305	1060	2	19.3	100:0	$\mathbf{A} = 100$
304	1060	2	17.9	100:0*	A = 99, (B) = 1
334	1120	1	16.9	95:5	(A) = 4, (B) = 11, G = 85 (1.548)
314	1100	2	18.7	95:5	A = 40, (B) = 5, G = 55 (1.545)
324	750	3	15.0	95:5*	A = 95, Q = 5
315	1100	2	12.5	90:10	(A) = 2, (B) = 3, G = 95 (1.541)
307	1060	2	19.3	90:10	A = 50, (B) = 1, G = 49 (1.521)
364	1050	2	14.0	80:20	A = 23, (B) = 2, G = 75 (1.530)
343	1040	2	17.8	70:30	$G = 100 (1.520), 12.5 \% H_{\circ}O, S.G. = 2.324$
362	940	- 2	17.0	60:40	$G = 100 (1.510), 11.7 \% H_{\bullet}O, S, G, = 2.280$
355	920	2	16.7	60 - 10	A = 3: G = 97 (1.516) 12.6% H ₂ O, S.G. = 2.26
254	020	2	16.0	55 + 45	G = 100 (1.513) 11.9% H ₂ O, S.G. = 2.26
220	920		10.0	55 • 45	$A = 1: C = 00 (1.512), 11.5 /_0 H_20, 0.0.5 = 2.20$
222	900	3	19.4	55.45	$A = 1, 0 = 33 (1.313), 12.0 \ /_0 = 120, 5.0. = 2.203 \pm 3$
359	850	Z	12.1	51:49	$G = 100 (1.504), 12.1 \% H_20, S.G. = 2.252$
361	940	z	20.4	50:50	$G = 100 (1.505), 11.2 \% H_2O, S.G. = 2.247$
358	850	2	19.6	50:50	$G = 100 (1.503), 12.0 \% H_2O, S.G. = 2.245$
318	800	3	29.7	50:50	$A = 2; G = 98 (1.506), 11.9 \% H_2O, S.G. = 2.26 \pm 2$
345	780	3	17.8	50:50	$A = 3; G = 97 (1.504), 10.0 \% H_2O$
348	765	4	17.9	50:50	$A = 4$; $G = 96$ (1.500), 12.8 % H_2O , S.G. = 2.250
367	1050	2	18.0	45:55	$G = 100 (1.503), 11.3 \% H_2O, S.G. = 2.224$
328	900	3	25.3	45:55	$G = 100$ (1.504), 12.5 % H_2O , $S.G. = 2.207 \pm 2$
357	850	2	18.1	45:55	$G = 100 (1.499), 11.2 \% H_2O, S.G. = 2.232$
319	800	3	14.9	45:55	$G = 98 (1.506), 11.8 \% H_2O, S.G. = 2.265 \pm 7; (Q) = 2$
349	765	4	26.4	45:55	(A) < 1; G = 97 (1.498), 11.8 % H ₂ O; Q = 3
350	765	4	18.2	45:55*	$G = 93 (1.499), 10.4 \% H_aO: Q = 7$
329	900	3	17.8	40:60	$G = 100 (1.500), 12.3 \% H_{\bullet}O, S, G_{\bullet} = 2.200 + 4$
356	850	2	18.0	40:60	$G = 100 (1.498), 11.6 \% H_{\bullet}O, S, G = 2.21 + 1$
351	765	- A	21 4	40 : 60	G = 90 (1.498), Q = 10
395	750	т 2	17.6	40.60*	$A = 40 \Omega = 60$
040 959	100	0 0	16 1	25 - 65*	G = 100 (1.495) 10.80/ II O S C = 9.96
000	940	4	10.1	00.00.	$G = 100 (1.433), 10.0 % H_2O, S.G. = 2.20$

Run	Tem-								
number	perature	Time	% H ₂ O in	An: Qz	Products ¹), estimated weight percent				
ANS.	$^{\circ}\mathrm{C}$	hours	capsule						
Experiments at 10 kb H_2O pressure — continued									
330	900	3	19.8	35:65	$G = 99$ (1.497), 12.2 % H_2O , S.G. = 2.21 ±1; $Q = 1$				
360	940	2	18.6	30:70*	$G = 100 (1.486), 12.0 \% H_2O$				
352	920	2	18.6	30:70*	$G = 97$ (1.490), 11.2 % H_2O , S.G. = 2.22; $Q = 3$				
331	900	3	22.9	30:70	$G = 95$ (1.490), 11.8 % H_2O ; $Q = 5$				
323	1032	2	21.0	10:90*	G = 95 (1.475), $Q = 5$				
326	750	3	20.1	5:95*	A = 5, Q = 95				
371	1060	2	39.5	0:100	G = 100 (1.479)				
376	1060	2	23.1	0:100*	G = 100				
365	1050	2	20.6	0:100	G = 15 (1.475), Q = 85				
366	1050	2	14.9	0:100*	G = 10, Q = 90				
380	1040	2	35.0	0:100*	$\mathbf{Q=100}$				

¹) A = anorthite, B = " β -alumina", C = cristobalite, G = glass (index of refraction in parentheses), Q = quartz, T = tridymite. Abbreviations in parentheses indicate quench or metastable phase.

* Crystalline starting materials used.

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