FLUID HISTORY FROM CRYSTAL CAVITIES IN RAPAKIVI, PYTERLAHTI, SOUTHEASTERN FINLAND

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KINNUNEN, KARI; LINDQVIST, KRISTIAN and LAHTINEN, RAIMO, 1987: Fluid history from crystal cavities in rapakivi, Pyterlahti, southeastern Finland. Bull. Geol. Soc. Finland 59, Part 1, 35-44.

The crystal cavities lined with quartz, feldspars, biotite, fluorite and a number of accessory phases occurring in the Haikonvuori pyterlite quarry in the Wiborg rapakivi massif were studied with mineralogical, petrographical and chemical methods. Primary fluid inclusions of quartz showed filling temperatures of 292°C to 342°C. The cavities as structures are considered to be miarolitic vugs of chamber pegmatites, but their filling is interpreted as largely secondary and deposited during at least three of the fluid stages present. The nature of the filling material was not reflected in the chemical composition of the pyterlites, but the presence of the cavities, irrespective of their filling, could be inferred from the La and Th depletion and from various petrographic characteristics indicating intensified fluid action. Thus, crystal cavities seem to provide ideal fluid traps for studies of the local fluid history of the crust.

Key words: quartz, fluid inclusions, topaz, molybdenite, miarolitic, rapakivi, chemical composition, Finland, Virolahti, Pyterlahti, Haikonvuori.

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Introduction

Rock cavities originally filled with fluid range in size from submicroscopic vacules and microscopic fluid inclusions via intergranular voids to giant fluid-filled spaces: vugs, pockets, cavities and eventually caves (see Richter and Simmons 1977; Fyfe et al. 1978; Roedder 1984). The larger cavities can be genetically considered as very large fluid inclusions, and in their pristine stage they may contain daughter minerals crystallized out from the fluid. However, because of their polycrystalline walls, the cavities have almost always leaked and are thus contaminated with later fluids present in the crust during uplift. Part of this long fluid history from cavity wall crystallization to supergene solutions present in the final weathering of the outcrops is reflected in the present filling: crystals of different generations are intermingled with various "clays".

In rapakivi granites these larger voids are present as small miarolitic vugs, which can be explained partly as enlarged intergranular voids into which the crystal faces of the rock-forming minerals protrude, and partly as larger crystal cavities characteristic of chamber pegmatites. In these now empty voids the rock-forming
minerals protrude into the cavity, where they constitute the wall material. Phases deposited from the original fluid form the daughter crystals. Several younger fluid types have wholly or partly dissolved earlier phases (especially the water-soluble ones) and deposited new secondary minerals and colloidal masses. In this respect the common quartz crystal cavities of Finnish rapakivi granites are of particular interest, because they contain evidence that several fluid types flowed into and out of the cavities after their formation together with the host rapakivi.

Only a few studies have been conducted on the quartz crystal cavities common in the Wi-borg rapakivi massif (see the literature in Härme 1980 b, p. 47), even though many of them contain interesting crystal specimens and hydrothermal ore minerals. The present study describes quartz crystal cavities associated with molybdenite and topaz crystals, recently found in the Haikonvuori pyterlite quarry (worked for dimension stones) in Pyterlahti, the municipality of Virolahti (Fig. 1). The crystal specimens were collected in 1981–84 from about ten cavities and crystal-lined fractures exposed during routine quarrying. Rock samples were collected from Haikonvuori and from two other granite quarries, Hevonniemi (with cavities) and Sysimäki (without known cavities), situated within the area of pyterlitic rapakivi in Virolahti (Fig. 1).

Cavities and crystallization

In Haikonvuori quarry the quartz crystal cavities had a narrow pegmatitic lining exhibiting the intergrowth textures (graphic granite) typical of granitic pegmatites. The mineral content

Fig. 1. A, Index map of Finland showing the Finnish part of the Wi-borg rapakivi massif (black) and the Pyterlahti area (arrow). B, simplified geological map of the Pyterlahti area (Härme 1980). Rapakivi quarries: 1. Sysimäki (without known cavities), 2. Hevonniemi (with cavities), 3. Haikonvuori (with abundant cavities). The rapakivi surrounding the pyterlite area (circled dots) is principally wiborgite (dots).

Fig. 2. Typical quartz crystal cavity in Haikonvuori quarry. Diameter ca. 3 dm. Most crystals removed.
depended on the size of the cavity, which varied from a few millimetres to some decimetres (Fig. 2). The smallest cavities lacked the minerals interpreted as contamination in the larger cavities; secondary late hydrothermal phases, supergene deposits from groundwaters and recent detrital sediments were deposited at the time the cavity was opened to a cave. The smallest cavities were usually composed only of the protruding wall minerals (feldspars, quartz, biotite and often fluorite), and can be interpreted as pure miarolitic cavities.

The analytical data (Tables 1, 2 and 3) show no significant chemical or mineralogical differences between the samples taken from different quarries (see Fig. 1). However, in the trace element composition there is a systematic decrease in the contents of La and Th from pyterlite sample 1 (without known cavities) through sample 2 (with cavities) to sample 3 (with many cavities). Light rare-earth elements (LREE) can act as compatible elements in very felsic magmas and diminish rapidly in abundance as differentiation progresses (Miller and Mittlefehldt 1982). The depletion of La could refer to the most differentiated part of the pyterlite cupola while indicating the presence of cavities. A few accessory minerals (allanite, monazite) have proved to be important reservoirs for LREE in felsic magmas (Mittlefehldt and Miller 1983;

| SiO₂ | Al₂O₃ | Fe₂O₃ (tot) | MgO | CaO | Na₂O | K₂O | MnO | TiO₂ | P₂O₅ | Sum |
|------|--------|-------------|-----|-----|------|-----|-----|------|------|------|
| 74.86 | 12.39 | 2.80 | 0.48 | 0.90 | 3.65 | 5.55 | 0.03 | 0.26 | 0.05 | 100.98 |
| 75.43 | 11.63 | 2.31 | 0.34 | 0.85 | 2.96 | 5.20 | 0.00 | 0.24 | 0.00 | 98.97 |
| 74.39 | 12.19 | 3.00 | 0.42 | 0.89 | 3.22 | 5.64 | 0.00 | 0.29 | 0.03 | 100.08 |
| 73.35 | 13.59 | 2.01 | 0.29 | 0.92 | 3.29 | 5.92 | 0.00 | 0.17 | 0.00 | 99.55 |
| 12.39 | 13.59 | 0.48 | 0.29 | 0.29 | 0.17 | 0.29 | 0.00 | 0.17 | 0.00 | 11.63 |
| 13.59 | 3.29 | 0.29 | 0.29 | 0.29 | 0.17 | 0.29 | 0.00 | 0.17 | 0.00 | 11.63 |
| 74.39 | 3.29 | 0.29 | 0.29 | 0.29 | 0.17 | 0.29 | 0.00 | 0.17 | 0.00 | 11.63 |
| 73.35 | 3.29 | 0.29 | 0.29 | 0.29 | 0.17 | 0.29 | 0.00 | 0.17 | 0.00 | 11.63 |

Table 1. Main component analysis of pyterlites and chamber pegmatite from Pyterlahti. Contents in weight percentages.

| Method | a | b | c | d |

Table 2. Trace element analyses of pyterlites and chamber pegmatite from Pyterlahti. Contents in ppm. | B | Bi | Co | Cu | Ga | Ge | La | Li | Mo | Nb | Ni | Pb | Sc | Sn | Sr | Th | V | W | Yb | Zn |
|-------|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| < 20 | < 20 | < 20 | < 20 | a | < 10 | < 10 | < 10 | < 10 | a | 2.3 | 2.5 | 2.3 | 2.1 | b | 4.4 | 5.1 | 5.4 | 4.4 | b | 24 | 24 | 25 | 28 | a |
| < 10 | < 10 | < 10 | < 10 | a | 180 | 120 | 82 | 49 | b | 50 | 51 | 51 | 62 | b | < 200 | < 200 | < 200 | < 200 | a | 0.0 | 0.8 | 0.3 | 0.0 | b |
| < 10 | < 10 | < 10 | < 10 | a | 2.4 | 2.8 | 2.9 | 0.5 | b | 9 | 8 | 9 | 8 | a | 78 | 83 | 70 | 78 | b | 36 | 31 | 26 | 14 | b |
| < 10 | < 10 | < 10 | < 10 | a | 5.7 | 5.4 | 6.3 | 5.8 | b | 6 | 4 | 7 | n.d. | c | 72 | 62 | 69 | 49 | b | 7 | 6 | 6 | 5 | a |
| < 200 | < 200 | < 200 | < 200 | a | 49 | 81 | 78 | 81 | b | 69 | 79 | 78 | 81 | b |

Samples as in Table 1.

Methods:

a. Optical emission spectrographic analysis
b. ICP-OES analysis after total dissolution (HF+AR+HClO₄)
c. Spectrophotometric determination using zinc dithioletricate

n.d.: not determined
Table 3. Petrographical analyses of the pyterlite samples listed in Tables 1 and 2.

|        | 1     | 2     | 3     |
|--------|-------|-------|-------|
| Quartz | M     | M     | M     |
| K-feldspar | M    | M     | M     |
| Plagioclase | M   | M     | M     |
| Biotite | m     | m     | M     |
| Chlorite | a    | a     | a     |
| Zircon | a     | a     | a     |
| Fluorite | a   | a     | a     |
| Apatite | a     | a     | a     |
| Haematite | a | a     | a     |
| Metamict | a   | a     | a     |
| Sericite | a    | a     | a     |
| Calcite | —    | a     | a     |
| Ilmenite | —   | a     | a     |
| Magnetite | —   | a     | —     |
| Chalcopyrite | — | a   | —     |
| Pyrite | —     | —     | a     |

M. major mineral, > 10 %
m. minor mineral, 5—10 %
a. accessory mineral
—. mineral not observed
1) other than zircon

The presence of cavities in pyterlite was inferred from petrographic features (Fig. 3): quartz-biotite symplectites, microcavities and widespread microclinitization indicating the action of fluids against the host rock. Augustithis (1973, Figs. 221—223) has described similar myrmekitic textures in biotite, which he attributed "to infiltration of quartz-forming solutions into the biotite". According to Vorma (1971), microclinitization in rapakivi granites indicates high fluid pressure favouring the transformation of primary orthoclase to maximum microcline. X-ray diffraction studies of alkali feldspar specimens from Haikonvuori quarry reveal the same trend (Table 4): the obliquity of the alkali feldspars increases towards the cavity.

Table 4. Structural types of the alkali feldspars in Haikonvuori quarry determined by X-ray diffraction methods according to Goldsmith and Laves (1954).

| Specimen | Structural type     | Obliquity |
|----------|---------------------|-----------|
| 1        | Orthoclase only     |           |
| 2        | Orthoclase only     |           |
| 3        | Maximum microcline  | 0.83      |
| 4        | Maximum microcline  | 0.84      |
| 5        | Maximum microcline  | 0.95      |

Specimens:
1. Alkali feldspar from an ovoid centre in pyterlite.
2. Alkali feldspar from pegmatite.
3. Alkali feldspar occurring together with molybdenite in cavities.
4. Alkali feldspar from pegmatitic rind surrounding the cavities.
5. Centre of a giant alkali feldspar crystal from a cavity.

Gromet and Silver (1983). Petrographic observations show that the metamict accessory crystals occur mainly as inclusions in the alkali feldspar ovoids in pyterlite sample 1 (without known cavities) but are enriched in the later biotite aggregates in pyterlite samples 2 and 3 (with cavities). The few samples leave this question to be answered in future work.

Fig. 3. Photomicrograph of the symplectitic texture bordering the biotite grain in pyterlite host rock in Haikonvuori. Clear grains are quartz and the pigmented ones alkali feldspar (mostly microclinitized). Note the numerous microcracks depicting the action of hydrothermal fluid(s). Transmitted light with one polarizer.
centre, indicating intensified action of the hydrothermal fluid medium in the crystallization process.

At Haikonvuori, the cavity crystallization was found to have begun with the formation of microcline and quartz as large crystals measuring up to several decimetres, together with albite laths, biotite, violet fluorite, platy haematite, black tourmaline, green beryl and topaz. The following minerals were deposited in the cavities and fractures during at least two hydrothermal stages: calcite (usually colourless) in three generations; pyrite as pyritohedron and as combinations of cube and pyritohedrons (with oscillatory growth); platy molybdenite; anhedral galena; vermicular chlorite; spherulitic marcasite and colloformic goethite.

Cavity minerals

Quartz

Quartz was the principal crystalline material of the cavities at Haikonvuori. The length of the crystals varied from a few millimetres to some decimetres. The habitus was long-prismatic and the crystals were ideal, the rhombohedral crystal planes being of almost equal size in each crystal. Most crystals lacked pyramidal faces, and the line connecting the nucleation centre to the present centre of gravity was in most crystals parallel to the c-axis. Individual quartz crystals were twinned according to the Dauphine law. This was seen from the natural etch markings on the prism and rhombohedral faces. Surface markings did not reveal the former presence of any late water-soluble, rare-mineral daughter crystals. Contact twins between quartz crystals were too rare to be determined for their twinning laws.

The quartz crystals varied from colourless to smoky. The darkest variety (morion) was found to contain 200–500 ppm Al (determined with optical emission spectrographic analysis). Doubly-polished plates revealed growth zoning in the quartz crystals, with smoky zones alternating with clear zones. In the outer parts of the crystals tourmaline needles occurred as primary mineral inclusions. The prism faces of the quartz crystals were nearly always etched to a matt gloss, but the rhombohedral faces were in places overlain by a later unetched quartz layer. Haematite globules were present on the contact plane of these two quartz types.

At room temperature, the primary and pseudosecondary fluid inclusions in the quartz crystals were composed of liquid, vapour and of four solid phases, three of which were anisotropic and transparent. One of these transparent phases shows its lowest refractivity index in the range of the inclusion brine, thus proving the presence of an unknown but water-soluble rare-mineral daughter crystal (cf. the lists in Phillips and Griffen 1981). The fourth solid phase was opaque, non-magnetic and under dark-field incident light, revealed red inner reflections indicative of haematite. Recrystallized (with necking down) pseudosecondary inclusion cavities occasionally contained halite cubes (isotropic and index of refraction within the range of those of quartz). The secondary fluid inclusions of hydrothermal type were composed of liquid, vapour and of two anisotropic and transparent crystal aggregates. Monophase secondary fluid inclusions were composed of liquid only, and they were observed together with chlorite and goethite spherules in the microfractures cutting all the earlier textures. The monophase secondary inclusions are probably supergene in origin.

During heating experiments (Leitz 350 and Chaixmeca stages) the primary inclusions were filled with liquid at 292°–342°C. Large inclusion cavities decrepitated at 320°–350°C. The secondary inclusions were filled with liquid at 134°–167°C.

Most of the primary and pseudosecondary inclusion cavities were in the shape of negative
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Fig. 4. Photomicrograph of a negative crystal in smoky quartz from Haikonvuori quarry. The primary fluid inclusion filling the cavity is composed of liquid, a vapour bubble and four solid phases (out of focus). Note the untwinned prism faces without striations and the curved rhombohedral ends in this negative crystal resembling in the main form (c/a ratio), its host quartz crystal. Transmitted plane-polarized light.

crystals consisting of short planar prism faces and weakly curved rhombohedral faces. The prism faces of the negative crystals did not have the twinning striations typical of quartz (Fig. 4).

The filling temperature range, 292°—342°C, of the primary fluid inclusions in the quartz from the Haikonvuori cavities is considered the lowest possible temperature of crystallization. The fact that the inclusion assemblages bear no visible indications of boiling can be interpreted as a sign that the fluid pressure exceeded 0.1 kb. At about this pressure the fluid inclusions with filling temperatures of 292°C to 342°C would have boiled according to the thermodynamic relations calculated by Haas (1971). Vorma (1971) has estimated that the lithostatic pressure during the formations of the Wiborg rapakivi massif was about 1 kb. If this figure is taken as the maximum fluid pressure during the pegmatitic stage, the estimated range (0.1—1 kb) raises the pressure-corrected temperature of crystallization to 312°—464°C with the nomograms by Potter (1977). The very existence of cavities makes 2 kb a likely maximum pressure in theory, because, as shown in the differential strain analysis by Feves et al. (1977), most open fractures would close mechanically at lighter lithostatic load.

The Al content of Haikonvuori quartz, 200—500 ppm, may offer a pressure-independent geothermometer. When the Al content is plotted on the diagrams of Dennen and Blackburn (1970), the minimum crystallization temperature obtained for Haikonvuori quartz rises to as much as 740°C. This is clearly excessive, as can be inferred from the fluid inclusion data and from the lack of high-quartz habitus in the earliest phantom generations. Very dark quartz (morion, see Webster 1983) tends to contain a few thousand ppm Al (Frondel 1962). According to laboratory synthesis, the smoky colour of quartz is caused by the colour centres formed by natural irradiation with gamma-rays on Al³⁺ ions substituting for Si⁴⁺ ions in the quartz lattice (Nassau 1983, pp. 190—192).

Topaz

An unusual feature of the Haikonvuori cavities was the occurrence of topaz as transparent colourless or brownish crystals (length generally 1—2 cm) and as translucent greenish fragments (length of largest 2 dm). It has been shown in experiments that, as with smoky quartz, natural irradiation with gamma-rays also causes the brownish color centres on colourless topaz (Nassau 1983, p. 200; Aines and Rossman 1986). The mineral inclusions consisted of tourmaline needles in the outer crystal zones and of spherulitic chlorite in the fracture fillings and on some crystal faces. No definitive primary fluid inclusions were found, although secondary fluid inclusions were common. Typical inclusions are needle-like tubes extending
from the prism faces to the inner crystal zones (Fig. 5). These inclusions may be interpreted as dislocations that were later hydrothermally etched to become optically visible tubes (cf. Fig. 9 in Carstens 1968; Fig 4 in Eadington and Wilkins 1980; Fig. 2–23 in Roedder 1984). Most of the transparent crystals are cleaved along the basal plane and generally etched. Typical specimens therefore revealed only striated prism faces and deeply etched fracture and cleavage surfaces.

In addition to the above topaz types, large (up to 5 cm) brownish and water-clear crystal fragments are said to have been found in the cavities of Haikonvuori. Unfortunately, we did not find any gem-quality material there and so are unable to confirm this information. However, we were able to borrow some of these crystals (rough and cut) for study. They contain goethite in the microfractures in a very specific spherule form. Interestingly, similar inclusions are described as typical of the brownish yellow topaz of Ouro Preto, Brazil, in the authoritative atlas on inclusions in gemstones from different localities compiled by Gübelin and Koivula (1986, pp. 202 and 389).

The crystal surfaces of the Haikonvuori topaz were covered with etch pits resulting from the corrosive action of a hydrothermal fluid (Fig. 6). On (120) prism faces the morphology of the etch pits corresponded to the triangular and trapezium-shaped types described by Joshi and Taku (1972); the large, semi-circular pits resulting from prolonged hydrothermal corrosion were not found. The matt gloss of the prism faces of the Haikonvuori topaz is due to the numerous etch pits.

**Molybdenite**

Molybdenite occurs as euhedral, hexagonal platelets about 0.2 mm thick and up to about 4 cm long. Molybdenite was found in the cavities only as secondary, late-hydrothermal filling material, usually together with calcite in the fractures associated with the cavities. The simultaneous growth of molybdenite and calcite
is demonstrated by the crystal growth figures seen on the pinacoid faces of the molybdenite platelets (Fig. 7). The most common texture is composed of pits resulting from the negative calcite crystal imprints, some of which are trigonal in shape. Corresponding positive forms were observed on calcite crystal plates. Natural etch pits were observed on pinacoid faces as triangular pits but they were only a few micrometres in diameter, testifying to the late hydrothermal genesis of molybdenite. Trace element analysis of molybdenite revealed the presence of Ag, Pb, Cu, Bi, Ni and Zn, each in the range of 10—100 ppm, and also of W and Re, but both below the detection limit, 200 ppm (AAS and OES methods, respectively).

Discussion

The quartz crystal cavities in the Wiborg rapakivi massif are usually interpreted as mioralitic (Wahl 1925, p. 60). There is a problem, however, in that not enough silica dissolves in the fluid to deposit the observed quantities of quartz in closed cavities, because in hydrothermal systems quartz is usually deposited as a result of a drop in temperature. A drop of 800° to 400°C is not sufficient for depositing more than 0.3 wt. % SiO₂. This estimate is based on the experimental data of Kennedy (1950). F and B in the system may increase the solubility of SiO₂, but only to about 1 wt. % (Pichavant 1979; Manning 1983). In the same way F and B increase the exsolution of H₂O from the pegmatitic melt (London 1986). It seems that the volume of fluid needed to produce the observed quantities of quartz and other silicate crystals should far exceed volume of the original cavity. In mioralitic crystallization the fluid inclusion assemblages of quartz should indicate the coexistence of fluid and a magma phase (devitrified glass inclusions together with ordinary fluid inclusions) and a high trapping temperature, 750°—800°C (Olsen and Griffen 1984). These features were not observed in the quartz of the Haikonvuori cavities or in their pegmatitic rim material. Moreover, the colour zoning of the Haikonvuori quartz crystals shows that the fluid environment was subject to pulses (PTX). Vorma and Paasivirta (1979) have interpreted the multizoned zircons of the Laitila rapakivi massif in western Finland as indicating changes in the H₂O pressure. Situated in the biotite aggregates altered by hydrothermal processes, multizoned zircons were also noted in the pyroelectric host rocks from Hevonniemi and Haikonvuori quarries (both with cavities).

The following stages are suggested for the cavity crystallization in Haikonvuori. First, the cavity spaces with their pegmatitic rim zone formed in a closed system under the influence of the autoclave effect (cf. Kosukhin 1978). As the temperature fell contraction fractures were formed. The cavities and fractures were then partly filled with material deposited from at least two hydrothermal and one supergene fluid types. The Haikonvuori cavities are thus considered primary but their filling is largely
secondary. Haapala (1977) drew the same conclusion from the microscopic, chlorite-filled miarolitic cavities of the Väkkärä rapakivi granite in southwestern Finland. Indications of pocket clays of supergene origin has recently been obtained from Californian pegmatites by Foord et al. (1986).

Quartz crystals are interpreted as having formed during the first hydrothermal stage. The temperature varied from 312° to 464°C according to fluid inclusion data, and the fluid pressure was most probably in the range 0.1—1 kb. Molybdenite and calcite were deposited from later hydrothermal fluids. During this stage some of the earlier crystals were fractured and etched. Fluid inclusions set the minimum temperature of this hydrothermal fluid at 137°—167°C. The latest supergene (?) fluid type is only found in monophasic inclusions and therefore is not suited to inclusion thermometry. Nevertheless, it could be shown that part of the chloritic material was deposited contemporaneously.

Chamber pegmatites, comparable to the Haikonvuori cavities in their mode of occurrence, mineral content and fluid inclusion types, have been described from the Ukraine by Zakharchenko (1971). These chamber pegmatites, located on the southwestern border of the Korostensky rapakivi massif, are worked for gem-quality topaz and beryl, fluorite and optical-grade quartz. The isotope ages of the Korostensky and Wiborg rapakivi masses are similar, being 1750 Ma (Semenenko et al. 1968) and 1700—1640 Ma (Vaasjoki 1977) respectively. According to Černý (1982), the chamber pegmatites in shallow granite masses are characteristic of their contact areas.

Acknowledgements. In 1981 Mr Seppo Leinikka, a teacher from Virojoki, brought some crystals from the cavities in Haikonvuori quarry to the Geological Survey of Finland for identification and provided the impetus for our investigation. The favourable attitude of Mr Paavo Vaarais of the Baltic Granite Co. towards our work made the collecting of field data and crystal specimens possible. The analytical skills of Mr Väinö Hoffren, Mr Gustaf Wansen and Mrs Mervi Wiik all of the Geological Survey of Finland, were indispensable and are gratefully acknowledged. We also thank Dr Matti Vaasjoki, likewise of the Geological Survey of Finland, for critically reading the manuscript and for suggesting many improvements.

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Manuscript received April 11, 1985