A CONTRIBUTION TO THE CRYSTAL CHEMISTRY OF MELANOPHLOGITE

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ABSTRACT

Melanophlogite from the second occurrence at Chvaletice, Bohemia, forms colorless cubes on lussatite in a vein cavity in a metamorphosed sedimentary pyrite-rhodochrosite deposit of Algonkian age. Fine growth zoning is parallel to the cube faces which represent bases of tetragonal sectors, composing each cube. The superstructural unit cell has space group $P4_3/nbc$, $a = 26.82$, and $c = 13.37$ Å. It breaks down by heating at 1,050°C to a cubic unit cell with space group $P4_3 2$ and $a = 13.4$ Å. The typical jet black color appears, but no carbon particles can be observed even at greatest magnifications. Emission spectrography indicates only Si in substantial quantity. Electron microprobe, activation, and microchemical analyses lead to a unit cell content that can be formulated as $46\text{SiO}_2\cdot3\text{C}_2\text{H}_5\text{O}_8\cdot\text{N}_2\cdot\text{H}_2\text{O}$. The sulfur content is much lower than that in the Sicilian melanophlogite. The clathrate structure, suggested by Kamb (1965) for melanophlogite, can be also applied for the Chvaletice mineral. Compounds of the minor elements, indicated by the formula, are framework-cavity guests and a probable cause of the tetragonal superstructure.

INTRODUCTION

The existence of melanophlogite, a cubic polymorph of silica, was confirmed only recently on material from the first occurrence of this mineral, Racalmuto in Sicily (Skinner and Appleman, 1963). Subsequent studies (Kamb, 1965; Appleman, 1965) have revealed that it is a clathrate-type compound, enclosing organic and other molecules in large polyhedral framework cavities.

A unique sample with melanophlogite was found by the mineral collector M. Duchoň in the Chvaletice deposit, eastern Bohemia, during the mining period 1964–65. Owing to its external appearance it was regarded as fluorite and kindly placed at the author’s disposal for study. Preliminary optical, spectrographic, and X-ray investigations revealed its identity with melanophlogite.

The mineral assemblage containing melanophlogite (Žák, 1967) forms a vein in the metamorphosed pyrite-rhodochrosite deposit of the Algonkian Ore-Formation (Svoboda and Fiala, 1951). Melanophlogite cubes (Fig. 1) overlie a one-millimeter of layer transparent bluish-white lussatite (Žák, 1968). Under magnification tiny lussatite spherulites are commonly observable on melanophlogite crystals. The lussatite layer encrusts colorless dolomite crystals, under which minute
pyrite crystals occur locally. Then follow layers of pinkish, concentrically-zoned rhodochrosite (X mm), pyrite (1 mm), and a fine grained pyrite-marcasite-apatite aggregate (5 mm). The lussatite is microscopically radial-fibrous and its medium index of refraction is 1.44. By X-rays a strongly disordered low cristobalite was established, alterable to an ordered low cristobalite after heating at 1,600°C for 10 hours (see Flörke, 1956).

**Physical Properties**

*Morphology and Optical Properties*

The cubes of melanophlogite are isolated or grow irregularly together.
Exceptionally, indications of penetration twins according to (111) were observed (see von Lasaulx, 1876). The surface of colorless transparent crystals is dull white and covered by a light brown limonite coating, easily dissolved by dilute acids. The whitish central (Žák, 1967) and other parts of crystal faces exhibit a remarkable skeletal growth (Fig. 2). Similar growth forms had been found in the Sicilian melanophlogite (von Lasaulx, 1876).

Microscopically, the melanophlogite powder is colorless and isotropic. By combination of immersion and refractometric methods, \( n_{Na}^{25} = 1.457 \) was determined. Sections parallel to the cube faces reveal a thin weakly anisotropic rim around a very weakly anisotropic to isotropic center. Similar observation was reported by Zambonini (1906) on melanophlogite from Giona, Sicily. The birefringence of the marginal zone of the Chvaletice mineral gradually decreases from the boundary with the core towards the cube faces. Thicker sections (\( \sim 1 \) mm) reveal both in the rim and in the core fine zoning, parallel to the cube faces (Fig. 3), and triangular sectors that resemble six tetragonal pyramids composing one cube of Sicilian melanophlogites.

Fig. 2. Skeletal structure of a melanophlogite cube face, as seen by scanning electron microscope. Gold coated. Photo by A. Klatt, Bochum.
Fig. 3. Fine growth zoning. Boundaries between two sectors and between the rim and the core (dark heavy line) are clearly visible. Section, parallel to a cube face, the anisotropic margin of which was polished off. Universal stage, polarizer only.

(Bertrand, 1880; Friedel, 1890; Skinner and Appleman, 1963). The sectors in the central parts of the crystals are very weakly birefringent with transitions to optical isotropy in places (Fig. 4). The extinction of the whole cube section is parallel to the cube edges. The optical axis of the anisotropic outer zone of each sector is perpendicular to \{001\}, which is indicated between crossed nicols by isotropy for the light propagating in this direction. No conoscopic figure can be obtained (see also Friedel, 1890; Zambonini, 1906). Finally, thin lamellae, differing in birefringence from their vicinity, or dark lines only, are seen parallel to \{110\} (Fig. 4). They intersect the rim and core boundary in some cases. The extinction of them is, again, conformable with that of the whole sector. No fluorescence was observed in the
Chvaletice melanophlogite in short (254 nm) and long-wave (366 nm) ultra-violet light.

Density

A density determination was made by suspension in Klein solution diluted by water, the liquid density being measured with a Mohr-Westphal balance. Three fragments, about one millimeter in size, including the marginal zone, were used. The limonitic coating was removed by washing with hot nitric acid and distilled water. The density determined at $21^\circ$C is 2.005 g cm$^{-3}$. 
Microhardness

Vickers' microhardness was determined by the Durimet (Leitz-Wetzlar) microscope. The load was 100 g, duration of the test 10 sec. A carefully polished section of melanophlogite, parallel to the cube face, and a nonoriented section of quartz from Chvaletice, mounted simultaneously in an epoxy resin block, were examined. Average values of ten measurements are 680 (range 649 to 724) kg mm\(^{-2}\) in melanophlogite and 1,330 (range 1,310 to 1,427) kg mm\(^{-2}\) in quartz. The Vickers' hardness number for melanophlogite corresponds approximately to 6.5 on the Mohs' scale (Zussman, 1967), which is in a good agreement with the value for the Sicilian material of 6.5–7 (von Lasaulx, 1876).

Thermal Behavior

After heating in air to a red glow for one minute or in an electric furnace at 800°C for three hours, the melanophlogite fragments became smoky brown in color. The previously anisotropic marginal zone became more intensively colored than the core, and turned isotropic. The intensity of the rim coloring increases towards the contact with the central part. In the Sicilian melanophlogite, observed on the heating stage (Zambonini, 1906), a colorless birefringent rim got nearly isotropic at 150°C in one case, and darkening of yellow-brown zones in the core occurred at different temperatures from 200 to 300°C. The color, zonal structure, and likely also the differences in darkening temperatures were ascribed to an organic pigment. After heating the Chvaletice mineral at 1,050°C for six hours, the crystal fragments, including both rim and core, turned jet black. They were opaque in thin section, with isotropic brown translucent or transparent edges. No heterogeneous carbon particles could be observed even at the greatest magnification of the microscope (1,600 X), contrary to the Sicilian melanophlogite (Skinner and Appleman, 1963). In powder preparations, translucent fragments show frequently brown and almost opaque narrow zones, rarely intersecting at an angle of about 120°. They can be correlated with the above-mentioned zoning and lamellae.

Infrared Absorption Spectrography

Absorption bands, characterizing unheated and heated melanophlogites, are presented in Figure 5. Non-aromatic hydrocarbons or their derivatives are suggested absorptions between 2,800 and 3,000 cm\(^{-1}\) (curve 1) disappearing after heating (curve 3), but proof is lacking. In some of other runs (e.g. curve 2) the mentioned absorptions were not
so clearly detectable, so that organic contamination could be present. Water molecules as framework cavity guests are uncertain, though the absorption at 1,630 cm\(^{-1}\) was also found by the Nujol technique in a sample dried at 120°C for 1 hour and immediately transferred into the Nujol paste. Water remaining adsorbed on the powder particles cannot be excluded.

**X-ray Crystallography**

The powder pattern (Table 1) contains a number of weak diffractions that cannot be indexed by a cubic 13 Å unit cell (Skinner and Appleman, 1963), and that indicate the presence of a superstructure with a doubled cell edge. This is confirmed by single crystal methods. Rotation photographs were made with a thin section (~0.2 mm) parallel to a cube face, after polishing and cutting off the anisotropic rim. This core section was prepared from the other half of the crystal, used for optical study (Figs. 3 and 4). The section was nearly
isotropic under universal stage observation. With rotation axes parallel to the plane of the section ([100] and [010]), the X-ray photographs are identical and show weak layer lines indicating doubled cell edges. With [001] rotation axis, only extremely weak reflections of these layer lines are obtained. Rotation photographs of a marginal zone section give similar results. The photographs indicate a common tetragonal supercell (axial ratio 2:2:1 relative to the 13 Å subcell) with vertical axis perpendicular to the plane of the thin section. This cell is confirmed and specified by equi-inclination Weissenberg photographs of the core section and cube edge fragments.

Survey of the Photographs Made

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<th>Rotation axes¹</th>
<th>Core section</th>
<th>Edge fragment</th>
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The photographs can be evaluated by the same tetragonal cell, the only difference being in the proportion of three orientations of intergrown cells in the X-rayed region. Whereas the edge fragments produce diffractions of a considerable intensity from two or three cell orientations with perpendicular c-axes, the core section demonstrates a prevalence of only one orientation of the cell. Therefore, the edge-fragment photographs are difficult to interpret in terms of a tetragonal cell, which is clearly confirmed by the core section photographs (Fig. 6). High-angle diffractions, e.g. 32.0.0 and 0.0.16 (indices relative to the 2:2:1 cell), demonstrate a difference of the corresponding interplanar spacings. The presence of the reflection 34.0.0 and the absence of an analogous 001 diffraction indicates doubling of the larger 13 Å subcell edge only. The photograph with the [001] rotation axis contains both 32.0.0 + 34.0.0 and 0.32.0 + 0.34.0 diffractions, with equal reflection angles in each pair of corresponding reflections.

The supercell dimensions \( a = 26.82 \pm 0.03 \text{ Å} \) and \( c = 13.37 \pm 0.02 \text{ Å} \).

¹ Indices relative to the prevailing orientation of the supercell.
² Comparable with second layer lines of photographs with [100] and [010] rotation axes.
Å were calculated from the 32.0.0 and 0.0.16 diffractions of a Weissenberg photograph (core section), calibrated by silicon (a = 5.43054 Å.

The photographs of the higher layer lines with [100] rotation axes display considerable differences in intensities of hkl and lkh reflections, in accordance with the tetragonality of the cell. The odd layer lines (exposures up to 100 hours) show weak diffractions with two odd indices (Fig. 7) relative to the axial ratio 2:2:2, confirming the 2:2:1 supercell (see also Pyatenko, 1967). No diffractions with three odd indices have been found.

The three orientations of the tetragonal supercell with perpendicular vertical axes produce splitting of high angle reflections, both in even and odd layer lines. The most striking are doubled reflections 32.0.0 + 0.0.16 in the photographs of the edge fragments. Weak reflections of a different cell orientation are obtained also in the photographs of the central section (Fig. 6). Some diffractions, e.g., 32.12.0, 12.0.16, and 32.0.6, should form triplets. Only doublets have been observed on the photographs. This can be explained by: (a) Only two prevailing orientations of the cell in the X-rayed region of the crystal, or, (b) In

Fig. 6. Zero layer h0l Weissenberg photograph, core section of the unheated melanophlogite. Note the 32.0.0, 34.0.0, and 0.0.16/∞ + ∞/reflections of the prevailing cell. Another orientation of the cell produces weak diffractions only: 32.0.0 ∞(arrow) and 0.0.16∞(arrow). Cu/Ni radiation, exposure 52 hours.
some other cases also by coincidence of two diffractions with a very small difference of Bragg angle. The h0l split reflections do not lie on central reciprocal lattice lines, as it seems from their postions in the photographs. Owing to the strongly pseudocubic character of the tetragonal 13 Å subcell and the great distortion of the reciprocal lattice geometry in the Weissenberg photographs, their maximum difference in position with respect to the central lattice lines is too small to be observed (~0.1 mm).
Table 1. Continued

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<sup>a</sup>Guinier-de Wolff camera, CuK<sub>α</sub> radiation, 40 kV, 25 mA, exposure 24 hours.

<sup>b</sup>Intensities are taken from a powder pattern, made without the internal standard

<sup>c</sup>Indices inconsistent with the space group P4<sub>2</sub>/nbc are left out.

<sup>d</sup>Found only in the powder pattern without the internal standard.

The reflection statistics (indexing relative to the tetragonal 2:2:1 supercell) lead to the space group P4<sub>2</sub>/nbc (Nuffield, 1966): hkl present in all orders, hk0 with h+k = 2n, hhl with l = 2n, and 0kl with k = 2n only.

The powder pattern (Table 1) was calculated for an average tetragonal supercell with a = 26.79 Å and c = 13.395 Å, based on a hypothetic cubic subcell with a = 13.395 Å. This subcell is similar in dimensions to the cubic cell of the heated melanophlogite.

**Twinning**

As indicated by optical and single-crystal X-ray study, the cubes of the Chvaletice melanophlogite are composed of six pyramids, whose apices meet at the center of the cube and bases represent the cube faces. The pyramids are formed by fine tetragonal tablets, growing parallel with their bases together (Fig. 3). Their vertical axis is common and perpendicular to the corresponding cube face. Similar
composite crystals were observed on melanophlogite from Sicily by Bertrand (1880), Friedel (1890), and Skinner and Appleman (1963). The adjoining pyramids and the lamellae in them probably represent some kind of twinning. Twinning of the adjoining individuals with \{110\} symmetry planes (relative to the cube) is not probable, owing to the tetragonal distortion of the 13 Å cubic cell, leading to a very complicated index of the symmetry plane. Twinning with \{201\} symmetry planes (relative to the tetragonal supercell) requires the c-axes of the individuals to meet at an angle of about 90°10'. No such deviation from perpendicularity could be observed in the X-ray single crystal photographs or by the microscope. In this connection it is interesting to note Friedel's (1890) observation of higher than 90° angles of the cube faces and Zambonini's (1906) observation of a tetrahexahedron of a complicated index, on the Sicilian melanophlogite crystals. A slight compression of cube corners towards the centre of the cube was observed on the Chvaletice crystals, too. The questions need further study.

**Heated Melanophlogite**

Small marginal-zone + core fragments were heated in air at 1,050°C for 12 hours. Powder and rotating crystal X-ray photographs (rotation axis [100]) revealed a breakdown of the superstructure to a cubic structure with \(a = 13.4\) Å. The superstructure reflections disappeared almost completely in the Guinier powder pattern and completely in the single crystal photographs. Equi-inclination Weissenberg photographs of zero and first layer lines (Fig. 7) display a diffuse character of split reflections. Different intensities of many reflections, when compared with the corresponding ones of the unheated mineral, are observable. The reflection survey records the presence of \(hkl\), \(hhl\), and \(0kl\) reflections in all orders and \(00l\) present with \(l = 2n\) only. Of the space groups \(P4_232\) and \(P2_13\), the group \(P4_232\) is indicated by the diffraction symmetry. The space group \(P4_232\) had been given for the Racalmuto melanophlogite by Skinner and Appleman (1963), but the space group \(Pm3n\) was suggested later (Kamb, 1965). Contrary to the Racalmuto melanophlogite with \(hhl\) present only if \(l = 2n\), the heated Chvaletice melanophlogite exhibits the \(hhl\) reflections in all orders (Fig. 8). Nevertheless, the \(hh1\) reflections are weak by comparison with other reflections and the corresponding \(hh1\) or \(h2h\) diffractions of the unheated melanophlogite have not been found.

**Chemical Composition**

Qualitative spectrographic analysis gave Si in substantial, Al, Cu,
MELANOPHLOGITE

Fig. 7. Third h3l(a) and first h1l(b) layer equi-inclination Weissenberg photographs of unheated and heated melanophlogites. Edge fragments. Note the weak 23.3.0, 313, and 616 reflections (arrows). Cu/Ni radiation, exposures 60 and 33 hours. Left upper corners: detail with longer exposures.

Mg, and Mn in insignificant (<0.1% percent), and Ag, Fe, and P? in trace quantities. JXA-3A electron microprobe tests for nitrogen and fluorine were negative, and only the sulfur test was slightly positive. A nitrogen content lower than 1 percent (Weinryb, 1967) probably cannot be detected under the experimental conditions used. Elements with \( Z \geq 11 \), except silicon, were not found.
Sulfur, silicon, and oxygen were determined by activation, carbon and hydrogen by microchemical methods from powdered samples, silicon and carbon by the electron microprobe.

Sulfur activation analysis (0.01 g, F. Kukula analyst) was carried out by nuclear reactor bombardment, combined with a chemical separation of the beta-active $^{35}$S. The standard was ammonium sulfate. A sulfur content $0.11 \pm 0.03$ percent was found.

Silicon and oxygen of the sample (0.03 g) were transformed into $^{28}$Al and $^{14}$N radioisotopes by fast neutrons, produced in a neutron generator (J. Bartošek and I. Kašparec analysts). The standards were quartz from Chvaletice and silica glass. The average value of three measurements is 45 percent (range 43 to 46) for silicon and 53 percent (range 51 to 56) for oxygen.

For silicon analyses with a Cambridge Geoscan electron microprobe, a graphite-coated polished section (from microhardness measurements), mica crystal, accelerating voltage 20 kV, sample current 20 nA, SiK$_\alpha$ radiation, and standard of quartz from Chvaletice were used. Carbon was measured with a Japanese JXA-3A probe (K. Stránský and A. Rek analysts), using a copper-coated section, electron beam diameter 1.8 $\mu$m, lead stearate crystal, 10 kV, 27 nA sample current, CK$_\alpha$ radiation, and standards of graphite and aragonite. The wavelength shift was respected (Castaing, 1960; Kohlhaas and Scheiding, 1969) and the carbon contamination reduced by a slow sample movement (100 $\mu$m per min.). A correction for the remaining carbon contamination was made by the carbon-free quartz standard, analyzed immediately after the melanophlogite. After corrections for background and dead time, Si = 43.6 and C = 0.3 (graphite standard) or 0.6 percent (aragonite standard) were obtained. These raw data were corrected for absorption (Birks, 1963; Springer, 1967) and atomic number (Springer, 1966). A model analysis (Table 2), based on analytical data and comparison of measured and calculated densities, was used for calculation of the corrections. The CK$_\alpha$ absorption coefficients for carbon (2,282) and calcium (68,502) had to be extrapolated from literature data (Henke, et al., 1957). To get the same result from both standards, an antagonistic change of absorption correction factors was necessary. Finally, Si = 44.2 and C = 0.8 $\pm$ 0.4 percent were obtained.

Carbon zonality: the mentioned carbon content was found in the central part of the melanophlogite section, parallel to the cube face at a certain distance from the surface. The anisotropic rim, about 0.2 mm thick, gave higher carbon contents. Seven analyses at different places of the margin with the electron beam moving from the cube plane to
the center of the section gave from 0.8 to 1.2 percent of carbon (average 1.0%), relative to the 0.8 percent carbon content of the central part. The carbon content was measured along 1 mm abscissae perpendicular to the zoning, the intensities of CKα radiation integrated in about 30 μm intervals. Rare anomalously high intensities in the core were attributed to section surface contamination. The optical picture of the transparent polished section used for analyses, was very similar to that depicted by Figure 4.

Microchemical analyses of carbon and hydrogen were made by a Perkin-Elmer C–H–N elemental analyser (J. Horáček analyst). The sample (0.006 g) was heated successively at 950°C for 26 min. and at 1,000°C for 23 min. in oxygen atmosphere. The elements were converted into carbon dioxide and water. The greenish yellow mineral powder after the analysis, when heated at 1,000°C for 2 hours in air afterwards, became greyish white, which indicated a nearly complete carbon liberation during the analysis. A mixture of the rim and core of the crystal gave in this way 1.06 percent carbon and 0.66 percent carbon.

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\[ \rho_{\text{meas.}} = 2.052 \text{ g cm}^{-3} \quad \rho_{\text{cal.}} = 2.06e \]
\[ \rho_{\text{meas.}} = 2.001h \quad \rho_{\text{cal.}} = 2.001h \]

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* B. L. Ingram analyst (Skinner and Appleman, 1963). Oxides were recalculated to elements.
  b Calculated from SiO₂ and SO₃.
  c Calculated by Kamb (1965).
  d By electron microprobe analysis.
  e By activation analysis.
  f Average of electron probe and microchemical analyses.
  g By microchemical analysis.
  h Calculated on the basis of the formula in text (4 x) and \( a = 26.82 \text{ Å} \), \( c = 13.37 \text{ Å} (V = 9,617.2 \text{ Å}^3). \)
hydrogen. After a correction for water and carbon dioxide (adsorption, inclusions), C = 1.0 and H = 0.6 percent.

**Crystallochemical Considerations**

A formula for the Chvaletice melanophlogite was calculated on the basis of the model analysis (Table 2), taking 46 silicon atoms (Kamb, 1965) in 1/4 of the tetragonal superstructural unit cell (a = 28.82, c = 13.37 Å):

\[ \text{Si}_{46}	ext{O}_{92} \cdot \text{C}_{2.17}\text{H}_{17.25}\text{O}_{5.43}\text{S}_{0.00} \cong 46 \text{ SiO}_2 \cdot \text{C}_2\text{H}_{17}\text{O}_5 \]

Six larger tetrakaidecahedral and two smaller pentagonal dodecahedral cavities in the 13 Å unit cell of melanophlogite (Kamb, 1965; Appleman, 1965) can accommodate guest molecules without chemical bounds to the silica framework. The chemical composition of the guest molecules has not been known exactly and was discussed by Kamb (1965). In the Chvaletice mineral, the following element combinations can be present in the guest molecules: C + H ± S ± O, H + O, C + O, and S ± O ± H. According to the preceding analytical data, non-aromatic hydrocarbons or their derivatives and water are probable and prevailing guests. Contrary to the Racalmuto melanophlogite (Skinner and Appleman, 1963), the sulfur content of the Chvaletice mineral, in accordance with its lower density and refractive index, is very low and probably unimportant for stabilization of the structure (see Kamb, 1965). The optical and thermal investigations of the Chvaletice melanophlogite especially indicate zoning of the crystals. Somewhat different physical and chemical properties are observed in the rim and the core of the crystals. No differences have been detected by X-rays, but the carbon content was found higher in the rim than in the core. Differences in occupancy of the larger and smaller framework cavities, in number of molecules in one cavity, and in frequency or chemical composition of the carbon compound guest molecules might be responsible for the property changes.

The idea of a lower than cubic symmetry of melanophlogite (Bertrand, 1880; Friedel, 1890; Kamb, 1965) was confirmed on the melanophlogite from Chvaletice. A tetragonal distortion of the cubic 13 Å cell, connected with the origin of the tetragonal superstructure, is most probably due to the guest molecules (Kamb, 1965). The importance of these guests can be seen from heating experiments. By decomposition of the organic molecules the superstructure was destroyed and the 13 Å cubic cell originated.

The Chvaletice melanophlogite originated most probably under low-temperature and low-pressure conditions, as indicated by its paragen-
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esis. It crystallized in the last stage of metamorphic hydrothermal vein formation by Alpine paragenetic processes. The source of carbon was country rock sediments: Graphite schists and fine-grained rhodochrosite with graphite.

Identification of the guest molecules and a detailed knowledge of the crystal structure (Appleman, 1965) are of utmost importance and the goals of future work on melanophlogite, making its laboratory synthesis easier.

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