New data on mineralogy of the Výskok Kámen deposits near Krásno, Slavkovský les area, Czech Republic

Nové poznatky o minerállogii ložisíc Výskok Kámen u Krásna (Slavkovský les, Česká republika)

(32 figs, 11 tabs)

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Detailed mineralogical study focused on two deposits located on slopes of the Výskok Kámen hill near Krásno in the proximity of the town of Horní Slavkov in Czech Republic. The open pit exploiting industrial feldspar material (mainly feldspathites and aplitic granites) offered new finds of minerals in the Slavkovský les. The following minerals were identified: autunite, bertrandite, beryl, goethite, hematite, koechlinlite, kolbeckite, meta-autunite, powellite and scheelite. In the nearby abandoned greisen deposit a phosphate mineral association was observed consisting of cacoxenite, leucophosphate, vivianite, wavelite and minerals of the chalcosiderite-turquoise and strengite-variscite series. In addition to them dickite and goethite were identified. The genesis of these mineral associations is discussed.

Keywords: greisen; supergene minerals; phosphates; molybdates; wolframates; Výskok Kámen near Krásno; Horní Slavkov; western Bohemia; Czech Republic

Introduction

The two studied deposits are both localized on the slopes of the elevation 819.4 m, west and northwest of Krásno, about 7 km south-southwest of Horní Slavkov in western Czech Republic. The first one is an abandoned Sn-W greisen deposit and the second one is feldspathite (feldspathite = leucocratic alkali feldspar syenite) body exploited presently as ceramic feldspar deposit. The greisen deposit was uncovered by an open pit (Fig. 1), directly at the NW edge of the Krásno village. During a recultivation the quarry was filled with various rock material. Thus the supergene* mineralization can be now observed only on a limited scale at the outcrop of a silicified fault zone exposed above the former quarry (Fig. 2). The feldspar deposit (called as Výskok Kámen – feldspar, in the following text) is exploited by a quarry (Fig. 3) with several levels (Fig. 4), located on the northern slope of the elevation 819.4 m, about 0.5 km west to northwest of the Krásno village.

Geological situation of the Výskok Kámen deposits

The greisen stock at Výskok Kámen is positioned southwest of the Schnöd stock, in extension of a line connecting the Schnöd and Huber granite stocks. It is located in an intrusion of a younger suite granite, i.e., classified with the Younger Intrusive Complex granite (YIC granite). The envelope to this granite body consists of migmatised biotite paragneiss, grading to medium-grained, advanced migmaticites. A marginal pegmatite (stockscheider) is often present at the granite contact. The stock has an oval shape in plan, but the southwestern flank is cut off by a fault zone, uplifted and strongly eroded (Fig. 5). A breccia-textured quartz gangue, stained by iron oxides, fills the fault zone (Fig. 6). Thus, only a low core part of the stock, composed of aplitic granite with greisen relics, is preserved (Beran 1999).

Coarse-grained topaz-mica greisen builds the remaining part of the stock and downwards it passes to greisenized granite. The greisen body is composed of several mineral zones with a total thickness of 40 to 60 m. In the central part, the greisen reaches upwards up to the

* note by referee: the term supergene is used in a relatively loose sense, in agreement with common practice in mineralogical literature. Some publications on ore deposits use the term specifically for a certain part of the oxide zone (zone of supergene enrichment).
granite-gneiss contact. The outcrop of the stock at the present surface has approximately 200 by 300 m size, disregarding the uplifted southwestern part. At the margins the greisen body is split to several smaller parts which gradually wedge-out and pass to greisenized granite. The greisenized granite directly below greisen is strongly hydrothermally altered, mainly by kaolinization. Toward the deeper levels, the alteration dies out and the rock passes into a fresh Li-mica-topaz granite. Still to deeper levels, about 150 m below surface, the stock is composed of a leucocratic topaz granite, which gradually evolved from ordinary granite via the disappearance of Li-mica. The Sn-W mineralization at Vysoký Kamen is formed by cassiterite and subordinate wolframite impregnation in greisen and greisenized granite. Miner-

Fig. 2 Relics of open pit at greisen deposit Vysoký Kámen near Krásno after recultivation (photo J. Sejkora 2002)

Fig. 3 Quarry at the Vysoký Kámen deposit near Krásno (feldspar) (photo J. Berger 2001)

Fig. 4 Quarry at the Vysoký Kámen deposit near Krásno (feldspar) (photo J. Sejkora 2002)
mica content and rare pegmatites are exploited as a feldspar raw-material with a low content of iron oxides.

The majority of studied samples was collected at the lower level of the Vysoký Kámen feldspar quarry, near a branch of the fault zone filled in by jasper, quartz and clay minerals. A lenticular body of "vein-type greisen", up to 3 m wide, trending NE-SW and dipping 75 to 80° SE was also observed there. This body has a core of re-crystallized quartz with wolframite, molybdenite, bismuth etc., and an outer zone of muscovite, quartz, apatite, beryl and clay minerals formed by greisenization of coarse-grained feldspar. The body has an outer kaolinized rim. It is probable that the rock is rather a greisenized pegmatite than a vein-type greisen.

Methods of mineral identification

Surface morphology of the samples was studied using optical microscope Nikon SMZ 1500 in combination with digital camera Nikon DXM1200F for microphotography in incandescent light. Details of surface morphology were studied in secondary electron images with the electron scanning microscopes JEOL JSM T-20 (Z. Mach; Research Institute of Ceramics, Karlovy Vary – Březová) and Tesla BS 340 (A. Gabašová and J. Sejkora, Czech Geological Survey, Prague).

The X-ray powder diffraction analysis was used for identification of minerals. To minimize complicated shape of background due to classic glass sample holder, the samples studied were placed on the surface of flat silicon wafer from alcoholic suspension. Step-scanned powder diffraction data were collected using Philips X’Pert MPD diffractometer with a metallo-ceramic copper tube which was operated at high-voltage of 40 kV and tube current of 40 mA. A graphite secondary monochromator has been used to produce CuKα,α radiation. In special cases a film methods – Gandelofi camera 114.6 mm (CoKα radiation) were used. The results were processed by using X-ray analysis software ZDS for DOS (Ondruš 1993), Bede ZDS Search/Match ver. 4.5 (Ondruš – Škála 1997); unit-cell parameters were refined by program FullProf (Rodriguez – Carvajal 2005).

We used energy dispersion analyser Tracor-Northern for qualitative chemical analysis (R. Rybká, Czech Geological Survey, Prague). Additional quantitative chemical analyses were made with the electron microprobe Cameca SX 100 (J. Sejkora and R. Škoda, Laboratory of electron microscopy and microanalysis of Masaryk University and Czech Geological Survey, Brno). Studied samples were mounted in epoxy resin discs and polished. The polished surfaces were coated with carbon layer 250 Å thick. Wavelength dispersion mode and operating voltage 15 kV were used in all analyses. Used beam current and diameter reflect stability of analyzed phases under electron beam. Stable phases were analyzed under 20 nA current and 2 µm beam size. Less stable and highly hydrated minerals were analyzed at conditions of 10–4 nA and 10–30 µm beam diameter. For smaller aggre-
gates (< 10 μm) of unstable minerals the beam diameter was as large as possible and applied beam current was only 1–2 nA. The sequence of elements determination during analyzing was operatively changed according to chemical composition. Volatile and major elements were analyzed prior to stable, minor, and trace elements. Higher analytical totals of minerals containing higher amount of hydroxyl or crystalline water were generally caused by escape of water under high vacuum, well documented by a collapse of crystals or by electron beam heating during analysis, causing escape of water, documented as intensively bright spots after analysis. Lower analytical totals of some samples were primarily caused by the porous nature or by poorly polished surface of soft or cryptocrystalline minerals.

To minimize peaks overlapping, the following analytical and lines were selected: Kα lines: Al (TAP, sanidine), As (TAP, InAs), Cu (TAP, diopas), P (PET, fluorapatite), Ca (PET, andradite), S (PET, barite), Fe (LIF, andradite), Zn (LIF, ZnO); L8 line: Sr (PET, SrSO4); L8 line: Ba (PET, benitoite); M8 line: Bi (PET, metallic Bi). Peak counting times (CT) were 10 to 20 s for main elements and 30 to 60 s for minor and trace elements. CT for each background was ½ of peak time. When only one background and appropriate slope had to be used, the background CT corresponded to peak CT. Applicable minor to trace elements were preferentially analyzed using high sensitivity large LPET and LLIF crystals. Raw intensities were converted to the concentrations using automatic PAP (Pouchou – Pichoir 1985) matrix correction software package.

Infrared spectra were recorded with PYE-UNICAM-PHILIPS SP3-300 and Perkin Elmer 521. Spectra were measured from KBr tablets in the range of 4000–200 cm⁻¹, time 7 minutes (O. Žilková, ÚVZÚ, Plzeň).

Minerals identified at the Vysoký Kámen (feldspar) deposit

Autunite Ca(UO2)2(PO4)2·10–12 H2O and meta-autunite Ca(UO2)2(PO4)2·6 H2O

Paulič (1990) reported autunite as light yellow powdery aggregates passing to tetragonal crystals with pearly lustre, up to 3 cm in size. Autunite was observed in a steeply dipping fracture within the greisen, locally healed by finely crystalline quartz associated with goethite.

Meta-autunite occurs through the whole profile and at all levels of the quarry. The most common form is that of fine and inconspicuous impregnations in fractures of leucocratic granite and of greisen. Yellowish mixture with clay minerals in cracks and fissures represents another frequent form of meta-autunite occurrence. Imperfect tetragonal tabular crystals or crystalline crusts of light yellow or yellow colours are relatively rare. In a short-wave and long-wave UV light it shows very strong yellow green fluorescence. The unit-cell parameters (Table 1) correspond to the published data.

| Table 1 Unit cell parameters of meta-autunite (tetragonal P4/nmm) |
|---------------------------------------|--------------------------|
| this paper | Makarov – Ivanov (1960) |
| a [Å] | 6.974(1) |
| c [Å] | 8.451(2) |
| V[Å³] | 411.0 |
| | 408.9 |

Bertrandite BeSi₂O₅(OH)₂

Bertrandite forms inconspicuous, thick tabular crystals up to 3 mm in size. They occur in greisen cavities on the crystals of quartz, rarely in the cavities after leached beryl in association with clay minerals, apatite and exceptionally with wolframite. Although the amount of leached beryl is notable at this locality, bertrandite is a rare mineral. It was also observed as aggregates of transparent acicular crystals up to 2 mm in size deposited on imperfect crystals of pink-beige apatite. Bertrandite has been observed close to corroded beryl only rarely, possibly due to relatively high mobility of Be ions. The unit-cell parameters (Table 2) correspond to published data.

| Table 2 Unit cell parameters of bertrandite (orthorhombic Cmc₂) |
|---------------------------------------|--------------------------|
| this paper | Giuseppetti et al. (1992) |
| a [Å] | 8.720(1) |
| b [Å] | 15.2785(9) |
| c [Å] | 4.5076(1) |
| V[Å³] | 608.93 |
| | 607.0 |

Beryl Be₃Al₂Si₂O₈

Beryl is a fairly common mineral in the greisen, as granular aggregates or perfectly formed crystals in vugs of greisen and quartz. The crystals are grey-brown, grey-blue, light blue, white, or rare colourless translucent up to 12 cm long and 2.5 cm wide. Some aggregates form layers up to several dm long, with the grains which were hydrothermally strongly corroded (Beran 1999). Granular beryl aggregates of yellowish or greenish colour are also very frequent enclosing in places imperfect bluish and grey green prismatic crystals or radiating crystal groups up to 15 cm long. The occurrences of beryl were confirmed by X-ray powder diffraction data.

Goethite FeO(OH)

It occurs as spheroidal radiating aggregates and small red-brown crystals in association with koehlinite-russellite, autunite, meta-autunite and powellite in the cavities of the quartz gangue. It is the youngest mineral in the mineral associations described. Goethite was identified by X-ray powder diffraction data.

Hematite Fe₂O₃

Hematite occurs in radiating spheroidal aggregates composed of microscopic red brown acicular crystals. The
aggregates are on average 5 mm in diameter and very similar to goethite aggregates. This type of hematite was deposited in some cavities in quartz and in fractures in the greisenized granite. Hematite along with goethite belong genetically to the youngest minerals at the deposit. Hematite was identified X-ray powder diffraction data.

**Koechlinite (BiO)₂MoO₄**

Koechlinite is a common mineral in the “greisen” body forming usually powdery coatings and infilling of cavities after leached molybdenite. Imperfectly foliated, ductile crystals up to 0.1 mm are rare. The locality provided relatively rich samples with yellow koechlinite coatings several cm² in size. Koechlinite was formed by decomposition of powellite and Bi was provided by native bismuth, observed as relics near some koechlinite occurrences.

Unit-cell parameters of koechlinite (Table 3) are in a good agreement with published data. Quantitative chemical analyses of koechlinite (Table 4) show, in addition to minor As and P contents, also minor W. This indicates that the mineral has the composition close to the end-member of the isomorphic series koechlinite (Mo) – russellite (W). Its chemical composition corresponds to the following empirical formula, based on six oxygen atoms: (BiO)₂.₀₆(Mo₀.₈₁W₀.₁₉P₀.₀₆As₀.₀₁O₄). Beran (1999) reported an occurrence of intermediate members of the russellite–koechlinite series from the same greisen body. However, no analytical data support this information. The infrared spectrum of koechlinite from Vysoký Kámen is given in Table 5 and in Fig. 7. Bands at 320 and 720 cm⁻¹ correspond to deformation vibration of the BiO⁺ ion. Bands at 320, 440, 795, 835, 1000 and 1070 cm⁻¹ correspond to vibration of the (MoO₄)²⁻ anion.

**Kolbeckite ScPO₄·2H₂O**

Kolbeckite was found in several samples only as blue or grey blue crystals (Fig. 8) rarely up to 2 mm long, of...
shortly prismatic morphology (Fig. 9). It occurs in quartz cavities in association with hematite, apatite and in the proximity of corroded beryl crystals. In a single case kolkbeckite was found as an overgrowth on scheelite in a quartz cavity. Scandium for crystallization of kolkbeckite was probably mobilized from wolframite which may contain up to 0.1 wt. % Sc.

X-ray diffraction pattern of a single crystal of kolkbeckite (Table 6) corresponds to the published data for this mineral from Fairfield, USA (Mrose - Wappner 1959). Some differences in the intensities of diffraction maxima were probably caused by small quantity of material available for study. Identification is confirmed by microprobe analysis showing the presence of Sc and P as the major elements.

<table>
<thead>
<tr>
<th>Vysoký Kámen</th>
<th>Fairfield, Utah, USA*</th>
</tr>
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<tbody>
<tr>
<td>L_a</td>
<td>d_a</td>
</tr>
<tr>
<td>30</td>
<td>6.94</td>
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<td>30</td>
<td>5.20</td>
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<td>2.08</td>
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<tr>
<td>10</td>
<td>1.877</td>
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<tr>
<td>10</td>
<td>1.822</td>
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</tbody>
</table>

Gandolfi camera 114.6 mm, CoKα radiation.
* kolkbeckite from Fairfield, Utah, USA (Mrose - Wappner 1959).

Meta-autunite Ca(UO$_2$)$_3$(PO$_4$)$_2$. 6 H$_2$O

See autunite.

Powellite CaMoO$_4$

Two generations of powellite occur at the studied locality. The older generation forms grey brown to dark brown flaky aggregates up to several cm long which are the pseudomorphs after molybdnite. This generation of powellite fills small cavities in quartz after leached molybdnite. Large flakes of the first generation of powellite are cemented by finely crystalline light brown powellite of a younger generation. The younger generation of powellite forms druses of imperfect bipyramidal crystals (Fig. 10) of brown to grey brown colour with glassy to greasy luster (Fig. 11). The crystals which deposited on quartz are exceptionally up to 3 mm in size. Druses of imperfect tabular crystals of powellite 1–2 mm in size are rare. The first generation of powellite was partly altered to a mixture of koechlinite-russellite, so that these minerals are regularly observed in a joint occurrence. Unlike koechlinite-russellite, powellite shows light yellow fluorescence in short-wave UV light; however, powellite of the second generation exhibits a weak fluorescence.

Unit-cell parameters were calculated for the younger generation of powellite (Table 7) and show a good correspondence with the published data. The quantitative chemical analyses (Table 8) show minor abundances of Mg, W, P and As. The empirical formula based on four oxygen atoms is Ca$_{1.02}$(MoO$_4$)$_{0.96}$(PO$_4$)$_{0.04}$. The infrared spectrum of powellite is given in Table 9 and Fig. 12.

<table>
<thead>
<tr>
<th>Table 7 Unit cell parameters of powellite (tetragonal I4$_1$/a)</th>
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<tr>
<td>this paper</td>
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<tr>
<td>a [Å]</td>
</tr>
<tr>
<td>c [Å]</td>
</tr>
<tr>
<td>V [Å$^3$]</td>
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Table 8 Chemical composition of powellite from Vysoký Kámen (in wt. %)

<table>
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<tr>
<th></th>
<th>Vysoký Kámen</th>
<th>theoretical composition</th>
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<tr>
<td>CaO</td>
<td>28.65</td>
<td>28.04</td>
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<tr>
<td>MgO</td>
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<tr>
<td>P₂O₅</td>
<td>1.41</td>
<td></td>
</tr>
<tr>
<td>As₂O₅</td>
<td>0.27</td>
<td></td>
</tr>
<tr>
<td>MoO₃</td>
<td>68.74</td>
<td>71.96</td>
</tr>
<tr>
<td>WO₃</td>
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<td></td>
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<tr>
<td>total</td>
<td>99.18</td>
<td>100.00</td>
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Table 9 Infrared absorption spectrum of powellite from Vysoký Kámen

<table>
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<th>[cm⁻¹]</th>
<th>*</th>
<th>[cm⁻¹]</th>
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<tr>
<td>3400</td>
<td>vw, b</td>
<td>805</td>
<td>vs, b</td>
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<td>1640</td>
<td>vw, b</td>
<td>695</td>
<td>w, sh</td>
</tr>
<tr>
<td>1095</td>
<td>m, sh</td>
<td>535</td>
<td>m</td>
</tr>
<tr>
<td>1030</td>
<td>s</td>
<td>470</td>
<td>m</td>
</tr>
<tr>
<td>1010</td>
<td>s</td>
<td>430</td>
<td>m</td>
</tr>
<tr>
<td>905</td>
<td>m, sh</td>
<td>320</td>
<td>s</td>
</tr>
<tr>
<td>860</td>
<td>s, sh</td>
<td></td>
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</table>

* Character of absorption maxima:
  sh – shoulder; b – broad. Intensity scale:
  vs – very strong; s – strong; m – medium;
  mw – medium weak; w – weak; vw – very weak.

The measured absorption bands at 320, 430 and 470 cm⁻¹ correspond to ν₁ and ν₂ deformation vibrations of (MoO₃)²⁻, a wide high-intensity band at 805 cm⁻¹ corresponds to ν₁ valence vibrations of (MoO₃)²⁻ and another band at 860 cm⁻¹ corresponds to ν₁ vibrations of the MoO₃²⁻ anion. Very weak bands at 1640 and 3400 cm⁻¹ are caused by adsorbed water.

**Scheelite CaWO₄**

It is a fairly common accessory in disseminated grains to 1 mm, associated with beryl, wolframite, koechlinite, apatite and other minerals. Exceptionally, scheelite occurred as well-formed grey crystal 1 cm in size, deposited in a quartz cavity and intergrown with minute kolbeckite crystals (Beran 1999).

**Supergene minerals identified at the Vysoký Kámen (greisen) locality**

**Cacoxenite (Fe₂,Al₂₅)(PO₄)₁₇O₆(OH)₁₂ · 75 H₂O**

Cacoxenite belongs to relatively common minerals at the locality. It forms coatings and aggregates in fractures of a quartz breccia in the Vysoký Kámen fault zone and in the drill cores from a depth up to 40 m below the present level.

It occurs as radiating aggregates up to 2 mm in diameter, composed of very fine crystals of golden yellow colour (Fig. 13) deposited in fractures in greisen and quartz.

![Fig. 12 The infrared spectra of powellite from Vysoký Kámen – feldspar, Krásno.](image)

![Fig. 13 Yellow crystalline crusts of cacoxenite, Krásno (greisen) near Horní Slavkov, width of figure 12 mm; photo J. Sejkora, Nikon SMZ1500.](image)
The aggregates are often corroded and at margins they show alteration to oxy-hydroxides of iron (*limonite*). The mineral was identified by X-ray powder diffraction data.

**Minerals of the chalcosiderite CuFe$_2$[PO$_4$(OH)$_2$]$_4$·4H$_2$O – turquoise series CuAl$_4$[PO$_4$(OH)$_2$]$_4$·4H$_2$O**

Minerals of this series are common at the locality, particularly as groups of spheroidal aggregates up to 1 mm in diameter, in fractures of a greisen, in quartz breccia and quartz veins.

Semi-spherical aggregates occur isolated (Fig. 14), or in small groups grading to thin crusts with irregular to botryoidal surface (Fig. 15). The aggregates are bright emerald to bluish green, probably in dependence on Al/Fe ratio. The surface of the aggregates is formed by platy crystals (Figs 16 and 17), or rare pale blue aggregates (Fig. 18) of prismatic crystals with acute terminations, which belong to the minerals close to the turquoise end-member (Figs 19 and 20).

The quantitative chemical analyses show various members ranging nearly within the whole isomorphous series (Fig. 21) and indicating chemical heterogeneity of stud-

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**Fig. 14 Spherical chalcosiderite aggregates, Krásno (greisen) near Horní Slavkov, width of figure 20 mm; photo J. Sejkora, Nikon SMZ1500.**

**Fig. 15 Crusts formed by hemispherical aggregates of mineral of turquoise group, Krásno (greisen) near Horní Slavkov, width of figure 30 mm; photo J. Sejkora, Nikon SMZ1500.**

**Fig. 16 Hemispherical chalcosiderite aggregates; Vysoký Kámen (greisen), Krásno, width of figure 125 μm; SE photo M. Fíkar.**

**Fig. 17 Groups of chalcosiderite crystals; Vysoký Kámen (greisen), Krásno, width of figure 125 μm; SE photo Z. Mach.**
ied aggregates (Fig. 22). Examples of compositionally heterogeneous chalcosiderite deposited on the crusts of crystallized turquoise were found (Fig. 23). Chemical composition of selected members of the series with empirical formulas is shown in Table 10. The powder X-ray data agree with the PDF2 database.

**Table 10** Chemical composition of members of the chalcosiderite – turquoise isomorphous series from the Vysoký Kámen (greisen) deposit (in wt. %)

<table>
<thead>
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<td>CuO</td>
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<td>CaO</td>
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<td>FeO</td>
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<td>20.22</td>
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<td>Al₂O₃</td>
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<td>22.18</td>
<td>31.97</td>
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<tr>
<td>P₂O₅</td>
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<td>31.08</td>
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<td>As₂O₅</td>
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<td>0.14</td>
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<td>SO₃</td>
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<tr>
<td>H₂O*</td>
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<td>14.38</td>
<td>15.53</td>
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<td>102.57</td>
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</tr>
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</table>

* H₂O is calculated from stoichiometry, assuming the ideal formula CuM₃[(PO₄)OH]₂·4H₂O (M = Fe, Al) and charge balance. Empirical formulas were calculated on the basis of 28 (O,OH).
1. theoretical composition of chalcosiderite CuFe₃[PO₄(OH)]₂·4H₂O
2. composition showing the highest iron content, corresponding to empirical formula:
(Cu₃Fe₃₋ₓAlₓ)[PO₄]₂(OH)₂·4H₂O, x = 0.40 where H₂O = 4.00H₂O
3. chalcosiderite showing an intermediate composition corresponding to the following empirical formula:
(Cu₃Fe₃₋ₓAlₓ)[PO₄]₂(OH)₂·4H₂O, x = 1.40 where H₂O = 4.00H₂O
4. turquoise showing an intermediate composition corresponding to the empirical formula:
(Cu₃Fe₃₋ₓAlₓ)[PO₄]₂(OH)₂·4H₂O, x = 2.40 where H₂O = 4.00H₂O
5. specimen showing the highest recorded Al-content, corresponding to empirical formula:
(Cu₃Fe₃₋ₓAlₓ)[PO₄]₂(OH)₂·4H₂O, x = 3.40 where H₂O = 4.00H₂O
6. theoretical composition of turquoise CuAl₃[PO₄(OH)]₂·4H₂O

Fig. 18 Spherical turquoise aggregates, Krásno (greisen), width of figure 8 mm; photo J. Sejkora, Nikon SMZ1500.

Fig. 19 Spheroidal aggregate of turquoise prismatic crystals with acute termination; Vysoký Kámen (greisen), Krásno, width of figure 450 μm; SE photo M. Fikar.

Fig. 20 Detail of turquoise crystals with acute termination; Vysoký Kámen (greisen), Krásno, width of figure 100 μm; SE photo M. Fikar.

Fig. 21 Fe – Al (mol. % of components) in minerals of the isomorphic series chalcosiderite – turquoise, Vysoký Kámen (greisen).
Goethite FeO(OH)

Goethite occurs mostly in fractures, dominantly in association with phosphates of the turquoise-chalcosiderite series, as spherical radiating aggregates of small red brown crystals. It belongs to the youngest minerals at the deposit. The mineral was identified by X-ray powder diffraction data.

Leucophosphate K(Fe,Al)_2(PO_4)_2 OH·2 H_2O

It occurs in a hydrothermally silicified tectonic zone in the upper part of the greisen quarry. Leucophosphate forms yellow green to grey yellow crystalline coatings and crusts on quartz. Less common are the aggregates of small, transparent and highly lustrous prismatic crystals grown on small quartz crystals (Fig. 24). Also found were spheroidal greenish aggregates up to 3 mm in diameter, showing radiating internal structure with drusy crystals on their surface (Fig. 25). Leucophosphate is document-

Dickite Al_2Si_2O_5(OH)_4

Dickite is a frequent mineral at the deposit. The oldest generation of dickite formed during the late stages of hydrothermal mineralization. However, some dickite samples could have been of supergene origin. It forms beige white soft aggregates, deposited on a crust of light blue turquoise. Dickite coatings several cm² in size cover quartz crystals in the cavities or turquoise-chalcosiderite crusts. The mineral was identified by X-ray powder diffraction data.
Table 11 Infrared absorption spectra of leucophosphite from Vysočany Kámen

<table>
<thead>
<tr>
<th>[cm⁻¹]</th>
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<tbody>
<tr>
<td>3430</td>
<td>s</td>
<td>795</td>
<td>w</td>
</tr>
<tr>
<td>3310</td>
<td>s</td>
<td>775</td>
<td>vw</td>
</tr>
<tr>
<td>2950</td>
<td>vw</td>
<td>630</td>
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<td>2850</td>
<td>vw</td>
<td>590</td>
<td>s</td>
</tr>
<tr>
<td>1640</td>
<td>m</td>
<td>550</td>
<td>m</td>
</tr>
<tr>
<td>1460</td>
<td>w</td>
<td>460</td>
<td>s</td>
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<td>1380</td>
<td>w</td>
<td>425</td>
<td>s</td>
</tr>
<tr>
<td>1150</td>
<td>s,sh</td>
<td>360</td>
<td>m</td>
</tr>
<tr>
<td>1070</td>
<td>vs</td>
<td>300</td>
<td>w</td>
</tr>
<tr>
<td>990</td>
<td>vs</td>
<td>232</td>
<td>m</td>
</tr>
<tr>
<td>840</td>
<td>m</td>
<td>208</td>
<td>m</td>
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</tbody>
</table>

* Character of absorption maxima:
  sh – shoulder; b – broad. Intensity scale:
  vs – very strong; s – strong; m – medium;
  mw – medium weak; w – weak; vw – very weak.

Fig. 26 The infrared spectra of leucophosphite from Vysočany Kámen (greisen), Krásno.

Minerals of the strengite FePO₄·₂H₂O – variscite AlPO₄·₂H₂O series

Members of the variscite-strengite series occur along fractures and in the cavities of greisen as minute grey white coatings with a botryoidal surface and a radiating-fibrous internal structure. Grey and light violet spheroidal aggregates 1 to 2 mm in diameter (Fig. 27) and with a drusy surface, composed of pyramidal crystals 0.1 mm

Fig. 27 Yellowish spherical aggregates od strengite, Krásno (greisen), width of figure 10 mm; photo J. Sejkora, Nikon SMZ1500.

ed from the association with the minerals of the turquoise-chalcosiderite and variscite-strengite series. The mineral was identified by X-ray powder diffraction data. The infrared absorption spectrum of leucophosphite is shown in Table 11 and Fig. 26.

Fig. 28 Light violet, radiating spheroidal aggregate of strengite; Vysočany Kámen (greisen), Krásno, width of figure 2 mm; SE photo M. Fíkar.

Minerals of the strengite FePO₄·₂H₂O – variscite AlPO₄·₂H₂O series

Fig. 29 Detail of the surface of spheroidal strengite aggregate composed of pyramidal crystals; Vysočany Kámen (greisen), Krásno, width of figure 300 μm; SE photo M. Fíkar.

Fig. 30 Spheroidal aggregates of pyramidal variscite crystals deposited on quartz crystals; Vysočany Kámen (greisen), Krásno, width of figure 5 mm; SE photo A. Gabašová.
long (Figs 28 and 29), occur in association with leuco-
phosphate and chalcosiderite. The morphology of the
variscite crystals is shown in Figs 30 and 31. The min-
erals of this isomorphous series have been identified in
addition to X-ray powder diffraction also by semi-quant-
tative chemical analyses. All the samples analyzed con-
tain Al and Fe. Pure end members of the series are un-
known at this locality.

**Turquoise CuAl₃[PO₄(OH)]₄·4 H₂O**

See minerals of the series chalcosiderite-turquoise.

**Variscite AlPO₄·2 H₂O**

See minerals of the strengite-variscite series.

**Vivianite Fe₅(PO₄)₃·8 H₂O**

Vivianite was observed rarely in cavities, deposited on
white quartz grading to amethyst in the fractures of jas-
per and in the greisen in the southern part of the quarry.
It forms typical radiating aggregates composed of dark
blue green to black green crystals (Fig. 32) with acute
termination, which are up to 3 mm long. Vivianite oc-
curs in association with cacaoxenite, minerals of the tur-
quoise-chalcosiderite series and strengite-variscite. Vi-
vianite was identified by X-ray powder diffraction.

**Wavellite Al₉(PO₄)₃(OH,F)₉·5 H₂O**

Wavellite occurs in association with other phosphates as
grey white coatings with a fibrous internal structure. Ex-
ceptionally it forms radiating aggregates with older tur-
quoise or chalcosiderite in their centres. Wavellite be-
longs to the youngest phosphates at the deposit. It was
identified by X-ray powder diffraction.

**Interpretation of supergene mineral assemblages at the Vysoký Kámen deposit**

**Vysoký Kámen (feldspar)**

Weathering processes, which took place at this locality
and resulted in formation of supergene minerals, repre-
sent several distinct sets of conditions. In a general way,
they include a range of processes from probably late hy-
drothermal to supergene (recent). Leaching and decom-
position of beryl, molybdenite and wolframite belong to
an early episode of alteration. The resulting supergene
minerals, including bertrandite, powellite (in pseudomor-
phs after molybdenite) and scheelite belong to the oldest
ones. They are followed by a younger generation of pow-
ellite and by kolbeckite. Koechinlite-russellite, autunite
and meta-autunite belong to the latest stages of supergene
alteration. Koechinlite-russellite replaced powellite; Bi
necessary for crystallization of this mineral was provid-
ed by native bismuth. There are indications for several
generations of koechinlite-russellite, autunite and meta-
autunite. Thus, an early generation of autunite crystals
was observed in cavities with quartz. The youngest gen-
eration of meta-autunite occurs relatively often as an in-
conspicuous phase in aggregates of clay minerals. Younger
koechinlite-russellite occurs in small cavities after leached
molybdenite. The youngest mineral in this association is
goethite, observed in some vugs and along fractures.
Vysoký Kámen (greisen)

A relatively simple association of phosphates formed in a breccia zone, partly heated by quartz, and in neighboring parts of fractured greisen. Leucophosphate, as the oldest mineral, and members of the variscite-strengite series are probably related to a closing stage of hydrothermal alteration. Younger phosphates of the turquoise-chalcociderite series occur in fractures of greisens and quartz gangue. The complicated compositional zoning in minerals of the variscite-strengite and turquoise-chalcociderite series indicates significant changes in chemical composition of solutions, including especially variation in the Al/Fe\(^{3+}\) ratio. Cacaoxenite, vivianite and goethite, as the youngest members of this mineralization, indicate absolute dominance of Fe\(^{3+}\) above Al\(^{3+}\) during the closing stages of supergene mineralization at this locality.

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References


Nové poznatky o mineralogii ložisek Vysoký Kámen u Krásna (Slavkovský les, Česká republika)

Podrobnejší mineralogický výzkum byl zaměřen na dvě ložiska lokalizovaná na svazích kóty 819,4 m Vysoký Kámen u Krásna (u horního Slavkova, Česká republika). V materiálu z činného lomu těžícího živcovou surovinu (zejména feldsparty jako součást přírodního mladšího intruzivního komplexu) byly zjištěny autunit, bertrandit, beryl, goethit, hematit, kockelinit, kolbeckit, meta-autunit, powellit a scheelit. Na opačněm greisenovém ložisku byla vedle dicitu a goethitu zjištěna minerální asociace fosfátů zastoupená kacoxenem, leukofosfitem, vivianitem, wavelitem a minerály série chalcociderit-týrkys. V práci je podána i stručná interpretace vzniku popisovaných minerálních asociací.