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Autor(en): Meinhold, K.D. / Frisch, Thomas

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## Manganese-Silicate-Bearing Metamorphic Rocks from Central Tanzania

By K. D. Meinhold\*) and Thomas Frisch\*\*)

With 3 figures and 2 tables in the text

Abstract. The Precambrian Konse Series includes isolated bodies of manganese-rich metasedimentary rocks, which were formed under amphibolite facies conditions in the Usagaran orogeny. Piemontite is a member of a variety of parageneses, providing a further example of its stability in a facies of higher grade than the greenschist or glaucophane schist facies. In piemontite-hornblende rock the amphibole is an optically positive magnesio-hornblende. Viridine with 8.50% Mn<sub>2</sub>O<sub>3</sub> is a major component of schist, in which it coexists with ferrian muscovite, manganoan phlogopite, spessartite, oligoclase An<sub>23-27</sub>, quartz, hematite, minor piemontite and accessory minerals. The garnet has the composition Spess<sub>76.0</sub> Py<sub>20.2</sub> Gross<sub>1.7</sub> Alm<sub>1.6</sub> Ti-Andr<sub>0.5</sub> and, as far as is known, is the most magnesian spessartite yet described.

#### INTRODUCTION

The metamorphic manganese-silicate rocks described below were discovered in the course of mapping by the German Geological Mission Tanganyika of the "Bundesanstalt für Bodenforschung (Hannover)" during the period 1963 to 1965. A brief petrographic description of these rocks forms part of a detailed report on their geologic setting by MEINHOLD (1968 and in press). Chemical analysis of the chief minerals of these unusual rocks provides the basis of the following more comprehensive description.

The first-named author is responsible for the field, optical and X-ray work, the second author for electron microprobe analyses.

<sup>\*)</sup> Institut für Geologie der Technischen Hochschule München. Present address: Bundesanstalt für Bodenforschung, Postfach 54, 3 Hannover-Buchholz, West Germany.

<sup>\*\*)</sup> Mineralogisch-Petrographisches Institut der Universität München, Luisenstrasse 37, 8 München 2, West Germany. Present address; Department of Geology, University of Alberta, Edmonton 7, Alta., Canada.

#### GEOLOGIC SETTING

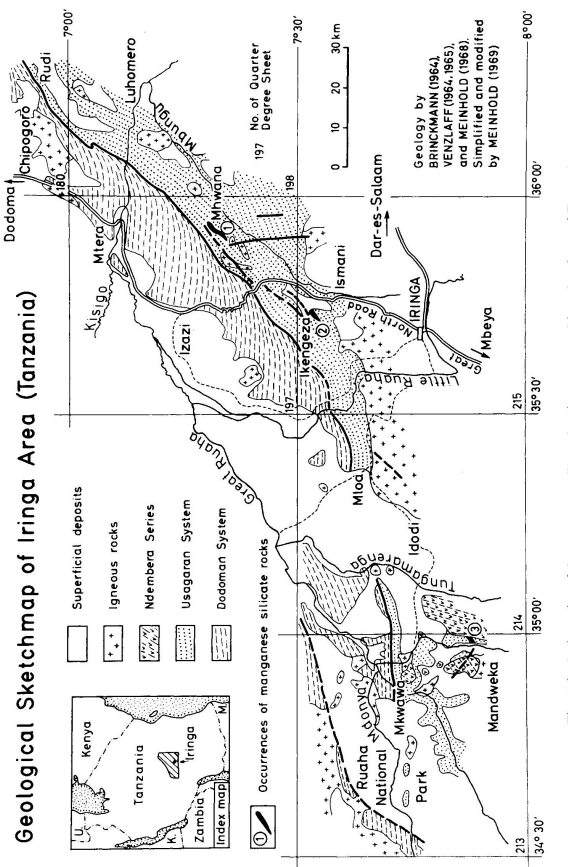
Manganese-silicate-bearing rocks comprise a part of the metasedimentary sequence of the Konse Series, which represents the base of the Precambrian Usagaran System in the area of Tanzania mapped by the German Geological Mission. The Konse Series is exposed in a curved belt, about 200 km long, extending from Rudi to Ruaha National Park, north of Iringa (Fig. 1). It lies unconformably on the migmatitic gneisses of the Archean Dodoman System, with the contact disturbed by a northwest-trending thrust fault.

The Konse Series begins with quartzites and conglomerates, which are overlain, in irregular succession, by marbles, calc-silicate rocks, calcareous mica schists, quartzites, quartz schists, ferruginous quartzites, mica schists, graphite schists, paragneisses, and amphibolites. Concordantly above the Konse Series lie interbedded para- and orthogneisses and amphibolites ("Gneiss-Amphibolite Series" of MEINHOLD, in press). South of the Mbungu River, northeast of Ismani (Fig. 1), banded gneisses and massive garnet or cordieritesillimanite gneisses (the "Amphibolite-free Gneiss Series") occur above the Gneiss-Amphibolite Series.

These three rock series are assigned to the Usagaran System (MEINHOLD, 1968 and in press). They were folded and metamorphosed together during the Usagaran orogeny, in which the amphibolite facies, locally even the granulite facies, was attained and migmatization occurred in places. The rocks of the Konse Series, the Gneiss-Amphibolite Series and the lower part of the Amphibolite-free Gneiss Series belong to the Barrovian facies series, whereas those of the upper parts of the Amphibolite-free Gneiss Series belong to the Abukuma type (cf. WINKLER, 1967).

Towards the end of the Usagaran orogeny, the Konse Series was faulted and thrust over the Dodoman Shield. The associated cataclasis coupled with invading hydrothermal solutions led to retrograde metamorphism of large parts of the Konse Series and smaller portions of the Gneiss-Amphibolite Series. Nevertheless, relics of the original high-grade metamorphism are everywhere preserved.

Manganese-silicate-bearing rocks have been observed in three places within the Konse Series (Fig. 1). The largest exposure, at Mwhana (Fig. 1, No. 1), is composed of quartz-viridine-mica schists and piemontite-quartz schists, in which lenses only a few meters thick of piemontite-amphibole rocks are intercalated. Interbeds of piemontite- and manganoan phlogopite-bearing marbles and of piemontite- and spessartite-bearing calc-silicate rocks occur locally. The exposure in the Mwhana Hills is about 130 m wide; to the northeast it wedges out rapidly, whereas to the southwest it splits into individual bodies 50 m thick, separated by marbles, calc-silicate rocks, quartzites, gneisses





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and mica schists. Seven kilometers farther southwest, the manganese-silicate rocks finally wedge out in the Mhumbirisa Hills.

The exposure of quartz-viridine-mica schists east of Ikengeza (Fig. 1, No. 2) is but a small lens with a maximum thickness of 10 m. The associated rocks comprise quartzite, calc-silicate rocks and paragneisses.

The Mandweka occurrence (Fig. 1, No. 3) was only cursorily examined. Here, piemontite-bearing quartz-viridine-mica schists are associated with metasedimentary rocks that may represent a facies repetition of the Konse Series within the Gneiss-Amphibolite Series or a saddle-type structure.

During the Usagaran orogeny, the rocks of all three occurrences were metamorphosed in the staurolite-almandine or the kyanite-almandine-muscovite subfacies of the almandine-amphibolite facies (Barrovian facies series) (cf. TURNER and VERHOOGEN, 1960; WINKLER, 1967). Retrograde metamorphism did not affect the rocks of the Mwhana occurrence but it is clearly evident at the other two localities.

#### ANALYTICAL PROCEDURE

Analyses of minerals were performed on separated grains mounted in "Araldite" embedded in plastic, except in the case of spessartite, for which a polished thin section was used. All samples were coated with beryllium to make them electrically conducting. The electron probe used was an A.R.L. EMX machine.

Analytical conditions were as follows: accelerating potential 15 kV, sample current 0.02 to 0.08 microampere, X-ray spot size 1 to 10 microns, integration time on scalers 10 to 40 sec. Pulse height discrimination was used for all elements. A minimum of four spot determinations was made on each of at least four grains of each mineral. Intensities so obtained were averaged and corrected for background and mass absorption, using the methods of SMITH (1965) and the absorption coefficients of HEINRICH (1966). When two standards were used for the same element in any one sample, the two "apparent" concentrations obtained never varied by more than ten per cent and their mean was used in the correction procedure.

A variety of natural and synthetic silicates was used as standards. For Si and Al in all minerals except spessartite, sillimanite (checked for minor element content and found to be essentially pure Al<sub>2</sub>SiO<sub>5</sub>). For Si, Ti, Al, Fe, Ca, and Mg in spessartite, natural pyrope (MASON, 1966, Tab. 1) and synthetic pyrope glass (prepared by M. J. O'Hara). For Ca, Fe and Mg in all minerals except spessartite and hematite, Ca-rich clinopyroxenes R 2397 (using microprobe values of SMITH, 1966, Tab. 3) and E-108 (AOKI and KUSHIRO, 1968, Tab. 3). For Fe, Ti and Mg in hematite, ilmenite K 13–131.8 (ANDERSON, 1968, Tab. 3). For Ti, Na and K in all minerals except spessartite and hematite, kaersutite with TiO<sub>2</sub> 4.38%, Na<sub>2</sub>O 2.60% and K<sub>2</sub>O 2.05% (B. Mason, unpublished analysis). For Mn in all minerals, Mn-cummingtonite No. 4 (KLEIN, 1968, Tab. 1).

Accuracy is estimated at 2 to 4% of the amount present for major elements and 5 to 10% for the minor ones.

	1	2	3	4	<b>5</b>	6	7
SiO <sub>2</sub>	38.1	52.1	35.8	38.8	38.5	45.1	n.d.
TiO,	0.06	0.20	0.03	0.13	0.81	0.76	1.13
$Al_2O_3$	23.6	7.30	53.2	21.7	18.6	32.7	0.42
Fe <sub>2</sub> O <sub>3</sub> <sup>a</sup>	11.0	3.72	2.90		2.06	3.90	97.7
FeOa				0.69			
Mn <sub>2</sub> O <sub>3</sub> b	1.90		8.50				
MnOb		0.53		35.0	1.72	0.25	1.75
MgO	0.19	21.7	0.23	5.27	23.2	1.45	0.06
CaO	23.3	12.4	0.02	0.76	n.d.	0.02	n.d.
Na <sub>2</sub> O	n.d.	0.97	n.d.	n.d.	0.43	1.44	n.d.
K₂Ô	n.d.	0.15	$\mathbf{n.d.}$	n.d.	9.30	8.94	n.d.
Total	98.2	99.1	100.7	102.4	94.6	94.6	101.1

Table 1. Microprobe analyses of minerals from the manganese-silicate-bearing rocks

<sup>a</sup> total iron <sup>b</sup> total manganese n.d. = not detected

#### Structural formulae

Basis $(0 =)$	12	23	20	<b>24</b>	22	<b>22</b>
Si	3.018	7.104	4.000	6.002	5.438	6.085
Al <sup>4</sup>		0.896			2.562	1.915
	3.02	8.00	4.00	6.00	8.00	8.00
Al <sup>6</sup>	2.199	0.284	7.007	3.958	0.526	3.287
Ti	0.005	0.016		0.037	0.085	0.081
$\mathrm{Fe}^{3}$	0.657	0.377	0.242		0.221	0.389
${\rm Fe}^2$				0.093		
$Mn^3$	0.114		0.725			
$Mn^2$		0.057		4.581	0.204	0.032
Mg	0.024	4.408	0.040	1.217	4.887	0.292
Ca	1.975	1.811		0.130		
Na		0.262			0.119	0.373
K		0.032			1.680	1.540
(A)	4.98	7.24	8.01	10.02	7.72	5.99

1 = Piemontite

te { from piemontite-amphibole rock, Mwhana

2 =Amphibole 3 =Viridine

3 =Virialite 4 =Garnet

5 = Mn-phlogopite from quartz-viridine-mica schist, Mhwana

6 = Muscovite

7 = Hematite

#### MINERALOGY

The following mineral assemblages were observed in the manganese-silicatebearing rocks of the Konse Series:

- 1. Quartz Mn-phlogopite muscovite piemontite ore (– tourmaline zircon) (retrograde: scapolite sphene).
- 2. Quartz biotite spessartite piemontite ore (– tremolite) (retrograde: tremolite sphene).
- 3. Quartz muscovite piemontite biotite (– ore garnet) (retrograde: sericite-chlorite-clinozoisite).

Table 2.	X-ray	powder	data	for	piemontite,	magnesio-hornblende,	viridine	and	muscovite
				(cf.	. Tab. 1). C	$u \ K_{lpha}$ radiation			

				u			
Piemon	tite	Mg-Horn	blende	Viridi	ine	Muscovit	e
d (Å)	I	d (Å)	Ι	d (Å)	Ι	d (Å)	Ι
5.0122	25	9.0175	10	6.1626	7	9.9051	95
4.1912	$\frac{10}{37}$	8.3860	81	5.5691	100	4.9676	34
4.1678	40	5.0748	4	4.5483	33	4.4622	4
3.4847	54	4.9130	4	4.2467	5	3.8734	3
3.4009	17	4.7410	6	3.9413	13	3.7200	3
3.2063	15	4.5071	8	3.7047	4	3.6656	<b>5</b>
3.0598	$\overline{17}$	4.1990	14	3.7017	<b>5</b>	3.4820	<b>5</b>
2.9153	19	3.8834	3	3.5365	17	3.3285	12
2.9005	85	3.4502	<b>5</b>	3.5063	16	3.3091	100
2.8202	<b>46</b>	3.3731	12	3.3432	<b>82</b>	3.1951	<b>5</b>
2.6914	37	3.2663	<b>25</b>	3.0867	4	3.1839	<b>6</b>
2.6773	42	3.1142	100	2.7843	95	2.9857	7
2.6603	21	2.9340	14	2.6961	7	2.8589	<b>5</b>
2.5989	37	2.7978	<b>26</b>	2.5902	<b>5</b>	2.7910	4
2.5321	<b>46</b>	2.7313	9	2.5128	<b>5</b>	2.6977	9
2.4024	63	2.7009	16	2.4793	<b>20</b>	2.5931	3
2.2783	35	2.5916	15	2.4237	4	2.5643	6
2.1661	<b>42</b>	2.5432	6	2.3635	12	2.5169	6
2.1176	25	2.3791	<b>5</b>	2.2816	14	2.4859	8
2.0723	15	2.3386	11	2.2651	7	2.1984	3
1.8841	19	2.3224	8	2.1841	<b>34</b>	2.1721	3
1.8739	21	2.2973	4	1.9584	6	1.9868	33
1.6417	17	2.1591	7	1.8178	8	1.6570	2
1.6369	<b>23</b>	2.0420	<b>5</b>	1.8104	6	1.6423	3
1.6258	17	2.0136	8	1.7647	4	1.4347	3
1.5888	100	1.9968	5	1.6709	4	1.4211	3
1.5769	17	1.8878	12	1.6039	6	1.3489	6
1.4576	19	1.8433	3	1.5470	5	1.2422	3
1.4387	15	1.8104	5	1.5363	8		
1.4097	21	1.6857	4	1.4933	14		
1.3941	31	1.6466	7	1.4847	5		
		1.6342	5	1.3937	14		
		1.5813	54	1.3104	8		
		1.5466	3	1.3020	4		
		1.5154	4	1.2919	5		
		1.4999	3	1.2008	<b>5</b>		
		1.4666	3				
		1.4543	3				
		1.4351	$\frac{27}{3}$				
		1.3718 1 2522	3				
		$1.3523 \\ 1.3082$	3 3				
		1.2940	$\frac{3}{2}$				
		1.2340	4				

- 4. Quartz viridine muscovite Mn-phlogopite spessartite plagioclase hematite (– piemontite – tourmaline – apatite – zircon – zoisite) (retrograde: ore – sericite – serpentine – quartz – albite – epidote).
- 5. Piemontite amphibole (– ore).
- 6. Carbonate Mn-phlogopite muscovite tremolite diopside (– piemontite – ore) (retrograde: tremolite – quartz – scapolite).
- 7. Quartz diopside tremolite Mn-phlogopite ore (– piemontite tourmaline – apatite) (retrograde: tremolite – epidote – sphene – carbonate).

Rocks of this type have not previously been reported from Tanzania. In western Tanzania, rocks termed "gondites" (manganiferous ironstones) by McCONNELL (1950, p. 31) contain spessartite as the only Mn-silicate, as well as manganese ores.

Minerals investigated by optical, X-ray and/or chemical (electron probe) methods are: piemontite, viridine, spessartite, Mn-phlogopite, muscovite, hematite, amphibole, tourmaline, and plagioclase.

#### 1. Piemontite

Piemontite was found in marbles, calc-silicate rocks, piemontite-quartz schists, quartz-chlorite-mica schists, quartz-viridine-mica schists, and piemontite-amphibole rocks. Only in the piemontite-quartz schists and in the piemontite-amphibole rocks does piemontite form a major constituent, in generally subhedral crystals, commonly stretched in "b". Pleochroism is intense: X = yellow, Y = pale violet pink, Z = red. Zonal structure, in which cores of crystals are paler than margins, is particularly common in the pied-montite-amphibole rocks. The piemontite of the calc-silicates has a paler colour than that of the other piemontite-bearing rocks.

Piemontite is optically positive in all the rocks studied. According to TRÖGER (1967, p. 311), this property would place the mineral in the  $Mn^3$  – rich part of the piemontite series, but chemical analysis (Table 1, Col. 1) indicates a low content of trivalent Mn.

The piemontite is commonly closely intergrown with ore and mica (Mnphlogopite and muscovite). In the quartz-viridine-mica schists it has in part grown along cleavage planes in muscovite, and in the piemontite-amphibole rocks it forms abundant inclusions in amphibole. Grains stretched in "b" display alignment with the schistosity and here and there are bent in small folds. In the same rocks piemontite may also occur in non-aligned crystals or in granular aggregates. Crystallization of the piemontite was evidently syn- to postkinematic with respect to the main deformation (MEINHOLD, in press).

The occurrence of diopside and, above all, the assemblage quartz-viridinemuscovite-oligoclase  $An_{23-27}$  are indicative of the almandine-amphibolite facies. In addition, the piemontite-bearing rocks are associated with rocks whose pre-retrograde mineralogy clearly indicates recrystallization in the amphibolite facies (MEINHOLD, in press). Textural relations show that the piemontite formed during the primary, amphibolite-facies metamorphism, not in the greenschist facies, which was reached during retrograde metamorphism. Whereas most piemontite previously reported belongs to the greenschist or glaucophane schist facies, piemontite in the amphibolite facies has recently been found in several other occurrences (SMITH and ALBEE, 1967, FUCHS, 1968, ROY and PURKAIT, 1968). In the piemontite analysis (Tab. 1, Col. 1), both the iron and the manganese are considered to be in the trivalent form. Microprobe wavelength scanning of the  $L_{\alpha}$  and  $L_{\beta}$  peaks of Fe (cf. SMITH and ALBEE, 1967, p. 191) confirmed that at least the majority of the iron is present as Fe<sup>3</sup>. Mn intensities in the L-spectrum were too low to provide unequivocal information as to the valence state of this ion.

Counts for manganese and iron varied antipathetically in some of the piemontite grains, doubtless reflecting the zoning that is visible optically. Maximum and minimum concentrations of  $Mn_2O_3$  recorded were about 3% and 1%, of Fe<sub>2</sub>O<sub>3</sub> about 11.5% and 9%. As a large number of spots were analyzed, the piemontite analysis given is considered to be a reliable bulk analysis.

The piemontite analyzed, from piemontite-amphibole rock at Mwhana, is characterized by a low Mn<sup>3</sup> content. MIVASHIRO and SEKI (1958, p. 427ff.) contend that this feature is typical of piemontites "from low-grade schists and associated rocks from areas where the temperature of metamorphism was so low that biotite was not formed". However, the Tanzanian piemontite, as well as the piemontite from San Gorgonio Pass, California (SMITH and ALBEE, 1967), all of which formed in the amphibolite facies, fall in the lowtemperature field of MIVASHIRO and SEKI'S Al<sup>3</sup>-Fe<sup>3</sup>-Mn<sup>3</sup> diagram (Fig. 2). Temperatures prevailing in the amphibolite facies are well over 500° C, accord-

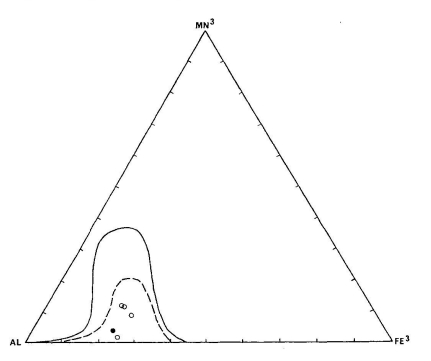


Fig. 2. Al-Mn<sup>3</sup>-Fe<sup>3</sup> diagram of piemontites from Tanzania (this study) – solid circle – and San Gorgonio Pass, California (SMITH and ALBEE, 1967) – open circles. Composition fields after MIYASHIRO and SEKI (1958, Fig. 4): the broken line marks the upper limit of piemontites formed at low temperature, in the greenschist and glaucophane schist facies; the solid line demarcates the field of piemontite formed at higher temperature.

ing to WINKLER (1967, Fig. 40). It appears that piemontites of similar composition may form at various temperatures ranging from low to high.

#### 2. Amphibole

Intercalated with the quartz-viridine-mica schists at Mwhana (Fig. 1, No. 1) are massive reddish rocks, composed entirely of piemontite, yellow amphibole and a little ore (Assemblage 5). The amphibole occurs interstitially and in schlier-like zones between piemontite-rich layers. The amphibole crystals are aligned in general but in places have grown parallel to the schistosity. Inclusions of piemontite in the amphibole are common. Growth of the amphibole is assigned to a postkinematic stage of recrystallization.

The amphibole exhibits weak pleochroism from colourless to pale yellow. On the basis of paragenesis and optical properties  $-2V_X = 91^{\circ}$  to  $98^{\circ}$ ,  $Z \wedge c = 16^{\circ}$  to  $23^{\circ}$ , length slow – MEINHOLD (1968) considered the amphibole to be manganese cummingtonite (Fig. 3). Chemical analysis (Tab. 1, Col. 2) shows it to be magnesio-hornblende (classification of LEAKE, 1968, Fig. 2): Ca and Al are high, Mn is low, and Al takes the place of a silicon atom in the Z position. However, magnesio-hornblendes investigated up to now are optically negative.

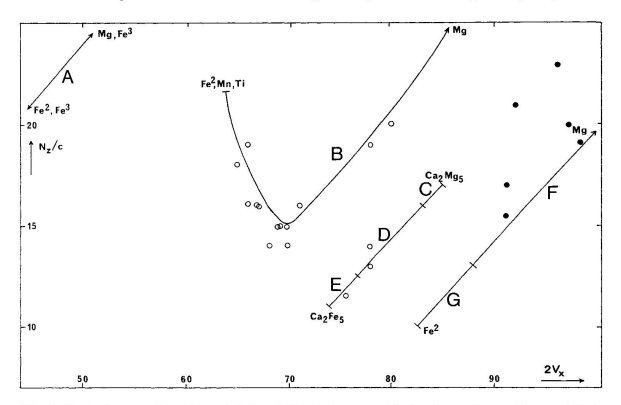


Fig. 3. Optical properties of amphiboles. Solid circles: amphiboles from piemontite-amphibole rocks, Mwhana; open circles: mean values of amphiboles from amphibolites of the Usagaran System. Determinative curves after TRÖGER (1956) and HELBIG (1969). A = Hastingsite series, B = Hornblende series, C = Tremolite, D = Actinolite, E = Fe-actinolite, F = Cummingtonite, and G = Grunerite.

Perhaps a solid solution series exists between the optically negative magnesiohornblendes and the optically positive pargasites, An X-ray diffraction study of the Tanzanian amphibole (Tab. 2) did not aid in resolving the problem but, as there is a lack of X-ray data on such amphiboles, results are presented herein.

#### 3. Viridine (manganandalusite)

Viridine, while limited to the quartz-viridine-mica schists (Assemblage 4), which are found in all three localities of the manganese-silicate rocks (Fig. 1), is a major constituent of these schists. Pleochroism is vivid: X = yellowish green, Y = grass green, Z = golden yellow. Isomorphous overgrowths on crystals are less intensely coloured than the primary viridine. HEINRICH and COREY (1959) state that the unusual pleochroism is the result of the copresence of Mn<sup>3</sup> and Fe<sup>3</sup>.

The viridine porphyroblasts, which are green in hand specimen and up to fist size, grew largely synkinematically. They are anhedral and commonly deformed into *s*-shapes and may contain helicitic inclusions of quartz, hematite, Mn-phlogopite and muscovite. Small, rounded spessartites are also included in many viridine porphyroblasts. Deformation related to small folds suggests that crystallization of viridine began early in the deformation history. Quartz and mica have penetrated along fractures and in embayments. During retrograde metamorphism the viridines were granulated and partly to wholly replaced by sericite, with concomitant formation of ore.

The pattern of the "L" peaks of Fe and Mn in the viridine shows that these ions are trivalent. The  $Mn_2O_3$  content, 8.50% (Tab. 1, Col. 3), is the highest yet reported for viridine, but it seems likely that the specimens from Yakutia, U.S.S.R. (SERDYUCHENKO, 1949, SHABYNIN, 1948), are richer still in trivalent Mn if, as HEINRICH and COREY (1959, p. 1268) suggest, they contain Mn<sup>3</sup> rather than Mn<sup>2</sup> as claimed by the Russian workers.

Substitution of aluminium by manganese and iron must obviously extend the stability field of andalusite. Andalusite is typically restricted to the hightemperature, low-pressure Abukuma-type facies series. Viridine, on the other hand, occurs not only in that facies series, alongside cordierite and sillimanite (SERDYUCHENKO, 1949, SHABYNIN, 1948), but also with kyanite in the almandine-amphibolite facies of Barrovian type (BACKSTRÖM, 1896, HEINRICH and COREY, 1959). The Tanzanian occurrence also belongs to the latter facies series.

The X-ray powder data of the analyzed viridine (Tab. 2) deviate from both the data of HEINRICH and COREY (1959, Tab. 2) and those given in the ASTM card 18-36. The discrepancy is probably due to differences in chemical composition.

#### 4. Garnet

Small (up to 4 mm diameter), colourless garnets are abundant in some of the quartz-viridine-mica schists (Assemblage 4) at Mwhana and in the Mhumbirisa Hills (Fig. 1, No. 1). The garnets formed largely prekinematically and have been granulated and strung out along the foliation planes. They are very irregularly distributed as single crystals, commonly included in viridine, Mnphlogopite and muscovite, and as aggregates of grains. Hematite and rutile occur as intergrowths with garnet. Beginning retrograde metamorphism in the Mhumbirisa Hills has caused brownish clouding of garnet centres.

Tectonically stretched, in part granulated and skeletal garnets occur in a biotite-garnet-tremolite-piemontite quartzite schist (Assemblage 2) in the Mhumbirisa Hills. These are colourless to pale yellow and generally closely intergrown with ore. Piemontite, along with sphene, replaces some of the garnet at its margins.

Chemical analysis of garnet from quartz-viridine-mica schist is given in Table 1, Col. 4. It is spessartite of unusual composition, Spess<sub>76.0</sub> Py<sub>20.2</sub> Gross<sub>1.7</sub> Alm<sub>1.6</sub>Ti-Andr<sub>0.6</sub> (mol per cent). MATTHES (1961, p. 266ff.) experimentally investigated the miscibility of pyrope in spessartite and found that spessartite took up no more than 15 mol per cent pyrope in solid solution and that incorporation of Mg in spessartite was favoured by high temperature. The compositions of natural spessartites support MATTHES' results. While a significant amount of magnesia might be expected in spessartite formed in the amphibolite facies, the Tanzanian example is anomalously high in MgO. The MgO in the spessartite was standardized against an homogeneous natural pyrope (with MgO = 18.80%) and a synthetic pyrope glass (MgO = 30.0%). The "apparent" (uncorrected except for background) Mg as MgO concentrations obtained were 4.60% and 4.30%. The mean of these two values after their correction for mass absorption is 5.27%. The analytical total for the analysis is slightly high, 102.4%, but the structural formula is very satisfactory. Thus there seems to be no compelling reason to reject the analysis and the Tanzanian spessartite may be considered to extend the known miscibility field of spessartite-pyrope garnets.

Total manganese in the spessartite has been calculated assuming all the Mn to be divalent but some of it may be present as  $Mn^3$ , replacing Mg, which has an identical ionic radius (0.66 Å).

#### 5. Manganoan phlogopite

Manganese-rich phlogopite is a significant constituent of the piemontitequartz schists and the quartz-viridine-mica schists (Assemblages 1 and 4). The mineral is also present in marble and calc-silicate rocks (Assemblages 6 and 7), associated with the manganese-silicate rocks at Mwhana and in the Mhumbirisa Hills.

Nematoblastic Mn-phlogopite generally displays good parallel orientation in the schistosity planes, indicative of prekinematic crystallization. However, in the same specimens, Mn-phlogopite may occur as a synkinematic growth in pressure shadows around viridine and as postkinematic crystals that grew athwart the foliation. Muscovite has formed in cracks in the Mn-phlogopite.

The Mn-phlogopite is distinctly pleochroic, X = very light reddish yellow, Y = Z = orange to reddish brown. Chemically (Tab. 1, Col. 5), it compares well with the phlogopite of the piedmontite gneiss from Southern California (SMITH and ALBEE, 1967), except that the latter has up to nearly twice as much manganese as the Tanzanian specimen. In keeping with the findings of SMITH and ALBEE (1967, Tab. 1), we have assumed the manganese to be divalent. The manganese content of the Tanzanian phlogopite is higher than normal for phlogopite but not high enough for manganophyllite, which contains at least twice as much Mn (JAKOB, 1924).

#### 6. Muscovite

Pink muscovite is the major component of the quartz-viridine-mica schists (Assemblage 4). It also occurs in the piemontite-quartz schists and in the metacalcareous rocks. As only the muscovite of Assemblage 4 was studied in detail, further discussion is confined to this variety.

Muscovite forms plates up to one centimeter in size in the quartz-viridinemica schists. It is well aligned with the schistosity, and longer flakes are commonly bent in microfolds. Muscovite has also grown in pressure shadows around viridine and across the foliation. Like Mn-phlogopite, it appears to have begun crystallization prekinematically and ended it postkinematically, but textural relations indicate that it probably continued to form after the cessation of Mn-phlogopite growth. At Ikengeza and Mandweka (Fig. 1, Nos. 2 and 3) diaphthoretic deformation has caused crushing of muscovite and its replacement by sericite, whereby the original pink colour was lost.

The pink colour of the mica is suggestive of alurgite, but X-ray diffraction measurements (Table 2) revealed the 2 M structure of muscovite; alurgite has the 3 T structure (HEINRICH and LEVINSON, 1955).

The iron content – nearly 4% Fe<sub>2</sub>O<sub>3</sub> (Tab. 1, Col. 6), assuming all the iron to be trivalent – is unusually high for muscovite and is almost twice as high as that of the coexisting Mn-phlogopite. High iron (more than 6% Fe<sub>2</sub>O<sub>3</sub>) in pink muscovite and a similar unusual partitioning of iron between coexisting muscovite and Mn-phlogopite was found by SMITH and ALBEE (1967) in the Southern Californian piemontite gneisses.

#### 7. Hematite

Hematite is an abundant accessory mineral in many of the rocks, and grains are generally well aligned with the schistosity. The common presence of hematite included in viridine, mica, quartz and plagioclase attests to early growth of the hematite. Rims of rutile around hematite grains may be the result of exsolution.

Hematite included in viridine was analyzed (Tab. 1, Col. 7). Ti and Mn are significant but minor constituents.

#### 8. Other minerals

Small, idiomorphic tournalines with weak pleochroism (colourless to pale yellow to ochre) occur in the piemontite-quartz schists, quartz-viridine-mica schists and piemontite-bearing calc-silicate rocks (Assemblages 1, 4 and 7). The colour suggests dravite but the paragenesis makes the manganese variety, tsilaisite, a possibility.

Small, well-twinned (Albite law) crystals of plagioclase are fairly abundant in the quartz-viridine-mica schists. They are up to 0.3 mm in size, anhedral and have the composition  $An_{23-27}$ . During retrograde metamorphism, the plagioclase was partly replaced by clinozoisite, pistacite and sericite, resulting in a decrease in anorthite content to 18 to 20 mol per cent.

Some specimens of quartz-viridine-mica schists contain minor, fractured zoisite with the typical blue-grey interference colour. Small zircons are surrounded by pleochroic haloes in Mn-phlogopite. Apatite is also present in rounded, generally much-cracked grains.

#### ORIGIN OF THE ROCKS

The manganese-rich rocks of the Konse Series belong to a suite of metamorphic rocks of undoubted sedimentary origin. The probable parent material comprised clay-rich sandstones, greywackes and sandy clays with manganese contents considerably higher than normal. Sedimentation most likely took place in small, isolated bodies of water under more or less euxinic conditions. In the Usagaran orogeny these sediments recrystallized during regional metamorphism that attained amphibolite-facies grade.

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