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Phase equilibria in the silica-undersaturated part of the KAISiO₄ – Mg_2SiO_4 – Ca_2SiO_4 – SiO_2 – F system at 1 atm and the larnite-normative trend of melt evolution

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Abstract The evolution of nephelinitic melts in equilibrium with mica-bearing liquidus assemblages and melting relations have been studied on two silica-undersaturated joins of the KAlSiO₄ – Mg₂SiO₄ – Ca₂SiO₄ - SiO₂ - F system at atmospheric pressure by quench runs in sealed platinum capsules. Fluorine has been added to the batch compositions by the direct exchange of fluorine for oxygen $(2F^{-} = O^{2^{-}})$. The first join is the pseudo-ternary Forsterite - Diopside - KAlSiO₃F₂ system. Forsterite, diopside, F-phlogopite and leucite crystallisation fields and a fluoride-silicate liquid immiscibility solvus are present on the liquidus surface of the join. Sub-liquidus and sub-solidus phases include akermanite, cuspidine, spinel, fluorite and some other minor fluorine phases. The second system is the pseudobinary Akermanite – F-phlogopite join that intersects the Forsterite - Diopside - KAlSiO₃F₂ join. Akermanite, forsterite, diopside, F-phlogopite, leucite and cuspidine are found to crystallise on the join. Forsterite (fo) and leucite (lc) are related to F-phlogopite (phl) by a reaction with the fluorine-bearing liquid: fo + lc + 1 = phl, and the reaction proceeds until forsterite or leucite are completely consumed. The reaction temperature and resulting phase association depend on batch composition. Thus, leucite is not stable in the sub-solidus of the Akermanite - F-phlogopite join, but is preserved in a part of the Forsterite - Diopside - KAl- SiO_3F_2 system where forsterite reacts out, or does not

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crystallise at all. The phlogopite-in reaction has an important effect on the composition of the coexisting liquid. The liquids initially saturated in forsterite evolve to extremely Ca rich, larnite-normative residuals. The experimental data show that larnite-normative melilitolites can crystallise from evolved melilititic melts generated from "normal" melanephelinitic parental magmas *with no normative larnite*. The evolution towards melilitites requires fractionation of phlogopitebearing assemblages under volatile pressure.

Introduction

The experimental study is designed to model the effect of volatiles on the crystal fractionation path in ultramafic alkaline magmatic systems at low pressure. Of special interest is the link between the stability of liquidus phlogopite and the development of highly larnite normative melilitite melts. We have focused on phlogopite because it is the most abundant hydrous liquidus phase in ultramafic alkaline rocks. Phlogopite is common not only in potassic rocks, but also in plutonic complexes of sodium-rich ultramafic alkaline rocks and related carbonatites. In both K- and Na-rich systems it is often found in close association with melilite in Ca-rich melilititic rocks. In a study of the petrography and bulk chemical composition of alkaline dyke swarms in the Gardiner complex, Nielsen (1994) suggested that highly larnite normative melilitolites may form from late-stage residual melts by phlogopite, amphibole, clinopyroxene and Fe-Ti oxide fractionation from a melanephelinite parental melt under closed, volatile-rich conditions in shallow-level magma chambers.

Phlogopite crystallisation has previously been studied mainly in the Ca-free KAlSiO₄ – Mg_2SiO_4 – SiO_2 – $H_2O(F)$ system (Luth 1967; Modreski and Boettcher 1973; Wendlandt and Eggler 1980 a, b; Foley et al. 1986; Gupta and Green 1988; Melzer and Foley 1996). Melilite liquidus equilibria have mostly been studied at atmospheric pressure in dry systems, such as NCMAS (Pan

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and Longhi 1989, 1990; Soulard et al. 1992), from which phlogopite is absent. High-pressure melting experiments with phlogopite-bearing potassic rocks were reviewed by Edgar and Vukadinovic (1992) and Foley (1992). In most cases, however, compositions of glasses in run products were not analysed due to problems with quench phlogopite. The aim of this study is accordingly to obtain experimental data on phlogopite-melilite liquidus relationships at crustal pressures, which have not previously been documented experimentally and to investigate the suggested evolution towards larnite-normative residual melt composition in phlogopite-bearing systems at low pressure. For this purpose we have studied several joins in the silica-undersaturated part of the $KAlSiO_4 - Mg_2SiO_4 - Ca_2SiO_4 - SiO_2$ system, known as the kalsilite-based normative tetrahedron (Yoder 1986) with the addition of fluorine.

The kalsilite-based normative tetrahedron (Fig. 1) has long proved to be a useful model for the study and the graphic presentation of phase equilibria in potassium-rich igneous rocks. The system, however, has

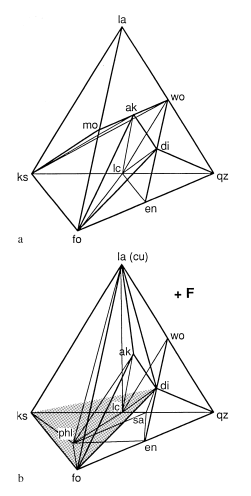


Fig. 1a, b Kalsilite-based normative tetrahedron. **a** volatile-free lowpressure conditions (Yoder 1986); **b** low-pressure fluorine-doped system at the conditions of phlogopite and cuspidine stability. *Shaded triangle* shows the Mg₂SiO₄ – CaMgSi₂O₆ – KAlSiO₃F₂ pseudoternary join. See the Appendix for abbreviations

broader application. The addition of water to the components of the tetrahedron at pressures > 0.7 kbar allows the study of the phlogopite-bearing phase assemblages. Phlogopite is a key phase in many petrological models, as it is the main reservoir for potassium, H₂O and a number of trace elements in the lithospheric mantle. Thus, studies of partial melts of phlogopitebearing mantle assemblages at elevated pressures are a necessity for understanding the origin of a broad spectrum of mantle-derived magmas. On the other hand, the crystallisation of liquidus phlogopite in shallow-level magma chambers at lower pressure may, as suggested by Nielsen (1994), have fundamental effect upon the crystal fractionation path of magmatic systems and result in the diversity of plutonic and volcanic rocks of ultramafic alkaline complexes.

Liquidus phlogopite can be stabilised in the kalsilitebased normative tetrahedron in two ways: (1) by applying water pressure >0.7 kbar; or (2) by adding fluorine by substitution of fluorine for oxygen $(2F^- = O^{2^-})$. We have chosen the latter method because fluorine-doped runs may be performed at 1 atm in sealed platinum capsules and such runs provide: (1) higher quenching rates; (2) the possibility to avoid quench phlogopite; (3) the formation of homogeneous glasses suitable for microprobe analyses. A further important point is that F-phlogopite thermal stability exceeds that of the OH-analogue by about 200 °C (Shell and Ivey 1969; Munoz 1984). The F-doped system thus enables the study of the extreme case of maximum phlogopite stability and its maximum effect on melt evolution.

Previous studies

Anhydrous phase equilibria in the kalsilite-based normative basalt tetrahedron at atmospheric pressure have been studied in detail for several decades (e.g. Schairer and Bowen 1938; Luth 1967; Gupta 1972). Experimental results on phase equilibria in the tetrahedron at anhydrous conditions and under H_2O pressure are reviewed by Yoder (1986) who showed that phlogopite takes part in a number of heteromorphic reactions and has an important effect on the topology of subtetrahedra within the kalsilite-based normative tetrahedron (Fig. 1).

Experimental studies at high pressure concentrated on the kalsilite-forsterite-quartz (KAlSiO₄ – Mg₂SiO₄ – SiO₂) border join. Phase relationships for this join have been investigated at different *P-T* conditions and in the presence of several volatile components: H₂O, CO₂, F (Luth 1967; Modreski and Boettcher 1973; Wendlandt and Eggler 1980a, b; Wendlandt 1984; Foley et al. 1986; Gupta and Green 1988). Much attention has been paid to the position of the fo + 1 = en + phl reaction point, where liquid is in equilibrium with the phase assemblage similar to that of the phlogopite-bearing upper mantle. Effects of fluorine on the phase equilibria in the kalsilite-forsterite-quartz system have been studied by Foley et al. (1986) and Melzer and Foley (1996). These investigations showed that the phlogopite liquidus field greatly expands with increasing fluorine content in the system.

A few high-pressure experiments have been carried out with Ca-bearing compositions. These include 1:1 (weight) diopside-phlogopite mixture (Kushiro 1970; Modreski and Boettcher 1973) in which the clinopyroxene-phlogopite association has been shown to be stable at pressures up to at least 32 kbar.

Experimental and analytical methods

The starting materials for the runs are mixtures of synthetic forsterite, diopside, akermanite, potassium aluminosilicate glass and chemical-grade AlF₃ (for the Mg₂SiO₄ – CaMgSi₂O₆ – KAlSiO₃F₂ join), or MgF₂ (for the Ca₂MgSi₂O₇ – KMg₃AlSi₃O₁₀F₂ join). Synthetic minerals and glasses were prepared from pure oxides and carbonates (SiO₂, Al₂O₃, MgO, CaCO₃, K₂CO₃) by mixing of appropriate proportions in a mortar under acetone and sintering in air at 950–1000 °C. Sintered mixtures were then crushed and heated again to temperatures between 1100 to 1400 °C, depending on the compound. Ready to use charges were kept dry in a desiccator. However, peralkaline glasses and some fluoride compounds used in reactant mixtures are hygroscopic and presence of some water in our charges can not be ruled out completely.

Quench experiments were carried out at 1 atm in a vertical electric furnace with a heater made of platinum wire. About 20 mg of fine powder were sealed in platinum capsules. The capsules were weighted before and after the runs to check for seal failure and fluorine loss. The temperature was controlled by Pt/PtRh30 thermocouple calibrated against the melting point of gold. The accuracy of temperature measurements is estimated to be ± 5 °C.

Run durations varied from 5 to 75 hours depending on the temperature and composition of the mixtures (see Table 1). We used two different heating procedures to check for equilibrium. In the majority of the runs starting mixtures were overheated up to about 100 °C above the run temperature and then slowly cooled to the run conditions. In high-temperature runs (>1200 °C) and in a few control runs at lower temperatures the run conditions were directly attained without overheating. Control runs show that the phase compositions of the run products are independent of the heating procedure. Nevertheless, runs with overheating usually produce larger crystals (above 10 microns) that are better suited for optical identification and electron microprobe studies. The samples were quenched in cold water. Mostly, quench rates were sufficiently high to avoid quench phlogopite and to produce homogeneous transparent glass.

Run products were studied optically in index oil. Crystal phases in some samples were determined by X-ray diffraction. Chemical compositions of crystal phases and glasses in selected samples were analysed by JEOL 7 Superprobe electron microprobe, Department of Geology, University of Copenhagen as described by Nielsen et al. (1997). A defocused beam (beam size 15–20 µm) was used for analyses of glasses to reduce K and F-loss. Crystallised globules of immiscible fluoride liquid were analysed by scanning method.

Experimental results

Four F-doped joins in the kalsilite tetrahedron were studied and include: the Forsterite – $KAlSiO_3F_2$ join, the Diopside – $KAlSiO_3F_2$ join, the Forsterite – Diopside – $KAlSiO_3F_2$ join and the Akermanite – F-Phlogopite join. Run conditions and phase composition of the run products are listed in Table 1. Liquidus surface

projection for the Mg₂SiO₄ – CaMgSi₂O₆ – KAlSiO₃F₂ join is presented in Fig. 2. The T-x phase diagrams for the $Ca_2MgSi_2O_7$ – $KMg_3AlSi_3O_{10}F_2$ and the CaMgSi₂O₆ - KAlSiO₃F₂ joins are shown in Figs. 3 and 4. Nine crystalline phases were identified in the run products: forsterite, diopside, leucite, akermanite, F-phlogopite, spinel, cuspidine, fluorite, and F-chondrodite (?). Some aluminofluorides (presumably K_3AlF_6) may be also present in F-rich compositions. Besides the crystalline phases, immiscible fluoride liquid is observed in the F-rich part of the Mg_2SiO_4 – $CaMgSi_2O_6$ – KAlSiO₃F₂ join (see Figs. 2 and 4). A vapour phase, though omitted in Table 1, may occur. Tiny $(5-10 \ \mu m)$ gas-filled vesicles are seen in some run products. They may suggest the presence of a free vapour phase during the runs, or a vapour phase formed during the quenching. In all cases, however, there is no indication (within the error of EMP analyses) of significant F-loss to a free vapour phase.

Crystalline phases

Forsterite

Forsterite forms small $(5-10 \ \mu\text{m})$ euhedral prismatic crystals and rounded oval grains. In the index oil it can in general be distinguished from diopside by smaller grain size and with confidence by parallel extinction. Analyses show that forsterite has nearly constant stoichiometric composition close to the ideal formula. The CaO content is 0.27–2.29 wt%.

Diopside

Clinopyroxene which, according to EMP data (Table 2), is close to pure diopside, usually occurs in the run products as large (about 40–50 μ m) elongated prismatic crystals. Representative EMP analyses in Table 2 show that Al substitutes for some Si in tetrahedral sites of clinopyroxene structure. This substitution is charge-balanced mainly by Al³⁺-Mg²⁺ exchange in M1 octahedral sites.

Leucite

High-temperature liquidus leucite occurs as medium-size $(20-30 \ \mu\text{m})$ perfect tetragontrioctahedral crystals. At lower temperatures it forms irregular grains. Oval glass inclusions are common in leucite. Under crossed nicols leucite is usually isotropic, but in some crystals complex twinning structures can be observed. The composition of leucite is stable and very close to the ideal formula. The CaO content does not exceed 0.22 wt%. Magnesium is also present in small amounts (0.07–0.23 wt% MgO) and appears to substitute for aluminium and silicon in tetrahedral sites.

2	5	n
2	J	υ

Table 1 Run conditions and run products. Abbreviations for components and phases are listed in the Appendix

Sample	Starting composition, wt%	Duration, h	T °C	Phase composition of run products
1	2	3	4	5
Forsterite – KAlSiC	D ₂ F ₂ ioin			
FKF-63	Fo80:KF20	5	1350	1 + fo
FKF-74		7.5	1300	1 + fo + phl
FKF-48		6.5	1250	1 + fo + phl
FKF-17		55	1200	1 + fo + phl + sp
FKF-1		75	1150	fo + phl fine-grain aggregate
FKF-32		60	1100	fo + phl fine-grain aggregate
FKF-64	Fo60:KF40	5	1350	1 + fo + phl
FKF-75		7.5	1300	1 + fo + phl
FKF-49 FKF-18		6.5 55	1250 1200	l + fo + phl + sp l + fo + phl + sp
FKF-2		75	1150	fo + phl fine-grain aggregate
FKF-33		60	1100	fo $+$ phl $+$ hu? fine-grain
FKF-65	Fo40:KF60	5	1350	1 + fo + phl
FKF-76		7.5	1300	1 + fo + phl
FKF-50		6.5	1250	1 + fo + phl
FKF-19		55	1200	1 + fo + phl
FKF-3		75	1150	1 + fo + phl
FKF-34		60	1100	1 + fo + phl
FKF-85	Fo30:KF70	7	1216	1 + phl
FKF-110		7	1130	1 + phl
FKF-77	Fo20:KF80	7.5	1300	1
FKF-51		6.5	1250	1 + phl
FKF-20		55	1200	1 + phl
FKF-35		60	1100	l + phl
FKF-95	E-00-KE10	18	1050	l + phl
FKF-86 FKF-96	Fo90:KF10	7 18	1216 1050	l l + phl
FKF-120		53	993	1 + phi 1 + phl
KF-143	KF	24	850	1 + pm
KF-144	KI	24	650	1 + few dendritic rystals
KF-145		24	650	1 + rare crystals
Diopside – KAlSiO	-F- ioin			2
DKF-87	Di80:KF20	7	1216	1
DKF-21	D100.1KI 20	55	1200	1 + di
DKF-5		75	1150	1 + di + lc
DKF-111		7	1130	1 + di + lc
DKF-36		60	1100	1 + di + lc + traces phl
DKF-97		18	1050	1 + di + lc + phl
DKF-53	Di60:KF40	6.5	1250	1
DKF-88		7	1216	1 + 1c
DKF-22		55	1200	1 + 1c + traces phl
DKF-112		7	1130	1 + 1c + phl
DKF-37		60 18	1100	1 + lc + phl + di
DKF-98		18 53	1050	1 + 1c + phl + di 1c + phl + di + fl
DKF-121 DKF-146	Di50:KF50	53 5	993 1170	lc + phl + di + fl l + lc
DKF-140 DKF-54	Di40:KF60	6.5	1250	1 + 1c
DKF-89		0.3 7	1230	$l_{s}^{1} + l_{f}$
DKF-23		55	1210	$l_{s}^{1_{s}} + l_{f}^{1_{f}} + l_{c}$
DKF-113		7	1130	$l_s^{r_s} + l_f^{r_f} + lc + phl$
DKF-38		60	1100	$l_s + l_f + lc + phl$
DKF-99		18	1050	$l_s + l_f + lc + phl$
DKF-122		53	993	$l_s + l_f + lc + phl$
DKF-133		50	950	1 + 1c + phl + fl
DKF-78	Di20:KF80	7.5	1300	1
DKF-55		6.5	1250	$l_s + l_f$
DKF-24		55	1200	$l_s + l_f$
		()	1100	$l_s + l_f$
DKF-39		60		
DKF-100		18	1050	$l_s + l_f + phl$
DKF-100 DKF-123		18 53	1050 993	$l_s + l_f + phl$ $l_s + l_f + phl$
DKF-100		18	1050	$l_s + l_f + phl$
DKF-100 DKF-123 DKF-134	e – KAlSiO ₃ F ₂ join	18 53 50	1050 993 950	$l_s + l_f + phl$ $l_s + l_f + phl$
DKF-100 DKF-123 DKF-134	e – KAlSiO ₃ F ₂ join Fo15:Di75:KF10	18 53	1050 993	$l_s + l_f + phl$ $l_s + l_f + phl$

Table 1 contd.

Sample	Starting composition, wt%	Duration, h	T °C	Phase composition of run products
1	2	3	4	5
FDKF-56		6.5	1250	1 + fo + di
FDKF-25		55	1200	1 + fo + di
FDKF-9		75	1150	1 + fo + di + phl
FDKF-40		60	1100	1 + fo + di + phl
FDKF-101		18	1050	1 + fo + di + phl + ?
FDKF-69	Fo55:Di25:KF20	5	1350	1 + fo
FDKF-80		7.5	1300	1 + fo
FDKF-57		6.5	1250	1 + fo + phl
FDKF-26		55	1200	1 + fo + phl
FDKF-10		75 60	1150	l + fo + phl + di
FDKF-41 FDKF-102		18	1100 1050	1 + fo $+ $ ph $1 + $ di 1 + fo $+ $ ph $1 + $ di $+ $ cu
FDKF-90	Fo40:Di40:KF20	7	1216	1 + 10 + phi + di + cd 1 + fo + phi
FDKF-114	1040.D140.K120	7	1130	1 + 10 + phi 1 + fo + phl + di
FDKF-70	Fo30:Di50:KF20	5	1350	1 + 10 + pm + m 1 + 10
FDKF-81	1000.0100.111 20	7.5	1300	1 + 10 1 + fo
FDKF-58		6.5	1250	1 + fo
FDKF-27		55	1200	1 + fo
FDKF-150		24	1190	1 + fo + phl
FDKF-147		5	1170	1 + fo + di + phl + lc
FDKF-11		75	1150	1 + fo + di + phl
FDKF-42		60	1100	1 + fo + di + phl + ak
FDKF-103		18	1050	fo + di + phl + ak + cu
FDKF-91	Fo15:Di65:KF20	7	1216	1 + fo
FDKF-151		24	1190	1 + fo + di
FDKF-148		5	1170	1 + di + lc + phl
FDKF-115		7	1130	1 + di + lc + phl
FDKF-104	E-10-D:(0-KE20	18	1050	1 + di + lc + phl
FDKF-152	Fo10:Di60:KF30	24 5	1190 1170	1 + 1c + phl
FDKF-149 FDKF-116		5 7	1170	l + lc + phl + di l + lc + phl + di
FDKF-71	Fo40:Di20:KF40	5	1350	1
FDKF-82	1 040.D120.K1 40	7.5	1300	1 + fo + phl
FDKF-59		6.5	1250	1 + 10 + phi 1 + fo + phi
FDKF-28		55	1200	1 + fo + phl
FDKF-12		75	1150	1 + fo + phl + sp
FDKF-43		60	1100	fo + phl + di + ak + traces lc
FDKF-105		18	1050	fo + phl + di + ak + cu + traces lc
FDKF-92	Fo30:Di30:KF40	7	1216	1 + phl
FDKF-117		7	1130	1 + phl + lc + di
FDKF-106		18	1050	1 + phl + lc + di + cu
FDKF-125		53	993	1 + phl + lc + di + cu
FDKF-131		50	950	phl + lc + di + ak + cu
FDKF-83	Fo20:Di40:KF40	7.5	1300]
FDKF-60 FDKF-29		6.5 55	1250 1200	1 + phl 1 + phl + lc + traces fo
FDKF-13		75	1150	1 + phi + lc 1 + phi + lc
FDKF-118		7	1130	1 + phl + lc
FDKF-44		60	1100	1 + phi + lc + di
FDKF-93	Fo10:Di50:KF40	7	1216	1 + lc + phl
FDKF-119		7	1130	1 + lc + phl 1 + lc + phl
FDKF-84	Fo20:Di20:KF60	7.5	1300	1
FDKF-61		6.5	1250	1 + phl
FDKF-30		55	1200	1 + phl
FDKF-14		75	1150	1 + phl
FDKF-45		60	1100	1 + phl + lc
FDKF-107		18	1050	1 + phl + lc
FDKF-126		53	993	l + phl + lc
FDKF-62	Fo10:Di10:KF80	6.5	1250	1
FDKF-94		7	1216	$l_s + l_f$
FDKF-31		55	1200	$l_s + l_f + phl$
FDKF-15		75	1150	$l_s + l_f + phl$
FDKF-46		60 18	1100	$l_s + l_f + phl$
FDKF-108		18 53	1050	$l_s + l_f + phl$ $l_s + l_s + phl$
FDKF-127 FDKF-135		53 50	993 950	$l_s + l_f + phl$ $l_s + l_f + phl$
1 DK1-133		50	200	$\mathbf{I}_{s} + \mathbf{I}_{f} + \mathbf{P}\mathbf{III}$

Sample	Starting composition, wt%	Duration, h	T °C	Phase composition of run products
1	2	3	4	5
Akermanite – l	F-phlogopite join			
AP-46	Ak80:Phl20	6	1350	1
AP-31		6	1300	1 + ak
AP-19		4	1250	1 + ak + fo
AP-13		9	1200	1 + ak + fo
AP-1 AP-62		5 9	1170 1150	1 + ak + fo 1 + ak + fo
AP-37		7	1130	1 + ak + 10 1 + ak + fo + di
AP-7		24	1100	1 + ak + 10 + di 1 + ak + fo + di + phl
AP-69		32	1075	1 + ak + fo + di + phl
AP-25		70	1050	ak + fo + di + phl + cu
AP-32	Ak70:Phl30	6	1300	1
AP-20		4	1250	1 + fo + ak
AP-14		9	1200	1 + fo + ak
AP-2		5	1170	1 + fo + ak
AP-63		9	1150	1 + fo + ak
AP-38		7	1130	1 + fo + ak + di
AP-8 AP-70		24 32	1100 1075	1 + fo + ak + di + phl 1 + fo + ak + di + phl
AP-26		70	1075	fo + ak + phl + di + cu
AP-48	Ak60:Phl40	6	1350	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
AP-33		6	1300	$\frac{1}{1 + fo}$
AP-21		4	1250	1 + fo
AP-52		5.5	1225	1 + fo
AP-15		9	1200	1 + fo + ak
AP-3		5	1170	1 + fo + ak
AP-64		9	1150	1 + fo + ak + lc
AP-43		7	1130	1 + fo + ak + lc + di
AP-9		24	1100	1 + fo + ak + lc + di + phl
AP-71 AP-27		32 70	1075 1050	1 + fo + ak + di + phl
AP-49	Ak50:Ph150	6	1350	fo + ak + phl + di + cu
AP-34	AK50.1 III50	6	1300	1 + 1 fo
AP-22		4	1250	1 + 10 1 + 10
AP-16		9	1200	1 + fo
AP-4		5	1170	1 + fo + 1c
AP-65		9	1150	1 + fo + 1c + ak
AP-44		7	1130	1 + fo + lc + ak + di
AP-10		24	1100	1 + fo + lc + phl + ak + di
AP-72		32	1075	1 + fo + phl + ak + di
AP-28	A 1-40-D1-160	70	1050	fo + ak + phl + di + cu
AP-50	Ak40:Phl60	6	1350	1 $1 + f_2$
AP-35 AP-23		6 4	1300 1250	$\begin{array}{c}1 + \text{ fo}\\1 + \text{ fo}\end{array}$
AP-53		5.5	1230	$1 + 10 \\ 1 + 10$
AP-17		9	1200	1 + 10 + 1c
AP-5		5	1170	1 + fo + lc
AP-66		9	1150	1 + fo + lc + phl
AP-41		7	1130	1 + fo + lc + phl
AP-11		24	1100	1 + fo + traces lc + phl + ak + di
AP-73		32	1075	1 + fo + phl + ak + di
AP-29	41 00 DI 100	70	1050	fo + ak + phl + di + cu
AP-75	Ak20:Phl80	5	1375	$\frac{1}{1+f_{0}}$
AP-67		4	1370	1 + fo 1 + fo
AP-36 AP-24		6 4	1300 1250	1 + 10 1 + 10 + 10
AP-54		4 5.5	1230	1 + 10 + phi 1 + fo + phl + lc
AP-18		9	1223	1 + 10 + phi + 1c 1 + fo + phi + 1c
AP-6		5	1170	1 + 10 + phi + 1c 1 + 10 + phi + 1c
AP-12		24	1100	1 + fo + phl + lc + ak
AP-74		32	1075	1 + fo + phl + ak
AP-30		70	1050	fo + phl + ak + di + cu

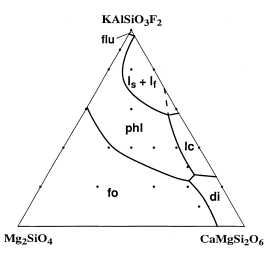


Fig. 2 Primary phase fields of the Mg_2SiO_4 – $CaMgSi_2O_6$ – KAlSiO₃F₂ pseudo-ternary join. *Points* indicate positions of reactant mixtures (starting compositions). For abbreviations see the Appendix

Akermanite

Early liquidus akermanite forms larger $(50-60 \mu m)$ tabular crystals. In near-solidus runs melilite occurs as irregular interstitial fillings and can only with difficulty be identified in the microscope.

The chemical composition of melilite in the run products (Table 3) is close to that of pure akermanite. Aluminium and potassium were detected in small amounts (1.36-1.82 wt% Al₂O₃ and 0.18-0.85 wt% K₂O) and they seem to enter melilite solid solution in the form of K-melilite (KCaAlSi₂O₇; K equivalent to sodamelilite) and gehlenite (Ca₂Al₂SiO₇) components.

F-Phlogopite

F-phlogopite usually occurs as hexagonal plates or scales that have a distinctive lustre visible to the naked

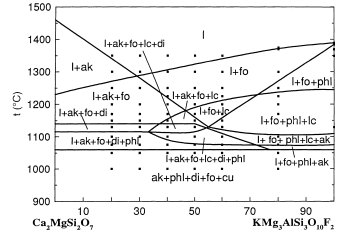


Fig. 3 T-x phase diagram of the Akermanite – F-phlogopite pseudobinary join; data from Table 1

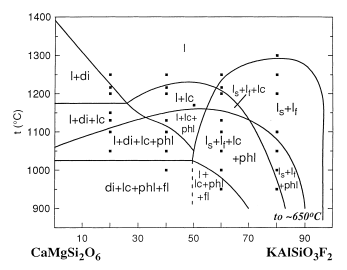


Fig. 4 T-x phase diagram of the Diopside – $KAlSiO_3F_2$ pseudobinary join; data from Table 1

eye in the run products. Small F-phlogopite needles appeared in a few runs and are interpreted to be quench crystals. Representative EMP analyses of F-phlogopite are listed in Table 4. Aluminium is only present in tetrahedral sites of the phlogopite structure. The Al/Si mole values vary (0.196–0.34) and correlate positively to the magnesium content. Thus, it is possible to suggest, that excess silicon in tetrahedral sites (relative to ideal formula) is counterbalanced by vacancies in octahedral sites and the main substitution mechanism for the phlogopite solid solutions is: $Mg^{VI} + 2Al^{IV} = 2Si^{IV} + \Box^{VI}$. Similar substitution has been described by Robert (1976) and Melzer and Foley (1996) for synthetic phlogopite in the SiO₂ – Al₂O₃ – MgO – K₂O – H₂O (F) system.

Spinel

Small amounts of spinel occur in some forsterite-rich runs as well-shaped isotropic octahedral crystals with very high refractive index. A single EMP analysis shows a pure Mg-Al phase (MgAl₂O₄). Spinel seems to disappear at lower temperatures due to the reaction with liquid.

Cuspidine

Cuspidine is a near-solidus and sub-solidus phase which is identified only by EMP. Analyses show that cuspidine contains MgO in essential amounts (2.61–3.36 wt%). The average composition of 14 analyses is: $(Ca_{3.7}Mg_{0.26} K_{0.02})Si_2O_7F_2$.

Fluorite and other minor phases

Fluorite was detected by XRF analyses and EMP in some sub-solidus leucite-bearing assemblages (see

3	5	4

Sample n	DKF 98 3	FDKF 42 2	FDKF 148 2	FDKF 115 2	AP43 2	AP9 4	AP10 4	AP11 3	AP29 2
SiO ₂	54.88	54.24	54.89	55.11	54.13	53.22	53.23	54.20	53.37
$Al_2 \tilde{O}_3$	0.58	0.75	0.84	0.25	1.15	2.30	2.14	1.79	2.51
MgO	19.17	19.22	20.65	20.60	19.44	18.89	18.89	18.70	18.47
CaO	26.50	25.62	24.16	24.19	26.24	26.06	25.87	25.37	25.29
K ₂ O	0.09	0.09	0.01	0.01	0.07	0.10	0.06	0.08	0.09
F	0.04	0.06	0.01	0.01	0.01	0.08	0.08	0.05	0.20
Sum	101.24	99.96	100.54	100.14	101.01	100.62	100.25	100.19	99.93
Atoms per	· 6 oxygens								
Si	1.96	1.96	1.96	1.98	1.94	1.92	1.92	1.95	1.93
Al	0.02	0.03	0.04	0.01	0.05	0.10	0.09	0.08	0.11
Mg	1.02	1.04	1.10	1.10	1.04	1.02	1.02	1.00	1.00
Ca	1.02	0.99	0.93	0.93	1.01	1.01	1.00	0.98	0.98
Κ	0.00	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00
F	0.01	0.01	0.00	0.00	0.00	0.01	0.01	0.01	0.02

Table 2 Representative compositions of clinopyroxene in wt% of oxides and atoms per formula unit (n number of analyses)

Table 3 Compositions of melilite in wt% of oxides and atoms per formula unit (n number of analyses)

Sample <i>n</i>	FDKF42 2	AP15 2	AP43 7	AP9 3	AP10 3	AP11 3	AP29 2
	2	-	,	5	5	5	2
SiO ₂	43.51	44.22	43.57	43.65	44.04	44.44	44.01
Al_2O_3	1.82	0.01	1.38	1.36	1.57	1.51	1.61
MgO	13.73	15.17	13.93	13.91	13.69	13.66	13.49
CaO	40.15	41.96	40.67	40.72	40.20	39.38	39.02
K ₂ O	0.68	0.18	0.62	0.76	0.70	0.59	0.85
F	0.19	0.01	0.03	0.01	0.01	0.01	0.08
Sum	100.07	101.51	100.18	100.39	100.21	99.59	99.06
Atoms per 7	oxygens						
Si	1.98	1.98	1.98	1.98	1.99	2.01	2.01
Al	0.10	0.00	0.07	0.07	0.08	0.08	0.09
Mg	0.93	1.01	0.94	0.94	0.92	0.92	0.92
Ca	1.96	2.02	1.98	1.98	1.95	1.91	1.91
K	0.04	0.01	0.04	0.04	0.04	0.03	0.05
F	0.03	0.00	0.00	0.00	0.00	0.00	0.01

Table 1). Its chemical composition is close to pure CaF_2 , but contains about 1.5 wt% MgF₂.

An unidentified anisotropic phase, presumably K_3AlF_6 , appears on the liquidus of the KAlSiO₃F₂ bulk composition at ca. 650 °C. It forms thin elongated crystals, or stubby plates and seems to have high bire-fringence.

Finally a representative of the humite group (F-chondrodite ?) was detected by a single EMP analysis in the sub-solidus assemblage of the Forsterite – KAlSiO₃F₂ join (see Table 1) and this phase may be present in trace amounts.

Topology of phase equilibria

Components and the phase rule

In a strict sense the studied joins are pseudo-binary and pseudo-ternary sub-systems of the seven-component Si - Al - Mg - Ca - K - O - F system. Nevertheless, the following assumptions allow to reduce the number of

components and to use three-dimension coordinates of the kalsilite-based normative tetrahedron (Fig. 1) for the graphic representation of phase equilibria and for the comparison to the relations established in volatilefree systems.

Firstly, the mass proportion of vapour, as mentioned above, seems to be negligible and the system may be treated as condensed. Thus, it is possible to present the components in the conventional oxide form and exclude the fluorine component $(2F^- = O^{2-})$. In effect, variations in F-O substitution and fluorine content of the phases will not show, and some F-bearing phases plot in the same positions as the corresponding F-free ones (for example, Ca₄Si₂O₇F₂ (cuspidine) and Ca₂SiO₄ (larnite). But the vast majority of phases differ in components other than F and O and retain a unique position within the tetrahedron.

Secondly, it is possible to combine Al_2O_3 and K_2O into a single kalsilite (KAlSiO₄) component, because K/ Al mole values for all starting compositions are equal to 1. It should, however, be noted that in some phases (such as spinel and in clinopyroxene, melilite and

Table 4 Representative phlogopite compositions in wt% of oxides and atoms per formula unit (*n* number of analyses)

Sample n	DKF 112 2	DKF 37 2	FDKF 147 2	DKF 121 2	FDKF 42 2	FDKF 103 2	FDKF 148 3	FDKF 125 2
SiO ₂	44.81	41.93	43.22	48.01	43.22	44.31	42.51	42.03
$Al_2 \tilde{O}_3$	10.05	12.06	11.24	7.98	11.53	10.32	11.69	11.62
MgO	28.32	28.50	28.62	27.63	28.68	28.13	29.12	28.45
CaO	0.18	0.11	0.15	0.01	0.15	0.17	0.09	0.42
K ₂ O	10.79	10.78	10.98	10.36	10.91	10.96	10.86	10.75
F	8.39	8.57	8.37	8.16	8.27	8.39	8.49	8.17
Sum	98.99	98.40	99.12	98.78	99.34	98.80	99.26	98.07
Atoms per	10 oxygens							
Si	3.13	2.97	3.02	3.31	3.01	3.10	2.98	2.97
Al	0.83	1.01	0.93	0.65	0.95	0.85	0.97	0.97
Mg	2.94	3.01	2.99	2.84	2.98	2.94	3.04	3.00
Ca	0.01	0.01	0.01	0.00	0.01	0.01	0.01	0.03
K	0.96	0.97	0.98	0.91	0.97	0.98	0.97	0.97
F	1.85	1.92	1.85	1.78	1.82	1.86	1.88	1.83

Table 4 (continued)

Sample n	FDKF106 2	FDKF 117 2	FDKF92 2	AP9 3	AP10 3	AP11 3	AP29 2	
SiO ₂	42.25	42.67	42.46	41.78	41.81	43.17	42.88	
Al_2O_3	11.02	12.13	11.78	12.08	11.99	12.14	12.01	
MgO	28.11	28.85	28.76	28.93	29.03	28.42	28.36	
CaO	1.44	0.09	0.14	0.17	0.15	0.21	0.23	
K ₂ O	10.55	10.84	10.94	10.88	10.60	10.95	10.99	
F	8.69	8.11	8.04	8.35	8.40	8.41	8.47	
Sum	98.47	99.32	98.78	98.76	98.52	99.85	99.46	
Atoms per 1	0 oxygens							
Si	3.01	2.96	2.97	2.94	2.95	3.00	3.00	
Al	0.93	0.99	0.97	1.00	1.00	0.99	0.99	
Mg	2.98	2.99	3.00	3.04	3.05	2.94	2.95	
Ca	0.11	0.01	0.01	0.01	0.01	0.02	0.02	
K	0.96	0.96	0.98	0.98	0.95	0.97	0.98	
F	1.96	1.78	1.78	1.86	1.88	1.85	1.87	

phlogopite solid solutions) K_2O and Al_2O_3 tend to behave as independent components. However, the variations in K/Al values during crystallisation are minor and can be neglected.

Data listed in Table 1 show that the number of equilibrium condensed phases (solids and liquids) on the joins at any given temperature never exceeds six. As discussed below, associations of six condensed phases are observed in narrow temperature intervals and they are very close to invariant isobaric equilibria. Thus, phase rule considerations suggest five effective components, and if vapour phase and variations in K/Al values are ignored, main phases and equilibria may be presented in the coordinates of kalsilite-based normative tetrahedron (Fig. 1).

Thermal barriers

Yoder (1986) has previously shown that phlogopite takes part in a number of heteromorphic relationships and a pattern of tie-lines and sub-tetrahedra within the kalsilite-normative tetrahedron changes drastically when

phlogopite stability is achieved. Important features of the volatile-free, and consequently phlogopite-absent, system (Fig. 1a), are thermal barriers close to the kalsilite-forsterite-akermanite (ks-fo-ak) and kalsiliteleucite-akermanite (ks-lc-ak) planes that separate the forsterite-saturated compositions from the highly larnite normative part of the system. In the volatile-free system silica-undersaturated compositions evolve through the fo-mo-ak-ks invariant reaction point further to the foak-lc-ks eutectic (Yoder 1986). The evolution of more silica-rich, but still undersaturated compositions to the right of the fo-ak-lc thermal barrier (Fig. 1a) terminates at the fo-ak-lc-di eutectic (see flow sheet for dry kalsilitebased normative tetrahedron presented by Yoder 1986). The stability of the F-phlogopite – cuspidine assemblage in the F-doped system breaks the ks-fo-ak thermal barrier and permits the liquid to evolve towards highly larnite normative residuals. Pure OH-analogue of cuspidine (custerite) does not appear to exist as a stable compound (Van Valkenburg and Rynders 1958). Similar to the F-doped system, the formation of OH-phlogopite and stabilisation of the phlogopite-monticellite association under H₂O-pressure result in the breakdown of the

ks-fo-ak thermal barrier. However, in the hydrated kalsilite-based tetrahedron no tie-line appears to connect phlogopite to the larnite apex (Yoder 1986).

Sub-solidus assemblages

All batch compositions on the Akermanite – F-phlogopite join crystallise into five-phase ak + phl + fo + di + cusub-solidus assemblage (Table 1 and Fig. 3). The join is situated between three stable sub-tetrahedra in the Fdoped normative tetrahedron (Fig. 1b) and all five phases of the sub-tetrahedra are present in sub-solidus (please notice that the join in fact is a part of the sevencomponent system).

The Forsterite – Diopside – $KAlSiO_3F_2$ plane cuts across the Akermanite - F-phlogopite join (see Fig. 1b) and thus the same five-phase sub-solidus assemblage is the final assemblage for many $Mg_2SiO_4 - CaMgSi_2O_6 -$ KAlSiO₃F₂ mixtures (see Table 1 and Fig. 2). But for the pseudo-ternary join this sub-solidus five-phase association is not unique and several other sub-solidus assemblages have been found on the join (Table 1). They include: phl+lc+di+ak+cu five-phase association (sample FDKF-131 in Table 1) and four-phase assemblage: di + lc + phl + fl which is present on the Diopside – KAlSiO₃F₂ border join (Fig. 4) and in adjacent forsterite-bearing starting compositions. Different sub-solidus assemblages within an isobaric join require thermal barriers, or reaction relations to be involved and this may be valid for the studied joins.

Six-phase isobaric equilibria with forsterite are plotted in Fig. 5 as a fragment of a flow sheet in coordinates of the KAlSiO₄ – Mg_2SiO_4 – Ca_2SiO_4 – SiO_2 – F system.

Reactions

The most important topological feature for the forsterite-bearing liquidus relationships (Fig. 5) is the sixphase l + fo + lc + di + ak + phl reaction "point" in which forsterite and leucite react with F-bearing melt to produce phlogopite:

$$fo + lc + l = phl \tag{1}$$

and the reaction proceeds until either forsterite, or leucite is completely consumed. The six-phase assemblage: 1+fo+lc+di+ak+phl has been found at 1100 °C in three batch compositions in the central part of the Akermanite – F-phlogopite join. On this join leucite is the phase to react-out. As the system in fact is a sevencomponent system (potassium and aluminium are in a strict sense independent components) the six-phase reaction assemblage is not an invariant point, but a reaction over a temperature range of about 30 °C. Traces of leucite are often preserved at temperatures below the reaction interval, as resorbed solid inclusions in large akermanite crystals and may be considered metastable,

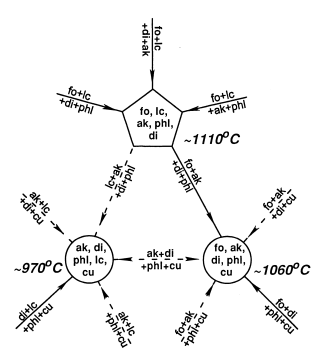


Fig. 5 Forsterite-saturated, low-silica part of a flow sheet for the $Mg_2SiO_4 - Ca_2SiO_4 - KAlSiO_4 - SiO_2 - F$ system at atmospheric pressure showing the six-phase invariant reaction point (*pentagon*) and two eutectics (*circles*). Experimentally determined invariant points and monovariant equilibria are shown in *solid lines*. For abbreviations see the Appendix

because the host effectively protects the remnants of leucite from the liquid and further resorbtion.

The reaction of forsterite and leucite with the F-bearing liquid takes place not only at the reaction "point". For some batch compositions it may start (and end) at higher temperatures along the three peritectic lines leading to the reaction point (see Fig. 5). For example, the resorbtion of leucite on the Akermanite – Fphlogopite join is observed to occur in equilibrium with forsterite, akermanite and diopside before phlogopite appears on the liquidus (Fig. 6). There are also batch compositions, e.g. Fo15:Di65:KF20 in which forsterite is consumed at 1175 °C while leucite remains on the liquidus and in the solidus assemblage together with diopside, leucite and phlogopite (see Table 1). Below the temperature of the 1+fo+lc+ak+phl+di reaction point the system may evolve either to the 1+ak+di + phl + lc + cu or to the l + fo + ak + di + phl + cu eutectics (Fig. 5) depending on which solid phase (forsterite, or leucite) is consumed first.

Other six-phase reaction points appear to be present in the forsterite-absent parts of the system. Six-phase equilibrium of di, lc, phl and fl with two immiscible liquids (silicate and fluoride) seems to exist close to the Diopside – KAlSiO₃F₂ border join at about 1050 °C (Fig. 4). It suggests a major homogeneous exchange reaction between species in the liquid:

$$CaMgSi_2O_6 + 2KAlSiO_3F_2 = CaF_2 + MgF_2 + 2KAlSi_2O_6$$
(2)

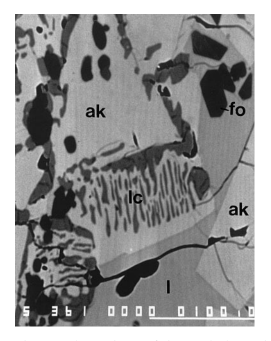


Fig. 6 Back-scatter electron image of the sample AP-43 showing leucite resorbtion in equilibrium with akermanite (*large, light euhedral crystals*), forsterite (*dark, small euhedral crystals*) and the melt (*light-grey regions*)

Phase equilibria on the Diopside – KAlSiO₃F₂ join imply that the right-hand assemblage of species is stable in the melt. In the presence of crystalline leucite and fluorite activities of KAlSi₂O₆ and CaF₂ components in the liquid approach 1. Thus, diopside can crystallise only from F-poor compositions where the CaMgSi₂O₆ component is in excess of reaction (2). Forsterite-free compositions that yield diopside crystallise completely at about 1030 °C to di+lc+phl+fl sub-solidus assemblage (see Table 1 and Fig. 4). Solidus temperatures for the F-rich part of the join from which diopside is absent are very low. The experimental data for this part of the system is insufficient for detailed discussions of topology.

Chemical evolution of the liquid

Electron microprobe analyses of quenched melts (glasses) in the charges on the studied joins provide direct information on the evolution of the coexisting melts. Thermal barriers, discussed in the previous section, result in different trends of melt evolution. Not all the numerous samples listed in Table 1 were studied by EMP. We have focused on the melts in equilibrium with the phase assemblages typical for igneous rocks and paid special attention to phlogopite-bearing assemblages of ultramafic alkaline complexes. Chemical compositions of representative glasses saturated with three or more crystalline phases are listed in Table 5.

Larnite-normative trend

For all studied batch compositions outside the silicate melt – F-melt immiscibility field MgO and possibly SiO_2 contents in the melts decrease and CaO content increases with falling temperature, while Al_2O_3 and K_2O concentrations show no significant variation. As a result, the evolved compositions of the residual melts are extremely larnite normative (Table 6).

The larnite-normative melt evolution is illustrated by three typical batch compositions in which forsterite is the first liquidus phase (Figs. 7 and 8). Weight proportions of the phases (Fig. 7) were calculated using massbalance equations based on EMP analyses of the run products. Common features of all three batch compositions are a strong decrease in proportion of the melt and a rapid growth of phlogopite at the expense of the melt, leucite and forsterite below the temperature of the phlogopite-in reaction (Eq. 1). For the Ak50:Phl50 batch composition the reaction takes place at the sixphase reaction point (see the previous section and the flow sheet in Fig. 5.), but for the other two compositions the reaction occurs at higher temperatures along the 1 + fo + lc + di + phl peritectic "line". The compositions Ak50:Phl50 and Fo30:Di50:KF20 represent the case where leucite is consumed completely in the reaction and forsterite remains, although in smaller quantities, in the assemblage until total crystallisation. For the Fo15: Di65:KF20 composition, on the contrary, forsterite is the phase to react out, and leucite is preserved in the sub-solidus. The latter batch composition differs from the other two by having fluorite in the sub-solidus assemblage instead of cuspidine.

The evolution of the melt in the three batch compositions is shown in Fig. 8 as concentrations of major components versus temperature and shows the effect of the phlogopite-in reaction. As crystallisation proceeds below the reaction temperature, SiO₂, K₂O and Al₂O₃ concentrations in the melt decrease, while CaO and F contents increase. The MgO content of the liquid decreases steadily within the whole crystallisation range and is not significantly affected by the reaction. Increase in CaO content of the liquid below the phlogopite-in reaction temperature is obviously related to intense phlogopite crystallisation, because F-phlogopite is rich in all the components except CaO (see Table 4). It should be noted that even akermanite crystallisation does not prevent the growth in CaO (e.g. batch composition Ak50:Phl50, Fig. 8). If leucite is preserved in the liquidus phase assemblage (Fo15:Di65:KF20 batch composition), fluorine initially introduced into the system in the form of the KAlSiO₃F₂ component accumulates in the residual melt in greater quantities (Fig. 8, see also increase in Fl norm in Table 6).

The compositions of residual melts for Ak50:Phl50 and Fo30:Di50:KF20 starting mixtures at 1100 °C are very similar (samples AP10 and FDKF42 in Table 5). The CIPW norms show that all residual melts have extremely high normative larnite (Table 6). The norms in

 Table 5 Selected compositions of glasses saturated in three or more crystal phases. Standard deviations are in brackets (n number of analyses)

Sample	T °C	п	Phases in	Compos	ition, wt%					
			equilibrium	SiO ₂	Al_2O_3	MgO	CaO	K ₂ O	F	Sum
FDKF	1100	3	fo + phl + di	41.19	8.42	12.53	26.54	4.65	6.34	96.99
41			*	(1.82)	(0.48)	(0.64)	(0.38)	(0.16)	(0.18)	
FDKF	1170	5	di + phl + lc	47.95	6.09	16.09	20.32	5.45	5.42	99.04
148				(0.14)	(0.18)	(0.18)	(0.40)	(0.08)	(0.18)	
FDKF	1130	4	di + phl + lc	42.19	4.86	13.34	28.74	3.72	9.53	98.37
115				(0.46)	(0.17)	(0.22)	(0.10)	(0.07)	(0.33)	
FDKF	1130	3	di + phl + lc	41.56	4.59	13.23	30.95	4.59	7.08	99.02
117				(0.10)	(0.22)	(0.14)	(0.19)	(0.16)	(0.18)	
DKF	1100	7	di + phl + lc	39.13	13.52	11.60	29.04	3.43	4.83	99.52
37			-	(0.67)	(0.24)	(0.34)	(0.34)	(0.11)	(0.28)	
DKF	1050	6	di + phl + lc	35.41	2.37	11.75	40.01	1.54	16.94	100.89
98			-	(0.27)	(0.16)	(0.24)	(0.34)	(0.11)	(0.24)	
FDKF	1050	5	di + phl + lc	39.14	4.69	8.85	35.80	3.53	12.80	99.42
104			-	(0.42)	(0.16)	(0.10)	(0.57)	(0.04)	(0.33)	
FDKF	993	2	di + phl + lc	37.74	6.59	18.00	25.29	6.20	9.12	99.10
125			+ cu	(1.48)	(0.66)	(0.69)	(1.95)	(0.37)	(0.11)	
DKF	950	4	phl + lc + fl	53.25	23.62	0.04	0.19	ì1.15	12.49	100.73
133			1	(0.27)	(0.28)	(0.03)	(0.14)	(0.65)	(0.55)	
FDKF	1170	7	fo + phl + di	45.48	5.99	16.68	22.03	5.15	5.72	98.64
147			$+ 1c^{T}$	(0.38)	(0.13)	(0.36)	(0.55)	(0.24)	(0.29)	
FDKF	1050	2	fo + phl + di	46.65	Ì1.91	6.21	21.85	7.04	7.43	97.97
102			+ cu	(1.76)	(0.50)	(0.33)	(1.27)	(0.43)	(0.67)	
AP43	1130	7	fo + ak + lc	41.87	5.28	15.02	28.63	5.43	6.04	99.73
			+ di	(0.17)	(0.14)	(0.56)	(1.17)	(0.13)	(0.18)	
AP44	1130	3	fo + ak + di	41.63	5.61	14.54	28.69	5.18	5.92	99.08
			+ 1c	(0.24)	(0.15)	(0.29)	(0.24)	(0.19)	(0.12)	
FDKF	1100	3	fo + ak	41.59	4.97	13.73	27.7	5.94	6.5	97.69
42			+ phl $+$ di	(0.40)	(0.42)	(0.24)	(0.49)	(0.18)	(0.39)	
AP11	1100	3	fo + ak +	40.65	3.48	15.21	29.32	6.17	7.79	99.34
••		-	phl + di	(0.41)	(0.07)	(0.21)	(0.23)	(0.13)	(0.02)	
AP9	1100	5	fo + ak + lc	40.30	3.47	14.91	28.90	6.70	7.78	98.78
/			+ di + phl	(0.40)	(0.42)	(0.24)	(0.49)	(0.18)	(0.39)	20.70
AP10	1100	6	fo + ak + lc	40.44	3.72	14.43	28.24	6.95	7.51	98.13
••		÷	+ di + phl	(0.52)	(0.21)	(0.24)	(0.56)	(0.25)	(0.16)	20.10

Table 6 CIPW norms of glasses (quenched liquids) plotted in Figs. 7 and 8. Abbreviations for normative components are listed in the Appendix

Sample	T ℃	Phases in	Normat	ive compo	sitions of g	lasses, wt%	0			
		equilibrium	Fl	Or	Lc	An	KS	Di	Fo	La
Fo30:Di50:KF	20									
Starting batch	compositio	on	8.67	15.49	12.07			25.95	37.81	
FDKF70	1350	fo	7.74	17.75	9.38	0.76		33.06	31.31	
FDKF127	1200	fo	9.53	21.45	11.56	1.50		38.88	17.09	
FDKF147	1170	fo + di + phl + lc	11.91		24.19	1.15		44.81	14.96	2.98
FDKF42	1100	fo + di + phl + ak	13.67		21.78		3.14	30.33	14.67	16.40
Fo15:Di65:KF	20	-								
Starting batch	compositio	on	8.67	15.49	12.08			40.95	22.81	
FDKF91	1220	fo	7.93	16.76	10.58	0.99		48.75	15.00	
FDKF151	1190	fo + di	9.78	25.65	9.79	1.55		37.80	15.44	
FDKF148	1170	lc + di + phl	11.24	10.27	17.44	0.53		47.63	12.88	
FDKF115	1130	lc + di + phl	19.91		17.52	2.31		51.88	6.82	1.57
FDKF104	1050	lc + di + phl	26.59		16.54	2.40		48.07		6.40
Ak50:Ph150		-								
Starting batch	compositio	on	9.27		25.91			19.85	31.50	13.47
AP22	1250	fo	8.09		27.86		0.05	23.15	23.54	17.31
AP4	1170	fo + lc	9.18		27.51	0.44		26.01	18.23	18.63
AP44	1130	fo + lc + ak + di	12.28		24.23			25.06	17.47	20.96
AP10	1100	fo + lc + ak +	15.73		16.23		8.15	26.61	17.02	16.27
		di + phl								

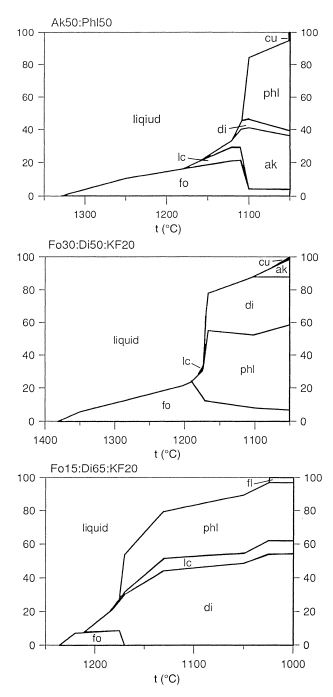


Fig. 7 Proportions of phases (in wt%) versus temperature for three representative batch compositions. The starting compositions (in wt%) are indicated *above* the plots. Phase proportions are calculated using mass-balance equations and EMP analyses of coexisting phases

Table 6 demonstrate that batch compositions Fo30: Di50:KF20 and Fo15:Di65:KF20 are examples of melanephelinitic compositions which initially are not larnite normative, but evolve to highly larnite normative (melilitie) residuals due to phlogopite crystallisation. In other words, they illustrate fractionation across the thermal barrier of the dry system to larnite-normative melilititic compositions under volatile pressure in accordance with the suggestions of Nielsen (1994) and Nielsen et al. (1997).

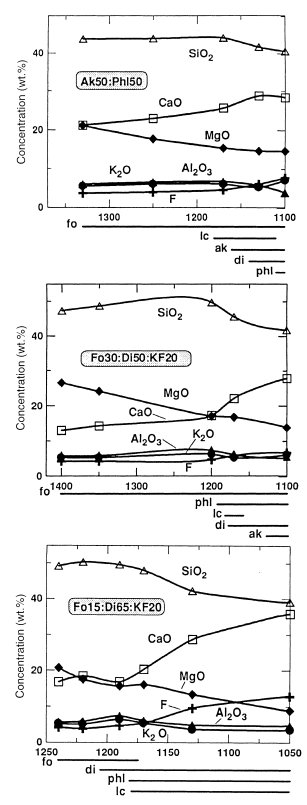


Fig. 8 Compositions of the remaining liquids (in wt% of oxides) versus temperature for the same batch compositions as in Fig. 7. *Lines below* the plots show crystallisation intervals for liquidus phases. For abbreviations see the Appendix

Homogeneous exchange reaction (2) seems to be a limiting factor for the larnite-normative trend of melt

evolution. In the F-rich part of the $Mg_2SiO_4 - CaMg_Si_2O_6 - KAlSiO_3F_2$ join where the $KAlSiO_3F_2$ component is in excess and neither diopside, nor forsterite are stable other trends are observed. For example, low-temperature residual liquids in equilibrium with leucite, phlogopite and fluorite are very Mg and Ca poor, SiO_2-rich and per-aluminous (sample DKF-133 in Table 5). Low K/Al mole values of these liquids are believed to result from crystallisation of phlogopite solid solutions with K/Al > 1 (due to the above-mentioned $Mg^{VI} + 2Al^{IV} = 2Si^{IV} + \Box^{VI}$ substitution). Liquid immiscibility is another phenomenon affecting the liquid evolution in the F-rich part of the system.

Silicate-fluoride liquid immiscibility

Immiscible fluorine-rich liquid occurs as spherical droplets, 20-100 microns in diameter, in the part of $Mg_2SiO_4 - CaMgSi_2O_6 - KAlSiO_3F_2$ join adjacent to the $CaMgSi_2O_6 - KAlSiO_3F_2$ border (Figs. 2 and 4). When quenched the fluorine-rich melt droplets crystallise to fine-grained aggregates and the conjugate silicate liquid often quenches to turbid, opacified glass. At lower temperatures silicates (leucite and/or phlogopite) start to crystallise in equilibrium with the immiscible liquids (see Table 1). Onset of the fluorite crystallisation together with leucite and phlogopite is a low-temperature limit for the liquid immiscibility (Fig. 4). The immiscibility field is located in a very F rich portion of the Mg₂SiO₄ – $CaMgSi_2O_6 - KAlSiO_3F_2$ join and is separated from the field of petrologically relevant compositions by the field of F-phlogopite. Thus, silicate-fluoride liquid immiscibility has little importance for applications discussed below. Details of phase equilibria and chemical composition of the immiscible liquids will be presented elsewhere.

Discussion

The application of the experimental results to natural magmatic systems requires examination of the intrinsic limitations. The simplified experimental system studied excludes major magmatic components such as ferrous and ferric iron, titanium, sodium, phosphorus and some phases common in natural magmatic systems are not present in the run products. It is also clear that the application is restricted to a limited, silica-undersaturated compositional range and low, crustal pressures. It can also be questioned to what extent the fluorinedoped system models the phase equilibria in natural volatile-bearing silicate melts. In general, H₂O rather than F is the main volatile component in natural magmas. Detailed discussion on the effect of the H₂O-F ratio on the crystallisation and properties of silicate melts is a broad field beyond the general scope of the study, but some effects are of great importance and outlined below.

Fluorine and water in silicate melts

Fluorine contents in the experimental batch compositions are much higher than those usually observed in igneous rocks and expected in natural magmatic melts. The F concentrations can only be compared to F concentrations in rare topaz-bearing rhyolites and ongonites with bulk-rock concentrations as high as 2-4.5 wt% (Congdon and Nash 1988; Kovalenko 1973). Melt inclusions in quartz suggest that fluorine content in the evolved silicic melts parental for these rocks may be even higher, up to 5.5 wt% (Carroll and Webster 1994). Very high bulk-rock F concentrations for mantle-derived mafic rocks are found in lamproites (up to 0.76 wt% F, Aoki et al. 1981; Edgar and Charbonneau 1991). These values are 3–4 times higher than the F concentrations in other mafic-ultramafic rocks and 6-8 times greater than in tholeiitic basalts. Thus, it must be expected that the F concentration in the silica-undersaturated mantle-derived magmatic melts is an order of magnitude lower than in the synthetic mixtures.

Water is believed to be the most abundant volatile component in magmas and despite the fact that F and H_2O (OH) play similar roles and generally substitute for each other in crystal structures there are many significant differences in their chemistry. The F-OH exchange influences the stability and physical properties of volatile-bearing crystalline phases and, as previously pointed out, the thermal stability of F-phlogopite significantly exceeds that of the hydroxyl analogue (Munoz 1984). Other F-bearing phases in the studied systems either do not have hydroxyl equivalents, e.g. cuspidine, or the hydrated phases have never been reported to crystallise from H₂O-bearing silicate melts.

The solubilities of H_2O and F in silicate melts have recently been reviewed by Carroll and Webster (1994) and McMillan (1994). Despite much controversy it is strongly suggested that fluorine forms complexes in aluminosilicate melts with network modifying cations, such as alkali and alkaline earths (Krigman and Kogarko 1981; Foley et al. 1986). This implies that F enhances the polymerisation of the melt and increases the activity of highly polymerised crystalline phases.

On the other hand, there is a strong evidence in favour of the depolymerising effect of water on the melt structure (McMillan 1994). In polymerised hydrated aluminosilicate melts linkages between tetrahedrally coordinated ions seem to be weakened (if not broken) by attachment of hydrogen to bridging oxygen and this may have important consequences for the crystallisation of the melts in H_2O -bearing systems.

Zeng and MacKenzie (1984) showed that H_2O decreases the stability of leucite and that the liquidus field of leucite shrinks dramatically under increasing water pressure. If the relative stabilities of phlogopite and leucite in analogue fluorine-doped and water-bearing systems can be compared it should be expected that both phases are less stable in hydrated systems, but the F-OH substitution may affect leucite stability to a greater

extent than the stability of phlogopite and this may change the topology of phase equilibria.

Effects of pressure

Akermanite is not stable at high pressure. Its stability field in the anhydrous systems is limited to a maximum pressure of about 15 kbar (Yoder 1986). Melzer and Foley (1996) demonstrated that leucite stability in the KAlSiO₄ - Mg₂SiO₄ - SiO₂ - F system depends on pressure and F content of batch compositions. Leucite was found to disappear from associations of liquidus phases at 10 kbar for 6 wt% F and 18 kbar for 3 wt% F. Neither leucite, nor akermanite are thus expected to be present at high pressures in phase associations of the kalsilite-based tetrahedron. However, instability of melilite and leucite will not prevent melts from evolving towards Ca-rich residuals. As shown above, the key phase for the larnite-normative trend is phlogopite and as long as the association fo + cpx + phl is stable the silica-undersaturated liquids may evolve towards the larnite apex of the normative tetrahedron (Fig. 1b).

Phlogopite liquidi are usually very steep on P-T phase diagrams for potassic rocks and in synthetic systems and it seems that pressure above 0.7 kbar has minor effect on the stability of phlogopite.

No compositional data appear to exist for glasses in equilibrium with forsterite, Ca-clinopyroxene and phlogopite in the water-bearing kalsilite normative tetrahedron at high pressure. Modreski and Boettcher (1973) report the composition of a glass on the diopside – phlogopite join (reactant mixture 1:1 by weight) at 1250 °C, 15 kbar, but at this temperature the glass is equilibrated with forsterite and diopside only, while phlogopite was found to crystallise at about 50 °C below. Even though, it is worth noticing that the glass is silica undersaturated and slightly larnite normative (0.9 wt% of normative larnite). Our results suggest that the onset of phlogopite crystallisation would drive the melt towards more larnite-rich compositions.

The topology of phase equilibria in the kalsilite-based normative tetrahedron with stable phlogopite (Fig. 1b) suggests that a plane close to the fo-di-phl join is critical for the larnite-normative melt evolution. Compositions above and to the left of this plane, or divide (Fig. 1b) would evolve towards larnite-normative residuals, whereas more SiO₂-rich and CaO-poor compositions below and to the right of the plane will not produce larnite-normative melts during fo+di+phl fractionation. Formation of diopside-enstatite solid solutions and the Mg^{VI} + 2Al^{IV} = 2Si^{IV} + \Box^{VI} substitution in phlogopite suggest that the critical plane should lie slightly below the fo-di-phl join. Previously published data on rock-melting experiments (see below) indicate that both trends do occur in natural multicomponent systems.

Applications

Evolution of the potassic magmas

The results of the experimental study can be applied to the generation and low-pressure evolution of group II ultrapotassic rocks, or kamafugites (classification introduced by Foley et al. 1987). Bulk compositions of kamafugites with leucite, melilite and phlogopite are characterised by low SiO₂ and Al₂O₃ and high CaO (Foley 1992; Foley et al. 1987). Kamafugite lavas from East Africa and other localities often contain micaclinopyroxenite and other phlogopite-bearing xenoliths believed to represent the samples of the highly metasomatized mantle source. This suggests that phlogopite plays a fundamental role for the origin and evolution of kamafugites.

The larnite-normative trend which is observed for the liquid in the F-doped system may help to explain the most distinctive chemical features of kamafugite melts: the SiO₂ depletion and the CaO enrichment. Liquidus phase equilibria in the model system suggest that partial melting of compositions in the vicinity of the Phl – Di join in the presence of stable phlogopite will produce silica-undersaturated larnite-normative melts. Melting experiments on a representative mantle-derived phlogopite-clinopyroxenite nodule composition reported by Lloyd et al. (1985) show that the larnite-normative trend is real for multicomponent water-bearing compositions. All partial melts produced in these experiments at 20 and 30 kbar in equilibrium with high-Ca clinopyroxene, phlogopite, ilmenite, apatite, magnetite and forsterite are larnite-normative. A series of isobaric runs at 30 kbar show that the CaO content in melts in equilibrium with phlogopite and clinopyroxene increase, while SiO_2 and K_2O concentrations decrease with falling temperature and decreasing degrees of partial melting. This is consistent with the observations in the model system and indicates that larnite-normative partial melts can be buffered by the residual clinopyroxene-phlogopite assemblage in the mantle source.

Natural glasses in upper mantle xenoliths from the volcanics of Gees, West Eifel (Edgar et al. 1989) provide further evidence for the genetic link between phlogopite stability in the mantle source and generation of larnite-normative kamafugite melts. Edgar et al. (1989) studied six xenoliths ranging from harzburgite to mica clino-pyroxenite. Glasses in association with phlogopite are larnite normative and differ from those surrounding and enclosed by spinel. All phlogopite related glasses show a strong negative correlation between SiO₂ and CaO similar to the phlogopite-saturated glasses in the studied F-doped system.

On the other hand, olivine and augite minettes studied by Esperança and Holloway (1987) and Righter and Carmichael (1996) provide examples of compositions with olivine, clinopyroxene and phlogopite on liquidus, that do not follow the larnite-normative evolution. The bulk compositions are more silica rich, Ca poor and orthoclase normative and plot below the fo-di-phl join in the kalsilite-based normative tetrahedron (Fig. 1b). Crystallisation of the ol-cpx-phl association from these compositions results in the increase of SiO₂ and decrease of CaO contents in the residual melts in agreement with the conclusions presented in the previous section.

Origin of the plutonic melilite-rich rocks

Phlogopite is abundant not only in potassic mafic silicaundersaturated systems, but also in more common sodium-dominated alkaline associations. Phlogopite accumulations in some ultramafic alkaline intrusions are even subjected to mining, e.g. Kovdor, Kola Peninsula. The experimental results and especially the development of residual melts in phlogopite-bearing systems (Fig. 8) imply that late-stage melilitolites of ultramafic alkaline complexes are low-pressure products of fractional crystallisation of phlogopite-bearing assemblages from a melanephelinitic parental melt. Other hydrous minerals, such as amphiboles, may play a role similar to that of phlogopite and also lead to increasing normative larnite.

The absence of the larnite-normative melt evolution in dry systems and the critical role of hydrous phases may also be illustrated by the experimental study of Gee and Sack (1988). In their experiments with natural melilitite lava compositions no increase in normative larnite in residual melts was observed despite larnitenormative starting compositions. Liquidus assemblages in the experiments of Gee and Sack (1988), however, include leucite, nepheline, clinopyroxene, spinel \pm melilite \pm olivine, and no phlogopite, which in the present study is shown to be the determining phase for the development towards highly larnite normative residuals in response to the increased volatile pressure.

In conclusion, the development of late-stage melilitolitic melts in ultramafic-alkaline complexes (e.g. Kovdor and Gardiner) is dependent on volatile content and the stability of phlogopite. Closed volatile-rich conditions during fractionation will favour a melilitic and possibly also a carbonatitic evolutionary trend as suggested by Nielsen (1994), whereas degassed and anhydrous conditions would favour developments towards more CaO-poor phonolitic residuals.

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Appendix

Abbreviations for phases:

ak akermanite cu cuspidine di diopside en enstatite fl fluorite flu unidentified fluoride phase, presumably K₃AlF₆ fo forsterite hu humite ks kalsilite la larnite lc leucite lf fluoride melt l_s silicate melt mo monticellite phl phlogopite sa sanidine qz quartz sp spinel wo wollastonite

Formulae for components:

Ak $Ca_2MgSi_2O_7$ An $CaAl_2Si_2O_8$ Di $CaMgSi_2O_6$ Fo Mg_2SiO_4 Fl CaF_2 KF KAISiO_3F_2 KS $K_2Si_2O_5$ La Ca_2SiO_4 Lc KAISi_2O_6 Or KAISi_3O_8 PhI KMg_3AISi_3O_{10}F_2 Wo CaSiO_3

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