Introduction

In September, 1965 the Geological Survey of Finland received a small boulder containing red corundum (Fig. 1) from a place called Paaraskalla in the Kittilä wilderness in Finnish Lapland. This sample was found by a school-boy, Mikko Tervo. During short periods in 1966—68 the area was studied for economic corundum deposits, and many tens of boulders and some outcrops containing corundum were found. The corundum occurs in strongly tectonized amphibolitic-hornblenditic rocks. When studying the Paaraskalla samples in the laboratory, sapphirine and kornerupine were identified. Kornerupine is a mineral new to Finland, and sapphirine has been described from only one locality (Rouhunkoski 1969). Because it was considered that the gravel at Paaraskalla could contain some precious corundum (ruby), panning was carried out on the spot. A great deal of corundum was obtained by panning the soft rock weathered during pre-glacial time and now located beneath the normal moraine. Many of the corundum crystals were weakly translucent and of a pink or red colour, but no precious corundum was found. Six crystals were polished by Mr. Tauno Paronen, but no star figures were visible. The amount of corundum in the rock is so small and the crystals so impure that quarrying for industrial corundum would not be economic. Nevertheless, the occurrence is interesting because of its mineralogy, chemistry and metamorphic facies.

The field work for this study was done by Haapala and Yletyinen. The chemical analyses were made by Siivola and Ojanperä, while Haapala is responsible for other mineralogical and petrological studies and the conclusions.
Geology

The corundum, sapphireine and kornerupine occur in medium-grained metamorphic mafic-ultramafic rocks with indistinct layered or gneissose structure. The mineral composition of the rocks is variable but in general they are amphibolite or, rarely, hornblendite. The main minerals are hornblende and plagioclase. In places hornblende forms phenocrysts up to several centimeters in diameter. Plagioclase is often concentrated in thin bands parallel to the foliation and to the elongation of the complex. However, plagioclase veinlets cutting the foliation have also been found. Plagioclase of the amphibolite-hornblendite formation may be almost fresh anorthite, but often it is strongly altered to sericite, clinozoisite and, in small amounts, to carbonate. The unaltered plagioclase was identified as anorthite by microprobe technique.

On the basis of thin section studies, the following main mineral associations were distinguished in the amphibolite-hornblendite formation:

1. Hornblende-plagioclase ± chlorite (most common type)
2. Hornblende-plagioclase-corundum ± chlorite
3. Hornblende-gedrite-sapphireine-corundum (-chlorite-plagioclase)
4. Hornblende-corundum-kornerupine-sericite-clinozoisite-chlorite (only one boulder)

Alteration products of plagioclase, and aluminous chromite are common accessories. The kornerupine-bearing sample also contains some dravite.

Corundum occurs as crystals detectable with the naked eye in only a few outcrops (Nos. 40, 41 and 44 in Fig. 2). Microscopically, small remnants of replaced corundum can be detected in outcrop 3. The corundum crystals often form rows of grains parallel to the foliation of the rock (Fig. 3). Boulders containing corundum can be traced over a long narrow zone parallel to the formation, part of which is visible in Fig. 2. This
Red corundum, sapphirine and kornerupine from Kittilä ... 223

Corundum

The corundum crystals found in the outcrops are generally about one centimeter or less in diameter, though crystals as large as 3—4 cm suggest that the corundum-bearing rock forms a conformable zone in the amphibolite-hornblende formation. Even in the richest part, the corundum content is only a few per cent. A hand specimen rich in corundum was polished and the corundum content (3.5 per cent) was determined with a planimeter.

A corundum-bearing sample (type 2) taken from outcrop 41 was analysed chemically by Pentti Ojanperä. The result was: SiO₂ 45.09, TiO₂ 0.18, Al₂O₃ 19.78, Cr₂O₃ 0.05, Fe₂O₃ 1.52, FeO 3.84, MnO 0.09, MgO 14.67, CaO 10.77, Na₂O 1.29, K₂O 0.26, P₂O₅ 0.01, CO₂ 0.00, H₂O + 2.24, H₂O— 0.04, total 99.83. Sp. gr. 3.03. The corresponding Niggli numbers are: si 88.64, al 22.95, fm 51.58, c 22.68, alk 2.78, mg 0.87, k 0.12, ti 0.27 and p 0.01.

The titanium content of this rock is much lower, and the alumina content higher, than in normal magmatic basic rocks (see e.g. Turner and Verhoogen, 1960).

On the western side of the amphibolite-hornblende formation is granite gneiss — a strongly tectonized rock with quartz, microcline, plagioclase, biotite and hornblende as the main minerals.

In the east, the amphibolite-hornblende formation is in contact with a finer grained banded amphibole gneiss-amphibolite formation. The banded structure is due to the concentration of quartz and plagioclase, and hornblende in different bands. Some quartz in these rocks occur as thin veins and is younger than the other main minerals. Hornblende is much darker than in the amphibolite-hornblende formation. Plagioclase is commonly strongly altered. In some thin sections almost fresh andesine, and areas in which andesine is replaced by albite and clinozoisite-epidote, can be seen. Titanite, calcite and opaque minerals occur in small amounts.

Corundum

The corundum crystals found in the outcrops are generally about one centimeter or less in diameter, though crystals as large as 3—4 cm


Fig. 3. Corundum-bearing amphibolite, outcrop 41. Corundum is visible as white roundish spots, some of which are marked with a small arrow. 1/10 of natural size. Photo Ilmari Haapala.
in diameter can be found. By far the largest and most striking corundum crystal is that of the original boulder sample (Fig. 1). The colour is normally pink, white or red, but under the binocular microscope also some deep blue corundum grains were detected in the material crushed for mineral separation. At the margins, the crystals are usually replaced by colourless optically positive chlorite (Fig. 4), and sometimes also by hornblende or sericite. The replacement is often almost complete so that only scattered remnants with a uniform orientation remain (Fig. 4).

Hornblende

In general, hornblende represents the most abundant mineral of the amphibolite-hornblendite formation, though in some samples the amount of plagioclase exceeds that of hornblende. The colour of hornblende is green in a hand specimen, but in thin section the mineral is colourless.

Hornblende was separated for chemical analysis from a sapphirine-bearing sample (40/IH/67) taken from outcrop 40. The physical properties of this material are given in Table 1. The unit cell dimensions were determined by the Buerger precession method using Zr-filtered Mo radiation. The film shrinkage was calibrated by recording silicon [111]-axis zero-level precession photographs on the same films. The specific gravity was determined with Clerici solution and a Westphal balance. Refractive indices were measured in Na light by the immersion method. A universal stage was used in measuring the optic angle. The same methods were also employed in determining the physical properties of gedrite, sapphirine and kornerupine.

On the basis of the formula calculated from the chemical analysis (Table 1), the mineral is magnesio-hornblende (Leake 1968). In the three-

TABLE 1
Chemical composition and physical properties of hornblende. Chemical analysis by Pentti Ojanperä

<table>
<thead>
<tr>
<th>Wt%</th>
<th>Number of ions on the basis of 24 (O, OH)</th>
<th>Physical properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>Si 6.690</td>
<td>Al²⁺ 1.130</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>Al⁴⁺ 0.892</td>
<td>Ti³⁺ 0.021</td>
</tr>
<tr>
<td>TiO₂</td>
<td>Cr⁺ 0.019</td>
<td>Fe³⁺ 0.123</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>Fe²⁺ 0.413</td>
<td>Mn⁰ 0.011</td>
</tr>
<tr>
<td>Fe₂O₃</td>
<td>Mg³⁺ 3.607</td>
<td>Ca 1.867</td>
</tr>
<tr>
<td>FeO</td>
<td>Na 0.312</td>
<td>K 0.030</td>
</tr>
<tr>
<td>MnO</td>
<td>OH 1.982</td>
<td>O 22.018</td>
</tr>
<tr>
<td>MgO</td>
<td>100.35</td>
<td></td>
</tr>
</tbody>
</table>

The unit cell:
Space group $\text{C2}/\text{m}$

$\alpha_a = 9.77 \pm 0.01\text{Å}$
$\beta_b = 17.97 \pm 0.02\text{Å}$
$\gamma_c = 5.30 \pm 0.005\text{Å}$
$\beta = 104^\circ 58' \pm 05'$
$V = 898.9\text{Å}^3$

Sp.gr. $\text{meas} = 3.086 \pm 0.005$
$D_{\text{calc.}} = 3.09 \text{g/cm}^3$

The optical properties:
$\alpha = 1.628$
$\beta_{\text{calc.}} = 1.638 \pm 0.002$
$\gamma = 1.652$
$2V\alpha = 101 \pm 3^\circ$
dimensional coordinate system proposed by Whittaker (1968) for the classification of amphiboles the mineral falls into the composition space centered on coordinates 211. The alumina content is remarkably high, reflecting the high alumina content of the rock as a whole. This mineral is biaxially positive and thus differs from most common hornblendes.

Hornblende from the kornerupine-bearing sample has refractive indices $\alpha 1.630$ and $\gamma 1.654$, suggesting a slightly lower Mg:Fe ratio than in the hornblende of the sapphirine-bearing sample.

Gedrite

Gedrite is found as long prismatic crystals in the sapphirine-bearing rock. The crystals may be several centimeters in length. Macroscopically, the mineral is beige in colour but in thin section it is colourless. It is very difficult to distinguish gedrite from hornblende in thin section, because both minerals appear to be colourless. For the universal stage measurements the gedrite grains had to be identified from a polished thin section by microprobe.

A partial chemical analysis was made from a polished thin section of the sample 40/IH/67 using the microprobe technique (Table 2). The ratio $\text{Mg}/(\text{Mg} + \text{Fe})$ is 84.7, i.e. somewhat lower than in hornblende (cf. Klein 1968). The optical properties are in good agreement with the diagrams for the relationships between these variables and the chemical composition of antophyllite-gedrite (Seki and Yamasaki 1957).

Sapphirine

Deep blue sapphirine is found in one outcrop (40) and in one boulder (7). In both cases, the rock has the mineral assemblage hornblende-gedrite-sapphirine-corundum (-chlorite-anorthite). Sapphirine is typically intergrown with hornblende (Fig. 5) or corundum, and is in many cases partially replaced by chlorite. Some larger sapphirine crystals are repeatedly twinned on.

From a crystal splinter of the sapphirine, $c$-axis zero-, first-, second-, and third-level Weissenberg photographs were taken with Ni-filtered Cu radiation, as well as $b$- and $[101]$-axis zero-, first-, and second-level precession photographs with Zr-filtered Mo radiation. The systematic extinctions ($hkl$ no conditions, $b0l$ absent if $b$ is odd, $0k0$ absent if $k$ is odd) agree with the requirements of the space group $Pnma$, recorded by Moore (1969) for Fiskenaesset sapphirine. The systematic extra-extinctions recorded for sapphirine from four localities by Fleet (1967) do not exactly fit the Kittilä material. After long expos-

### Table 2

<table>
<thead>
<tr>
<th>Wt %</th>
<th>Number of cations on the basis of 23 O.</th>
<th>Physical properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO$_2$</td>
<td>Si 6.28</td>
<td>The unit cell:</td>
</tr>
<tr>
<td>TiO$_2$</td>
<td>Al[4] 1.72</td>
<td>Space group $Pnma$</td>
</tr>
<tr>
<td>Al$_2$O$_3$</td>
<td>20.4</td>
<td>Mg 4.38</td>
</tr>
<tr>
<td>Cr$_2$O$_3$</td>
<td>trace</td>
<td>Fe 0.79</td>
</tr>
<tr>
<td>FeO</td>
<td>trace</td>
<td>Ca 0.12</td>
</tr>
<tr>
<td>MnO</td>
<td>trace</td>
<td>Na 0.29</td>
</tr>
<tr>
<td>MgO</td>
<td>21.8</td>
<td></td>
</tr>
<tr>
<td>CaO</td>
<td>0.8</td>
<td></td>
</tr>
<tr>
<td>K$_2$O</td>
<td>trace</td>
<td></td>
</tr>
<tr>
<td>Na$_2$O</td>
<td>1.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>97.7</td>
<td></td>
</tr>
</tbody>
</table>

The ratio $\text{Mg}/(\text{Mg} + \text{Fe}) = 84.7$.
Sapphirine was analysed from a polished thin section of sample 40/IH/67 by microprobe (Table 3). The silica content is a little lower and the alumina content somewhat higher than in the previous sapphirine analyses (Deer, Howie and Zussman 1963, Wilson and Hudson 1967, Lutts and Kopaneva 1968, Forestier and Lasnier 1969) suggesting a pronounced coupled substitution Al, Al for Mg, Si. The asymmetric formula unit contains $\text{Mg}_3\text{Fe}^{2+}_{0.14}\text{Fe}^{3+}_{0.20}\text{Al}_{3.94}\text{Si}_{14.45}\text{O}_{20}$.

**Kornerupine**

Kornerupine was recognised only in the original boulder sample found by Mikko Tervo (Fig. 1). The main minerals in this sample are hornblende, corundum, kornerupine, sericite, clinozoisite and chlorite. Fresh plagioclase (anorthite), tourmaline (dravite) and chromite occur in smaller amounts. Kornerupine occurs as pale bluish green crystals up to 1 cm in diameter. In thin section, the mineral is colourless and has indistinct {110} cleavage. The kornerupine crystals are veined and strongly replaced by chlorite and fine-grained mica (Fig. 6), and the associated plagioclase grains are in general

**Table 3**

Chemical composition and physical properties of sapphirine. Electron microprobe analysis by Jaakko Siivola

<table>
<thead>
<tr>
<th>Wt%</th>
<th>Number of cations on the basis of 80 O</th>
<th>Physical properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{SiO}_2$ 12.2</td>
<td>Si 5.80</td>
<td>The unit cell:</td>
</tr>
<tr>
<td>$\text{TiO}_2$ n.o.</td>
<td>Al 35.35</td>
<td>Space group $P2_1/\alpha$</td>
</tr>
<tr>
<td>$\text{Al}_2\text{O}_3$ 63.1</td>
<td>Cr 0.02</td>
<td>$a_0 = 11.28$</td>
</tr>
<tr>
<td>$\text{Cr}_2\text{O}_3$ 0.04</td>
<td>$\text{Fe}^{3+}$ 0.82</td>
<td>$b_0 = 14.42 \pm 0.01\text{Å}$</td>
</tr>
<tr>
<td>$\text{Fe}_2\text{O}_3$ 2.31</td>
<td>$\text{Fe}^{3+}$ 0.60</td>
<td>$\epsilon_0 = 9.95$</td>
</tr>
<tr>
<td>$\text{FeO}$ 1.51</td>
<td>Mg 13.53</td>
<td>$\beta = 125.28\pm 0.05^\circ$</td>
</tr>
<tr>
<td>MnO n.o.</td>
<td></td>
<td>$V = 1318\text{Å}^3$</td>
</tr>
<tr>
<td>MgO 19.1</td>
<td></td>
<td>Sp-gr$_{\text{mass}} = 3.518 \pm 0.005$</td>
</tr>
<tr>
<td>CaO n.o.</td>
<td></td>
<td>The optical properties:</td>
</tr>
<tr>
<td>Na$_2$O n.o.</td>
<td></td>
<td>$\alpha = 1.713$</td>
</tr>
<tr>
<td>K$_2$O n.o.</td>
<td></td>
<td>$\beta = 1.716 \pm 0.002$</td>
</tr>
<tr>
<td>Loss 0.371</td>
<td></td>
<td>$\gamma = 1.718$</td>
</tr>
<tr>
<td>of ign. 98.61</td>
<td></td>
<td>$2V_0 = 65 \pm 2^\circ$</td>
</tr>
</tbody>
</table>

3) Wet chemical determination by Pentti Ojanperä. n.o. not observed
completely saussurized. Idiomorphic twinned clinozoisite crystals are present and may be up to two millimeters long (Fig. 7). Remnants of strongly altered idiomorphic plagioclase crystals are common inclusions in kornerupine (Fig. 8). Unaltered plagioclase can be found only as enclosed, shielded crystals. A microprobe analysis of such a crystal showed it to be anorthite. Corundum may also occur as inclusions in kornerupine.

A series of precession photographs about all the crystallographic axes were taken to determine the space group and unit cell dimensions of kornerupine. The conditions for non-extinction of the reflections were: \( hkl \ b + k = 2n, \ b0l \ l = 2n. \) These data are compatible with space groups \( Cmcm, \ Cm2_1, \) and \( C2cm, \) conforming to the studies of Bartl (1965), McKie (1965) and Knorring et al. (1969). Moore and Bennet (1968) recorded space group \( Cmcm \) on the basis of a crystal structure analysis of the Mautia Hill kornerupine.

The Kittilä kornerupine was so strongly metamorphosed that it was not possible to obtain sufficient pure material for a complete wet chemical analysis. Therefore, the main part of the analysis was made by microprobe. \( H_2O^+ \) was determined using a Penfield tube from a small amount (60 mg) of carefully hand-picked material. Iron was determined by a wet chemical method and boron spectrochemically from material containing small amounts of alteration products as impurities. The amount of impurities was so small that their influence on the iron and boron contents is believed to be insignificant. However, the water content may be slightly high. According to the microprobe analysis, the total iron content calculated as FeO is 4.9%.

The boron content (Table 4) is somewhat lower than in other contemporary kornerupine analysis (cf. Knorring et al. 1969). Correspondingly, the silica content is slightly greater than usually, suggesting an unusually low substitution of B for Si. Moore and Bennett (1968) suggested the simplified formula \( Mg_3Al_6(Si, Al)_2O_{21}(OH) \)
TABLE 4
Chemical composition and physical properties of kornerupine. Electron probe micro-
analysis by Jaakko Siivola

<table>
<thead>
<tr>
<th>Number of ions on the basis of 88 (O, OH)</th>
<th>Physical properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wt %</td>
<td>Si 15.62</td>
</tr>
<tr>
<td>SiO₂ 31.7</td>
<td>B 0.85</td>
</tr>
<tr>
<td>B₂O₃ 1.0</td>
<td>Al 25.27</td>
</tr>
<tr>
<td>TiO₂ n.o.</td>
<td>Cr 0.13</td>
</tr>
<tr>
<td>Al₂O₃ 43.5</td>
<td>Fe²⁺ 0.85</td>
</tr>
<tr>
<td>Cr₂O₃ 0.33</td>
<td>Fe³⁺ 0.78</td>
</tr>
<tr>
<td>FeO 2.3</td>
<td>Mn 0.04</td>
</tr>
<tr>
<td>FeO 1.9</td>
<td>Mg 12.78</td>
</tr>
<tr>
<td>MnO 0.1</td>
<td>Ca 0.04</td>
</tr>
<tr>
<td>MgO 17.4</td>
<td>OH 4.93</td>
</tr>
<tr>
<td>CaO 0.07</td>
<td>O 83.07</td>
</tr>
<tr>
<td>Na₈O trace</td>
<td></td>
</tr>
<tr>
<td>K₂O n.o.</td>
<td></td>
</tr>
<tr>
<td>H₂O⁺ 1.5</td>
<td></td>
</tr>
<tr>
<td>99.80</td>
<td></td>
</tr>
</tbody>
</table>

1) Spectrochemical determination by Arvo Löfgren
2) Wet chemical determination by Pentti Ojanperä
n.o. not observed

for kornerupine. The formula calculated for the Kittilä kornerupine is Mg₃₋₉Fe²⁺⁸⁻₀₋₂Al₄₋₅Fe³⁺⁰₋₂Si₃₋₄B₂₋₅O₂₋₈(OH)₁₋₂.

A partial semiquantitative microprobe analysis of the colourless, optically positive chlorite associated with kornerupine gave SiO₂ 29.3, Al₂O₃ 26.6, FeO 6.8 and MgO 21.9 wt %. This indicates that the mineral is clinochlore. An analysis of tourmaline by a similar method gave SiO₂ 35.6, Al₂O₃ 35.6, FeO 2.0, MgO 10.8, CaO 1.5—1.8, and Cr₂O₃ 0.41 wt % showing the mineral to be dravite. Tourmaline often occurs at the margins of the corundum crystals and may penetrate them as thin veinlets. This suggests boron metasomatism.

Discussion

The literature dealing with the rare rock-forming mineral sapphirine is fairly comprehensive. Types of sapphirine and kornerupine occurrences and their stability relations are summarized by Vogt (1947), Sørensen (1955), Deer, Howie and Zussman (1963), Herd et al. (1969), Forestier and Lasnier (1969) and Schreyer and Seifert (1969). The most common type is composed of sapphirine-bearing metamorphosed basic-ultrabasic rocks of granulite terrains. The minerals commonly associated with sapphirine are orthopyroxene, hornblende, gedrite, phlogopite, plagioclase, cordierite, spinel, corundum and rarely kornerupine. Recently, sapphirine has also been described from rocks of the skarn type (Knorring 1967, Rouhunkoski 1968, p. 29).

In his study on the sapphirine occurrences, Sørensen (1955) discussed the stability relations of sapphirine. He concluded that sapphirine was formed in rocks rich in alumina and magnesia but poor in silica, which have been metamorphosed under the P-T conditions corresponding to high amphibolite facies or low granulite facies. His study largely conforms to the conclusions of Vogt (1947) and Ramberg (1948). According to Turner and Verhoogen, (1960) sapphirine is formed under conditions of hornblende-granulite subfacies. Herd et al. (1969) made a detailed study of the petrology of different sapphirine-bearing rocks of the Fiskensæset region in West Greenland. These rocks were derived from layered spinel-bearing ultramafic rocks which formed a minor part of chromite-bearing layered anorthosite complex. The comp-
lex has undergone hornblende-granulite facies
metamorphism followed by cordierite-amphi-
bole'ite facies metamorphism. Herd et al. dis-
tinguished enstatite, pargasite, gedrite and phlo-
gpite types of sapphirine-bearing assemblages,
which represent increasing degrees of Si, Ca, K
and H2O metasomatism related to shearing and
deformation. Herd et al. (op. cit, p. 41): »Sapphirine
was thus formed under hornblende-granulite
facies conditions, was stable in reduced
amounts under cordierite-amphibolite facies
conditions and was stable even in relatively calcium
rich assemblages where it was able to crystallise
under amphibolite facies conditions». Accordingly
they state that sapphirine is stable under
a wide range of P-T conditions, but within a
limited range of chemical environments.

According to the experimental studies of
Schreyer and Seifert (1969) of the system
MgO-Al2O3-SiO2-H2O assemblages containing
sapphirine, gedrite and kornerupine may be
formed at elevated pressures and intermediate
temperatures. Sapphirine, corundum and gedrite
appear as stable coexisting phases in the system
MgO-Al2O3-SiO2-H2O at a pressure of 14 kb
and temperature 850° (Schreyer 1968, Fig. 27).
At higher temperatures (810°—910°C, depending
on the pressure) gedrite breaks down to enstatite-
bearing assemblages. Boronfree kornerupine
appears to be stable at 8—13 kb and 850° ± 50°C.
Because of the greater complexity of natural
systems, however, the phase relations in the
system MgO-Al2O3-SiO2-H2O can be applied
only with reservations. For example, the Kittilä
material contains notable amounts of calcium,
iron and alkalies, which would change the PT-
conditions.

The corundum-sapphirine-kornerupine occur-
rence described in this paper is located just out-
side the great granulite complex of Lapland
(Mikkola 1937, Mikkola and Sahama 1936). Poor
exposures make it difficult to draw any detailed
conclusions concerning the genesis and petrology
of the amphibolite-hornblendite formation. In
every case it is apparent that the complex was
metamorphosed under at least two different P-T
conditions; (a) under high-grade amphibolite —
low-grade granulite facies, and (b) with slight
retrogression under greenschist facies. According
to Winkler (1967) gedrite is not stable under
granulite facies conditions. The sapphirine-
bearing assemblage (type 3, p. 222) corresponds
to a transition between the pargasite and gedrite
types of Herd et al. (1969).

Zakrutkin (1968) has shown that the compo-
sition of hornblende depends on the meta-
morphic facies. Plotted on his diagram based on
Al4 and Al6 contents, the analysed hornblende
from the sapphirine-bearing assemblage falls into
the amphibolite facies field but very near the
junction of the albite-apidote-amphibolite facies.
A similar result is obtained from Zakrutkin's
diagram based on the contents of (Fe3+ + A6) +
Ti and (Mg + Fe2+ + Mn). The high content
of Al6 suggests unusually high pressure during
metamorphism (Kostyuk and Sobolev 1969).

It is not clear what the rocks were before
the high-grade metamorphism. The Niggli
numbers (p. 223) suggest the formation to be
an "ortho-amphibolite". Compared with c and
al-alk the mg number is somewhat higher than
it should be in mixtures of normal pelites or
semipelites and dolomites or limestones (cf. Leake
1964). Also the occurrence of chromite as an
accessory mineral points to igneous origin. The
large hornblende phenocrysts are possibly
pseudomorphs after pyroxene. The high alumina
and magnesia contents may be derived from the
primary spinel-rich layers in the mafic-ultramafic
formation, as is the case in the Fiskenaeset
region (Herd et al. 1969). Spinel has been de-
scribed from some ultramafic rocks near to the
granulite complex of Lapland (Mikkola and Sa-
hama 1936). However, the data are insufficient
for excluding the possibility of metasedimentary
origin.

It appears that the main metamorphism took
place under high pressure and high temperature
conditions at or near the boundary between
amphibolite and granulite facies. Greenschist
In this sample unshielded plagioclase is completely replaced by sericite and clinozoisite, and kornerupine very strongly by chlorite. Thus, the retrograde metamorphism shows a tendency to destroy corundum, sapphirine and kornerupine.

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