



Title	Differentiation of melilite nephelinitic rocks in light of experimental study of the system Na ₂ O-CaO-MgO-Al ₂ O ₃ -TiO ₂ -SiO ₂
Author(s)	Onuma, Kosuke; Yagi, Kenzo
Citation	Journal of the Faculty of Science, Hokkaido University. Series 4, Geology and mineralogy, 17(3), 437-449
Issue Date	1977-02
Doc URL	http://hdl.handle.net/2115/36066
Type	bulletin (article)
File Information	17(3)_437-450.pdf



[Instructions for use](#)

DIFFERENTIATION OF MELILITE NEPHELINITIC ROCKS
IN LIGHT OF EXPERIMENTAL STUDY OF THE SYSTEM
 $\text{Na}_2\text{O}-\text{CaO}-\text{MgO}-\text{Al}_2\text{O}_3-\text{TiO}_2-\text{SiO}_2$

by

Kosuke Onuma and Kenzo Yagi

(with 1 table and 9 text-figures)

(Contribution from the Department of Geology and Mineralogy,
Faculty of Science, Hokkaido University, No. 1480)

Abstract

Differentiation of the melilite nephelinitic rock series is discussed in light of recent experimental data of the system $\text{Na}_2\text{O}-\text{CaO}-\text{MgO}-\text{Al}_2\text{O}_3-\text{TiO}_2-\text{SiO}_2$. The crystallization sequence in the join diopside-akermanite-nepheline- $\text{CaTiAl}_2\text{O}_6$ including principal constituent minerals in melilite nephelinite is particularly referred to construct an inferred differentiation trend. The trend is olivine nephelinite \rightarrow olivine-melilite nephelinite \rightarrow perovskite-bearing olivine-melilite nephelinite or melilite nephelinite \rightarrow perovskite-bearing melilite nephelinite \rightarrow corundum and perovskite-bearing melilite nephelinite. When this inferred trend is compared with the observation in natural melilite nephelinitic rocks, it is evident that this differentiation trend is now valid to elucidate the formation of the melilite nephelinitic rock series. The relationship between titaniferous pyroxene and perovskite is also discussed with the data of the join diopside- $\text{CaTiAl}_2\text{O}_6-\text{SiO}_2$.

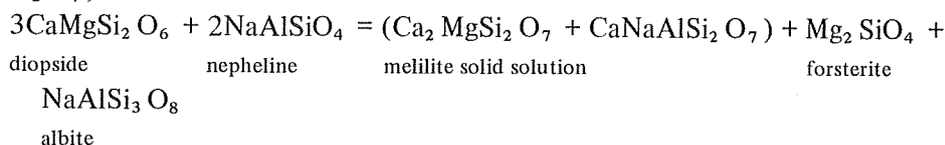
Introduction

The sodium-rich mafic and ultramafic volcanic rocks are characterized by higher Na_2O and TiO_2 and lower SiO_2 compared to rocks belonging to average basaltic rock series. The undersaturated character of these rocks are reflected by the presence of such minerals as nepheline, melilite, titanite and olivine. Some of the interesting rock types, belonging to this family include: nephelinite, melilite nephelinite, olivine-melilite nephelinite and olivine nephelinite, etc. The primary magma, responsible for the formation of these rocks, are believed by many petrologists to have been generated by partial melting of the upper mantle (Bultitude and Green, 1967, Green 1969, Kushiro, 1969, O'Hara, 1970, Bultitude and Green, 1971). However, the genesis and differentiation of these rock types under volcanic and sub-volcanic condition at shallow depths remain an interesting problem. Differentiation of these various rock type at 1 atm will be discussed in this paper in light of the experimental study of the system $\text{Na}_2\text{O}-\text{CaO}-\text{MgO}-\text{Al}_2\text{O}_3-\text{TiO}_2-\text{SiO}_2$.

The System $\text{Na}_2\text{O}-\text{CaO}-\text{MgO}-\text{Al}_2\text{O}_3-\text{SiO}_2$ *Expanded basalt tetrahedron*

First experimental work on the melilite nephelinitic rocks has been carried out by Bowen (1922) who showed the preliminary phase equilibrium diagram of the system diopside-nepheline in which the principal constituent minerals of this rock type were encountered. Bowen also pointed out that it is of importance to study the system $\text{CaO}-\text{MgO}-\text{SiO}_2$ -nepheline to elucidate the genesis and the differentiation of the melilite and nepheline-bearing rocks (Bowen, 1928, p. 164).

Later the system diopside-nepheline was studied in detail by Schairer et al. (1962) and olivine, clinopyroxene, melilite, and nepheline, all of which are main constituent minerals in the melilite nephelinitic rocks, were confirmed in the system. Olivine and melilite may be formed by the following reaction and melilite includes considerable amount of soda-melilite molecule ($\text{CaNaAlSi}_2\text{O}_7$).



Schairer and Yoder (1964) studied the system nepheline-forsterite-larnite-silica including melilite and nepheline composition as a comprehensive model of basaltic rock. Since this is an expanded model of the basalt tetrahedra proposed by Yoder and Tilley (1962), it is called expanded basalt tetrahedron, which includes all the mineral assemblages observed in natural rocks.

The silica-undersaturated part of this system is related to the melilite nephelinitic rock series and the relationship among the mineral assemblages in the system and the rock terminology was given by Schairer and Yoder (1964) as follows: olivine nephelinite, olivine melilitite, olivine-melilite nephelinite, melilite nephelinite, wollastonite-melilite-nephelinite, wollastonite nephelinite, and wollastonite-nepheline tephrite. There are three invariant points in the silica-undersaturated part of the quaternary system; two are reaction points and one is a eutectic point. At one reaction point forsterite is consumed by the reaction with liquid, and at the other melilite is consumed. Such a phenomenon has not yet been described from natural rocks, but the reaction involving melilite is important for the incompatibility of melilite and plagioclase in natural rocks. Further petrographic data on these minerals are expected.

The system diopside-akermanite-nepheline

Bowen (1928) stressed the importance of the system $\text{CaO}-\text{MgO}-\text{SiO}_2-$

nepheline. Onuma and Yagi (1967) studied the join diopside-akermanite-nepheline which is a part of this system. The liquidus diagram of the system diopside-akermanite-nepheline is shown in Fig. 1. The system has no invariant

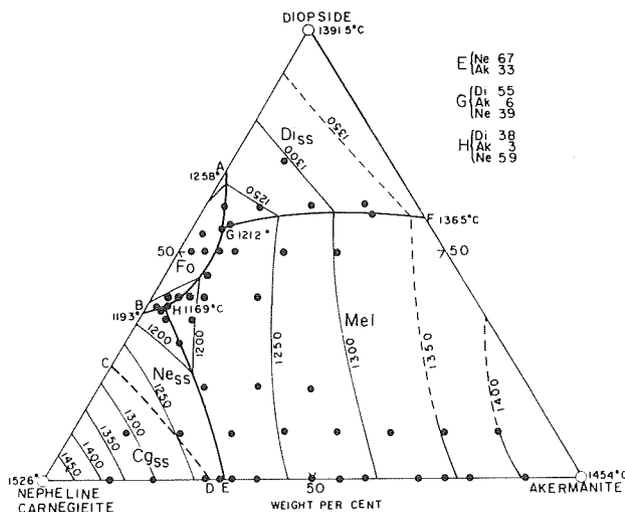


Fig. 1 Liquidus diagram for the system diopside-akermanite-nepheline (after Onuma and Yagi, 1967). Di_{SS} = diopside_{SS}, Fo = forsterite, Mel = melilite, Ne_{SS} = nepheline_{SS}, CG_{SS} = carnegieite_{SS} (ss: solid solution).

point and contains two piercing points G and H. G lies on the univariant line separating the primary phase volumes of melilite, forsterite and diopside in the system CaO-MgO-SiO₂-nepheline, and at this point forsterite, melilite, and diopside coexist with liquid. On the other hand H lies on the univariant line along which nepheline, forsterite, and melilite coexist with liquid. The position of these univariant lines in the system CaO-MgO-SiO₂-nepheline is shown in Fig. 2. Both univariant lines proceed towards the plane of CaO-SiO₂-nepheline with falling temperature. The fractional crystallization along these univariant lines produces a residual liquid rich in CaO and SiO₂. The two univariant lines cross each other at an invariant point in the quaternary system and the liquid coexists with forsterite, diopside, melilite, and nepheline at this invariant point I as shown in Fig. 3. This is also confirmed by Schairer and Yoder (1964). After forsterite is consumed at this point, crystallization proceeds towards another invariant point with falling temperature. During this crystallization the residual liquid becomes enriched in CaO and SiO₂, and finally wollastonite would crystallize out.

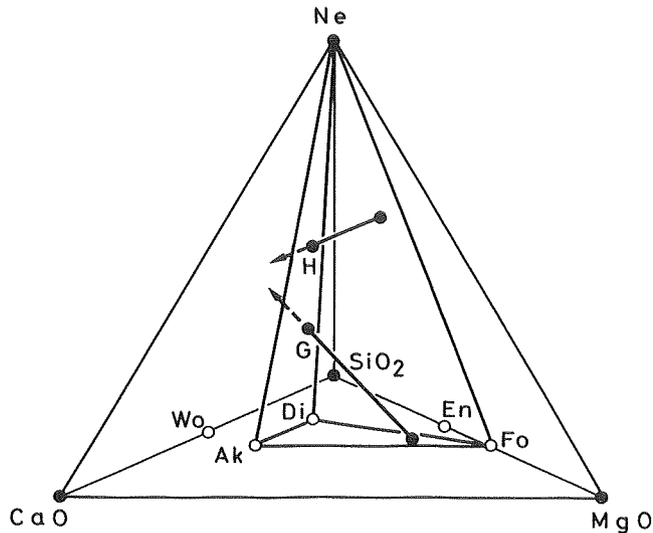


Fig. 2 Nepheline-CaO-MgO-SiO₂ tetrahedron showing two univariant lines and two piercing points in the join diopside-akermanite-nepheline.

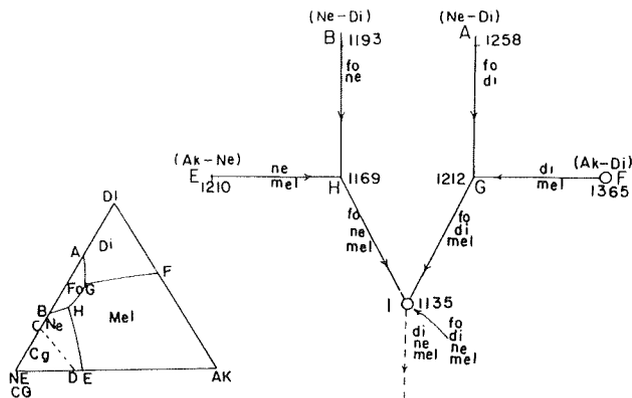
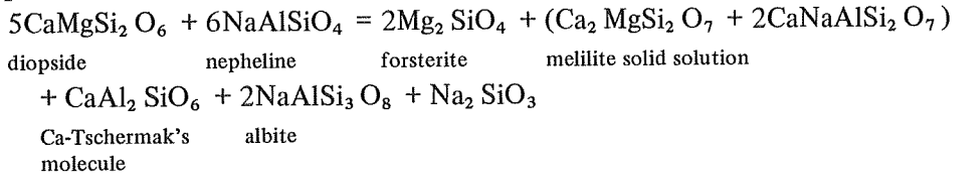


Fig. 3 Flow sheet of the system diopside-akermanite-nepheline showing an invariant point (reaction point) forsterite + diopside + nepheline + melilite + liquid (Onuma and Yagi, 1967).

In the system diopside-nepheline-akermanite the mineral assemblage at G corresponds to olivine melilitite and that at I to olivine-melilite nephelinite. After forsterite disappears by reaction at I, the mineral assemblage represents that of melilite nephelinite. This system is a part of five component system but sometimes the crystallization ceases on a univariant line, resulting a three phase assemblage because of the complex solid solution of the phases encountered in

the system. For example, Onuma and Yagi (1967) have inferred that the diopside contains $\text{CaAl}_2\text{SiO}_6$ (Ca-Tschermak's molecule) and melilite contains $\text{CaNaAlSi}_2\text{O}_7$ (soda-melilite molecule) from the reaction between diopside and nepheline.



The melilites crystallized near G and H are nearly isotropic, showing that soda-melilite is incorporated. Yoder (1964) has shown that natural melilite contains 1/3 of soda melilite molecule.

Possible differentiation course

On the basis of the experimental results of Schairer and Yoder (1964) and Onuma and Yagi (1967), the differentiation course of the melilite nephelinitic rock series are inferred as shown in Fig. 4. The wollastonite in the last product

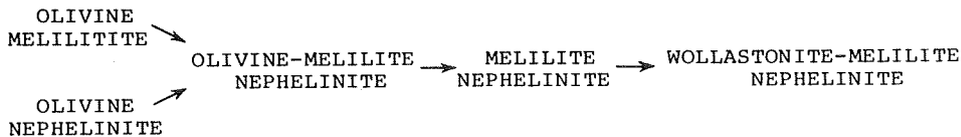


Fig. 4 Inferred differentiation course of the melilite nephelinitic rock series without taking TiO_2 into account.

might be replaced by perovskite or sphene when the residual liquid is rich enough in TiO_2 to produce one of these minerals. When the residual liquid is poor in TiO_2 , however, it is incorporated in clinopyroxene, and wollastonite together with nepheline is consumed to form soda melilite, which in turn enters into the structure of akermanite to form solid solution (Schairer and Yoder, 1964). Thus neither perovskite nor sphene is crystallized.

As indicated in the above statements, the role of TiO_2 in the melilite nephelinitic rock series is indeed very important in the differentiation of these rock series which is richer in TiO_2 than tholeiite rock series and ordinary alkaline rock series.

The System $\text{Na}_2\text{O-CaO-MgO-Al}_2\text{O}_3\text{-TiO}_2\text{-SiO}_2$ and the Differentiation of

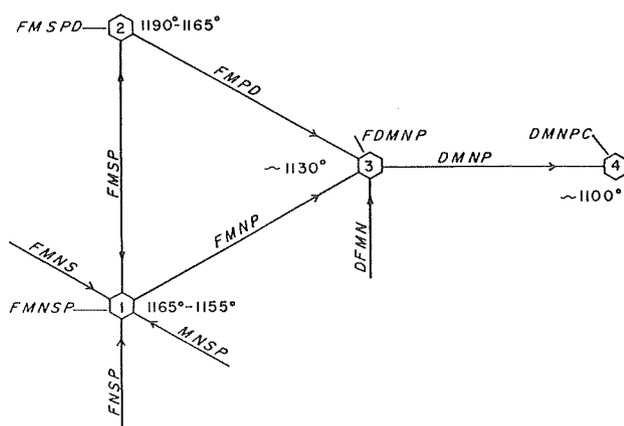


Fig. 6 Mutual relations of five-phase and six-phase assemblages in the system diopside-akermanite-nepheline- $\text{CaTiAl}_2\text{O}_6$ (after Yagi and Onuma, 1969). C = corundum and remainder abbreviations are same as in Fig. 5.

these experimental results. At V and Z in Fig. 5 both showing five phase assemblages liquids coexist with forsterite, spinel, melilite, and nepheline and with forsterite, spinel, melilite, and perovskite, respectively.

Natural rocks, corresponding to mineral assemblages 1, and 2 of Fig. 6 have not been found. The phase assemblage of DFMN (diopside + forsterite + melilite + nepheline + liquid) corresponds to olivine-melilite nephelinite and the assemblage at 3 corresponds to perovskite-bearing olivine-melilite nephelinite. At 3 forsterite is consumed and the assemblage between 3 and 4 indicates perovskite-bearing melilite nephelinite. The temperature falls towards 4 where corundum crystallizes out.

As mentioned above, the crystallization course which proceeds along DFMN through point 3 and along DMNP ultimately to 4 has a special petrologic significance. It is therefore interesting to analyze relationship between DFMN and the four-phase assemblage which is generated from the five-phase assemblage along DFMN. These five-phase assemblages were also confirmed as invariant points in the system diopside-akermanite-nepheline. In Fig. 7 a schematic flow diagram is given. The temperature at which a four-phase assemblage (three crystalline phases + liquid) transfers to a five-phase assemblage is not fixed and consequently DFMN is a reaction line along which forsterite reacts with liquid with falling temperature. In Table 1 crystallization sequences of selected mixtures in the system diopside-akermanite-nepheline- $\text{CaTiAl}_2\text{O}_6$ are given with the bulk composition of starting material. The data show that the crystallization sometimes ceases on the univariant line or divariant plane before the eutectic point is reached, because of the complex solid solution of

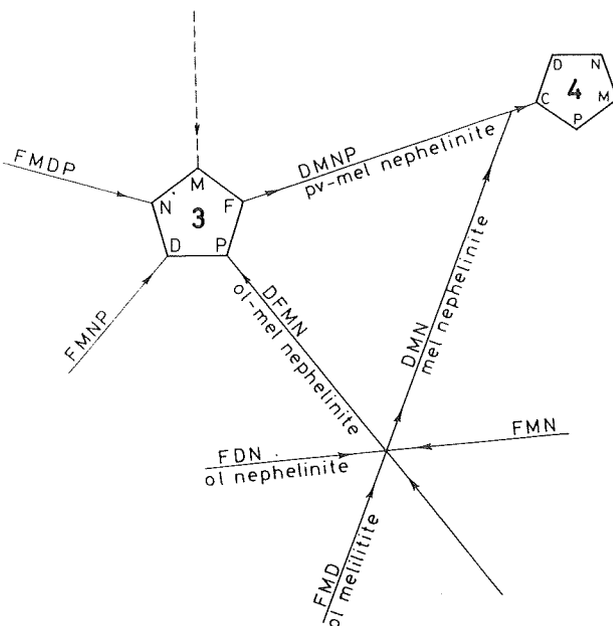


Fig. 7 Mutual relation of invariant, univariant, and divariant assemblages of a part of the system $\text{Na}_2\text{O}-\text{CaO}-\text{MgO}-\text{Al}_2\text{O}_3-\text{TiO}_2-\text{SiO}_2$ with corresponding rock names. Abbreviations are same as in Figs. 5 and 6.

the crystalline phases. On the basis of these data it is evident that if the liquid is rich in TiO_2 and poor in SiO_2 the crystallization proceeds along the DFMN until point 3 is reached, where perovskite joins the assemblage, while if the liquid is poor in TiO_2 and rich in SiO_2 , forsterite is consumed by the reaction before point 3 is reached, and the crystallization course leaves the DFMN and proceeds along the DMN (actually divariant plane), separating melilite, nepheline, and diopside (melilite nephelinite).

O'Hara and Biggar (1969) suggested that the melilite nephelinite is formed by the low-pressure extreme fractional crystallization of alkali olivine basalt magma. Recently Onuma and Yamamoto (1976) also indicated that the crystallization at depth does not lead to the melilite nephelinite but produce the olivine nephelinite and that the olivine melilitite and the melilite nephelinite are formed by the low pressure fractional crystallization of the magma. Considering these two discussions, it seems that the crystallization sequence obtained by the experimental data at atmospheric pressure is valid for the interpretation in the differentiation of the natural melilite nephelinitic rock series.

From the above discussions an inferred crystallization differentiation of the

Table 1 Crystallization sequence of the selected compositions in the system diopside-akermanite-nepheline-CaTiAl₂O₆.
(data are quoted from Yagi and Onuma, 1969)

	Di	Ak	Ne	Tp	Crystallization Sequence
(1)	40	10	40	10	L → Fo+L → Fo+Mel+L → Fo+Mel+Ne+L → Mel+Ne+Di+Pv+L → Mel+Ne+Di+Pv+Cor
(2)	75	5	5	15	L → Fo+L → Fo+Di+L → Fo+Di+Mel+L → Fo+Di+Mel+Pv+L → Di+Mel+Pv+Ne+L → Di+Mel+Pv+Ne
(3)	70	10	10	10	L → Fo+L → Fo+Di+L → Fo+Di+Mel+L → Fo+Di+Mel+Ne+L → Di+Mel+Ne+Pv+L → Di+Mel+Ne+Pv
(4)	70	10	15	5	L → Fo+L → Fo+Mel+L → Fo+Mel+Di+L → Mel+Di+Ne+L → Mel+Di+Ne

	(1)	(2)	(3)	(4)
SiO ₂	43.73	45.96	47.54	49.61
TiO ₂	3.36	5.04	3.36	1.68
Al ₂ O ₃	18.65	8.24	7.88	7.54
MgO	8.92	14.69	14.50	14.50
CaO	16.82	25.02	24.59	23.42
Na ₂ O	8.72	1.09	2.18	3.27

Di = diopside (CaMgSi₂O₆), Ak = akermanite (Ca₂MgSi₂O₇), Ne = nepheline (NaAlSi₃O₈), Tp = CaTiAl₂O₆, Fo = forsterite, Pv = perovskite, Mel = melilite, Cor = corundum, L = liquid.

melilite nephelinitic rock series is as shown in Fig. 8. Now the wollastonite in the wollastonite-melilite nephelinite is replaced by perovskite, supporting a suggestion by Schairer and Yoder (1964). Perovskite was reported from the natural olivine-melilite nephelinite and melilite nephelinite by Winchell (1947) and Spencer (1969). Spencer also described that perovskite is observed in the olivine-melilite nephelinite but not in the olivine nephelinite. This would support the inferred differentiation course of the melilite nephelinite rock series.

Although corundum has not been reported as a constituent of melilite

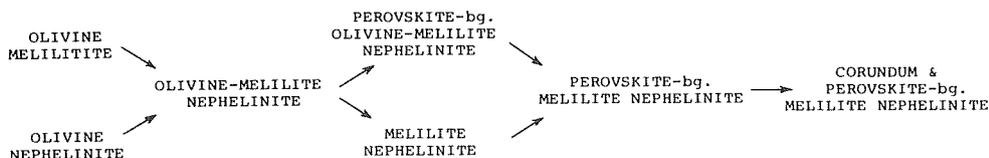


Fig. 8 Inferred differentiation course of the melilite nephelinitic rock series with taking TiO₂ into account.

nephelinite, occurrence of corundum megacrysts in nepheline basanite, hawaiite or related alkalic rocks from several localities in northern Queensland, Australia is worthy of notice (Stephenson, 1976). At Cheviot Hills, for example, rounded corundum crystals, sometimes up to 10 mm in diameter, are found associated with abundant, corroded megacrysts of anorthoclase. Genetic relation between the corundum and the host rocks has not yet been clarified, and the origin from the deep buried metamorphic rocks cannot be denied. It is, however, also possible to consider that corundum might have grown directly from the nepheline normative basaltic melt in the later stage under some special conditions, as supported by the results of the recent experiments.

Recently we had a chance to observe the olivine-melilite nephelinites from Eifel, West Germany. The rocks contain phenocrysts of olivine and titaniferous clinopyroxene and the groundmass always carries clinopyroxene, nepheline, melilite and perovskite (Detailed study on this rock will be published elsewhere by Yagi et al.). It is noticed that the phenocrystic olivine has corroded anhedral form and that olivine is absent and perovskite is always present in the groundmass. This observation leads to the following process during the formation of olivine-melilite nephelinite. The magma crystallizing olivine and clinopyroxene both of which are observed as phenocrysts (A-G in Fig. 3) was erupted. Then the crystallization course followed FMD and DFMN (Fig. 7), precipitating clinopyroxene, melilite, and nepheline, and finally reached the point 3, where perovskite joined in the groundmass. During this process olivine did not precipitate but decreased in amount by a reaction with liquid as experimental data showed. In nature, generally crystallization ceased at this point before the olivine is completely consumed, resulting in the formation of melilite nephelinitic rock always carrying olivine. Probably, the magma was cooled too quickly to follow further crystallization, because the most part of crystallization took place after the magma was erupted.

When the liquid is rich in SiO_2 sphene crystallizes instead of perovskite and the mineral assemblages would be different. The relationship between sphene and perovskite is controlled by activity of silica as will be stated in the following section.

Relation Between Perovskite and Sphene and the System $\text{CaO-MgO-Al}_2\text{O}_3\text{-TiO}_2\text{-SiO}_2$

Gupta et al. (1973) studied the system diopside- $\text{CaTiAl}_2\text{O}_6$ -silica. In this system there is an interesting relationship between pyroxene and associated titanium-bearing minerals, sphene and perovskite. The flow sheet given in Fig. 9 indicates that silica content increases from the upper part to the low part. Two

anorthite may be replaced by melilite in the silica-poor portion, and thus the assemblage may correspond to the natural assemblage (cf. olivine melilitite).

In the system diopside-akermanite- $\text{CaTiAl}_2\text{O}_6$ representing the silica-poor portion of the system $\text{CaO-MgO-Al}_2\text{O}_3\text{-TiO}_2\text{-SiO}_2$ Onuma and Yagi (1971) found the phase assemblage of diopside + forsterite + akermanite + perovskite + liquid (olivine melilitite) instead of diopside + forsterite + anorthite + perovskite + liquid which is encountered in the system diopside- $\text{CaTiAl}_2\text{O}_6$ -silica. These two joins may be connected by the divariant assemblage of forsterite + diopside + perovskite + liquid.

When sodium is added to the system, nepheline instead of pure anorthite, would appear in the silica-undersaturated portion, whereas albite-rich plagioclase would appear in the silica-rich portion of the system $\text{Na}_2\text{O-CaO-MgO-Al}_2\text{O}_3\text{-TiO}_2\text{-SiO}_2$. O'Hara and Biggar (1969) and Schairer and Yoder (1964) reported that in the system $\text{CaO-MgO-Al}_2\text{O}_3\text{-SiO}_2$ anorthite and melilitite coexist, showing inconsistency with the observation in natural rocks. However, the addition of Na_2O to the system changes the phase-assemblage and melilitite never coexist with plagioclase in the system $\text{Na}_2\text{O-CaO-Al}_2\text{O}_3\text{-SiO}_2$, indicating that a increase of component makes the system closer to the natural rocks.

Acknowledgement

We are grateful to the following persons of our department. Dr. A.K. Gupta for his critical reading of the paper in manuscript, Mr. S. Terada for his help in preparation of figures, and Mrs. S. Yokoyama for her typing the manuscript. Part of the cost for the present study was defrayed by the grant of Japan Society for Promotion of Science.

References

- Bowen, N.L., 1922. Genetic features of alnöitic rocks at Isle Cadieux, Quebec. *Am. J. Sci.* 5th Series, 3: 1-34.
- Bowen, N.L., 1928. *The evolution of the igneous rocks*. Princeton Univ. Press, Princeton, N.J., 332 pp.
- Bultitude, R.J. & D.H. Green, 1967. Experimental study at high pressures on the origin of olivine nephelinite and olivine melilitite nephelinite. *Earth & Planet. Sci. Letter*, 3: 325-337.
- Bultitude, R.J. & D.H. Green 1971. Experimental study of crystal-liquid relationships at high pressure in olivine nephelinite and basanite composition. *J. Petrol.*, 12: 121-147.
- Carmichael, I.S.E., J. Nicolls & A.L. Smith, 1970. Silica activity in igneous rocks. *Am. Mineral.*, 55: 246-269.
- Green, D.H., 1969. The origin of basaltic and nephelinitic magma in the earth's mantle. *Tectonophy.*, 7: 409-422.

- Gupta, A.K., K. Onuma, E.G. Lidiak & K. Yagi, 1973. Effect of silica concentration on the diopsidic pyroxenes in the system diopside-CaTiAl₂O₆-SiO₂. *Contr. Mineral. Petrol.*, 41: 333-344.
- Kushiro, I., 1960. Si-Al relation in clinopyroxenes from igneous rocks. *Am. J. Sci.*, 258: 548-554.
- Kushiro, I., 1969. Discussion of the paper "The origin of basaltic and nephelinitic magma in the mantle" by D.H. Green. *Tectonophy.*, 7: 427-436.
- O'Hara, M.J., 1968. The bearing of phase equilibria studies in synthetic and natural systems on the origin and evolution of basic and ultrabasic rocks. *Earth-Sci. Rev.*, 4: 69-133.
- O'Hara, M.J. & G.M. Biggar, 1969. Diopside + spinel equilibria, anorthite and forsterite reaction relationships in silica-poor liquidus in the system CaO-MgO-Al₂O₃-SiO₂ at atmospheric pressure and their bearing on the genesis of melilitites and nephelinites. *Am. J. Sci.*, 267-A: 364-390.
- Onuma, K. & K. Yagi, 1967. The system diopside-akermanite-nepheline. *Am. Mineral.*, 52: 227-243.
- Onuma, K. & K. Yagi, 1971. The join CaMgSi₂O₆-Ca₂MgSi₂O₇-CaTiAl₂O₆ and its bearing on the titanopyroxenes. *Mineral. Mag.*, 38: 471-480.
- Onuma, K. & M. Yamamoto, 1976. Crystallization in the silica-undersaturated portion of the system diopside-nepheline-akermanite-silica and its bearing on the formation of melilitites and nephelinites. *J. Fac. Sci., Hokkaido Univ., Ser. 4*, 17: 347-355.
- Sahama, Th. G. & A. Meyer, 1958. Study of the volcano Nyiragongo: Exploration du Parc National Albert, Inst. Parcs Nat. Congo Belge, 2, 1-85.
- Schairer, J.F. & H.S. Yoder, Jr., 1960. The nature of residual liquids from crystallization, with data on the system diopside-nepheline-silica. *Am. J. Sci.*, 258-A: 273-283.
- Schairer J.F., K. Yagi & H.S. Yoder, Jr., 1962. The system nepheline-diopside. *Carnegie Inst. Wash. Year Book*, 61: 96-98.
- Schairer, J.F. & H.S. Yoder, Jr. 1964. Crystal and liquid trend in simplified alkali basalt. *Carnegie Inst. Wash. Year Book*, 63: 65-74.
- Schairer, J.F. & H.S. Yoder, Jr., 1970. Critical planes and flow sheet for a portion of the system CaO-MgO-Al₂O₃-SiO₂ having petrological application. *Carnegie Inst. Wash. Year Book*, 68: 202-214.
- Smith, A.L., 1970. Spene, perovskite and coexisting Fe-Ti oxide minerals. *Am. Mineral.*, 55: 264-269.
- Spencer, A.B., 1969. Alkalic igneous rocks of the Balcones province, Texas. *J. Petrol.*, 10: 272-306.
- Stephenson, P.J., 1976. Sapphire and zircon in some basaltic rocks from Queensland, Australia. Abstracts of 25th Int. Geol. Cong. Sydney.
- Winchell, H., 1947. Honolulu series, Oahu-Hawaii. *Geol. Soc. Am. Bull.*, 58: 1-48.
- Yagi, K. & K. Onuma, 1967. The system CaMgSi₂O₆-CaTiAl₂O₆ and its bearing on the titanite. *J. Fac. Sci. Hokkaido Univ. Ser. 4*, 13: 463-83.
- Yagi, K. & K. Onuma, 1969. An experimental study on the role of titanium in alkalic basalts in light of the system diopside-akermanite-nepheline-CaTiAl₂O₆. *Am. J. Sci.*, 267-A: 509-549.
- Yoder, H.S., Jr., 1964. Soda melilite. *Carnegie Inst. Wash. Year Book*, 63: 87-89.
- Yoder, H.S., Jr. & C.E. Tilley, 1962. Origin of basaltic magma: An experimental study of natural and synthetic rock systems. *J. Petrol.*, 3: 342-532.
- Yoder, H.S., Jr. & I. Kushiro, 1972. Composition of residual liquid in the nepheline-diopside system. *Carnegie Inst. Wash. Year Book*, 71: 413-416.

(Received on Oct. 12, 1976)