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Crystal chemistry of diopsides from garnet Iherzolites (Cima Lunga/Adula Nappe, Central Alps)

by Paolo Nimis¹

Abstract

The first crystal chemical data on garnet pendotite clinopyroxene are presented through the study of ^a suite of diopsides from metamorphic garnet Iherzolites from the Cima Lunga/Adula Nappe, Central Alps (CL-Cpx).

CL-Cpx have low Al^{iV} occupancies and moderate, though variable, Na and \widehat{Al}^{γ} contents. Small Al^{IV} may be due to fractionation of Al₂O₃ into coexisting garnet \pm amphibole, and involves low T site volumes (2.218–2.226 Å³). Contraction of the almost Si-saturated tetrahedron favours expansion of adjacent M1 polyhedron and requires relatively low AI^{VI}. This produces relatively large cell volumes (435.6–438.3 \AA ³), despite the relatively high-pressure origin (garnet stability field).

The crystal chemical evolution of CL-Cpx is characterized by a rimward NaAl^{VI} depletion causing significant cell volume increase. Fe3+ appears not to be negligible in CL-Cpx and decreases as volumes increase. These changes are consistent with ^a decompressional evolution involving breakdown of garnet and formation of ^a spinel + amphibolebearing assemblage. The different NaAl^{VI} contents of the clinopyroxenes from the different localities may reflect either different original equilibration pressures or compositional effects, such as different $H₂O$ contents.

Petrographic features and mineral chemistry suggest that breakdown of garnet occurred at $T < 800$ °C (perhaps even lower than 700 °C) and $P < 16$ kbar.

Keywords: clinopyroxene, crystal chemistry, garnet peridotites, Cima Lunga/Adula nappe, Central Alps.

Introduction

Abundant crystal structural data available on C2/c clinopyroxene (Cpx; Fig. 1) (CARBONIN et al., 1991, and references therein) show that cation substitution mechanisms depend on either the physico-chemical conditions of formation or the intracrystalline constraints induced by geometrical variations of the coordination polyhedra. Detailed crystal chemical studies are therefore essential to understand the behaviour of Cpx in different petrogenetic environments.

Available data on peridotite Cpx are related to spinel- or garnet + spinel-bearing nodules (Dal NEGRO et al., 1984; CUNDARI et al., 1986; PRINCIvalle et al., 1989; Nimis, 1994), representing mantle melting residua, and to massif Iherzolites reequilibrated in the plagioclase facies (Nimis, 1994). In the present work the first crystal chemical data on garnet peridotite Cpx are presented through the study of ^a suite of diopsides from

metamorphic garnet Iherzolites from the Cima Lunga/Adula Nappe (Central Alps), hereafter belled CL-Cpx. Moreover, the structural refinement provides an independent evaluation of the chemical composition (Dal Negro et al., 1982) and may therefore be used to constrain site pancies and chemical data, which are crucial to geothermometric estimates. A list of abbreviations used in the text is given in table 1.

Geological and petrological outline

Four garnet peridotites from Cima di Gagnone (CG) , Verzasca Valley, Alpe Arami (AA) , northwest of Bellinzona, Switzerland, and M. Duria (MD), north of Lago di Como, Italy, were selected for crystal chemical analysis. These localities belong to the same tectonic unit, namely the Cima Lunga/Adula nappe.

Detailed geological and petrological outlines

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clinopyroxene(s);
Cima Lunga/Adula;
Alpe Arami;
M. Duria;
Cima di Gagnone;
temperature;
pressure;
atoms per formula unit;
calculated;
observed:
total:
mean electron density on s site;
cell and site volumes; Vc, VM1, VT, VM2
mean M1 cation-oxygen bond
length;
mean T cation-non bridging
oxygen (O1 and O2) bond
length;
sum of $(A^{13+} + Fe^{3+} + Cr^{3+} + Ti^{4+})$
in M1 site;
Al in tetrahedral (T site)
or octahedral (M1 site)
coordination.

Tab. 1 Legend to abbreviations.

were given e.g. by EVANS and TROMMSDORFF (1978), Fumasoli (1974), Ernst (1978), and TROMMSDORFF (1990). Briefly, the garnet peridotites outcrop as small lenses or macroboudins accompanied by a variety of rocks comprising leucocratic migmatitic gneisses, semi-pelitic gneisses, marbles and mafic dykes (eclogites and metarodingites). At AA and MD garnet develops as subhedral porphyroblasts. At CG elongate poikiloblastic garnet is confined to strongly deformed bands (PICCARDO et al., 1993) and overgrows preexisting folds marked by an amphibole + py roxenes \pm olivine-bearing assemblage.

The garnet peridotites are related to ^a regional eclogitic-facies metamorphism (HEINRICH, 1986). $P > 20$ kbar and $T \approx 800$ °C, consistent with the P -T derived for the associated eclogites, were estimated by Evans and Trommsdorff (1978). More severe conditions were inferred for AA by Carswell and Gibb (1980; 836-893 °C, 29.4- 33.0 kbar) and by Ernst (1981; 950 °C, 40 kbar). During the subsequent meso-Alpine amphibolitefacies metamorphism, garnet breaks down into ^a spinel + amphibole kelyphite and, in H_2O -rich domains, into late chlorite clots. Kelyphite around garnet is generally less developed at CG than at AA and MD, but is present in all the investigated specimens.

The close association of rocks pertaining to different lithospheric levels has been interpreted as ^a lithospheric mélange formed in ^a subduction

Fig. I Perspective view of C2/c pyroxene structure perpendicular to the (100) plane with atom and silicate chain nomenclature (Burnham et al., 1967) adopted in this paper.

zone through tectonic erosion (TROMMSDORFF, 1990), followed by tectonic exhumation and very rapid uplift (Vance and O'Nions, 1992). Evidence of Eocene/Oligocene age for the subductionrelated eclogitic metamorphism has been recently provided by GEBAUER et al. (1992) and by BECKER 1993). The possibility that AA garnet peridotites may have much older protoliths and may have experienced other high-pressure events before Eocene (at 1.2 Ga and 650 Ma) has been posed by GEBAUER et al. (1992) and by TROMMS-DORFF (1990).

Analytical methods

Fragments (0.1-0.2 mm) of Cpx grains were handpicked from ca. ¹⁵⁰ mm-thick rock sections, choosing optically homogeneous core and rim portions. Structural data (Tab. 2) were obtained through single-crystal X-ray diffraction on ^a mens AED II four-circle automated diffractometer with M o $K\alpha$ radiation monochromatized with ^a flat graphite crystal following procedures scribed by DAL NEGRO et al. (1982). EDS electron microprobe analyses of the same crystals used for structural refinement were carried out with an ETEC-AUTOSCAN system operating at ¹⁵ kV and ⁵ nA, equipped with an ORTEC EEDS spectrometer. A MAGIC program (COLBY, 1972) in the ORTEC MAGIC IVM version was used to

DIOPSIDES FROM GARNET LHERZOLITES (CENTRAL ALPS)

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 $\ast : I_{\rm hkl} \! > \! 3\sigma(I_{\rm hkl})$

 $\frac{1}{2}$

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Fig. 2 Cell volume vs M1 volume relationship in peridotite clinopyroxenes. Sp : spinel peridotite field (crosses) after DAL NEGRO et al. (1984), CUNDARI et al. (1986), and PRINCIVALLE et al. (1989). Pl: plagioclase peridotite field (stars) after Ntmis (1994). Solid lines: core-rim trends for Cima Lunga samples (cores: filled symbols; rims: open symbols). Error bars correspond to 2σ .

convert elemental X-ray counts into weight % oxides. Results are considered accurate to within 2-5 relative % for major elements and ca. 9% for minor elements. Neither inclusions nor significant zoning were detected in the examined crystal fragments. The program of Papike et al. (1974) was used to convert weight % oxides into atoms per formula unit and to make ^a first estimation of Fe3+ with the charge-balance method. Site partitionings were calculated according to Dal Negro et al. (1982). These authors showed that site partitioning can be constrained on the basis of <Ml-0> mean distance and mean electron sities on M2 and M1 sites, both derived from structural refinement, through the equations;

- (1) 18 Ca + 10 Na + 23 Mn + 24 Fe $_{M2}^{2+}$ + 10 Mg_{M2} = e_{M2}^-
- (2) 10 Mg_{M1} + 24 Fe²⁺_{M1} + 23 Fe³⁺ + 10 Al^{V1} + 21 Cr $+ 18$ Ti = e^{-}_{M1} ;
- (3) 2.081 Mg_{M1} + 2.126 Fe $^{2+}_{M1}$ + 2.03 Fe³⁺ + 1.93 $Al^{VI} + 2.01$ Cr + 1.99 Ti = <M1-O>.

Cation fractions are multiplied by their respective electrons in the former two equations, and by their octahedral cation-oxygen mean distance in the latter. This is particularly important for $Fe³⁺$ estimation, as Fe-O bond lengths are different depending on iron oxidation state $(Fe^{2+}-O)$ 2.126 Å; Fe³⁺-O = 2.030 Å; DAL Negro et al., 1982). This does not mean that the structural finement provides a direct measure of Fe^{2+}/Fe^{3+} partitioning. Nevertheless, good-quality chemical

analyses should fit the above equations within alytical uncertainties. Structural data will be tentatively used to constrain Fe³⁺ estimates in both Cpx cores and rims by selecting those partitionings that yield the best match between observed and calculated values of <Ml-0> and electron densities $\left(\langle M1-O \rangle_{\text{calc}} - \langle M1-O \rangle_{\text{obs}} \right) \leq 0.003$ Å; $(e_{\text{calc.}} - e_{\text{obs.}})_{M1+M2} < 0.3 e$; Tab. 3). These partitionings will be referred to as " constrained analyses ".

Crystal chemistry

Studies on Cpx from peridotites equilibrated in the spinel (DAL NEGRO et al., 1984; CUNDARI et al., 1986; Princivalle et al., 1989), plagioclase and garnet + spinel faciès (Nimis, 1994) showed that, in general, Cpx adapts itself to increasing pressure by reducing site and cell volumes. The largest variations involve the Ml site, whose volume is linearly correlated with the cell volume (Fig. 2). Substitution (Na Ca₋₁)_{M2} (Al^{V1} Mg₋₁)_{M1} is the main responsible for these volume reductions and is therefore strongly pressure-dependent.

CL-Cpx conform to the linear cell volume vs Ml volume relationship (Fig. 2). The Ml volume is correlated with the content of small-sized and high-charged R_{M1}^{3+} (= Al^{VI} + Fe³⁺ + Cr³⁺ + Ti⁴⁺), mainly Al^{VI} (Fig. 3). Substitution of Na⁺ for Ca²⁺ in M2 site balances the charge excess determined by substitution of R^{3+} for bivalent cations in M1 (Tab. 3). Geometrical variations of M2 and T polyhedra are small (Tab. 2), M2 being little fected by the main (Na Ca_{-1})_{M2} substitution occurring in the site, and Al^{IV} in the T site being very low and little variable (0.006-0.040 a.f.u.). A common rimward trend towards higher Ml and cell volumes is characteristic to all specimens, but Cpx from specimen 9-5-2A (Arami) are distinct for their smaller cell and Ml volumes (Fig. 2). The other samples show relatively high Ml volumes but have lower cell volumes with respect to plagioclase peridotite Cpx with similar M1, because of very small tetrahedron (VT = $2.218-$ 2.231 \AA ³). Fe³⁺ appears not to be negligible in the studied Cpx and decreases as volumes increase (Fig. 3). This is more evident if only "constrained analyses" are considered.

Relatively large Ml site of CL-Cpx is related to contraction of the almost Si-saturated T hedron (Fig. 4). Shortening of T-O1 and $T-O2$ $(**T-O**_{nbro})$ distances, which are the most sensitive to $(AI^{iv}Si_{-1})_T$ substitution, is counterbalanced by lengthening of $|M1-O\rangle$. A similar relationship, but at lower pressure and for higher $Fe²⁺_{M1}$ (i.e. longer <Ml-0>), occurs in Cpx from volcanic rocks (Dal Negro et al., 1989). Note that more

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Tab. 3 (cont.)

Fig. 3 (a) Al^{VI} vs M1 volume in peridotite clinopyroxenes. Fields as in figure 2. (b) $Fe³⁺$ vs M1 volume in Cima Lunga clinopyroxenes. Dashed line encloses "constrained analyses" (see text). All other symbols as in figure 2. Fe^{3+} values may vary over the entire range of the figure as a result of \pm 2 wt% relative inaccuracy of SiO₂ detection.

Na-rich suites (e.g. $9-5-2A$) show shorter <M1-O> distances (i.e. higher R_{M1}^{3+}) for the same value of $<$ T-O_{nbrg}>.

Evolution of clinopyroxene from Cima Lunga/Adula garnet lherzolites

The main chemical variations of CL-Cpx involve the exchange:

(4) $(Ca Na_{-1})_{M2} (Mg Al_{-1})_{M1}$.

This exchange is responsible for the compositional zoning from cores to rims. In exchange (4) , Fe³⁺ may appear instead of Al^{VI} , as its negative corre-

Fig. 4 <M1-O> vs <T-O_{non-bridging}> relationship in peri-
dotite clinopyoxenes. Inset: Al^N vs <T-O_{non-bridging}> (CL: this work). Le: Mt. Leura spinel peridotite field (crosses) after DAL NEGRO et al. (1984) . X: clinopyroxenes from basic volcanics (DAL NEGRO et al., 1989). Other symbols as in figure 2.

lation with VM1 suggests (Fig. 3). According to EVANS and TROMMSDORFF (1978), exchange (4) may be related to the formation of amphibole. Depletion of Fe³⁺ in Cpx rims may be instead related to the formation of spinel, which is present in the kelyphite replacing garnet as well as in small recrystallized grains near kelyphite. Accordingly, the reequilibrated NaAl^{VI}-poor Cpx rims are probably not in equilibrium with the original garnet-peridotite assemblage.

Smaller M1 and cell volumes (i.e. higher NaAl^{VI} contents) of Cpx 9-5-2A (Arami) suggest higher pressure conditions with respect to the other studied samples (Fig. 2). The same holds for Cpx cores with respect to rims in each specimen. It is also worth noting that Cpx from specimens 9-5-2A and 9-88, although both from the same locality, fall in clearly distinct fields (Fig. 2). A possible explanatory hypothesis is that: a) the rimward volume increase in the Cpx from each specimen was related to a pressure decrease, which produced a retrograde metamorphism recorded through the passage from a garnet- to a spinel + amphibole-bearing assemblage and through a rimward decrease of Na and Al^{VI} in Cpx ; b) differences in the NaAl^{VI} content of the Cpx from the different localities, and hence their respective positions in the Vc vs VM1 plot (Fig. 2), reflect either different equilibration pressures or compositional effects.

In particular, at CG the presence of a hydrous assemblage prior to garnet formation (PICCARDO et al., 1993) might partly explain the low jadeite content of Cpx cores, as Na and Al would have been fractionated by ^a coexisting amphibole. A higher original pressure for Arami garnet peridotites would be consistent with the estimates of CARSWELL and GIBB (1980) and ERNST (1981) (see also Becker, 1993).

Geothermometry

Previous studies suggested $T \ge 800$ °C for Cima Lunga/Adula garnet Iherzolites (e.g. Evans and Trommsdorff, 1978; Carswell and Gibb, 1980; Ernst, 1981). These estimates were mostly based on pyroxene solvus (Ca-transfer reaction).

As regards garnet-clinopyroxene $Fe²⁺-Mg$ exchange, KROGH's (1988) geothermometer, which matches best experimental data on garnet lherzo-(Brey and Köhler, 1990), yields significantly lower temperatures compared to previous garnet-Cpx geothermometers (\lt 800 °C; Tab. 4). Moreover, the present work points out that $Fe³⁺$ is not negligible in CL-Cpx (Tab. 3). Assuming Fe^{3+}/Fe_{tot} ratio 0.035 in garnet [mean(2) of Fett's (1989) Mössbauer analyses of Arami garnets] and 0.29 in Cpx (mean(13) of "constrained analy-" for CL-Cpx) temperature estimates would be further lowered by some $100 °C$ (Tab. 4). This is clearly a rough estimate, since Fe^{3+}/Fe_{tot} ratio may not be the same for the considered specimens (cf. Fig. 3b) and can only be calculated with large uncertainty, as both ferric and ferrous iron are very low in CL-Cpx (Tab. 3). Nevertheless, table ⁴ gives an idea of the possible bias that may arise from neglecting $Fe³⁺$.

The apparently higher closure temperature for Ca transfer (≥ 800 °C) relative to Fe²⁺-Mg exchange (< 800 °C) would be compatible with a retrograde metamorphic history (Saxena et al., 1986), consistently with the slower kinetics of the former (e.g. BRADY and McCALLISTER, 1983; Loomis et al., 1985).

Concluding remarks

 $CL-Cpx$ are diopsides with low Al^{IV} occupancies and moderate, though variable, Na and Al^{VI} contents. Low Al^{IV} may be due to fractionation of Al_2O_3 into coexisting garnet and (for CG) amphibole, and involves low T volumes. Contraction of the almost Si-saturated tetrahedron favours pansion of adjacent M1 polyhedron and requires relatively low R^{3+} . This may explain the relatively large cell volumes (435.6–438.3 \AA ³) of Cpx cores, despite their relatively high-pressure origin (garnet stability field).

Tab. 4 KROGH's (1988) garnet-clinopyroxene thermometry ($P = 25$ kbar) based on Evans and Trommsdorget is (1978) garnet-pyroxene pairs chemical analyses; (a) all iron as Fe²⁺; (b) Fe³⁺/Fe_{tot} = 0.29 in clinopyroxene and 0.035 in garnet (see text for further explanation).

	Spec.	а	
Arami	$9-5-2C$	767	660
Duria	264.3	764	659
Gagnone	160-4-8	717	617

The crystal chemical evolution of CL-Cpx is characterized by a rimward NaAl^{VI} depletion causing sensible cell volume increase. $Fe³⁺$ appears not to be negligible in CL-Cpx and decreases as volumes increase. These changes are consistent with a decompressional evolution volving breakdown of garnet and formation of a spinel + amphibole-bearing assemblage. The different NAA^{VI} contents of the Cpx from the different localities, and hence their respective positions in the VM1 vs VC plot (Fig. 2), may reflect either different original equilibration pressures or compositional effects, such as different $H₂O$ contents. Breakdown of garnet may have occurred at $T < 700$ °C. At such a low temperature, $P \approx$ ¹⁶ kbar would be sufficient to maintain ^a peridotite within the stability field of garnet (cf. O'Neill, 1981).

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