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Crystal-melt equilibria involving potassium-bearing clinopyroxene as indicator of mantle-derived ultrahigh-potassic liquids: an analytical review

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Abstract

Crystal-liquid equilibria, including phase relationships of minerals with silicate and/or carbonate melts, are reviewed in order to understand the occurrence of clinopyroxene with up to 2 wt.% K₂O (KCpx). This mineral occurs as inclusions in diamond from kimberlite pipes and in garnet from garnet-clinopyroxene potassium-poor silicate rocks intercalated with diamondiferous silicate-carbonate rocks of the Kokchetav Complex, northern Kazakhstan. The analysis of the available experimental data allowed estimation of the effect of P, T and compositional parameters on the equilibrium of KAlSi₂O₆ (in Cpx) = 1/4K₄Si₂O₆ + 3/4Al_{4/3}Si₂O₆ (in *melt*) in various silicate systems. A strong dependence of the K₂O partition coefficient both on pressure and SiO₂ and Al₂O₃ contents in the melt was identified and thermodynamically described. The resulting thermodynamic equation allows the calculation of pressure of the KCpx formation within interval 50–100 kbar for known melt compositions. A model for the formation of KCpx in deep-mantle potassium-rich carbonate-silicate magmas was derived using the available experimental and petrologic data. The formation of KCpx from any potassium-poor (K₂O < 0.2 wt.%) melt is impossible because the partition coefficient, $K_p^{Cpx/L} = K_2O$ in Cpx/K_2O of melt is much smaller then 1. The presence of Cpxwith 1 wt.% of K₂O as inclusions in Grt from potassium-poor rocks could be explained only by the crystallization of KCpx from potassium-rich silicate or silicate-carbonate magma in the presence of KCl brine at P > 70 kbar. This chemical zoning of potassium-bearing clinopyroxene suggests its crystallization together with feldspar from silicate melt during rapid ascent of melt toward the Earth's surface from depth about 200 km. Since Tschermak-type substitution has not been observed in either pyroxene, we suggested that feldspar resulted from the peritectic reaction $KAlSi_2O_6+[SiO_2]_{L/f}=KAlSi_3O_8$ at relatively shallow mantle levels. A hypothetical phase diagram for the system Ca(Mg,Fe)Si₂O₆-KAlSi₂O₆ is presented. © 2002 Elsevier Science B.V. All rights reserved.

Keywords: Mineral equilibria; Potassium-bearing clinopyroxene; Inclusions; Mantle; Melting relations

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1. Introduction

A possible way to understand the sources of potassium in the Earth's mantle is to consider the origin of potassium-rich rocks, such as fergusites, leucite basalts, leucitites, etc., as well as kimberlites and lamproites. Kushiro (1980) showed that an increase of pressure causes an appreciable displacement of the cotectic toward kalsilite in the system Mg₂SiO₄-KAl-SiO₄-SiO₂. That should assist the formation of potassium-rich magmas at mantle pressure. Addition of water into the system results in a decrease of the liquidus temperature, but does not change the effect of pressure on expansion of the primary crystallization fields of olivine and orthopyroxene at the expense of kalsilite (Perchuk, 1987). Thus, pressure (P) is the most important parameter that regulates the formation of highly potassic magmas. Such magmas are characteristic for rifts (e.g., eastern Africa, the Rhein trough, and northern Baikal) and occur locally in some fold belts (e.g., the Pamirs) and subduction zones (northern Indonesia and Italy). However, ultra-high pressure minerals such as diamond, coesite, potassium-bearing clinopyroxene, etc., are not found as phenocrysts in these rocks. That implies that the generation of the initial magmas occurred in depths less than 100 km. According to the data on melt inclusions (Kogarko et al., 1991; Solovova et al., 1996; Nielsen et al., 1997), such magmas originate in the upper mantle, but beyond the diamond stability field (Wyllie and Lee, 1998).

Microinclusions of potassium-bearing clinopyroxene (KCpx) and fluids are common in diamonds from kimberlite pipes (e.g., Reid et al., 1976; Bishop et al., 1978; Navon et al., 1988; Harlow and Veblen, 1991; Israeli et al., in press). The presence of such inclusions created no theoretical problems until the data on inclusions of KCpx from the extremely potassium-poor silicate and calc-silicate rocks of the Kokchetav Complex, northern Kazakhstan, appeared (Sobolev and Shatsky, 1990; Perchuk et al., 1995, 1996). These problems are as follows. Firstly, according to experimental data, *KCpx* is stable at $P \gg 40$ kbar and 1200-1500 °C (Table 1). Therefore, its occurrence in crustal metamorphic environments is hard to explain. Secondly, the K₂O partitioning coefficient between KCpx and a melt of any composition is always less then 1.0. Thus, Cpx cannot be richer in K₂O than the rocks in which it occurs, unless it is residue of some primary K-rich melt, subsequently removed from the rocks. Thus, the occurrence of KCpx implies unusual circumstances. Perchuk and Yapaskurt (1998) proposed that at mantle

Table 1 Run conditions for the synthesis of potassium-bearing clinopyroxenes in the model and natural systems

Reference	<i>T</i> (°C)	P (kbar)	System	Na2O (wt.%)	K ₂ O (wt.%)		
				System before run	System before run	Cpx ^a	
Alumino-silicate systems							
Erlank and Kushiro (1970)	1000 - 1450	15-30.5	Di-Phl, Kfs-An ₅₀ Fo ₅₀	0	?	0.05 - 0.14	
Shimizu (1971)	1400, 1350	38 - 100	$Di_{50}Fo_{13.5} + An_{25.5}Phl_{10}$	0	1.2	0.42 - 2.2	
Shimizu (1974)	1100 - 1200	15 - 30	a model system $Di + An + K_2O$	0	2.2	< 0.01	
Luth (1992)	1200-1500	50-110	model systems $Di + KAlSi_2O_6$ and $Di + KCrSi_2O_6 \pm H_2O$?	?	0.13-2.0	
Edgar and Vukadinovic (1993)	1250-1500	50 - 60	armalcolite-phlogopite lamproite	0.96	8.84	0.12 - 1.67	
Mitchell (1995)	1000 - 1700	40 - 70	sanidine-phlogopite lamproite	1.25	11.59	0.34 - 1.86	
Luth (1995, 1997)	1250-1600	30-170	a model system Phl-Di	0	5.6	0.13 - 1.2	
Edgar and Mitchell (1997)	900-1200	70 - 120	armalcolite-phlogopite lamproite	0.96	8.84	0.3 - 2.59	
Okamoto and Maruyama (1998)	900-1200	26 - 70	eclogite + Ms^{b}	?	?	0.0 - 0.23	
Tsuruta and Takahashi (1998)	1100-2050	10 - 125	high-K basalt	2.85	1.46	0.01 - 1.03	
Wang and Takahashi (1999)	1400-2350	25-250	high-K basalt	2.64	1.31	0.0 - 1.9	
Carbonate-silicate systems							
Harlow (1997)	1300 - 1700	50-140	a model system $Cpx + K_2CO_3$ (KHCO ₃)	?	?	1.09 - 4.51	
Matveev et al. (1998)	1000 - 1600	70	a model system $K_2Mg(CO_3)_2 + Di + Grs;$	0	?	0.56-4.83	

?-data unknown.

^a After run.

^b Ms/rock ratio in the starting material was not given.

conditions *KCpx* could crystallize at liquidus of a potassium-rich carbonate–silicate liquid. The miscibility of the carbonate and silicate liquids decreases as pressure drops (e.g., Lee and Wyllie, 1997), while crystallizing *KCpx* could be captured by any latter mineral, for example garnet.

In this paper, all available experimental and natural data are reviewed in order to:

- 1. find general dependencies of potassium substitution in *Cpx* on the external parameters and the melt composition;
- deduce an equation for the dependence of the potassium partition coefficient between silicate melt and *Cpx*, and to calculate *P* and *T* for *KCpx* formation in mantle conditions;
- preliminarily calculate the high-temperature phase relations of in system CaMgSi₂O₆-KAlSi₂O₆;
- evaluate the possibility of *KCpx* crystallization from potassium-rich mantle liquids using the compositions of melt, fluid, and brine inclusions in diamonds from kimberlitic pipes;
- explore the possibility of the peritectic reaction, producing the clinopyroxene-K-feldspar aggregates in *Cpx* grains from quartz-free rocks of the Kokchetav Complex (Perchuk et al., 1996) and some diamondiferous eclogites from South Africa (Reid et al., 1976).

2. A review of the existing experimental data on the equilibria involving *KCpx*

Because of the structure of Cpx, the coupled substitution of large potassium cation into the M2 site, (Mg, Fe)^{M1}Ca^{M2} \Leftrightarrow Al^{M1}K^{M2}, is impossible at low pressure (Harlow, 1996). Therefore, all experimental studies on the *KCpx* synthesis were performed at very high pressures (see Table 1). The experimental studies cover a wide spectrum of system compositions, from alumino-silicate (Erlank and Kushiro, 1970; Shimizu, 1971, 1974; Edgar and Vukadinovic, 1993; Mitchell, 1995; Edgar and Mitchell, 1997; Luth, 1992, 1995, 1997; Okamoto and Maruyama, 1998; Tsuruta and Takahashi, 1998; Wang and Takahashi, 1999) to carbonate–silicate systems (Harlow, 1997; Matveev et al., 1998).

2.1. Alumino-silicate systems

Experiments on the crystallization of KCpx in alumino-silicate systems have been conducted in both model and natural systems. Erlank and Kushiro (1970) attempted to synthesize KCpx from the mixtures of *APhl* or *Kfs* with diopside or $An_{50}Fo_{50}$ at 1000–1450 °C and 15–30.5 kbar. The maximal K₂O content in diopside coexisting with *Fo* and glass did not exceed 0.14 wt.% at 21 kbar and 1250 °C. At 70 kbar, *KCpx* was absent in the run products.

Shimizu (1971) studied the pressure dependence of the K_2O content in Cpx in the sub-solidus system *Di*₅₀*Fo*_{13.5}*An*_{26.5}*APhl*₁₀ at 38 kbar (1350 °C), 72 kbar (1350 °C) and 100 kbar (1400 °C). The maximal K_2O in Cpx was 2.2 wt.% at 100 kbar and 1400 °C. Later, Shimizu (1974) studied the partitioning of potassium and some minor elements (Sr, Ba, Rb, Cs) between Cpx and melt in the system Di-An-Ab at 15-30 kbar and 1100-1200 °C. The concentrations of K, Sr, Ba, Rb, Cs in the system were similar to those of high-K basalts (for example, 2.2 wt.% K₂O). Despite the low K₂O concentrations (maximum is 0.012 K₂O wt.% at 25 kbar and 1100 °C), the partition coefficient of K_2O , $K_{\rm p}$, between KCpx and $L_{\rm Si}$ steadily increased from 0.0014 at 15 kbar to 0.0026 at 30 kbar. Shimizu's (1971, 1974) experiments clearly demonstrated that K_2O becomes more compatible in *Cpx* with increasing pressure.

In order to determine a maximal solubility of potassium in clinopyroxene, Luth (1992) carried out experiments in quasi-binary joins CaMgSi₂O₆-KAl-Si₂O₆ and CaMgSi₂O₆-KCrSi₂O₆ at 50-110 kbar and 1200-1500 °C. Following Shimizu's (1971, 1974) conclusions, Luth (1992) found that in the first system concentration of K₂O in Cpx increased from 0.13 wt.% at 50 kbar to 0.61 wt.% at 110 kbar at 1300 °C. In the later paper, Luth (1995) reported even 3 wt.% of K₂O in clinopyroxene in the system CaMg-Si₂O₆-KAlSi₂O₆. However, author did not mention conditions of the synthesis of such highly potassic clinopyroxene. In addition, Luth (1992) experimentally confirmed the effectiveness of Cr in stabilization of K in the clinopyroxene structure that was previously described by Harlow and Veblen (1991).

In order to study the relative stability of *Phl* and *Cpx*, Luth (1995, 1997) studied the system *Phl–Di* at 30-170 kbar and 1250-1600 °C. According to his

data, the assemblage $Cpx-Grt-Ol-L_{\rm Si}$ is stable from the liquidus to the solidus of the system within the *P* range 60–150 kbar. At lower *P*, this assemblage is replaced by $Cpx-Ol-L_{\rm Si}$. The maximal K₂O content in Cpx (1.3 wt.%) was obtained at 170 kbar and 1600 °C, but at $P \le 50$ kbar the K₂O content in Cpx did not exceed 0.3 wt.%. The $K_{\rm p}^{Cpx/L}$ increases systematically with *P*, whereas a *T* dependence of the $K_{\rm p}^{Cpx/L}$ was not observed.

Most experimental studies on the KCpx synthesis in natural systems were conducted using potassium-rich mantle rocks such as lamproites. Edgar and Vukadinovic (1993) investigated the conditions of the KCpx crystallization in a melt of armalcolite-phlogopite lamproite from Smoky Butte. The experiments were carried out at 50 and 60 kbar in the T interval 1250– 1500 °C. All run products contained glass. Taking into account earlier experiments of Edgar et al. (1992), Edgar and Vukadinovic (1993) constructed a phase diagram including melting relations in the intervals 10–60 kbar and 900–1500 °C. They found that Cpxwith the highest K_2O content (1.58–1.67 wt.%) formed at 1250-1200 °C and 60 and 50 kbar in the assemblage $Grt + Phl + Ru + Cos + L_{Si}$. With increasing temperature, Phl disappears first from the assemblage, being followed by Cos. Cpx was the only liquidus phase at 50 kbar and 1400 °C. At 60 kbar and 1500 °C, Grt and Cpx were both stable on the liquidus. Grt disappeared below 50-45 kbar, being replaced by Cpx + Opx and Cpx + Phl. Edgar and Vukadinovic (1993) showed that at high K₂O content in the system and P > 50 kbar, relatively low T(< 1300)°C) was most favorable for the crystallization of KCpx. However, Harlow (1997) called attention to the dependence of the K₂O concentration in Cpx from experiments of Edgar and Vukadinovic (1993) on its enstatite-ferrosilite content, which is negatively correlated with temperature. According to Harlow (1997), an increase of the enstatite-ferrosilite content in KCpx causes a decrease in the molar volume of Cpx, and this is the only reason for the T dependence of the K_2O content in Cpx in the experiments of Edgar and Vukadinovic (1993). Later, Edgar and Mitchell (1997) studied the same system at 70-120 kbar and observed no dependence of K_2O content in Cpx on temperature. Edgar and Mitchell (1997) found that KCpx was stable with garnet at the liquidus within the whole P interval. Opx and Ol appeared only at P < 50 kbar. The K₂O content of *KCpx* varied from 0.3 wt.% at 1500 °C and 80 kbar to 2.58 wt.% at 1500 °C and 120 kbar. Edgar and Mitchell (1997) found that the K₂O content of clinopyroxene decreased in presence of K-bearing phases (phlogopite, K–Ti silicates, K–Ba phosphates) near the solidus (1100–1300 °C).

Mitchell (1995) used sanidine-phlogopite lamproite from Leucite Hills for his experiments at 40-70 kbar and 1000-1700 °C, and found that both Grt and Cpx were present at the liquidus at 1600 °C and above 45 kbar. Coesite, K-Ti silicate, and K-Ba phosphate joined the assemblage below 1400-1500 °C, while *Phl* and K-richterite were stable at 1100 °C. KCpx was the only liquidus phase at P < 45 kbar. The K_2O content in Cpx increased with P reaching 1.86 wt.% at 70 kbar and 1400 °C. A dependence of the K₂O content in Cpx on temperature was not observed. Sanidine was found at the solidus at P < 55 kbar and T < 1250 °C. Its formation was accompanied by a significant decrease in the K₂O content in Cpx. For example, at 1200 °C, the K₂O content in Cpx decreased from 1.03 wt.% at 60 kbar to 0.55 wt.% at 50 kbar.

Tsuruta and Takahashi (1998) and Wang and Takahashi (1999) studied the melting relations of anhydrous alkalic basalt (JB1, $K_2O = 1.31$ wt.% and $Na_2O = 2.64$ wt.%) at pressure up to 200 kbar and 1100–2400 °C. They showed that KCpx is the liquidus phase at 10-70 kbar. At P>70 kbar, Cpx has been replaced by Grt. The following crystallization sequence was observed with decreasing pressure from 10 to 70 kbar: (1) Cpx; (2) Grt (P < 40 kbar) or coesite (P > 40 kbar); (3) San at temperature 1200–1400 °C and P < 60 kbar; (4) the assemblage Cpx + Grt + San + Cpx + Grt + GrCos + Rt at 1000–1200 °C. An inflection in the solidus within 60-75 kbar was documented. The slope of the solidus line changes from 5°/kbar to 40°/kbar, and subsequently flattens at P above 100 kbar to almost 2°/kbar. This behavior resulted from the appearance of San which is not stable at P > 60 kbar. The K₂O content in Cpx increased over 1 wt.% in the P interval studied. The K₂O content in Cpx was observed to distinctly decrease in presence of melt. KCpx with up to 1.03 wt.% K₂O was found to be stable when it was the only potassium-bearing phase. A temperature dependence of the K_2O content in Cpx was not observed as well (Tsuruta and Takahashi, 1998).

Okamoto and Maruyama (1998) analyzed KCpx at the eclogite solidus obtained for two different samples from the Dabie Shan Complex (Eastern China). Muscovite was added to one sample, and its K₂O content was 9.32 wt.%. The K₂O content of KCpx increased at 900 °C from 0.07 wt.% at 30 kbar to 0.23 wt.% at 70 kbar. A similar pattern was observed at 1000 °C. A minor increase of the K₂O content in Cpx with temperature was reported.

Konzett and Ulmer (1999) experimentally explored the stability of hydrous potassic phases (phlogopite and K-richterite) at 40–95 kbar and 800–1400 °C in both the analogue system KNCMASH and in natural lherzolite with addition of 5 wt.% of phlogopite or 10 wt.% of K-richterite. Konzett and Ulmer (1999) found that *Cpx* was stable within the all P-T interval studied, but the K₂O content in *Cpx* depended critically on the phase assemblage. The K₂O content abruptly decreased to ~ 0.2 wt.% in the presence of phlogopite and to 0.45 wt.% in the presence of K-richterite and/or melt. The K₂O content in *Cpx* from pure lherzolite without any additions of K₂O did not exceed 0.05–0.1 wt.%. The composition of *KCpx* at given *P* and *T* depended only on the bulk K₂O content.

2.2. Carbonate-silicate systems

Only two experimental studies are known on the KCpx crystallization in carbonate-silicate systems. Harlow (1997) investigated the partitioning of K_2O between Cpx and coexisting carbonate melt in the system CaMgSi₂O₆-NaAlSi₂O₆-NaCrSi₂O₆-K₂CO₃-KHCO₃ at 50–110 kbar and 1300–1700 $^{\circ}$ C, with addition of small amount of quartz and enstatite to it. The choice of such a complex system was based on the results of a study of chromium-bearing KCpx inclusions in diamond from some kimberlite pipes of South Africa and Botswana (Harlow and Veblen, 1991). The maximal K₂O contents in Cpx, 4.39 and 4.51 wt.%, were observed at pressure 100 kbar for the system Di₅₀Ko₅₀-K₂CO₃. As mentioned above, the presence of chromium assisted to the potassium substitution in Cpx. Under the same P-T conditions, Crfree KCpx was two to three times poorer in K_2O . An X-ray examination study in Cpx in the run product showed a dependence of its molar volume on the K₂O content. A similar dependence has been previously found for natural Cpx (Harlow and Veblen, 1991).

This dependence explains the high K₂O content in the Cr-bearing *Cpx* in comparison to the Cr-free *Cpx*, as well as the influence of sodium on the potassium solubility in *Cpx*. This study also confirmed the dependence of K_p on *P*, while the *T* dependence was not observed. The most important result of Harlow's study is the confirmation of the low K_p in the system $KCpx-L_c$, which varies from 0.03 to 0.071. In other words, *Cpx* with K₂O > 1 wt.% can crystallize in equilibrium with the carbonate liquid containing 14–33 wt.% K₂O at any given *T* and *P* within 1300–1700 °C and 50–110 kbar intervals.

Matveev et al. (1998) investigated the stability of KCpx at 70 kbar and 1500 and 1600 °C on the liquidus of the carbonate-silicate system K₂Mg(CO₃)₂-K₂CO₃-*Di*-*Grs*, i.e., a synthetic analogue system of the garnet-clinopyroxene-carbonate diamondiferous rocks of the Kokchetav Complex. Beside *KCpx*, the run products contained pyrope-grossular *Grt*, dolomite, and homogeneous silicate glass. The maximal K₂O content in *Cpx* was 4.83 wt.%, i.e., the highest K₂O concentration documented so far.

3. Discussion of the observed potassium solubility in *Cpx*

Although the experiments reviewed above cover a wide range of P and T, most of the starting compo-

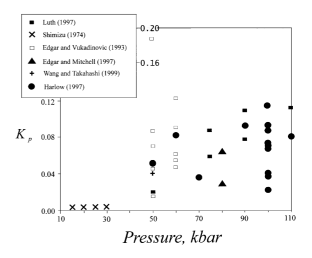


Fig. 1. The variation of the K_2O partition coefficient (K_p) between clinopyroxene and melt according to available experimental data.

sitions did not include simple binary systems, which would allow to constrain specific P-X or T-X phase diagrams involving *KCpx*. The data by Luth (1992) on

the join CaMgSi₂O₆-KAlSi₂O₆ are very confined. This limits the applicability of the data for a rigorous analysis of the potassium solubility in *Cpx*. Never-

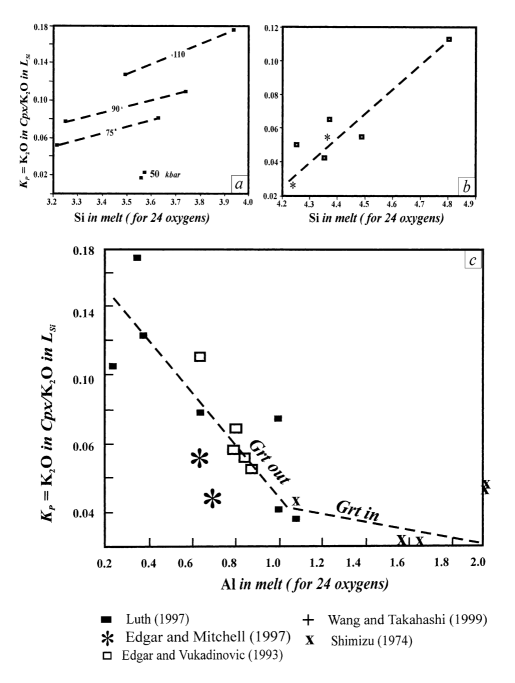


Fig. 2. The variation of the K_2O partition coefficient (K_p) between clinopyroxene and melt with SiO₂ and Al₂O₃ in the silicate melt according to available experimental data (see text for more details).

theless, a semi-quantitative evaluation of the different parameters controlling the *KCpx*-melt equilibrium is possible.

(1) The K partitioning between Cpx and melt is P dependent in any bulk composition. However, it is impossible to calculate the dependence of $K_p^{Cpx/L}$ on P even approximately because of the appreciable scatter in the available data (Fig. 1). Nevertheless, this scatter likely results from the very strong influence of the melt and clinopyroxene compositions on $K_p^{Cpx/L}$.

(2) $K_p^{C_{px/L}} \ll 1.0$ for any composition within a wide range of *P* and *T*.

(3) The nature of the *T* dependence of K_2O content in *Cpx* is ambiguous (Mitchell, 1995; Luth, 1997; Edgar and Mitchell, 1997; Tsuruta and Takahashi, 1998).

(4) Existing experimental data indicate that the clinopyroxene from carbonate-silicate melts is richer in K_2O when compared with the clinopyroxene from alumino-silicate melts. Even at $P \approx 200$ kbar, the K_2O concentration in *Cpx* crystallizing from alumino-silicate melts never exceeds 3 wt.% while the K_2O content in *Cpx* above 4 wt.% is observed in carbonate-silicate melts (Harlow, 1997; Matveev et al., 1998). Nevertheless, new experimental studies may change this conclusion.

(5) *KCpx* either does not crystallize or its K₂O content is extremely low in the K₂O-poor aluminosilicate systems (K₂O < 1 wt.%; Okamoto and Maruyama, 1998; Konzett and Ulmer, 1999). In contrast, *KCpx* with about 0.5 wt.% may crystallize even at significantly lower *P* in the relatively K₂O-rich systems, such as lamproites (Edgar and Vukadinovic, 1993; Mitchell, 1995; Edgar and Mitchell, 1997).

(6) The K_2O contents in *Cpx* depend both on the bulk composition and the mineral assemblage of a system, e.g., K_2O in *Cpx* abruptly decreases in the presence of K-bearing phases (e.g., Luth, 1995, 1997; Tsuruta and Takahashi, 1998).

(7) $K_p^{Cpx/L}$ depends directly on the SiO₂ (Fig. 2a and b) and inversely on the Al₂O₃ contents (Fig. 2c) of the alumino-silicate melt.

(8) The reciprocal substitution $(Mg,Fe)_{M1}Ca_{M2} \Leftrightarrow (Al,Cr,Fe^{3+})_{M1}K_{M2}$ is the most efficient scheme for potassium incorporation into Cpx (Luth, 1992; Edgar and Vukadinovic, 1993; Harlow, 1997).

(9) The potassium solubility in Cpx is controlled by its composition, which in turn optimizes the molar

volume (Harlow, 1997). Cr-bearing diopside is the most favored, while omphacite is least favored for potassium (Harlow and Veblen, 1991; Luth, 1992; Harlow, 1997).

4. Thermodynamic treatment of the available experimental data for the *KCpx* equilibria in alumino-silicate systems

The available experimental data show that the K_2O content in *Cpx* is very low to produce correct thermodynamic model for the *KCpx* solid solution. However, existing experimental data allow the derivation of a simplified thermodynamic model for the system CaMgSi₂O₆-KAlSi₂O₆ for low K₂O concentrations.

Let us consider the equilibrium of *KCpx* with the alumino-silicate melt (L_{Si}) by taking into account the dependency of $K_p^{Cpx/L}$ on *P* and the SiO₂ content in the melt (Figs. 1 and 2). This equilibrium can be expressed as:

$$\begin{aligned} \text{KAlSi}_2\text{O}_6(KJd \ in \ CPx) \\ &= 1/4\text{K}_4\text{Si}_2\text{O}_6 + 3/4\text{Al}_{4/3}\text{Si}_2\text{O}_6(in \ L_{Si}), \end{aligned} \tag{1}$$

The Gibbs free energy of this reaction is

$$\Delta G_{(1)}^{0} = \Delta H_{(1)}^{0} - T \Delta S_{(1)}^{0} + P \Delta V_{(1)}^{0} + RT \ln[(a_{K}^{L})^{1/4} (a_{AI}^{L})^{3/4} / a_{KJd}^{Cpx}]$$
(2)

where $a_{\rm K}^{\ L}$ and $a_{\rm Al}^{\ L}$ -activities of the K₄Si₂O₆ and Al_{4/} ₃Si₂O₆ components of the silicate melt, respectively; $L=L_{\rm Si}$ and $a_{KJd}^{\ Cpx}$ is activity of the KAlSi₂O₆ component in the clinopyroxene solid solution; $\Delta G_{(1)}^0$, $\Delta H_{(1)}^0$, $\Delta S_{(1)}^0$ and $\Delta V_{(1)}^0$ -free energy, enthalpy, entropy and volume differences for reaction (1).

Using the multi-site solution model, the mixing energy of the $K^{M2}Al^{M1}Si_2{}^{T}O_6$ (*KJd*) component in clinopyroxene is expressed as

$$RT\ln(a_{KJd}^{Cpx}) = RT(a_{KJd}^{id}) + G_{KJd}^{e}$$
$$= RT[X_{K}^{M2}X_{Al}^{M1}(X_{Si}^{T})^{2}] + G_{KJd}^{e}, \qquad (3a)$$

where X_i are mole fractions of components in corresponding sites and G^{e}_{KJd} — an excess free energy of the

No.	No. Run No.		P (kbar)	Mineral assemblage	Compositional parameters							References
					KCpx				Melt			
					K ^{M2}	Ca ^{M2}	Na ^{M2}	Al^{M1}	X _K	X_{Si}	X _{Al}	
1	1106	1600	110	Ol+Grt+Cpx+X*	0.037	0.811	0	0.050	0.090	0.449	0.047	1
2	1163	1600	110	Ol + Grt + Cpx	0.046	0.804	0	0.053	0.065	0.504	0.044	1
3	1211	1550	90	Cpx + Ol + Grt	0.050	0.831	0	0.042	0.115	0.480	0.026	1
4	1212	1500	90	Cpx + Ol + Grt	0.037	0.846	0	0.041	0.118	0.419	0.129	1
5	1213	1550	75	Cpx + Ol + Grt	0.027	0.803	0	0.046	0.084	0.466	0.081	1
6	1208	1450	75	Cpx + Ol + Grt	0.027	0.862	0	0.034	0.131	0.414	0.118	1
7	1178	1500	50	Cpx + Ol	0.009	0.844	0	0.049	0.130	0.457	0.139	1
8	1179	1400	50	Cpx + Ol	0.011	0.834	0	0.059	0.121	0.459	0.128	1
9	_	1400	50	Cpx + Grt + Cos	0.014	0.446	0.337	0.570	0.085	0.619	0.139	2
10	_	1300	80	$Grt + Cpx + Cos + Rt + X^*$	0.046	0.513	0.214	0.244	0.207	0.545	0.080	3
11	_	1300	60	Cpx + Grt + Cos + Phl + Ru	0.062	0.531	0.017	0.308	0.138	0.600	0.083	4
12	_	1400	60	Cpx + Grt + Cos	0.0317	0.475	0.199	0.239	0.145	0.561	0.102	4
13	-	1500	60	Cpx + Grt	0.0278	0.452	0.191	0.246	0.138	0.531	0.105	4
14	_	1350	50	Cpx+Grt	0.0203	0.422	0.141	0.195	0.119	0.544	0.109	4
15	_	1300	50	Cpx + Grt + Cos	0.0277	0.403	0.146	0.197	0.106	0.546	0.109	4
16	_	1100	15	Ċpx	0.0004	0.937	0.024	0.066	0.070	0.522	0.207	5
17	_	1100	20	Cpx	0.0004	0.904	0.042	0.132	0.053	0.535	0.216	5
18	_	1100	25	Ċpx	0.0005	0.873	0.071	0.191	0.072	0.549	0.221	5
19	_	1200	25	Cpx	0.0004	0.894	0.046	0.105	0.071	0.559	0.232	5
20	P194	1200	30	Cpx + Grt + Cos	0.0005	0.532	0.333	0.509	0.035	0.679	0.175	6
21	S222	1400	50	Cpx + Grt + Cos + Kfs	0.011	0.513	0.329	0.522	0.036	0.696	0.179	6
22	S303	1900	100	Cpx+Grt	0.009	0.378	0.453	0.575	0.016	0.527	0.138	6
23	S257	1900	75	Cpx+Grt	0.009	0.411	0.344	0.576	0.021	0.522	0.150	6
24	S270	1700	60	Cpx+Grt	0.003	0.422	0.270	0.555	0.030	0.535	0.152	6
25	S228	1700	50	Cpx	0.002	0.478	0.238	0.455	0.020	0.500	0.158	6
26	P182	1500	30	Cpx	0.002	0.530	0.173	0.343	0.022	0.531	0.164	6

Table 2 Available experimental data on compositions of KCpx and the alumino-silicate melt used in the regression

References: (1) Luth (1997), (2) Wang and Takahashi (1999), (3) Edgar and Mitchell (1997), (4) Edgar and Vukadinovic (1993), (5) Shimizu (1974), (6) Tsuruta and Takahashi (1998).

X*-diverse K-bearing phases (Luth, 1995, 1997; Edgar and Mitchell, 1997).

KJd component in the clinopyroxene solid solution. G_{KJd}^{e} reflects a very limited solubility of *KJd* in clinopyroxene that corresponds to the strong non-ideality of the solid solution. In addition to the *Di*, *Hed*, *Jd* and *KJd* end-members, synthetic clinopyroxenes contain a significant amount of the *En-Fs* component (up to 20–30 mol%). Regarding the Mg-Fe-Ca-Na-K mixing in the M2 site, we tentatively used the fivemember regular model for G_{Cpx}^{e} to derive G_{KJd}^{e}

$$G_{Cpx}^{e} = \sum_{i=1}^{c-1} \sum_{j \neq i}^{c} W_{ij} X_i X_j$$
(3b)

where c = 5; *i*, *j*-components in the M2 site (Mg, Fe, Ca, Na, K), and W_{ij} -interaction parameters, which

could be computed by the regression of the experimental data.

Using similar approach, the mixing energy of the $K_4Si_2O_6$ component in the alumino-silicate melt is expressed as

$$RT\ln(a_{\rm K}^L) = RT\left[\left(X_{\rm K}^L\right)^4\right] + \left(G_{\rm K}^{\rm e}\right)^L \tag{3c}$$

and for the $Al_{4/3}Si_2O_6$ component

$$RT\ln(a_{\rm Al}^L) = RT\left[\left(X_{\rm Al}^L\right)^{4/3}\right] + \left(G_{\rm Al}^{\rm e}\right)^L \tag{3d}$$

Thus,

$$RT\ln\left[\left(a_{\rm K}^{L}\right)^{1/4}\left(a_{\rm Al}^{L}\right)^{3/4}\right) = RT\ln\left[\left(X_{\rm K}^{L}\right)\left(X_{\rm Al}^{L}\right)\right] + \left(G^{\rm e}\right)^{L}$$
(3e)

where X_i^L is mole fraction of *i*-component in the melt and $(G^e)^L$ —a total excess free energy of the melt species, whose expression and parameters could be fitted by regression of the experimental data. The mole fraction of a component *i* in the melt is $X_i^L = i/$ (Si + Ti + Al + Cr + Fe + Mn + Mg + Ca + Na + K), where symbols of elements denote their quantities in the melt formula per six oxygens. This approach allows the treatment of a wide range of the melt compositions (Table 2).

As a result, the best fit for 26 experimental data points (Table 2) at the least square error $r^2 = 0.934$ was obtained with the following values of Eq. (2):

 $\Delta H^0_{(1)} = -24317.3(\pm 6937) \text{ J/mol},$ $\Delta V^0_{(1)} = 0.219(\pm 0.106) \text{ J/mol/bar},$ $W_{\text{Si}} = 1.716(\pm 0.5) \text{ J/mol/bar},$

 $W_{\rm KCa} = 63881.44(\pm 9495) \text{ J/mol},$

Eq. (2) reproduces experimental pressure values and $X_{\rm K}^{Cpx}$ within ± 9.1 kbar and ± 0.012 , respectively, in the intervals T=1100-1900 °C and P=15110 kbar. The equation also illustrates the dependence of reaction (1) on *pressure*, *melt composition*, *and Cpx composition*. The positive value $\Delta V_{(1)}^{0} =$

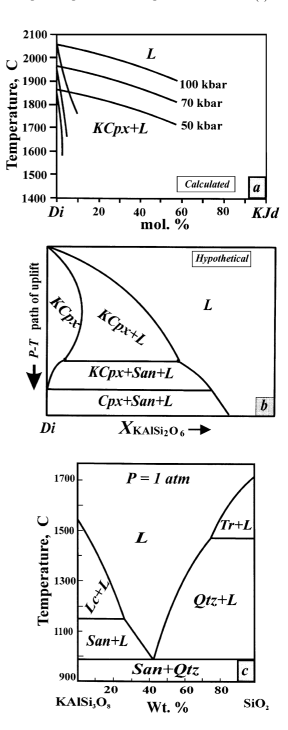


Fig. 3. Calculated (a) and hypothetical (b) diagrams for melting in the system Di-KJd and their comparison with the diagrams of melting in the systems KAISi₃O₈–SiO₂ (diagram c, after Schairer and Bowen, 1955). The diagram (a) was calculated using Eq. (2) and data by Gasparik (1996) on diopside melting. Diagram (b) reflects change in phase relations with decrease of temperature taking into account that *P* and *T* form an uplift path (pseudo-isobaric diagram) and illustrates the possible existence of a peritectic point, similar to that in the system KAISi₃O₈–SiO₂ (diagram c).

0.219 (\pm 0.106) J/mol/bar reflects the observed dependence of the potassium content in Cpx in equilibrium with the alumino-silicate melt on pressure. Compositional factors can amplify or moderate this effect without change of its sign. The calculated $W_{\rm Si} = 1.716(\pm 0.498) * P(\text{bar})$ J/mol reflects the observed positive correlation of K2O in clinopyroxene with SiO_2 in coexisting melt (Fig. 2a,b). Fig. 2c shows some dependence of K_2O in Cpx on Al_2O_3 in the coexisting melt. However, in contrast to SiO₂, Al_2O_3 in the melt strongly correlates with pressure and, therefore, concentration of Al₂O₃ in the melt is already regarded in Eq. (2). Eq. (3b) shows that addition of any component (i.e., Jd, En, Fs) into the Di(Hed)-KJd solid solution results in decrease of the potassium solubility in clinopyroxene, as well.

We used Eq. (2) and the data by Gasparik (1996) on diopside melting for calculation of the hightemperature isobaric relation in the system Di-KJd, which is a portion of the system Di-Kfs. Fig. 3a illustrates the major features of the KCpx-liquid equilibria. Low-KAlSi2O6 clinopyroxene precipitates first from a melt at high temperature. It becomes richer in the KJd component with decrease of temperature at any pressure. However, solubility of the KJd component in clinopyroxene drastically drops with decrease of pressure. As pressure decreases, both clinopyroxene and liquid compositions displace toward the alkali-poor region of the system. By now, there are no data on KCpx equilibria at temperatures below 1600–1500 °C. However, we assume that there should be a "pseudo-peritectic" point at pressures below 60-50 kbar which produces K-feldspar and potassium-free clinopyroxene (Fig. 3b). This conclusion follows from both the experimental and natural data. Experiments by Mitchell (1995), Tsuruta and Takahashi (1998) and Wang and Takahashi (1999) show that the potassium content of clinopyroxene abruptly drops (by 0.5-1 wt.%) in presence of K-feldspar. Similar conclusion has been done from the study of K-feldspar blebs in clinopyroxene from garnet-clinopyroxene rocks of the Kokchetav Complex (see below, Perchuk et al., 1996). The proposed peritectic reaction is analogous to the reaction of an incongruent melting of sanidine in the system San-Qtz at P=1 atm (Fig. 3c, Schairer and Bowen, 1955).

5. Immiscibility in the carbonate-silicate melts and the stability of *KCpx*

Liquid immiscibility in sodic carbonate-silicate systems was experimentally investigated (Koster van Groos and Wyllie, 1968, 1969, 1973; Freestone and Hamilton, 1980; Perchuk and Lindsley, 1982; Lee and Wyllie, 1997, 1998; Wyllie and Lee, 1998), following the discovery of sodic carbonatitic lavas (e.g., Dawson et al., 1987; Dawson, 1989) and sodic carbonate inclusions in minerals from alkalic rocks (e.g., Kogarko et al., 1991; Solovova et al., 1996; Nielsen et al., 1997). Freestone and Hamilton (1980) found that a miscibility gap in the system SiO₂-Al₂O₃-TiO₂-CaO-FeO-MgO-Na₂CO₃-H₂O at 1250 $^{\circ}$ C expands with the increase of P from 2 to 7.6 kbar, mostly at expense of displacing of the field of the silicate melt toward the alkali-poor region. This conclusion has been experimentally tested by Wyllie and Lee (1998) in the pseudo-ternary system [SiO₂, Al₂O₃]-Na₂O-(Ca,Mg,Fe)O. Wyllie and Lee (1998) found, however, complete miscibility at P>25 kbar and 1300-1500 °C (se also Lee and Wyllie, 1998). Fig. 4 demonstrates systematic decrease of the silicate melt field with the increase of P or the decrease of Al/ Si ratio in the melt.

Carbonatites often accompany alkali-rich basic and ultra-basic magmatism in continental platforms, but only locally in oceanic islands (e.g., Tenerife). This observation has been explained by a model of carbonation of basaltic magmas (Perchuk, 1971), whereby an increase of the Na₂O and CO₂ activities in deepseated fluid drives the reaction:

$$\begin{split} \text{NaAlSi}_{3}\text{O}_{8}(Ab) + \text{CaFeSi}_{2}\text{O}_{6}(Hed) \\ &+ (1.5\text{Na}_{2}\text{O} + \text{CO}_{2})_{\text{fluid}} \\ &= 2\text{NaAlSiO}_{4}(Ne) + \text{NaFeSi}_{2}\text{O}_{6}(Aeg) \\ &+ 2\text{CaCO}_{3}(Cal). \end{split}$$

to the right. Owing to carbonate-silicate liquid immiscibility, the increase of Na₂O and CO₂ activities leads to the formation of the association *ijolite* (Ne+Aeg)+carbonatite (Cal) instead of the usual basalt or gabbro (*Pl+Hed*). An experimental study of the system NaAlS₃O₈-CaAl₂Si₂O₈-Na₂CO₃-H₂O

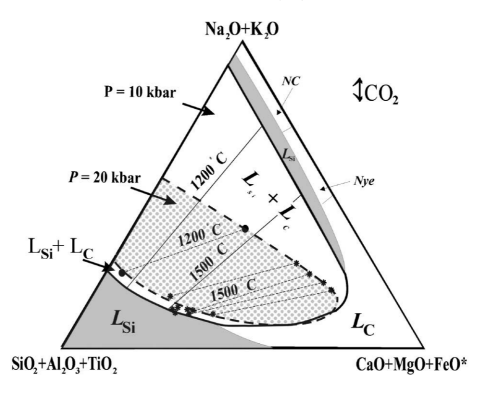


Fig. 4. Melting relations for the pseudo-ternary system $[SiO_2-Al_2O_3-TiO_2]-(Ca,Fe,Mg)O-(K,Na)_2CO_3$ (of the Freestone and Hamilton's, 1980 projection from the CO₂ apex) illustrating the immiscibility of carbonate-silicate liquid in the sodic system at 10 kbar (solid) and in the potassic system at 20 kbar (dashed). P=10 kbar. The system $SiO_2-Al_2O_3-CaO-Na_2CO_3$ (Lee and Wyllie, 1998); tie-line 1200 °C corresponds to the same system at P=20 kbar in the presence of H₂O (Perchuk and Lindsley, 1980a, 1982). P=20 kbar. The system $[SiO_2-Al_2O_3]$ -(Ca, Mg)O-(K₂CO₃, showing carbonation of garnet peridotite melt at T=1500 °C in the presence of H₂O (Perchuk and Lindsley, 1980a, b, 1982). A strong effect of P on the mutual solubility of silicate melt and carbonate liquid and, consequently, an abrupt widening of the miscibility gap toward alkali-rich melts (see text) is apparent. Nve—nyereriet stability field (Lee and Wyllie, 1998).

(Koster van Groos and Wyllie, 1973) confirmed the immiscibility of carbonate and silicate melts. Later, Perchuk and Lindsley (1982) studied reaction (4) in the system NaAlS₃O₈-CaAl₂Si₂O₈-Na₂CO₃-K₂CO₃-CaCO₃-H₂O using plagioclase glasses of compositions $An_{90}Ab_{10}$ and $An_{90}Ab_{10}$ (in mol%) at 10 and 20 kbar and 1200-1500 °C. Alkali and CO₂ activities were fixed by Na, K and hydrate salts, as well as by $C_2H_4O_4 \cdot 2H_2O$ and $C_3H_4(OH) \cdot 3Na_2CO_3 \cdot 5H_2O$. The experiments showed that the melt split into two silicate liquids and one carbonate-silicate liquid. The shaded area in Fig. 4 denotes the $L_{\rm Si}-L_{\rm c}$ miscibility gap at 20 kbar in the potassic system. Compositions of the carbonate-silicate liquid were calculated on the basis of the compositions of a starting mixture and the silicate melt after the experiment (Perchuk and Lindsley, 1982). The slopes of the tie-lines are gentler in comparison with the sodic system. That indicates that the silicate melt depleted in alkalis can coexist with potassium salt melts (K_2CO_2 or KCl), whereas the potassium partition coefficient for the two liquids must be larger than 20 in favor of the salt melt. In this case, a high potassium activity is buffered in the silicate melt by the coexisting salt melt and may provide the conditions for the crystallization of *KCpx* from the silicate melt.

Koster van Groos and Wyllie (1973), Lee and Wyllie (1997, 1998), Wyllie and Lee (1998) as well as Freestone and Hamilton (1980) not only confirmed the immiscibility in the Na–Ca–carbonate–silicate systems, but also provided phase diagrams.

In contrast, the system K₂O-CaO-SiO₂-Al₂O₃-CO₂ was not studied systematically. The nature and the behavior of ultra-potassic salt melts at high P has been studied only superficially. Perchuk and Lindsley (1982) studied the carbonation of garnet peridotite melt in the system Mg3Al2Si3O12-CaMgSi2O6-K₂CO₃-H₂O at 1500 °C and 20 kbar in order to find evidence for the carbonate-silicate immiscibility. Glass globules of phlogopite composition included in Ca-Mg-K carbonate matrix were found after the experiments. As expected, the carbonate fraction extracted Ca and partially Mg from the silicate melt (Fig. 4). Other information on immiscibility of potassium-rich alumosilicate and carbonate-silicate melts is very restricted and non-systematic. Harlow (1997) observed rare silicate globules in the experiments on potassium exchange between K₂CO₃ melt and KCpx at 100 kbar. However, Matveev et al. (1998) did not observe such carbonate-silicate immiscibility in the system K₂Mg(CO₃)₂-K₂CO₃-Di-Grs at 70 kbar and 1500–1600 °C.

Besides the preliminary study of Perchuk and Lindsley (1982), liquid immiscibility in potassic carbonate-silicate systems has not been studied experimentally. Perchuk and Yapaskurt (1998) suggested that the observation of Freestone and Hamilton (1980) can be applied for potassic systems: the miscibility gap abruptly narrows and moves to the alkali-poor region with the increase of pressure. Therefore the increase of pressure expands the field of the potassic carbonate liquid and contracts that of the silicate liquid.

The above review shows that the most experimental studies of the carbonate-silicate systems were carried out to solve the problem of genesis of carbonatites. From mid-1990s, Jones et al. (1995), Litvin et al. (1997), Pal'yanov et al. (1998, 1999), Sokol et al. (1998, 2000) and Borzdov et al. (1999) showed that the alkali-carbonate and the alkali-carbonate-silicate systems are appropriated media for synthesis of large diamond crystals. In addition, investigations of phase relationships in the potassium-bearing carbonic-alumino-silicate systems seems to be the right way to understanding the origin of ultra-high pressure rocks with diamond, *KCpx*, phlogopite, etc.

6. A review of the natural data

6.1. Inclusions of KCpx in diamonds and garnets

There are numerous reported occurrences of KCpx as inclusions in diamonds from eclogite and peridotite nodules in kimberlites and lamproites (e.g., McGregor and Carter, 1970; Moore and Gurney, 1985; Ricard et al., 1989; Harlow and Veblen, 1991; Stachel et al., 2000). Prinz et al. (1975) pointed out a wide variation of K_2O in Cpx inclusions in diamonds (0.02–0.87) wt.%). Reid et al. (1976) found up to 0.29 wt.% of K₂O in Cpx from diamondiferous eclogites. Having summarized all known data on Cpx composition from mantle xenoliths and inclusions in diamonds, Bishop et al. (1978) showed that Cpx from eclogites was richer in potassium, than Cpx from lherzolites. The K_2O content in Cpx from eclogites varies from 0.05 to 1.62 wt.%, whereas it is less then 0.37 wt.% in lherzolites. Based on the correlation of Na₂O in garnet with K_2O in coexisting Cpx, Bishop et al. (1978) confirmed strong dependence of the K₂O content in *Cpx* on pressure. Stachel et al. (2000) described *KCpx* $(0.53-1.44 \text{ wt.}\% \text{ K}_2\text{O})$ coexisting with Na₂O-bearing majoritic garnets (6.313-6.558 Si atoms in garnet formula per 24 oxygen) from inclusions in diamonds from the Kankan region (Guinea). All these data unambiguously proved the ultra-high pressure origin of natural KCpx.

Perchuk et al. (1996) reported about 1.2 wt.% K₂O in *Cpx* from diamond-free coarse-grained garnet-clinopyroxene silicate rocks (Group A) of the Kumdy-Kol microdiamond mine in the Kokchetav Complex, Kazakhstan. Even higher content of K₂O, up to 1.5 wt.%, was measured in *Cpx* from garnet-clinopyroxene fine-grained carbonate–silicate rocks (Group B) (Sobolev and Shatsky, 1990; Perchuk et al., 1995; Shatsky et al., 1999). Average chemical compositions of the rock groups are:

	SiO_2	TiO ₂	Al_2O_3	Cr_2O_3	FeO	MgO	CaO	Na ₂ O	K_2O	P_2O_5	CO_2
A	54.7	0.23	8.44	0.08	4.17	11.80	16.02	4.02	0.12	0.25	4.65
В	56.4	0.24	8.84	0.12	3.51	9.88	13.35	2.16	1.81	0.26	21.78

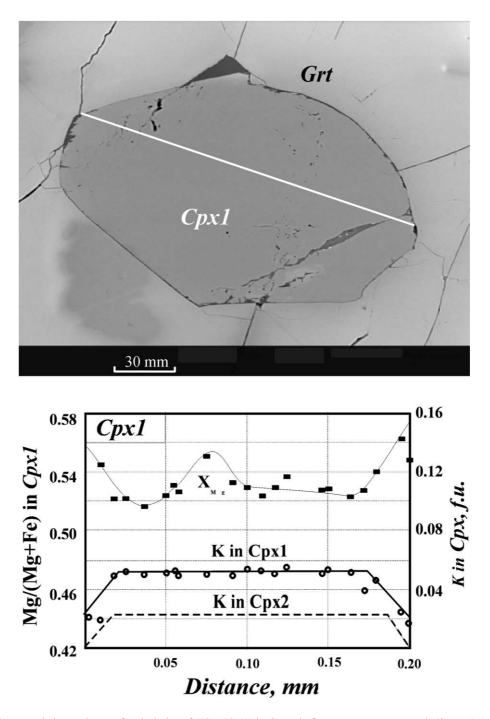


Fig. 5. A back-scattered electron image of an inclusion of KCpx (Cpx1) in Grt grain from pyroxene–garnet rock (Group A, see text) of the Kumdy-Kol mine in the Kokchetav Complex (N. Kazakhstan) and a microprobe profile (solid) across this grain (Perchuk et al., 1996; Perchuk and Yapaskurt, 1998). The profile of the potassium content in matrix Cpx (Cpx2) containing Kfs blebs from Fig. 6 is shown for comparison (dashed). Electron microscope CamScan.

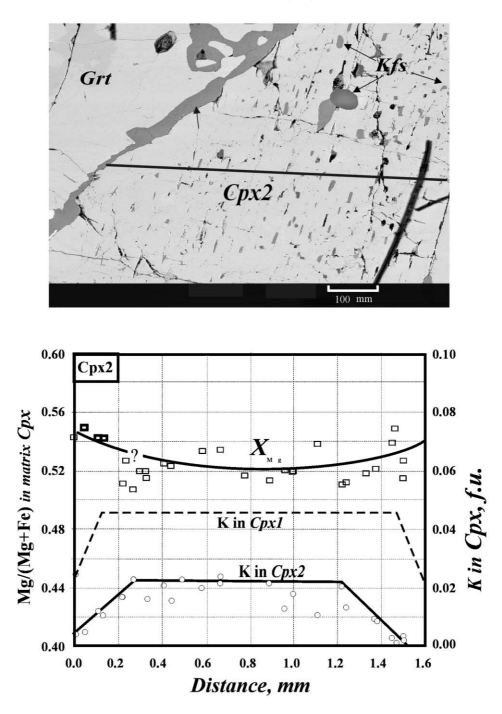
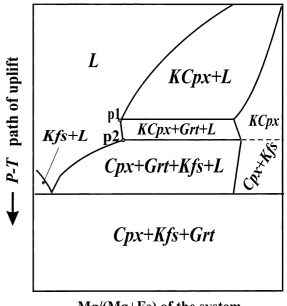


Fig. 6. A back-scattered electron image of the matrix Cpx (Cpx2) with Kfs blebs in the core from the diamond-free K-poor pyroxene–garnet rock (Group A, see text) of the Kumdy-Kol mine in the Kokchetav Complex (N. Kazakhstan) and a microprobe profile across this grain. The analyses are made with defocused beam in order to obtain an integrated bulk composition in every point (Perchuk et al., 1995; Perchuk and Yapaskurt, 1998). The profile of the potassium content in Cpx1 inclusion in garnet from Fig. 5 is shown for comparison. Electron microscope CamScan.

These undeformed rocks form interlayers, boudins, and lenses in strongly foliated garnet-biotite diamondiferous gneisses and schists and alternate with the eclogite boudins (Lavrova et al., 1999). The relationships of KCpx, potassium-free clinopyroxenes, and Kfs in the rocks of Group A are the most interesting. The rocks are composed of large (locally up to 7 cm) grains of garnet and clinopyroxene, accessory sphene, apatite, calcite, and secondary quartz, hornblende, K-feldspar, epidote, and calcite (Perchuk et al., 1995, 1996). Garnet contains microinclusions of KCpx (Cpx1) whose K2O concentrations can reach 1.2 wt.% (see the top of Fig. 5). The bottom of Fig. 5 illustrates the distribution of potassium (f.u.) and X_{Mg} across a ~ 200 μ m wide inclusion of KCpx in the Grt. The K₂O content in the core of the Cpx1 inclusion is uniform. At its edges, K_2O symmetrically drops from 0.055 (~ 1.2 wt.%) to 0.02 f.u. (~ 0.4 wt.%). The X_{Mg} profile is more complicated. According to Perchuk and Yapaskurt (1998), it reflects a sequence of mineral precipitation from a melt during uplift and cooling. Being the liquidus mineral, Cpx1 precipitated first (X_{Mg} decreases), and subsequently Grt forms (X_{Mg} increases) at lower T and P. The cores of large matrix Cpx in the rocks (Cpx2) contain tiny, less then 10 μ m, Kfs blebs (Fig. 6). The bottom of Fig. 6 shows a compositional profile across Cpx2 obtained with a defocused electron beam. Similar to Cpx1, the integrated K_2O content in core in Cpx2 is constant, about 0.02 f.u., but symmetrically decreases to zero in the rims (Perchuk et al., 1995; Perchuk and Yapaskurt, 1998). Reid et al. (1976) described similar Kfs blebs in potassium-free Cpx in diamondiferous eclogite from African kimberlite pipes (South Africa, Botswana, Tanzania). The sodium content in Grt from these eclogites reached 0.15 wt.% and is positively correlated with Ti (substitution Grs-Na $Ca_2[Ti,Al]Si_3O_{12}$). The jadeite content in coexisting Cpx varies from 20 to 60 mol%. Taking this into account, Reid et al. (1976) suggested that the crystallization of Grt and Cpx from a melt occurred at a depth of about 180-200 km. A subsequent uplift of eclogite toward the Earth's surface caused the exsolution of jadeite from Cpx. Partial melting of the rock during the rapid uplift led to K₂O partitioning into the melt, which was subsequently preserved as the Kfs blebs in Cpx. Luth (1997) proposed that the



Mg/(Mg+Fe) of the system

Fig. 7. Hypothetical pseudo-isobaric T-X ($X=X_{Mg}=Mg/Mg+Fe$) diagram of melting for the pseudobinary system Grt-KCpx, deduced from relationships between Cpx1, Cpx2, Grt, and Kfsand their compositions and zoning, shown in Figs. 5 and 6 (after Perchuk and Yapaskurt, 1998).

formation of blebs is related to exsolution from *Cpx* by the reaction:

$$3\text{KAlSi}_2\text{O}_6 + 2\text{Ca}_{0.5}\text{AlSi}_2\text{O}_6$$

= 3KAlSi_3O_8 + CaAl_2SiO_6, (5)

or

$$KCpx + Esk = Kfs + Tsc.$$

In contrast, KCpx (Cpx1) from clinopyroxene– garnet rocks of the Kokchetav Complex does not contain the Ca-eskolaite end-member. Therefore, reaction (5) is unable to explain the formation of the *Kfs* blebs in *Cpx2* (Perchuk et al., 1995, 1996). On the basis of phase relationships and chemical zoning of both the *Cpx1* inclusions in *Grt* and the matrix *Cpx2* with the *Kfs* blebs (Fig. 6) from the pyroxene–garnet rocks of Group (A), Perchuk and Yapaskurt (1998) showed that reaction (6)

$$KAlSi_2O_6 + [SiO_2]_{melt} \Rightarrow KAlSi_3O_8, \tag{6}$$

or

$$KJd$$
 (in Cpx) + $[SiO_2]_{melt} = Kfs$,

operates during uplift and cooling of the potassiumrich carbonate-silicate melt. In addition to Fig. 3b illustrating peritectic point in the system diopsidesanidine, Fig. 7 demonstrates hypothetical non-isobaric $TX (X=X_{Mg}=Mg/Mg+Fe)$ diagram of melting for pseudobinary system Grt-KCpx deduced from the data of detailed microprobe profiles of Cpx1 (Fig. 5) and Cpx2 (Fig. 6) from the Kokchetav Grt-Cpxrocks.

6.2. Non-crystalline inclusions in diamonds

In order to determine the source and the depth of melt generation and its primary composition, the most useful information can be obtained from analyses of non-crystalline inclusions in minerals from carbonatites and in diamonds from kimberlite and lamproite pipes. Despite the successful experiments on diamond crystallization in alkali or alkali-earth carbonate and carbonate-silicate systems (Jones et al., 1995; Litvin et al., 1997; Pal'yanov et al., 1998, 1999; Sokol et al., 1998, 2000; Borzdov et al., 1999), analysis of fluid and melt inclusions in minerals from carbonatites (e.g., Kogarko et al., 1991; Solovova et al., 1996; Nielsen et al., 1997) does not support deep-seated origin of majority of carbonatitic melts. The absence of diamond in carbonatites, even as inclusions in phenocrysts, indicates a relatively shallow depth of their origin, i.e., less then 100 km. Nevertheless, diamonds are common in kimberlites and lamproites. It is now evident that diamonds from kimberlites and lamproites contain inclusions of mantle minerals, fluids, silicate melts and brines. Non-crystalline inclusions in diamonds (Dawson et al., 1987; Navon et al., 1988;

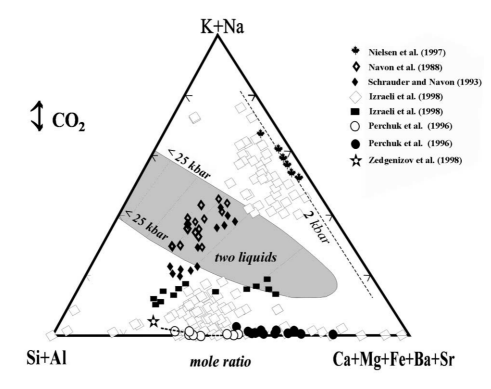


Fig. 8. The Freestone and Hamilton's (1980) projection from $CO_2 + Cl$ apex showing a comparison of the compositions of fluid inclusions in diamonds (all open and solid diamonds and rectangles) from some African kimberlite pipes with the composition of some fluid inclusions in carbonatites (maples), bulk compositions of Grt-Cpx (open circles) and calc-silicate rocks (solid circles) from the Kokochetav complex (Kazakhstan). Asteriks represent the composition of the silicate melt from Fig. 10. See text for further explanation.

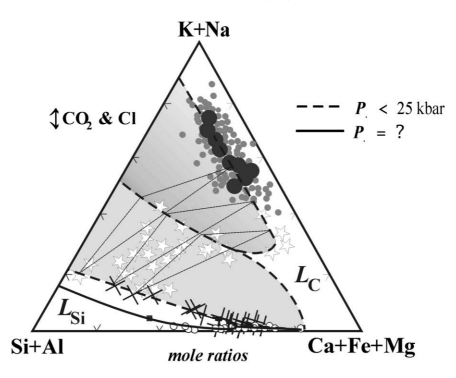


Fig. 9. Plot of fluid (open and solid rhombs) and brine compositions from inclusions in diamonds, some kimberlites and lamproites onto the Freestone and Hamilton's (1980) projection from CO_2 . Large solid diamonds—Brines: black circles—average compositions of the brine in the nine diamonds, small gray circles-analyses of the individual inclusions in all nine diamonds. Stars—average compositions of fluid inclusions in fibrous diamonds. Crosses—lamproites, xs—kimberlites. Solid rectangle = asterisk from Fig. 8, open circles = silicate and calc-silicate magmatic rocks from Kumdy-Kol microdiamond main, the Kockchetav complex (northern Kazakhstan).

Dawson, 1989; Chen et al., 1992; Schrauder and Navon, 1993, 1994; Schrauder et al., 1993, 1996; De Corte et al., 1998; Izraeli et al., 1998, 1999, 2001; Zedgenizov et al., 1998; Navon, 1999; Navon and Izraeli, 1999) indicate pressure of their entrapment of about 40-70 kbar (e.g., Navon, 1991), i.e., to a depths interval from 120 to 200 km.

6.3. Fluid inclusions in diamonds

Very few data exist on composition of fluid inclusions in diamond because of detection and analytical problems. According to Navon et al. (1988), Chen et al. (1992) and Izraeli et al. (2001), only three major groups of fluid inclusions are present in diamond: (1) silicate-bearing of diverse composition; (2) aqueous-carbonic fluid and (3) brine. However, because of their small size, measurements on such inclusions are imprecise and some analyses correspond to mixtures of inclusions. In Fig. 8, a portion of the fluid-melt inclusions (open and solid diamonds) corresponds approximately to equal mixtures of all three constituents (Navon, personal communication). The diagram also shows a preliminary two-phase immiscibility region of the fluidized silicate melt in absence of the salt melt presumably at pressure of above 20– 25 kbar and T>1000 °C. All inclusions are characterized by K/Na ≥ 1 as exemplified by the data (wt.%) from Navon et al. (1988) and Schrauder and Navon (1994):

#	1	2	3	4	5	6	7	8	9	10	11	12	13
Na ₂ O	2.4	2.6	2.2	3.0	2.3	3.5	3.8	3	2.9	2.3	2.4	2.7	3.4
K ₂ O	18.7	21.4	22.2	18.4	18.1	29.7	17.7	23.7	20.8	19.4	21.1	25.2	19.3

This large K/Na ratio is able to explain coexistence of diamond with KCpx rather than omphacite. De Corte et al. (1998) showed that microdiamonds from metamorphic rocks of the Kokchetav massif are similar to those of kimberlitic pipes. On the other hand, analyzing the infrared spectra of microinclusions in microdiamonds from garnet grains of the rocks, De Corte et al. (1998) observed predominance of the aqueous inclusions containing solid carbonate minerals. That indicates a high oxygen fugacity in the C–O–H fluid during the diamond formation.

6.4. Brine inclusions in diamond

Chen et al. (1992) was the first to report an occurrence of concentrated KCl inclusions in diamond. Similar inclusions were subsequently found in diamonds from kimberlite pipes Koidu (Sierra-Leone), Koffiefontein (South Africa) and Udachnaya (Russia) (Fig. 9; Navon and Izraeli, 1999; Izraeli et al., 1999, 2001). The average composition of the brine from these inclusions is (wt.%): $30-40 \text{ H}_2\text{O}$, 30-34(Na,K)Cl, 23-25 (Fe,Ca,Mg)CO₃, $3-4 \text{ SiO}_2$ and 5% of an undetected chloride (to account for the systematic excess of the observed chlorine). Izraeli et al. (2001) suggested an average formula for this fluid: (K,Na)₈(Ca,Fe,Mg)₄SiO(CO₃)₄Cl₁₀(H₂O)₂₈₋₄₄. They assumed that the fluid inclusions were mixtures of two fluids, i.e., a KCl–NaCl brine and a Ca, Na, Si, Fe, Mg-rich carbonate fluid. This conclusion is supported by the direct correlation of the alkali-earth component with $X_{CO_3}^{cl}$.

6.5. Inclusions of silicate melt in diamond

Navon et al. (1988), Schrauder and Navon (1994), and Izraeli et al. (2001) investigated silicate melt inclusions in diamond from kimberlites of Botswana and Zair (Table 3). The major characteristics of the melt inclusions are very low aluminum contents ($Al_2O_3 < 6$

Table 3

Selected analyses on metallic oxides, and H₂O/CO₂ ratios in fluid inclusions trapped by diamonds (Navon et al., 1988; Schrauder and Navon, 1994)

1994)	TW/NI	OTD	INVAL	THEFT	TAVAT	OTD	OTD	CTD	OTD	CDD	CDD	CDD	CDI
Sample #	JWN 110	CTP 6268	JWN 108	JWN 106	JWN 89	CTP LB	CTP MM1	CTP L0	CTP L6	GRR 1504	GRR 1508	GRR 1517	GRF 1518
Region	Botsw	ana				Zair	Zair			No dat	ta		
Points*	42	4	70	16	27	2	2	5	2	8	10	3	2
SiO ₂	41.3	31.9	23.9	28.4	29.4	34.6	40.4	41.2	43.3	42.3	42.4	30.3	42.4
TiO ₂	4.6	4.2	4.6	4.9	4.8	2.1	2.9	2.4	2.5	2.6	2.7	3.4	2.9
Al ₂ O ₃	4.9	2.9	2.5	2.2	2.9	5.6	4.5	6.1	5.4	4.9	4.9	5.3	4.4
FeO	11.1	15.7	15.2	14.9	16.1	4.9	7.2	5	5.6	11.1	6.1	5	8
MgO	5.1	5.7	10.9	10.6	8.2	2.3	4.6	2.8	3.8	4.6	3.6	4.3	4.9
CaO	6.8	10.5	13.5	12.6	13.4	12.3	13.9	10.7	10.6	7.8	11.9	18.7	9.8
Na ₂ O	2.4	2.6	2.2	3.0	2.3	3.5	3.8	3	2.9	2.3	2.4	2.7	3.4
K ₂ O	18.7	21.4	22.2	18.4	18.1	29.7	17.7	23.7	20.8	19.4	21.1	25.2	19.3
Total **	94.9	94.9	95.0	95.0	95.2	95.0	95.0	94.9	94.9	95.0	95.0	94.9	95.1
H ₂ O , ppm	64	407	99	38	128	140	165	191	619	118	269	241	:115
CO ₂ , ppm	120	600	250	75	420	66	89	79	135	44	133	139	70
X ^{fl} _{H2O} ***	0.6	0.5	0.5	0.6	0.4	0.8	0.6	0.8	0.8	0.8	0.7	0.8	0.8

* Number of inclusion analyses for each diamond crystal.

** Average major oxide compositions (wt.%) of the volatile-free fractions of the inclusions, normalized to the sum of eight rock-forming oxides equal to 95%.

** $X_{H_2O}^{fl} = H_2O/H_2O + CO_2$, molar fraction of H_2O in a fluid.

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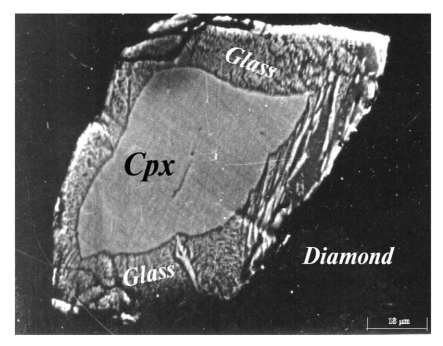


Fig. 10. Omphacite grain surrounded by a silicate glassy matrix; an inclusion in diamond from the Satykanskaya kimberlite pipe, Yakutia (Zedgenizov et al., 1998). *P* of 81–83 kbar for 1000–1800 °C was calculated with Eq. (2).

wt.%), wide variations of SiO_2 content (from 24 to 58 wt.%), and high alkalinity with significant predominance of potassium over sodium. The following nine examples (wt.%) clearly illustrate the conclusions of Izraeli et al. (2001) on extremely potassic fluid: Data on inclusions in diamonds from kimberlite pipes from Sierra-Leone, South Africa and Yakutia (Izraeli et al., 2001) indicate that the inclusions of silicate melt are accompanied by the inclusions of highly concentrated KCl brines and diverse mineral

	1	2	3	4	5	6	7	8	9
Na ₂ O	4.6 ± 2.7	1.9 ± 2.4	1.1 ± 1.2	9.9 ± 6.9	4.4 ± 1.8	2.8 ± 4.3	2.5 ± 0.8	2 ± 0.9	3.7 ± 2.6
K ₂ O	28.8 ± 4.9	31.9 ± 3.4	36.5 ± 1.3	29.8 ± 1.8	26.7 ± 3.2	39 ± 5.7	33.7 ± 4.2	30.6 ± 4.2	31.4 ± 4.4
K ₂ O/Na ₂ O	6.26 ± 1.81	16.8 ± 1.4	33.2 ± 1.2	3 ± 0.26	$6{,}07 \pm 1.8$	13.9 ± 1.3	13.5 ± 5.2	15.3 ± 4.7	8.5 ± 1.7

The studied inclusions are also characterized by relatively low X_{CO_2} (CO₂/(H₂O + CO₂)) that varies from 10 to 50 mol% (Navon et al., 1988; Schrauder and Navon, 1994). The Ca, Mg and Fe contents in the inclusions correlate directly with CO₂. The aqueous inclusions are richer in Al₂O₃ and SiO₂, while the carbonate inclusions are enriched in alkali, alkali-earth components, and P₂O₅. Elevated concentrations of chlorine are also evident.

inclusions. Mineral inclusions, associated with the ultra-potassic melts and fluids, belong to either eclogite (Grt-Cpx) or peridotite (Grt-Cpx-Ol) assemblages.

Zedgenizov et al. (1998) published data on a potassic-bearing clinopyroxene included in diamond from the Satykanskaya kimberlitic pipe. The core of this inclusion composed of KCpx is surrounded by glass (solidified fluidized silicate melt) (Fig. 10). The compositions of these phases are the following:

	SiO ₂	TiO ₂	Al_2O_3	Cr_2O_3	MnO	FeO	MgO	CaO	Na ₂ O	K_2O
KCpx	54.7	0.23	8.44	0.08	0.06	4.17	11.8	16.02	4.02	0.12
Melt (?)	56.4	0.24	8.84	0.12	0.01	3.51	9.88	13.35	2.16	1.81

The presence of such inclusion implies crystallization of KCpx within the diamond field stability. Despite the very low K₂O contents in the inclusion, the melt is by 15 times richer in potassium than the coexisting pyroxene. These data allows estimation of pressure for KCpx crystallization using Eq. (2): P=81-83 kbar within the wide temperature interval, from 1000 to 1800 °C. These values correspond to the depth of about 250 km.

Thus, the available data suggest that in the upper mantle, at ~ 40–50 kbar and ~ 1100–1500 °C, minerals can coexist with three liquid phases: aqueous-carbonic fluid, silicate melt and brine. Immiscibility in the carbonate-brine system has not been proven experimentally. Pure carbonate melt inclusions were also not found (the brine portion contains only rare carbonates). Compositions of all liquids are plotted on diagram of Fig. 9. The fluid phase is not distinct, because of the difficulties in its identification. However, the boundaries of the miscibility gap for fluid at P > 25 kbar can be outlined from Fig. 4. They are indicated in Fig. 9 by dashed lines, whereas the tie-lines connect the possible compositions of coexisting liquid phases, i.e., the silicate melt (kimberlite/ lamproite proposed by Israeli et al., in press), the fluid, and the brine. In accordance with Fig. 4, it can be assumed that the miscibility gap disappears or appreciably narrows, displacing toward alkali-poor silicate melts. Such compositions correspond to the pyroxene-garnet rocks of the diamond-bearing Kumdy-Kol area of the Kokchetav Complex in northern Kazakhstan.

7. Conclusions

The reviewed experimental and natural data indicate the following phase relationships involving *KCpx* and melt:

(1) *KCpx* of at least 1 wt.% of K₂O may crystallize on the liquidus of relatively high-potassic silicate and carbonate-silicate melts at P > 70 kbar.

(2) K_p depends mostly on *P* and the SiO₂ content of the melt, but $K_p < 1$ for any bulk composition.

(3) The existing experimental data on dependence of K_p on pressure and the melt compositions were used to derive an equation for the *P* calculations from the compositions of *KCpx* and the coexisting silicate melt.

(4) The data on fluid inclusion from natural diamonds suggest that the deep-seated magma splits at P < 50 kbar into three liquids, i.e., silicate melt, a CO_2-H_2O -potassium-rich fluid, and a potassium-rich brine.

(5) The *KCpx* inclusions (K₂O>1.2 wt.%) in garnet from ultra-high *P* potassium-poor rocks (K₂O<0.2 wt.%) from the Kokchetav massif could have resulted from the crystallization of pyroxene on the liquidus of a silicate magma, which mixed or coexisted with potassium-rich brine at depth of about 250 km. Liquid immiscibility between the silicate melts and brine occurring during uplift resulted in a decrease of alkali activity in the silicate melt and a decrease of K₂O content in *Cpx*. The *Kfs* blebs observed in cores of coarse-grained clinopyroxenes may have resulted from the peritectic reaction KCpx+melt=Di+Kfs at $P \sim 50-45$ kbar during the uplift.

Mineral and phase abbreviations used in the text

Abbreviation	Formula	Name of mineral
Ab	NaAlSi ₃ O ₈	albite
Aeg	NaFeSi ₂ O ₆	aegerine
Alm	Fe ₃ Al ₂ Si ₃ O ₁₂	almandine
An	CaAl ₂ Si ₂ O ₈	anorthite
APhl	KMg ₃ AlSi ₃ O ₁₁	anhydrous
		phlogopite
Cal	CaCO ₃	calcite
Cpx		clinopyroxene
		solid solution
Cos	SiO ₂	coesite
Di	CaMgSi ₂ O ₆	diopside
En	$Mg_2Si_2O_6$	enstatite
Esk	Ca _{0.5} AlSi ₂ O ₆	Ca-Eskola pyroxene
Fs	Fe ₂ Si ₂ O ₆	ferrosilite
Fo	Mg ₂ SiO ₄	forsterite
Grs	Ca ₃ Al ₂ Si ₃ O ₁₂	grossular
Grt		garnet solid solution
Hed	CaFeSi ₂ O ₆	hedenbergite
Jd	NaAlSi ₂ O ₆	jadeite
KCpx		potassium-bearing
		clinopyroxene
Kfs	KAlSi ₃ O ₈	K-feldspar
KJd	KAlSi ₂ O ₆	fictive K-jadeite
Ko	NaCrSi ₂ O ₆	kosmochlor
Ks	KAlSiO ₄	kalsilite

Ms	KAl ₃ Si ₃ O ₁₀ (OH) ₂	muscovite
Nc	Na ₂ CO ₃	Na-carbonate
Ne	NaAlSiO ₄	nepheline
Ol	-	olivine solid solution
Opx		orthopyroxene solid solution
Phl	KMg ₃ AlSi ₃ O ₁₀ (OH) ₂	phlogopite
Pl		plagioclase solid
		solution
Prp	$Mg_3Al_2Si_3O_{12}$	pyrope
Qtz	SiO ₂	quartz
Rt	TiO ₂	rutile
San	KAlSi ₃ O ₈	sanidine
Tsc	CaAl ₂ SiO ₆	Ca-Tschermak
		pyroxene
L_{c}		carbonate liquid
L_{Si}		silicate liquid
fl		fluid

Thermodynamic symbols used in the text

Т	tomporatura
-	temperature
P	pressure
X_i^j	mole fraction of an <i>i</i> -component in a mineral
	<i>j</i> ,
$N_i = 100$	$X_i \mod \%$
a_i	activity of an <i>i</i> -component
Kp	partition coefficient
K _r	reaction constant
$\Delta G_{ m r}^{\ 0}$	free energy difference in reaction r
$\Delta H_{\rm r}^{0}$	enthalpy difference in reaction r
$\Delta S_{\rm r}^{0}$	entropy difference in reaction r
$\Delta V_{\rm r}^0$	volume difference in reaction r
G_i^{e}	excess free energy of mixing of an i-
	component
R	gas constant (8.31 J/mol/K)

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