K in clinopyroxene at high pressure and temperature: An experimental study

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Abstract

Clinopyroxene (Cpx) is a principal upper-mantle phase for concentrating large cations but has not been viewed as a major crystal-chemical reservoir for K because K⁺ is considered too large to enter the largest site, M2, in the pyroxene structure. Accumulating data from high-pressure conditions indicate this inference is incorrect, so multianvil experiments have been performed to evaluate maximal K solubility in Cpx at high pressure. End members and mixtures of diopside, jadeite, and kosmochlor have been mixed with K_2CO_3 , KHCO₃ or both in welded platinum capsules and heated typically for 24 h in the range of 5 to 14 GPa and 1200 to 1700 °C. These experiments produced K-rich Cpx in solid solutions by means of a fictive Kcpx component (KCrSi₂O₆ or KAlSi₂O₆). The maximum K₂O content obtained is 4.7 wt% in a Cpx (Di₃₈Ko₃₉Kcpx₂₂En₁) formed from a 50:50 Di + Ko mixture at 10 GPa, 1400 °C. K uptake and partitioning is dependent positively on P, complexly on Cpx composition but not demonstrably on T. $C_{PX/Iiq}D_{K_2O}$ is in the range of 0.03–0.1 and $C_{PX/Iiq}D_{Na_2O}$ varies from 0.5 to 5, although the variations for each with Cpx composition are different. In diopside, Kcpx increases are always accompanied by increases in Nacpx, and cooperative Nacpx solution is necessary for Kcpx solution in the compositional systems examined. K appears to be accommodated in the M2 site of the Cpx structure by two types of spatial averaging: a large average M2 site, as in the case of Di, ameliorates the fit, but local accommodation by size averaging with a smaller M2 occupant, presumably Na, appears necessary, suggesting that the polyhedral compressibility of Na and K are large in comparison with Ca. In application to Cpx inclusions in diamond, the data here imply that a chromium diopside with ~ 1 wt% K₂O forms in the presence of a C-rich melt with 15–28 wt% K₂O.

INTRODUCTION

Cpx, an important mineral of the upper mantle and crust, was not considered a major reservoir for K because the preponderance of compositional data for pyroxenes of all kinds show only trace K content. The crystal-chemical reasoning is that K^+ [$R(^{[8]}K^+) = 1.51$ Å; Shannon 1976] is too large to enter the largest site, M2, in the pyroxene structure. Harlow and Veblen (1991) have shown that K⁺ at levels of at least 0.07 cations per 6 O atoms (1.5 wt% K_2O) can reside in the crystal structure of Cpx included in diamond and, thus, have confirmed the accuracy of some analyses of mantle Cpx with substantial K₂O (Prinz et al. 1975, McCandless and Gurney 1989, Rickard et al. 1989, Sobolev et al. 1991a) and more recently in Cpx inclusions from ultradeep [or ultra-high pressure (UHP)] metamorphic rocks (Sobolev et al. 1991b, 1994). Harlow and Veblen (1991) reasoned that high pressure and a K-rich environment were required to yield the K-rich pyroxenes found in diamonds. Because diopside, with a cell volume (V_c) of ~439 Å³, has larger average M2 (and M1) coordination polyhedra than omphacite ($V_c = -425 \text{ Å}^3$) or paticularly jadeite ($V_c = \sim 401 \text{ Å}^3$), they reasoned diopside should be a better host for K. However, for re-0003-004X/97/0304-0259\$05.00

placement of Ca by K a coupled exchange to balance charge such as

$$\mathbf{K} + {}^{[6]}\mathbf{M}^{3+} \leftrightarrow \mathbf{Ca} + (\mathbf{Mg}, \mathbf{Fe}) \tag{1}$$

is required and observed in all natural samples. Because the cell volume and size of the M2 coordination polyhedron also increases with increasing M1 site occupant (Oberti and Caporuscio 1991; Cameron and Papike 1980), Harlow and Veblen (1991) proposed that K uptake should be favored in solid solutions incorporating a relatively large trivalent M1 cation, such as Cr^{3+} . These predictions have needed experimental testing.

Recently, long after the pioneering work of Shimizu (1971), a few experimental studies have produced K-rich Cpx at high pressure, e.g., Doroshev et al. (1992), Harlow (1992), Luth (1992), Edgar and Vukadinovic (1993). Here are reported experiments designed to yield maximum K_2O content in Cpx otherwise in the diopside-jadeite-kosmochlor (-enstatite) field. The goals also include testing the above mentioned hypotheses and examining the partition coefficients, pyroxene crystal-chemistry, and phase assemblages in a Cpx-potassium carbonate system.

TABLE 1. Microprobe analyses of pyroxene starting materials

		Weight percent oxides								
	1	2	3	4	5					
SiO ₂	55.50	59.31	59.61	53.53	60.42					
TiO₂	0.03	0.02	0.01	0.06	0.00					
AI_2O_3	0.04	1.10	0.44	0.00	25.39					
Cr_2O_3	0.00	0.01	0.04	33.56	0.00					
Fe ₂ O ₃ *	0.08	0.00	0.22	0.15	0.12					
FeO	0.04	0.21	0.00	0.00	0.12					
MnO	0.02	0.02	0.01	0.00	0.02					
MgO	18.60	39.40	40.11	0.00	0.06					
CaO	26.00	0.00	0.00	0.04	0.20					
Na₂O	0.03	0.01	0.01	13.72	15.45					
K₂O	0.00	0.00	0.00	0.00	0.00					
Total	100.34	100.09	100.45	101.06	101.54					
		Cations f	or 6 atoms							
Si	1.996	1.982	1.986	2.001	2.00					
^{4]} AI	0.002	0.018	0.014	0.000	0.000					
SUM T	1.997	2.000	2.000	2.001	2.00					
Ti	0.001	0.000	0.000	0.002	0.000					
^{6]} AI	0.000	0.025	0.003	0.000	0.99					
Cr	0.000	0.000	0.001	0.992	0.000					
Fe ^{3+*}	0.002	0.000	0.006	0.004	0.003					
Fe	0.001	0.006	0.000	0.000	0.003					
Mn	0.001	0.001	0.000	0.000	0.000					
Mg	0.997	1.963	1.992	0.000	0.003					
Ca	1.002	0.000	0.000	0.002	0.007					
Na	0.002	0.001	0.001	0.994	0.99					
K	0.000	0.000	0.000	0.000	0.000					
Total	4.003	3.997	4.003	3.996	4.00					

Notes: 1 is 39849, Wakefield Di; 2 is 100634, Zabargad Hi-Al En; 3 is 100634, Zabargad Low-Al En; 4 is Synthetic Kosmochlor; 5 is 33399 Manzanal Jd.

* Fe3+ determined by cation sum and charge balancing.

MATERIALS, EXPERIMENTAL TECHNIQUE, AND ANALYSIS

Materials

Starting materials in the experiments were mixtures of ground reagent SiO₂, pyroxenes (most grains ≤ 100 mesh), K₂CO₃, or KHCO₃. Carbonates were chosen as a K source because of their low melting temperature and their potential importance as components in some diamond-forming environments (Navon et al. 1987). Experiments involved exchanging K between carbonate and pyroxene on the Di-Jd and Di-Ko joins (either as solid solution or mixtures). Some experiments included small amounts of En, to examine minor Di-En solution on K uptake, and silica was added to one set of experiments at the suggestion of J.R. Smyth to avoid subsilicic (<2 Si per 6 O) Cpx. Natural pyroxene samples include diopside (AMNH 39849, Wakefield, Quebec, Canada), jadeite (AMNH 33399, Manzanal, Guatemala), and enstatite (AMNH 100634, Zabargad Island, Egypt). Kosmochlor was synthesized in a procedure intermediate to that of Ikeda and Yagi (1972) and Frondel and Klein (1965), where a sodium chromium silicate glass was ground and crystallized at 800 °C for 96 h in a muffle furnace (see Table 1 for analyses).

High-pressure experimental procedure

Experiments were performed in multianvil apparati using 600-2000 ton presses in the laboratory at Lamont



FIGURE 1. Drawing showing our latest experimental multianvil setup of octahedral assembly with $LaCrO_3$ heater, platinum capsule and off-set Tc at its base.

Doherty Earth Observatory, following the uniaxial design of Walker et al. (1990) and Walker (1991). Truncated tungsten carbide cubes in the assembly were both 25 and 16 mm in dimension, the latter employing tool-steel shims to match the assembly size of the former. The pressure-medium octahedra, to match cubes with 6 or 8 mm truncated edge lengths, were made of fired castable ceramic with integral fin gaskets and drilled for insertion of a solid cylindrical LaCrO₃ heater with a \sim 3.18 mm I.D. Potassium-carbonate is extremely reactive and mobile, so it must be encapsulated with other reactants in platinum capsules. Approximately 20 mg total of $Cpx + K_2CO_3$ in equal amounts were packed into a cylindrical platinum capsule and dried for 20 min at ~150 °C before welding the capsule. Experiments with KHCO₃ were not dried because the intent was to add H₂O to enhance reaction and diffusion rates. The capsule with an alumina liner was placed in contact with a D-type (W3%Re/W25%Re) transversely mounted thermocouple (Tc) so that the central hot-spot was ~ 1 mm away in the bottom of the capsule (see Fig. 1). Experiments using two thermocouples, one at the center of the assembly and the other at the far end of the capsule, have measured gradients of $\sim \leq 200$ °C with the central Tc at 1700 °C. Consequently, the Tc temperature at 1 mm below center is estimated to be 50° below the hottest point in the capsule and is comparable to the average temperature in the capsule; the total gradient in the capsule is 100 °C. Experiments were brought up to P, then brought up to T, kept at a temperature for normally 24 h, then quenched, and finally depressurized. Some experiments were superheated by $\sim 100^{\circ}$ for 1 h before lowering to the final temperature. Other aspects of the experimental procedure, including pressure calibration that is considered accurate to \pm 1/4 GPa, are similar to those described by Johnson and Walker (1993) and Walker (1991). Conditions for experiments cited here are given in Table 2.

Analytical technique

Processed octahedral assemblies were imbedded in epoxy and sawn in half with a 4 mil diamond blade. One

TABLE 2.	Conditions f	for experiments
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	<i>P</i> (GPa)	<i>T</i> (°C)	Duration(h)	Phases	Experiment				
Di _{aa} Ko _a Qtz ₅ +									
K ₂ CO ₂	5	<1500	29	Cpx, uvar, mgchr, glass	GG404				
K,CO,	6	1400	23	Cpx, uvar, mgchr, glass	BB330				
K ₂ CO ₃	7	1500	24	Cpx, uvar, esk, glass	BB109				
K ₂ CO ₃	9	<1700	9	Cpx, uvar, glass	BB243				
KĤCŎ ₃	10	>1200	25	Cpx, K-wad, devit. glass	BB206				
K ₂ CO ₃ *	10	1500	24	Cpx, K-wad, devit. glass	BB470				
K ₂ CO ₃	10	1500	21	Cpx, devit. glass	GG477				
K ₂ CO ₃ **	11	1500	14	Cpx, KMgSil, Fo, glass	GG307				
K ₂ CO ₃	11	1700	24	Cpx, devit. glass or carb	GG299				
K ₂ CO ₃	12	1500	45	Cpx, esk, devit. glass or carb	BB85				
K ₂ CO ₃	14	1500	26	Cpx, devit. glass or card	BB89				
			Jd+						
K _a CO _a	11	1300	100	Cpx. carb	GG229				
K₂CO₃	10	1500	24	Cpx, carb, glass	GG405				
			Ko⁺						
K ₂ CO ₂	11	1300	24	Cpx. carb	GG227				
K₂CO₃	10	1500	27	Cpx, K-wad, esk, KSiCr ₃ O ₇ , carb, glass?	GG415				
			Di₅₀Jd₅₀	+					
K _a CO _a	10	1500	26	Cpx, K-wad, carb	GG416				
KHCO ₃ + K ₂ CO ₃	10	<1400	14	Cpx, K-cym?, carb	BB308				
	10	1450	26	Cpx, glasses	GG499				
$KHCO_3 + K_2CO_3^*$	11	1450	24	Cpx, glasses	GG568				
			Di ₅₀ Ko ₅₀	+					
K2CO2**	10	1450	24	Cpx, K-wad, esk, carb	GG486				
K ₂ CO ₂	10	1500	24	Cpx, K-wad, KSiCr ₂ O ₂ , glass, carb	GG412				
KHCO ₂ + K ₂ CO ₂	10	1400	21	Cpx, K-wad, esk, carb	GG421				
KHCO ₃	11	1500	22	Cpx, K-wad, esk, glass	TT48				
			Di ₇₀ Ko ₂₀ Er	n_{20}^+					
KHCO ₃	10	1300	24	Cpx, K-wad, KMgCrsil, carb	BB211				
K ₂ CO ₃	11	1400	27	Cpx, K-wad, KMgCrsil, uvar, glass	GG389				

Notes: carb = carbonate, esk = Cr_2O_3 , K-cym = KAlSi_3O_8 · H_2O, K-wad = $K_2Si_4O_8$, KMgCrsil = new KMgCr-silicate, mgchr = Mg_2CrO_4, uvar = uvarovite, devit = devitrified.

* Temperature was higher by 50° for the first hour; > or < indicates that, based on the final geometry (e.g., Tc position or compaction texture of the LaCrO3 heater as not symmetrical) the appropriate capsule temperature may be higher or lower (probably by \leq 50°).

** Temperature was higher by 100° for the first hour.

half of each experiment was polished on diamond-impregnated laps with mineral spirits or oil for study of textures and compositions. The other half was kept in reserve for X-ray diffraction and ATEM/HRTEM analysis. Samples were stored in a drying oven at 60 °C, however, unless embedded in epoxy the carbonate phases eventually hydrate and effloresce. The polished samples were observed in incident and reflected light and imaged using a Zeiss DSM950 SEM with BSE and Link AN100 EDS. Microprobe analysis was carried out for constituent major elements except C and O using an upgraded ARL-SEMQ operating at 15 kV and 12.5 nA sample current. Natural minerals were used as standards, and the CIT-ZAF correction scheme according to Armstrong (1989) was employed. A focused beam was used for all phases except devitrified glasses and fine-grained carbonate for which a 5–10 μ m beam was traversed for > 10 spots across the sample. Corrections for unanalyzed CO₂ and H₂O were made by adding a specified missing quantity (in the proportion of carbonate and bicarbonate used in the charge) to the average of a group of analyses so that the adjusted average total equaled 100% (having taken everything into account in the ZAF correction).

RESULTS

Cpx and glass or carbonate, are the dominant products, although other phases occur in minor amounts (Table 2). Phase segregation by thermal compaction (Lesher and Walker 1988) is conspicuous and common because of thermal gradients in the capsules. Carbonate or melt seeks the hot sides of the charge, and Cpx (and other crystalline phases) seek the colder center and ends (Figs. 2 and 3a); whereas an hourglass thermal profile should be expected in the capsule, inconsistent electrical properties of our LaCrO₃ result in variable compaction geometries. K-rich Cpx forms as overgrowths and recrystallizations, which was inferred from sharp compositional boundaries of overgrowth K-rich Cpx on relict cores and from compaction texture (Fig. 4). In some experiments, Cpx forms either tooth-shaped protuberances into carbonate (akin to feldspars in pegmatites), usually when compaction is incomplete, or new prisms extending from Pt or compacted Cpx into glass or carbonate, probably because of decreasing temperature or pressure creep from thermal pressure in the hydraulic system. In some experiments, there are immiscible carbonate-rich and silica-rich glasses (e.g.,

FIGURE 2. Texture of experiment GG405 (Jd + K_2CO_3): Compaction and recrystallization of jadeite (dark gray) in center of capsule (black is epoxy, replacing plucked jadeite grains), finer unrecrystallized jadeite mixed with K_2CO_3 at bottom, and formation of immiscible carbonate (devitrified blobs) and silicate liquids at hot sides of Pt capsule.

GG477 and GG405; Fig. 2), and in a mixed carbonatebicarbonate experiment (GG499) a devitrified, void-rich sphere occurred in a silicate glass, presumably an immiscible carbonate-rich and H₂O-rich liquid. A result of larger thermal gradients and total Cpx recrystallization is thermally induced composition profiles (see Fig. 3). The generally smooth boundary between melt and carbonate and Cpx is an isotherm, and compositions were measured, if possible, adjacent to this boundary. Compositional data for glasses, carbonate, and non-Cpx phases will be reported in a separate treatment of the solubility characteristics and solution modeling.

The other crystalline experimental products typically exhibit distinctive habits: eskolaite (Cr_2O_3) forms hexagonal plates, garnet forms dodecahedra, potassium-wadeite $(K_2Si_4O_9)$ and forsterite form rhombic prisms or blebs, magnesiochromite and $KSiCr_3O_7$ forms blebs, and a phase with the formula $K_{12-x}Mg_{11+2x}(Cr,Al)_{2-x}Si_{14}O_{48}$ forms a boundary between Cpx and glass (complete information on new phases will be reported elsewhere).

K in Cpx

These experiments have produced K contents in Cpx exceeding amounts previously cited, and the compositions of Cpx products are appropriate to pyroxene crystal chemistry. K enrichments are measurable under all conditions attempted except when no potential balancing $M1^{3+}$ cation is present, further substantiating pyroxene stoichiometry. The cation totals are invariably 4 per 6 O atoms, suggesting no vacancies, even in the H-bearing experiments, and K + Na is balanced by ^[6]Al + Cr (Fig.

3c). Tetrahedral substitution of Si by Cr is not observed, but some Al substitution occurs at higher temperatures (GG416 in Table 3). Maximum content achieved so far is ~ 4.7 wt% K₂O at 10 GPa in GG412 and GG421 (see Tables 3 and 4) with a molar Kcpx component of 22% [here Kcpx = KCrSi₂O₆ but in general it will be used to represent K(Cr or Al)Si₂O₆]. Whereas high-resolution transmission electron microscopy has not as yet been conducted on products, several X-ray diffraction studies (on GG421 and TT48) by both powder and single-crystal methods show only sharp diffractions of C2/c Cpx without any other diffraction artifacts or phases. Results of these diffraction studies including structures will be presented elsewhere.

Experiments with K_2CO_3 produce the greatest K enrichments in Cpx for a given *P* and *T* simply because of partitioning response, but these compositions are usually restricted to rims on unreacted starting crystals. Higher temperature experiments produce greater recrystallization (and melting and carbonate-silicate liquid immiscibility) but lower K content in the Cpx (in part because of dilution of K_2O in the increased volume of melt and carbonate). Addition of H by means of KHCO₃ to the charge produces large (> 100 µm) homogeneous Cpx crystals, probably in response to increased fluxing and recrystallization, but maximum K content can be diminished by the dilution of K in the carbonate.

Several important features of K uptake in Cpx are evident from the experiments. First, K uptake depends on the composition of Cpx involved, which appears to reflect the role of cell volume, in part. This aspect is clearest with the isomorphous exchanges of KNa-1 in jadeite and kosmochlor; in experiments at comparable conditions (GG415 and GG405) the level of K is two to three times greater in kosmochlor, with the larger V_c (~419 vs. ~401 Å³) and M1 polyhedron (10.55 vs. 9.37 Å³), than in jadeite. Second, however, in the Cpx mixtures studied here, maximum K uptake occurs in intermediate solid solutions of Di-Jd or Di-Ko rather than in compositions closer to diopside, which should have greater $V_{\rm e}$. Third, Kcpx solution into diopside is accompanied by Jd or Ko solution in a sympathetic but not necessarily correlated manner (Na:K≠1:1 or some other fixed ratio; see Fig. 3d and Table 4). These last two observations suggest (1) an average structure effect, that is some amelioration of fitting K into the crystal by mixing Na on the M2 site or Al or Cr and Mg on the M1 site or (2) deficit volume of mixing along the Di-Jd or Di-Ko join at high P. As for this latter possibility, data at ambient conditions for Di-Jd solid solutions (Rossi et al. 1983) show no departure from ideal mixing. Finally, a major problem in interpreting K-uptake P, T systematics for diopsidic Cpx is the compositional effect (describe above); local recrystallization of admixed pyroxenes yields different Di contents and thus different Kcpx as well. Nevertheless the data demonstrate (most persuasively for the $Di_{86}Ko_9Qtz_5 + K_2CO_3$ experiments and Cpx with about the same Di content, see Table 4) that K uptake, and perhaps partitioning, is dependent positively on P. The dependence







1.0

0.9

FIGURE 3. (a) BSE image of experiment GG299 that used a long narrow capsule and probably produced a thermal gradient > 200 °C. The bright boundary is the Pt capsule; the pyroxene with convex form has coalesced at the cold end of the capsule due to thermal compaction and quenched carbonate-silicate glass fills center and hottest portion (beyond edge of photo). (b) Pyroxene compositions from probe traverse from cold end to carbonate contact in GG299; compositions show thermally induced zoning from K-Cr-rich cold end to Di-rich hot end. Some of the compositional variation is due to surface cracks and some presumably to incomplete recrystallization. (c) and (d) are plots of Cpx cations per 6 O atoms vs. K + Na and K cations, respectively, for a complete suite of analyses on GG299 Cpx. The term $M^{3+} + 2Ti + {}^{6i}Al - {}^{4i}Al$ (where $M^{3+} = Cr^{3+} + Fe^{3+}$) represents the charge balancing cation sum to balance the alkalis if Cpx crystal chemistry is to be maintained. Regression lines show in (c) show good agreement with ideal stoichiometry.



FIGURE 4. Pyroxene composition profile across a single grain showing the sharp boundary between Di core and Ko- + Kcpx-rich overgrowth in experiment GG389.

on T is unclear from most of these experiments largely because of a lack of T variation among the data (which is in turn due to minimal reaction kinetics at lower T and excessive melting at high T) and in some part because of a lower T accuracy from gradients and Tc junction variations. The thermal diffuse zoning in GG299 and a few comparisons among the data argue for K uptake depending inversely on T.

The average structure effect has been observed in a natural K-rich Cpx ($Ca_{0.80}K_{0.073}Na_{0.023}Mg_{0.95}Fe_{0.06}Cr_{0.07}$ Al_{0.02}Si_{1.99}O₆; Harlow 1996), an inclusion in a Koffiefontein diamond. In the average structure of this Cpx, 10% (Fe + Mg) in M2 (with radii 0.92 and 0.89Å, respectively) balances the 7% K (with radius 1.51Å) yielding an average M2 polyhedron only slightly larger than in diopside, 25.89 vs. 25.76 Å³, respectively. Moreover, the substitution of Fe + Mg into M2 is charge balanced in M1 by Mg, which is larger than Al or Cr, improving the fit for large occupants in M2 further (see Harlow 1996). Experiments with the mixture $Di_{70} + Ko_{20} + En_{10}$ were conducted to reproduce similar Cpx, but a new KMgCr silicate and K-rich Cpx without any significant En content resulted. Nonetheless, if this kind of effect operates in the

experimental K-rich Cpx of intermediate Na-Ca composition, Mg is not the size-averaging cation.

Partitioning

Another purpose of these experiments is to estimate the partitioning between K-rich media and K-Cpx as a minor component in Di-Jd or Di-Ko solid solutions appropriate to the formation of K-rich Cpx in diamonds and, indeed, other environments. K has always been treated as a trace element in Cpx, with a silicate melt partition coefficient, $^{Cpx/liq}D_{K}$, in the range of 10⁻⁴ at low pressure to 10⁻² at 2– 3 GPa (Hart and Dunn 1993; Shimizu 1974). The results here (Table 4) show that $C_{px/liq}D_{K}$ (for an admittedly carbonate-rich melt) is greater at high pressure than previously measured and, from the few coexisting immiscible carbonate and silicate melts, it is generally greater than $^{Cpx/carb}D_{K}$. Moreover, generally the data demonstrate D_{K} is a function of Cpx composition. There is not a systematic relationship for D with the K₂O content of the glass-carbonate or for any single group of experiments. Strictly, the data indicate that there is a maximum saturated Kcpx content in a particular pyroxene at a given P and T in the experiments, whereas the K content of the glass-carbonate varies according to bulk composition and the phase assemblage. The plateau of D values, particularly in the intermediate Di-Nacpx, indicates the maximum for D is near 0.1 at 10 GPa and 1500 °C. However, the Cpx solutions are actually controlled by components, so a more complex formulation is necessary, particularly for the complex Di-Ko-K₂CO₃ system, and is the subject of further examination. Finally, the data also demonstrate the nearly compatible nature of Na in diopsidic Cpx with Cpx/ $^{liq}D_{Na}$ values between ~0.5 and 1, but Na is compatible in more sodic Cpx when values of D reach values of about 10 in jadeite. Again, D_{Na} varies with the composition of the Cpx.

DISCUSSION

An essential issue in this study is the mechanism of solubility of K in Cpx that is enabled by high pressure. The effects of high pressure on crystal chemistry are usually viewed in terms of phase transformations in which coordination number increases, such as ^[4]Al (in albite) \rightarrow ^[6]Al (in jadeite) or ^[4]Si (in pyroxene) \rightarrow ^[6]Si (in silicate perovskite). It is not surprising that observations of K in Cpx from mantle xenoliths were viewed skeptically. Many of such samples were shown to contain exsolved and contaminant K-rich phases, and the coordination number of K in Cpx (presumably 8-7) is not larger than in the contaminant phases (14-12 in phlogopite; 9-8 in sanidine). However, a less well-studied relationship of crystal chemistry and pressure is differential ionic or polyhedral compressibility as it affects substitutions. Larger, lower-charge (low field strength-low FS) ions have relatively greater polyhedral compressibility in comparison with smaller, higher charge (high FS) ions (Hazen and Finger 1982; Hazen 1988) and generally greater linear bond compressibility (Kudoh et al. 1992 and Table 5

 TABLE 3.
 Microprobe analyses of experimentally produced K-rich pyroxene

	Weight percent oxides									
	1	2	3	4	5	6	7	8	9	10
SiO ₂	55.53	54.29	53.96	59.68	53.30	55.21	57.48	52.94	53.03	54.92
TiO ₂	0.04	0.00	0.01	0.00	0.00	0.02	0.00	0.01	0.00	0.05
AI_2O_3	0.11	0.00	0.15	24.93	0.17	11.25	13.37	0.09	0.14	0.38
Cr_2O_3	3.44	6.10	6.80	0.01	34.05	0.01	0.00	17.92	19.45	8.85
Fe ₂ O ₃ *	0.18	0.03	0.18	0.18	0.00	0.08	0.02	0.00	0.00	0.00
FeO	0.00	0.00	0.00	0.07	0.00	0.00	0.00	0.00	0.00	0.00
MgO	16.95	14.82	14.20	0.19	0.05	10.72	9.06	8.57	7.96	13.94
CaO	22.87	20.49	19.66	0.35	0.02	14.57	12.59	10.90	10.18	18.43
Na₂O	0.60	0.80	0.99	14.42	12.72	4.35	6.23	4.74	4.95	1.73
K₂O	1.58	2.50	3.09	1.09	1.56	3.10	2.60	4.39	4.51	3.13
Total	101.30	99.04	99.05	100.92	101.87	99.32	101.35	99.57	100.22	101.44
				Cation	ns for 6 oxyg	en atoms				
Si	1.996	2.007	2.003	2.000	1.990	1.969	1.990	1.996	1.991	1.993
^[4] AI	0.004	0.000	0.000	0.000	0.008	0.031	0.010	0.004	0.006	0.007
SUM T	2.000	2.007	2.003	2.000	1.998	2.000	2.000	2.000	1.997	2.000
Ti	0.001	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.001
^[6] AI	0.000	0.000	0.007	0.985	0.000	0.442	0.536	0.000	0.000	0.010
Cr	0.098	0.178	0.200	0.000	1.005	0.000	0.000	0.534	0.577	0.254
Fe ^{3+*}	0.005	0.001	0.005	0.005	0.000	0.002	0.000	0.000	0.000	0.000
Fe	0.000	0.000	0.000	0.002	0.000	0.000	0.000	0.000	0.000	0.000
Mg	0.908	0.817	0.786	0.009	0.003	0.570	0.467	0.482	0.445	0.755
Ca	0.881	0.812	0.782	0.013	0.001	0.557	0.467	0.440	0.410	0.717
Na	0.042	0.057	0.071	0.937	0.920	0.301	0.418	0.346	0.360	0.122
К	0.072	0.118	0.147	0.047	0.074	0.141	0.115	0.211	0.216	0.145
Total	4.007	3.991	4.000	3.997	4.001	4.014	4.004	4.014	4.006	4.003

Notes: Values are for an individual analysis that is typical (within 1 S.D.) for a group of at least 6 analyses. 1. BB470: Di+Ko+Qtz+KC, 1500 C, 10 GPa. 2. GG299: Di+Ko+Qtz+KC, 1700 C, 11 GPa. 3. BB89: Di+Ko+Qtz+KC, 1500 C, 14 GPA. 4. GG405: Jd+KC, 1500, 10 GPa. 5. GG415: Ko+KC, 1500, 10 GPa. 6. GG416: Di₅₀Jd₅₀+KC, 1500 C, 10 GPa. 7. GG499: Di₅₀Jd₅₀+KC+KBC, 1450 C, 10 GPA. 8. GG412: Di₅₀Ko₅₀+KC, 1500 C, 10 GPa. 9. GG421: Di₅₀Ko₅₀+KC+KBC, 1400 C, 10 GPa. 10. GG389: Di₇₀Ko₂₀En₁₀+KC, 1400 C, 11 GPa. KC = K₂CO₃; KBC = KHCO₃. * Fe³⁺ determined by cation sum and charge balancing.

where the interpretation has been extended to K^+ and Na⁺). The complete compressibility data for Cpx do not exist and predictions for structures as complex as in Cpx are, indeed, extrapolations. Nevertheless, the inference is that with increasing pressure a large ion like K⁺ or Na⁺ will behave more like relatively smaller ions Ca²⁺ or Mg²⁺. Given sufficient charge-balancing trivalent cations, with increasing pressure the large ions should have increasing solubilities in Ca or Mg sites where the coordination number (in this case, 7–8 in the phases stable at high-*P*) is near or below the lower limit for the large ions at ambient conditions; temperature should have a reciprocal effect. The experimental data for Kcpx solution in Di, Jd, and Ko are consistent with differential compressibility based on polyhedral values in Table 5.

The experimental results may point to the significant compressibility of Na in Cpx by means of its role in Kcpx solubility in diopside. The size of K appears to be accommodated in the M2 site by spatial averaging in the only structure studied so far, as noted above, and Nacpx content varies sympathetically in diopsidic Cpx produced experimentally. Thus, a smaller Na⁺ at high pressure would provide a mechanism of size averaging for K⁺ to fit into M2 of diopside. Following this hypothesis, a limit is reached because the volume decrease due to Nacpx mixing in diopside ($Vc \approx 439$ Å³ for Di, 401 Å³ for Jd, and 419 Å³ for Ko) reduces the Kcpx solubility as compositions become dominated by Nacpx.

If Na⁺ fulfills the accommodating role for Kcpx solu-

tion in Di at high pressure, Na⁺ must be smaller than Ca²⁺ at pressure. For tabulated radii at ambient conditions where Na^+ is larger than Ca^{2+} (1.18 vs. 1.12 Å). The observed radii in Cpx at ambient conditions are more in line with the high pressure implications; the average M2-O bond lengths in Jd and Di at ambient conditions, 2.469 and 2.498 Å respectively (Cameron, et al. 1973), yield radii of 1.09 Å for Na⁺ and 1.12 Å for Ca²⁺ in coordination with ^[iv]O. Using these bond lengths from Cpx and values of β_c from Table 5, the calculated average bond lengths at 10 GPa are 2.389 Å for jadeite vs. 2.424 Å for diopside, only 0.006 Å greater than the difference at ambient conditions. The polyhedral compressibilities, β_{n} , indicate a much greater difference between the alkalis and Ca²⁺. The contrast in observed polyhedral volumes $V_{p(0)}$ at ambient conditions and calculated values at 1500 °C and 10 GPa, $V_{p(P,T)}$, (see Table 5) could indicate both the size-accomodating behavior of Na at pressure and the greater compatibility of K in the M2 polyhedron of Cpx. However, actual configurations, bond lengths and polyhedral volumes, in high pressure structures of Nacpx and experimentally grown K-rich Cpx are needed to test these hypotheses.

These experimental results, which indicate that Kcpx content depends on high pressure and is correlated positively with *P* and perhaps negatively with *T* for a coexisting K-rich melt, can be compared with some data for natural Cpx. K contents in Cpx from diamonds where *P* > 5 GPa, $T = 1000^{\circ}-1400$ °C (Harlow and Veblen 1991)

TABLE 4. Compositions and partition coefficients (D) for K₂O and Na₂O

			K₂O	(wt%)		Na ₂	O (wt%)			
Starting mixtures/ Cpx composition	<i>P</i> (GPa)	<i>T</i> (°C)	Срх	Carb or glass	D (K ₂ O)	Срх	Carb or glass	D (Na ₂ O)	Matrix	Experiment
				Di	₀Ko₀Qtz₅ +	K ₂ CO ₃				
Di ₈₆ Ko ₅ En ₄ Kcpx ₄ Di ₈₈ Ko ₅ Kcpx ₅ En ₂ Di ₈₈ Ko ₅ Kcpx ₅ Di ₈₂ Kcpx ₁₁ Ko ₄ En ₂ Di ₆₀ Kcpx En *	5 6 7 9	<1500 1400 1500 <1700 1200	0.9 0.9 1.0 2.4 2.5	18(1) 11(1) 28(3) 25(2) 37(2)	0.050 0.082 0.036 0.096 0.068	0.75 0.75 0.66 0.55 1.6	0.9(1) 0.52(2) 0.4(1) 0.9(1) 1.4(2)	0.83 1.4 1.7 0.61 1.1	glass glass glass glass carb	GG404 BB330 BB109 BB243 BB206
Di ₈₈ Kcpx ₇ Ko ₄ En ₁ ** Di ₇₄ Kcpx ₁₆ Ko ₈ En ₁ ** Di ₇₄ Kcpx ₁₆ Ko ₇ En ₁ Di ₈₅ Kcpx ₃ Ko ₄ En ₃ † Di ₈₅ Kcpx ₈ Ko ₄ Di ₈₇ Kcpx ₈ Ko ₄ En ₁ Di ₈₅ Kcpx ₆ Ko ₆ En ₁ Di ₇₅ Kcpx ₈ Ko ₆ En ₁	10 10 11 11 12 12 14 14	1500 1500 1500 1500 1700 1500 1500 1500	1.6 3.6 2.1 1.8 1.7 1.7 2.2 3.2 3.5	48(1) 48(1) 30(3) 36(2) 37(1) 34(3) 34(3) 51(2) 51(2)	0.033 0.075 0.070 0.050 0.046 0.050 0.065 0.063	0.6 1.1 1.0 0.60 0.6 0.6 0.8 0.9 0.9	$1.7(1) \\ 1.7(1) \\ 1.5(3) \\ 0.9(1) \\ 1.4(9) \\ 1.1(3) \\ 1.1(3) \\ < 0.02 \\ < 0.02$	0.35 0.65 0.67 0.67 0.43 0.55 0.73 45	glass/carb glass/carb glass/carb devit glass devit glass devit glass devit glass devit glass devit glass	BB470 BB470 GG477 GG307 GG299 BB85 BB85 BB85 BB89 BB89
D1771 (OPX171 (O6	14	1000	0.0	51(2)	'ld + K C	0.0	<0.02	-10	ucvit glass	BB00
Jd ₉₉ Kcpx₁ Jd ₉₆ Kcpx₄	11 10	1300 1500	0.14 1.0 1.0	? 46(2) 26(1)	0.022 0.039	14.8 14.2 14.2	? 3.5(9) 2.3(1)	4.1 6.2	carb carb glass	GG229 GG405 GG405
Ko ₉₆ Kcpx₄ Ko ₉₂ Kcpx ₈	11 10	1300 1500	0.86 1.6 1.6	? 57(3) 44(2)	0.028 0.036	13.0 12.8 12.8	? 2.2(3) 9.3(4)	5.8 1.4	carb carb glass?	GG227 GG415 GG415
				[Di ₅₀ Jd ₅₀ + K	22 CO 3				
$\begin{array}{l} Di_{53}Jd_{31}Kcpx_{13}Ct_3En_1\\ Di_4Jd_{90}Kcpx_6 \end{array}$	10 10	1500 1500	2.9 1.2	44(2) 44(2)	0.066 0.027	4.5 13.5	3.6(4) 3.6(4)	1.2 3.8	carb carb	GG416 GG416
				Di₅₀Jo	I₅₀ + KHCO	₃ + K ₂ CO ₃				
$\begin{array}{l} Di_{49} Jd_{37} Kcpx_8 Ct_4 En_2 \\ Di_{32} Jd_{58} Kcpx_7 Ct_2 En_1 \\ Di_{46} Jd_{40} Kcpx_{10} Ct_2 \dagger \\ Di_{56} Jd_{36} Kcpx_5 Ct_1 \dagger \end{array}$	10 10 10 10	<1400 <1400 1450 1450	1.9 1.6 2.3 1.2	62(2) 62(2) 25(1) 33(1)	0.031 0.026 0.092 0.036	5.5 8.6 6.0 5.0	0.8(2) 0.8(2) 2.0(1) 2.8(2)	6.9 10. 3.0 1.8	carb carb glass glass	BB308 BB308 GG499 GG568
				Di₅₀Ko	0 ₅₀ + K ₂ CO ₃	± KHCO₃				
Ko ₆₄ Di ₂₃ Kcpx ₁₂ En ₁ † Di ₄₃ Ko ₃₄ Kcpx ₂₂ En ₁ Di ₃₈ Ko ₃₉ Kcpx ₂₂ En ₁	10 10 10	1450 1500 1400	2.4 4.6 4.6 4.7	33(2) 53(1) 40(3) 51(2)	0.073 0.087 0.12 0.092	8.7 4.6 4.6 5.5	2.0(4) 4.5(1) 4.3(1) 1.2(2)	4.4 1.0 1.1 4.6	glass carb glass carb	GG486 GG412 GG412 GG421
Ko55Di27Kcpx16En2	11	1500	3.2	39(2)	0.082	7.5	2.5(1)	3.0	glass	1148
				Di70K020	$En_{10} + K_2C$	$O_3 \pm KHCC$	D ₃			
$\begin{array}{l} Di_{72}Kcpx_{16}Ko_{13}\\ Di_{71}Kcpx_{14}Ko_{13}\\ Di_{70}Ko_{12}En_2Kcpx_{16} \end{array}$	11 11 10	1350 1450 1300	3.5 3.2 3.3	60(2) 47(3) 47(3)	0.058 0.068 0.062	1.8 1.8 1.7	0.8(2) 2.0(3) 1.6(3)	2.25 0.9 1.06	carb carb carb	GG389 GG389 BB211

Note: ? indicates that compositions could not be determined.

* lone experiment using KHCO₃ instead of K₂CO₃. ** Temperature was higher by 50° for the first hour; < and > as in Table 2. † Temperature was higher by 100° for the first hour.

TABLE 5. Comparison of linear compressibilities and polyhedral compressibilities (Mbar-1)

	β_{c}	β _o	β _p (O)	$\beta_{p}(F)$	$\beta_{p}(Cpx)$	α_{p}	$V_{\rm p(0)}$	$V_{p(P,T)}$
Cr	0.14	0.14	0.43			1.69	10.51	10.31
AI	0.16	0.14	0.42			2.65	9.37	9.33
Mg	0.22	0.21	0.62-0.67	1.0	0.95	4.25	11.85	11.39
Ca	0.30	0.30	0.91-1.18	1.16	0.99	4.42	25.76	24.72
Na	0.33*		3.13	2.22	Jd	3.74	24.58	20.18
					Ko	3.50	25.36	20.75
К	0.30*		3.7	3.413		3.74	36.40	25.30

Notes: β_c = calculated linear bond compressibility and β_o = observed linear bond compressibility data from Kudoh et al. (1992); $\beta_p(O)$ = polyhedral compressibility for Various O polyhedra and $\beta_p(F)$ = polyhedral compressibility for F in fluorides from Hazen and Finger (1982); $\beta_p(Ox)$ data: Di from Levien and Prewitt (1981); α_p = polyhedral thermal expansivity for Cpx (Cameron et al. 1973) except K which is from Hazen and Finger (1982) for 8-fold coordination; $V_{p(0)} = polyhedral volume in Cpx except for K for which the microcline value is used (Smyth and Bish 1988); <math>V_{p(P,\eta)} =$ the calculated polyhedral volume at 1500 °C and 10 GPa using α_p and β_p (Cpx) for Mg and Ca, $\beta_p(O)$ for Cr & AI, and $\beta_p(F)$ for K and Na (which are less than $\beta_p(O)$) for Na and K). * The equation $\beta_c = 0.217 [\Delta r / \Delta \log(z/n)]^2$ (Kudoh et al. 1992) was used with radii (*r*) from Shannon (1976).

range as high as 1.5 wt% K₂O in the Di₈₀ Cpx noted above. K-rich diopside (1.1–1.5 wt% K₂O, foremost as Di₈₈Kcpx₅Jd₃Cts₂Fs₂) is included in garnet grains from UHP metamorphic rocks from Kazakhstan with estimates of $P = \sim 4$ GPa, $T = \sim 1000$ °C, (Sobolev and Shatsky 1990; Sobolev et al. 1991b). These conditions are potentially subsolidus but in a very limited way show a similar relationship as these experiments.

Only limited prior experimental data exist for comparisons. Earlier experiments showed a small increase of K in diopside in synthetic basaltic melts containing between 1.2 and $\sim 3 \text{ wt}\% \text{ K}_2\text{O}$ as an anhydrous or oxvphlogopite component: ≤0.017 wt% K₂O at 32 kbar and 1450 °C yielding $^{Cpx/liq}D_{\kappa}$ of 0.0013 (Erlank and Kushiro 1970) and 0.26 wt% K₂O at 10 GPa and 1400 °C (Shimizu 1971). These values are very low compared with the data presented here, and the difference may be due to the absence of Na in the starting material. Thus even though Al is available for charge compensation of K⁺ in Cpx, the absence of Na in its perceived role as a size compensator in the M2 site may prevent the formation of K-enriched Cpx. Other experiments by Erlank and Kushiro (1970) using omphacite + anhydrous phlogopite or diopside + potassic richterite component at 24-25 kb and 1000 °C resulted in 50 ppm or less K in Cpx, in spite of the presence of Na in the Cpx (no compositions reported). It is possible that at 1000 °C the reactions were too sluggish or zones of K-rich Cpx too small to be observed.

The results of Edgar and Vukadinovic (1993) for Cpx in lamproite partial melt at 1200 to 1500 °C and 5 to 6 GPa are comparable to the results here. Maximum K₂O content of ~1.6 wt% K₂O is produced at lowest T and highest P in Di₅₇Jd₁₅₋₁₉ Cpx, and Kcpx content drops off simultaneously with increasing T, En content, and decreasing K₂O content of the coexisting melt. All these factors should tend to decrease Kcpx content based on the observations made here. The partition coefficients $C_{px/liq}D_{\kappa}$ of Edgar and Vukadinovic (1993) show a somewhat higher maximum value, 0.19 at 1200 °C and 5 GPa, but the value decreases dramatically with increasing T. This highest value may represent a closer convergence of the melt composition to the optimum for Kcpx solution. The dramatic changes in $^{Cpx/liq}D_{K}$ in their experiments undoubtedly manifest the effect of the En content (decreased V_c) of Cpx on its capacity to accept a Kcpx component. Consequently, their results appear consistent with those reported here but should only be applied to Cpx of the compositions they report.

Because Kcpx solid solution in diopsidic Cpx appears to depend on Nacpx solid solution as well, the pressure dependencies of Nacpx-Di miscibility are an important factor in the formation of K-rich Cpx. The pressure dependence of jadeite stability and Di-Jd solubility is well known (Gasparik 1985), such that low pressure formation of KAlSi₂O₆ component in diopsidic Cpx is not expected. For conditions of high P/T metamorphism, recent data for metasomatic omphacite coexisting with phengite formed at ~ 400 °C and 6–10 kbar showed K₂O contents only at the detection limit, ~ 0.01 wt% (and so were not reported in Harlow 1994), consistent with other data for comparable conditions. One infers that the values of D for omphacite-phengite or omphacite-K-feldspar are much lower at these conditions than for Cpxmelt or Cpx-carbonate at 10 GPa: 10⁻³ vs. 10⁻¹. Kosmochlor-diopside solutions may be a better subject for lower pressure evaluation as kosmochlor is stable at ambient pressure even though such Cpx is restricted to mantle xenoliths and rare metasomatic high P/T environments (Harlow and Olds 1987). Unfortunately, the experimental data for the Di-Ko system are contradictory. Partial miscibility of Ko in Di is reported by Ikeda and Yagi (1972) at 1 ATM, a maximum content of Ko_{23.1} at 1140 °C (probably as low as 1050 °C) that decreases upon increasing T. On the other hand, complete miscibility is reported by Yoder and Kullerud (1970) at 0.2 GPa, and yet Vredevoogd and Forbes (1975) state that Ko solubility decreases substantially below 1 bar values at 2 GPa and that Di is insoluble in Ko. Di-Ko solubility may be nearly complete at 10 GPa and 1500 °C, on the basis of observations presented here. Clarifications are required in the Ko-Di system before inferences can be made about KCrSi₂O₆ component solubility in diopsiderich members at lower pressures.

IMPLICATIONS FOR THE MANTLE

These results show that diopsidic pyroxene may be more useful for measuring the K content in the mantle than K-rich minerals. Mica and amphibole appropriate to mantle composition are not stable at high P; they undergo major coordination-changing or dehydration reactions with depth in the mantle, breaking down between 30 and 200 km (1-6 GPa at 1000 to 1300 °C; Sudo and Tatsumi 1990). Potassium-feldspar is a stable host for K only up to ~7 GPa and 1500 °C (Yagi et al. 1994) and was found only rarely in eclogitic xenoliths and diamond inclusions (Schulze and Helmstaedt 1988). The higher pressure phases K₂Si₄O₉ wadeite and KAlSi₃O₈ hollandite are stable at high P but have never been recovered in natural samples. Whereas K-rich phases may be exceedingly rare or absent (or missed in thin section) in typical mantle rocks, Cpx is not likely to be missed, may be the fundamental solid host for K in most of the upper mantle, and should be a very good monitor of K activity in the mantle.

The values of $^{Cpx/liq}D_{\kappa}$ measured here, although reflecting an unusual composition, can be used along with the data of Edgar and Vukadinovic (1993) to put some limits on K contents in coexisting liquids. K-rich Cpx found as inclusions in diamonds results from high pressure and high K activity: the data here imply that a diopside with ~ 1 wt% K₂O forms in the presence of a C-rich meltfluid with 14 to 33 wt% K₂O; for omphacite a slightly less K₂O-rich melt-fluid would be needed. Very K-rich environments are indicated for these Cpx found in diamonds.

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