Selenide Mineralization in the Příbram Uranium and Base-Metal District (Czech Republic)

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Abstract: Selenium mineralization in the Příbram uranium and base-metal district (Central Bohemia, Czech Republic) bound to uraninite occurrences in calcite hydrothermal veins is extremely diverse. The selenides—antimonselite, athabascaite, bellidoite, berzelianite, brodtkorbite, bukovite, bytízite, cadmoselite, chaméanite, clausthalite, crookesite, dzharkenite, eskebornite, eucairite, ferroselite, giraudite, hakite, klockmannite, naumannite, permingeatite, příbramite, sabatierite, tiemannite, and umangite were found here, including two new mineral phases: Hg-Cu-Sb and Cu-As selenides. Those selenides—and in some cases their sulphidic equivalents—are characterized using wavelength-dispersive spectroscopy, reflected light, powder X-ray diffraction, single crystal X-ray diffraction, Raman spectroscopy, and electron backscatter diffraction. The selenide mineralization in the Příbram uranium district is bound to the border of the carbonate-uraninite and subsequent carbonate-sulphidic stages. Selenides crystallized there at temperatures near 100 °C in the neutral-to-weakly-alkaline environment from solutions with high oxygen fugacity and a high Se2/S2 fugacity ratio.

Keywords: selenides; chemical composition; crystal structure; wavelength-dispersive spectroscopy; X-ray diffraction; Příbram

1. Introduction

The Bohemian Massif belongs to those areas on a global scale having anomalously abundant occurrences of selenides. This is also true with respect to the number of determined selenide species. The first occurrence of selenides in the Bohemian Massif was recognized by Johan [1] at the Černý Důl uranium deposit in the Krkonoš Mountains. Later, large accumulations of selenides were found at the West-Moravian uranium deposits [2–4] and at the Předbořice uranium deposit in Central Bohemia [5–7]. Additional and important selenide occurrences were reported in Western Bohemia—Oldřichov near Tachov [8,9] and Ústaleč [10,11]. The Zálesí uranium deposit [12–15], the Jelení vrch [12], and the Bílá Voda [16] occurrences were discovered in the Rychlebské hory Mountains. The rich association of selenides from the uranium ore district Niederschlema-Alberoda in the Saxonian part of the Krušné hory Mountains was also studied in detail [17–23]. Recently, selenide occurrences tied to uranium-vanadium mineralization in the Prachovice quarry [24] and those associated with minor uranium mineralization at fluorite-baryte Moldava [25] and Běstvina [26] deposits were studied.
Selenide occurrences outside uranium mineralization are only sporadic in the Bohemian Massif. Worthy of mention are rare finds of naumannite and tiemannite in the heavy mineral fraction of sediments near Svoboda nad Úpou [27]; Mo-selenides and clausthalite in black schists near Hromnice [28]; clausthalite and Ag–Cu selenides in stromatolites at Kokšin [29]; inclusions of clausthalite in wolframite from the Sn-W Krupka deposit [30]; and Bi-sulfoselenotellurides in products of burning of coal-mine dump at Radvanice near Trutnov [31].

The exceptional character of selenide occurrences in the Bohemian Massif is underlined by 18 newly-discovered selenide mineral species, out of a total of ca. 90 known selenide minerals. The new minerals include bukovite [32] and sabatierite [33] from the Bukov locality; belloïdoite from Habří [34]; bohdanowiczite from Kletno in Poland [35]; pousiaite [8] and součekite [9] from Oldřichov near Tachov; krut’alite [36] and petrovicite [37] from Petrovice; fischesserite [38], hakite [39], permingeatite [40], milotaite [41], and petříčekite from Předbořice [42]; mgriite [43] and schlemaite [44] from Schlema in Germany; litochlebite from Záleší [45]; and příbramite [46] and bytizite [47] from Příbram.

At the uranium and base-metal Příbram ore district (the most important hydrothermal uranium district in the Czech Republic), clausthalite was originally reported as rare and local in its occurrence [48]. The first relatively important selenide occurrence was described on the basis of a preserved reference sample by Litochleb et al. [49]. The extraction of material from mine dumps at some Příbram uranium mines (in particular, shafts 16, 11A, and 9) provided the opportunity during the period of 2005–2015 to monitor selenium minerals in the mine dump material and obtain more samples for a complex mineralogical study of selenide mineral assemblages. The first partially-published results include clausthalite in association with arsenolamprite [50] and cuprostitite [51], permingeatite [52], S-rich antimonelite [53], hakite [54], příbramite [46], brodkorkbite [55], and bytizite [47]. A comprehensive study of the selenide mineralization at the Příbram uranium base-metal ore district was the topic of a Ph.D. thesis of Škácha [56], and forms the basis of this paper. Empirical formulae of selenides and some associated sulphides are shown in Table S1.

2. The Příbram Uranium and Base-Metal Ore District

2.1. History of Mining

Two main ore districts are known in the Příbram ore area (Central Bohemia, Czech Republic). They include the Březové hory base-metals district and the Příbram complex uranium and base-metals district (Figure 1), referred to as the “Příbram uranium district” in the following text. The Březové hory base-metal district—located in the western part of the area—was known from at least the 13th century, but the main mining activity took place from 1810 to 1978. A total of 22 million tonnes of extracted ore contained 3837 tonnes of silver, 517,961 tonnes of lead, and 70,300 tonnes of antimony [57].

The history of the easterly-positioned Příbram uranium district started in 1947 with the discovery of the first uranium anomalies. Mining started as early as 1948 with the opening of the first shafts (1 and 2, Vojna). In 1950, uranium exploitation started on a large scale. In succession, 26 shafts were constructed, which opened eight main deposits of the uranium district (Třebško, Kamenná, Lešetice, Brod, Jerusalem, Háje, Bytíz, and Skalka). The deposits used to be historically designated as “vein nodes”.

Minerals 2017, 7, 91
Uranium mining culminated there in 1975, when this ore district became the main producer of uranium ores in the former Czechoslovakia. In the following year, the final depth of 1838.4 m was reached in shaft 16 (Háje). At that time, this was the deepest ore mine, vertically, in Europe. Owing to a change in economic and political conditions in the former Czechoslovakia in 1989, the extraction of uranium ores in this ore district was terminated in 1991. The following numbers indicate the remarkable extent of the mine works in the Příbram uranium ore district: 23 km of vertical shafts, 2188 km of horizontal adits, and 300 km of chutes were created in an area of 57.6 km² over 44 years (1948–1991). In the process, more than 2500 hydrothermal (especially carbonate) veins were exposed; the uranium mineralization occurred in 1641 veins, base metal mineralization in 35 veins, and finally, monomineralic silver mineralization in 19 veins [57]. The total production of 48,432 t of pure U metal represented 49% of Czechoslovak production since 1945. The parallel mining of base metals and silver from veins produced more than 6100 tonnes of Pb, 2400 tonnes of Zn, and 28 tonnes of Ag [58]. These data qualify the Příbram uranium district for competition with the most important hydrothermal vein uranium districts on a worldwide scale.

2.2. Geological Situation

Both of the main ore districts (Březové Hory base-metals district and the Příbram uranium district) are situated near the southeastern contact of Barrandian sedimentary rocks of the Neoproterozoic and Lower Cambrian age, with the western margin of granitoids of the Central Bohemian Plutonic Complex. The Příbram uranium district is located tightly in the exocontact of the Central Bohemian Plutonic Complex in metasedimentary rocks of the Barrandian Unit. The uranium hydrothermal vein mineralization is located in a zone of tectonically-deformed Neoproterozoic rocks 1–2 km wide and 25 km long, including the Příbram antiform [58]. A predominant part of uranium mineralization (96%) occurred in veins located in the Dobříš Group of Neoproterozoic age with a predominance of claystones, siltstones, and sandstones. The underlying Davle Group of tuffs and tuffites was encountered by two structural boreholes and by deep levels of mines in the Bytíz area.
Lower Cambrian sediments—including conglomerates overlying the Dobříš Group—were penetrated by numerous mines. This concerns the upper parts of the central and northeastern part of the ore district and at greater depth in the northern part of the vein Bt4. All rock types are affected by contact metamorphism caused by granitoids of the Central Bohemian Plutonic Complex [59]. Granitoids are represented here by the so-called marginal-type and Blatná-type granitoids. Mineralized veins wedge out upon reaching plutonic rocks.

There are four main mineralization stages generally identified in the Příbram uranium district [59]: (I) siderite-sulphidic; (II) calcite; (III) calcite-uraninite; and (IV) calcite-sulphidic. The oldest siderite-sulphidic (stage I) is developed on a smaller scale in comparison with the neighbouring Březové Hory district. The younger calcite stages are characterized by notably lower temperatures, and are more abundant; calcite generations were used in separating mineralization stages. The calcite stage (II) is characterized by pre-ore calcite DK and calcite K1. In the calcite-uraninite stage (III)—carrying the main part of the industrial uranium mineralization (uraninite, coffinite, and U-bearing anthraxolite)—calcite types K2–K4 are present. In the last calcite-sulphidic stage (IV), post-ore calcite K5 appears.

The calcite-uraninite (stage III) and the youngest calcite-sulphidic (stage IV) stages are important for selenide mineralization. Selenide mineralization occurs in close association with uraninite of the calcite-uraninite mineralization, but selenides are always younger than uraninite. It is uncertain if selenides in Příbram originated in the conclusion of the calcite-uraninite stage, or at the beginning of the following calcite-sulphidic stage, which is mentioned by some authors [55].

The age of the uranium mineralization obtained by the U-Pb dating of uraninite showed the middle Permian ages of 275 ± 4 and 278 ± 4 Ma [60]. Uraninite from the nearby Březové Hory base-metal ore district has similar ages (269.8 ± 20.3 and 263.2 ± 8.9 Ma) [61]. According to the fluid inclusions study [62], the carbonate veins carrying uraninite in the Příbram uranium district formed at low-temperatures (80–140 °C) from low-salinity fluids. Other data indicate that fluids instrumental in the deposition of the uraninite-carbonate mineralization derived from local meteoric waters and evolved in the process of migration through upper crustal rocks [62].

3. Methods of Research

3.1. Reflected Light Microscopy

A Zeiss Stemi 2000C optical microscope (Jena, Germany) with a Canon 5DSR digital camera (Tokyo, Japan) (in the Příbram Mining Museum) was used for the study of sample morphology in reflected light. Samples for microscopy and the following laboratory study were prepared as polished sections 2.54 cm across, mounted in resin, and polished with diamond suspensions in the Diatech s.r.o. company, Prague, Czech Republic. Over 250 polished sections representing all types of vein filling were used.

Optical properties of minerals in reflected light were studied using a Nikon Eclipse ME600 microscope (Minato, Japan) at the National Museum, Prague, Czech Republic. Over 250 polished sections representing all types of vein filling were used.

3.2. Quantitative Analysis of Chemical Composition

Chemical analyses were performed using a Cameca SX100 electron microprobe (Gennevilliers Cedex, France) (National Museum, Prague; P. Škácha, J. Sejkora and I. Macek as analysts) operating in WDS (wavelength-dispersive spectroscopy) mode (25 kV, 20 nA, and 2 µm wide beam). The following standards and X-ray lines were used to minimize line overlaps: Ag (AgLα), Au (AuMα), Bi (BiMβ), CdTe (CdLα), Co (CoKα), chalcopyrite (CuKα), FeS2 (FeKa, SKα), HgTe (HgMα), NiAs (NiKα, AsLα), PbS (PbMα), PbSe (SeLα), PbTe (TeLα), Sb2S3 (SbLα), Ti(BrI) (TlLα), and ZnS (ZnKα). Peak counting times were 20 s for all elements and one-half of the peak time for each background. Contents of the above-listed elements (which are not included in tables) were analyzed quantitatively, but with
 contents below the detection limit (ca. 0.01–0.04 wt % for individual elements). Raw intensities were converted to the concentrations of elements using automated “PAP” [63] matrix-correction software.

3.3. X-Ray Powder Diffraction Methods (PXRD)

In the first stage, small fragments of samples carrying selenides were dissolved in hydrochloric acid. Insoluble residues were washed with water and neutralized with Na$_2$CO$_3$. Small selenide aggregates submerged in water were mechanically separated according to colour and morphological properties of individual minerals.

Selected monomineral subsamples were used for diffraction data using a Bruker D8 Advance X-ray powder diffractometer (National Museum, Praha, Czech Republic) with a LynxEye semiconductor detector, using CuK$_\alpha$ radiation (40 kV, 40 mA). Powder mounts were prepared with acetone suspension on a plate of monocrystalline silicon. X-ray data were recorded in a step-scanning manner (step 0.01°, time 30 s/step of the detector; total time of experiment more than 50 h). Positions of individual diffraction maxima were refined with a Pearson VII profile function in the ZDS program for DOS [64]. Indices hkl were assigned based on tabulated theoretical values calculated from available structural data with the Lazy Pulverix program [65]. Lattice parameters for individual minerals were refined using Burnham program [66].

3.4. Single-Crystal X-ray Diffraction Methods (SXRD)

Single-crystal diffraction experiments were done using a Rigaku Oxford diffraction SuperNova diffractometer (Tokyo, Japan) equipped with an Atlas S2 CCD detector. Mirror-monochromatized MoK$_\alpha$ radiation ($\lambda = 0.71073$ Å) from a microfocus X-ray source was utilized. Unit-cell parameters were refined from the diffraction data employing least-squares algorithm within the CrysAlis Pro program package (Rigaku).

3.5. Electron Backscatter Diffraction (EBSD)

For checking the identity of the crystal structure of selected minerals with a presumed structure model, the EBSD method (electron backscatter diffraction) was used. The surface of the sample in the polished section was treated by chemical polishing, using amorphous SiO$_2$ in suspensions with successively acidic and alkaline pH.

Measurements were conducted with a Tescan Vega (Brno, Czech Republic) scanning electron microscope (Faculty of Science, Charles University, Praha), operated by M. Racek. The apparatus was equipped with a NordlysNano EBSD detector (Oxford Instruments, Abingdon-on-Thames, UK). The following operating conditions were used: accelerating voltage of 20 kV, beam current ca. 7 nA, the area of the analysed sample (diameter of the area for signal collecting) less than ca. 1 $\mu$m, and tilted by $70^\circ$ from horizontal. Data obtained were registered and processed using Channel5 software (Oxford Instruments, Abingdon-on-Thames, UK); data for the structural model (unit-cell parameters, space group, and atoms and their co-ordinates in the elementary cell) were downloaded from the crystal structure refinements cited in the discussion of the individual minerals below.

3.6. Raman Spectroscopy

Raman spectra of mineral phases in polished sections were collected in the range 30–3500 cm$^{-1}$ using a DXR dispersive Raman spectrometer (Thermo Scientific, Waltham, MA, USA) mounted on a confocal Olympus microscope (National Museum, Prague, Czech Republic). The Raman signal was excited by a green 532 nm diode-pumped solid-state laser and detected by a CCD detector. The experimental parameters were: 50× objective, 1 s exposure time, 1000 exposures, 900 lines/mm grating, 50 $\mu$m slit spectrograph aperture, and 0.5 mW laser power level. The instrument was set up by a software-controlled calibration procedure using multiple neon emission lines (wavelength calibration), multiple polystyrene Raman bands (laser frequency calibration), and standardized
white-light sources (intensity calibration). Spectral manipulations were performed using Omnic 9 software (Thermo Scientific).

4. Results

4.1. Characterization of Selenide Occurrences in Příbram

Růžička [48] recorded the first occurrence of selenium mineralization in the Příbram uranium district by describing clausthalite from vein Bt4 at the Bytíz deposit. Later on, selenides were identified in documentation samples (deposited in the National Museum, Praha) from vein Bt 22B, 4th level, north [49]. In the course of the continued research of documentation, a sample carrying uraninite was located with another sample containing small accessory aggregates of selenides. This sample came from the Bt 4 vein, 8th level, north, corresponding to the depth of 450 m. Due to the sampling period, the documentation set did not contain samples from deeper levels of the deposit. This might indicate that the selenium mineralization is developed only in the uppermost part of the district. However, the considerable distribution of the selenide mineralization in the dumps at shafts 16 and 11A, and the overall relative abundance of material with selenides, supports the alternative of a deeper vertical extent of the selenide mineralization—at least at the Bytíz deposit. Several additional samples with selenides were found at the dump of shaft 9 (Jerusalem), used for the exploitation of veins in the upper part of the Jerusalem deposit.

Samples used in the present study were collected at the dumps at shafts 9 (Jerusalem), 16 (Háje near Příbram), and 11A (Bytíz). Dumps at shafts 16 (Figure 2A) and 11A (Figure 2B) were recently intensively processed by the Ecoinvest Company for the production of crushed aggregate, in part as a construction material for the D3 highway. Continued exploitation and technical separation of dispersed radioactive material made sample collection from the lower parts of dump 16 (Háje near Příbram) and parts of the dump 11A (Bytíz) possible.

![Figure 2. Dump of the shaft (A) 16-Háje in 2007 and (B) 11A-Bytíz in 2017. Photos by P. Škácha.](image-url)

The possible occurrence of material with selenide mineralization was checked at some other dumps using a sensitive scintillation detector. This included dumps at some other shafts of the Příbram uranium district (Figure 1)—in particular, shafts 2 (Vojna), 4 (Lešetice), 3C (Kamenná), 5, 6, 15 (Brod), and 19 (Dubenc). No additional selenide occurrences were found in the course of this field work. This negative result does not necessarily prove the absence of selenide, as the recultivated surface of the dumps presents adverse conditions for sample collection. Local radioactive anomalies on the surface of the dumps were subject to controls and material collection at the time of mine operation, and later attracted the attention of private mineral collectors.

At the Březové Hory ore district, bordering at the west of the Příbram uranium district, no selenide occurrences were found, although some local occurrences of uranium minerals are known. A single indication of a possible presence of selenides in this ore district is the report of supergene copper and lead selenate munakataite in several samples collected at a dump of the Lill mine [67].
4.2. Characteristics of Individual Minerals of the Selenide Association

4.2.1. Antimonselite Sb$_2$Se$_3$

Antimonselite was found at the dump of the mine 16, Háje near Příbram [53]. It forms euhedral acicular crystals up to 1.5 × 0.1 mm (Figure 3). The crystals are enclosed in calcite gangue in association with uraninite, hakite, claustralite, Se-rich chalcostibite, tiemannite, permingeatite, luzonite, tennantite, dzharkenite, and Příbramite (Figure 4). Antimonselite is among the youngest minerals in the studied polished sections. The crystals are brittle, with perfect (010) cleavage. In the reflected light, antimonselite is white and shows bireflection. Under crossed polarizers, it is distinctly anisotropic. X-ray powder diffraction data for Příbram antimonselite are given by Škácha et al. [53], including refined unit-cell parameters: $a = 11.7437(6)$, $b = 3.9358(7)$, $c = 11.5846(8)$ Å, $V = 535.41(1)$ Å$^3$. Based on single-crystal X-ray data, Škácha et al. [53] solved the crystal structure of the studied antimonselite. Based on the data (3353 reflections), refined unit-cell data were obtained using the least-squares algorithm of CrysAlis Pro [68] as follows: $a = 11.7156(3)$, $b = 3.9514(11)$, $c = 11.5645(3)$ Å, $V = 535.36(15)$ Å$^3$, $Z = 4$. The obtained volume of the unit-cell of antimonselite from Příbram is compared with the published data for members of the stibnite-antimonselite solid solution in Figure 5.

![Figure 3. Antimonselite prismatic crystals partly etched from calcite. Field of view (FOV) 2.0 mm. Photo by P. Škácha.](image3)

![Figure 4. Antimonselite prismatic crystals and their rhombic sections in association with yellow chalcopyrite and grey tetrahedrite. Reflected light, one polarizer. Field of view 210 μm. Příbram, a dump of the shaft 16. Photo by P. Škácha.](image4)
Among cations, the dominant Sb is substituted by minor Cu contents below 0.02 apfu.

The average empirical formula of Příbram antimonselite calculated on the basis of 5 synt., [72] synt., [71] synt., [70] deposit 504, China, [73] synt., [72] deposit 504, China, [74] synt., [72] is (Sb$_{2.06}$Cu$_{0.02}$)Σ$_{2.08}$ (Se$_{2.67}$S$_{0.25}$)Σ$_{2.92}$.

In the course of the chemical study of newly-polished sections of antimonselite (Table S2), minor sulphur contents in the range of 0.03–0.16 apfu were found in addition to the dominant Se. Škácha et al. [53] determined slightly higher sulphur contents in the range 0.31–0.56 apfu. Similar S contents have been reported for some samples from Chinese localities La’ermo and Qiongmo by Liu et al. [75]. The study of a synthetic stibnite-antimonselite series [72] shows a complete miscibility (Figure 6). Among cations, the dominant Sb is substituted by minor Cu contents below 0.02 apfu.

The average empirical formula of Příbram antimonselite calculated on the basis of 5 apfu for 16-point analyses (including those in Škácha et al. [53]) is (Sb$_{2.06}$Cu$_{0.02}$)Σ$_{2.08}$ (Se$_{2.67}$S$_{0.25}$)Σ$_{2.92}$.

**Figure 5.** Se contents (at% of anion) vs. the unit-cell volume graph for the members of the stibnite-antimonselite solid solution. PXRD: X-ray powder diffraction; SXRD: single crystal X-ray diffraction [69–74].

**Figure 6.** Se vs. S (apfu) graph of antimonselite from various localities.
4.2.2. Athabascaite Cu$_5$Se$_4$

In the studied assemblage, athabascaite is a relatively rare mineral. It forms up to 300 µm long anhedral aggregates composed of very thin lamellar crystals (Figure 7). Owing to its blue color, the mineral is very similar to berzelianite. Athabascaite is intergrown with umangite, and a local intergrowth with eucarite occurs. In addition, athabascaite is closely associated with Hg-hakite and uraninite. At the type locality Martin Lake Mine, Saskatchewan, Canada, athabascaite occurred as inclusions in umangite and as pseudomorphs after umangite was enclosed in gangue calcite. Several of the lath-shaped crystals show remnant cores of umangite that is finer-grained than the surrounding material, suggesting that the athabascaite is an alteration product of former umangite [76]. Umangite cores enclosed in athabascaite were also observed in samples from Prˇibrˇam (Figure 7).

![Figure 7.](image-url) Whitish to dark blue (pleochroism colors) aggregate of athabascaite with creamy white eucarite and very small violet umangite grains. The dark brown uraninite rim surrounds the periphery. Reflected light, one polarizer. Field of view 210 µm. Prˇibrˇam, a dump of the shaft 16. Photo by P. Škácha.

In reflected light, athabascaite is grey-blue, and shows distinct bireflectance with color ranging from yellow-grey to blue-grey. Anisotropy is strong with colors changing from cream white to dark grey.

The chemical composition of athabascaite from Prˇibrˇam (Table S3) is represented by cation/anion ratio in the range 1.10–1.37 (based on 44 analyses). The studied mineral grain is probably forming very fine intergrowths with other Cu selenides or sulphides, and this may be the reason for the observed non-stoichiometry. The cation part of the formula contains minor Ag (up to 0.14 apfu), and locally, Fe (up to 0.36 apfu), in addition to the dominant Cu. The anion part features a significant SeS$^{-1}$ isomorphism; sulphur contents attain up to 1.14 apfu (Figure 8)—i.e., contents higher than values reported for this mineral in the literature [14,49,76,77]. Additionally, a locally-significant Te content up to 0.16 apfu was rarely observed. The empirical formula for studied athabascaite on the basis of 9 apfu (average of 44 point analyses) can be expressed as $(\text{Cu}_{4.98}\text{Fe}_{0.06}\text{Ag}_{0.03})_{\Sigma5.07}(\text{Se}_{3.12}\text{S}_{0.80}\text{Te}_{0.01})_{\Sigma3.93}$. 


4.2.3. Bellidoite Cu$_2$Se

Bellidoite is an extremely rare selenide so far reported only from Habří (Czech Republic), Tumiñico (Argentina), El Dragón (Bolivia), and Jiguanzui and Jilongshan (China) [34,78–80]. Its occurrence in the Příbram uranium district is in the samples with abundant berzelianite, umangite, and eucairite, confirmed by X-ray powder diffraction.

In reflected light, bellidoite is creamy white (Figure 9). Anisotropy was not observed.

Figure 8. Se vs. S (apfu) graph of athabascaite from various localities.

Figure 9. Bellidoite (Be) euhedral grains in association with bluish berzelianite (Bz). Reflected light, (A) one polarizer, and (B) partly-crossed polarizers. Field of view 210 µm. Příbram, from a dump of shaft 16. Photo by P. Škácha.

Bellidoite identification in polished sections is made difficult at first sight by the similarity of the optical properties of bellidoite to those of berzelianite in small sizes of grains and their intergrowths. In addition, berzelianite and bellidoite often form very close intergrowths, and the size of individual grains is under 1 µm. Eucairite—and especially berzelianite—are rather abundant in the studied assemblage.

The X-ray powder diffraction data of bellidoite from Příbram (Table S4) agree very well with the published data of this mineral phase [34]. Due to the unknown crystal structure of bellidoite,
the experimental pattern (selected maxima of bellidoite in a mixture with berzelianite and eucalrite) was indexed on the base of a tetragonal ($P4_2/n$) unit-cell with $a = 11.52$ and $c = 11.74$ Å, proposed by Montreuil [34]. The refined unit-cell parameters of bellidoite from Příbram are $a = 11.607(7)$, $c = 11.74(1)$ Å, and $V = 1582(2)$ Å$^3$.

The chemical composition of Příbram bellidoite may be partly affected by its intimate intergrowth with berzelianite (Figure 10). The cation part of the formula contains Ag (0.00–0.02 apfu) and Fe (0.00–0.06 apfu); in anion parts, only a minor S content was observed (0.00–0.01 apfu) (Table S5). Very low S content is typical for this phase according to the published data [34]. The empirical formula of Příbram bellidoite (from an average of 109 analyses) based on 3 apfu is $\text{Cu}_{1.96}(\text{Se}_{1.00}\text{S}_{0.03})\Sigma_{1.03}$.

![Figure 10. Se vs. S (apfu) graph for bellidoite from Příbram.](image)

4.2.4. Berzelianite $\text{Cu}_{2-x}\text{Se}$

Berzelianite belongs among the most abundant selenides at the deposit. It forms typical blue to blue-green anhedral aggregates in calcite gangue. Berzelianite aggregates rarely exceed 2 mm. The mineral was observed as inclusion in uraninite and as abundant rims separating carbonate generations. It was frequently identified in the proximity of massive uraninite veins, in thin veinlets in the adjacent carbonate, or directly in uraninite (Figure 11). The presence of berzelianite in carbonate-uraninite vein filling deposited at dumps is indicated by abundant malachite coatings formed by the recent weathering of berzelianite. Berzelianite aggregates often show replacement by umangite, starting in the center of aggregates. Overgrowth of berzelianite aggregates by chalcopyrite is rare (Figure 12). The association of berzelianite with Hg-hakite is typical (Figure 13). In reflected light, isotropic berzelianite has a steel grey-blue color. X-ray powder data for berzelianite (Table S6), as well as the refined parameters of the unit-cell (Table S7), correspond well to the data published for this mineral.
Figure 11. Spherical aggregates of uraninite (white) overgrown by later berzelianite (grey). Backscattered electron (BSE in the further text) photo. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.

Figure 12. Earlier berzelianite (blue) with umangite (violet) inclusions replaced by later chalcopyrite (yellow). Reflected light, one polarizer. FOV 525 μm. Příbram, a dump of shaft 16. Photo by P. Škácha.

Figure 13. Berzelianite aggregates (dark grey) in Hg-hakite (light grey) in association with uraninite (white). BSE photo. Příbram, a dump of shaft 16. Photo J. Sejkora and P. Škácha.
The X-ray powder diffraction data were obtained for two types of berzelianite from Příbram (Table S6). The pattern of berzelianite (S-poor) agrees very well with the data calculated from the crystal structure information [81]; the positions of diffraction maxima of S-rich berzelianite are shifted to lower 2θ angles. The refined unit-cell parameters are given in Table S7; the increased content of S results in a decrease of the berzelianite unit-cell volume.

Study of the chemical composition of S-poor berzelianite (Table S8) and S-rich and S + Te-rich berzelianite (Table S9) results in cation/anion ratios in the range of 1.62–1.94 (Figure 14). This value may include—besides non-stoichiometry of the mineral—the effect of intergrowth with other Cu-selenides on a sub-microscopic scale. We can distinguish three chemical types of berzelianite: normal S-poor berzelianite, S-rich berzelianite, and S, Te-rich berzelianite. The most common are S-poor berzelianite and much rarer S-rich berzelianite, in association with arsenic and arsenolamprite, or with chalcocite and S-rich umangite was found. S, Te-rich berzelianite was found only in one case in association with Te-tiemannite. For all three types, the cation part of the formula (except the dominant Cu) frequently contains minor silver (up to 0.03 \( \text{apfu} \)), and in the case of S-poor berzelianite and S-rich berzelianite, local Fe content up to 0.07 \( \text{apfu} \) and Hg content up to 0.004 \( \text{apfu} \) were found. The anion part exhibits isomorphic substitution of Se by S (up to 0.51 \( \text{apfu} \)) in the case of S-rich berzelianite and rarely by Te (up to 0.06 \( \text{apfu} \)), together with S (up to 0.19 \( \text{apfu} \)). Sulphur contents in berzelianite from various localities are reported in Figure 14. For the first time, such samples, containing up to 1.24 S \( \text{apfu} \), were reported from Martin Lake (Canada) by Harris et al. [82]. Litochleb et al. [49] mentioned elevated sulphur contents up to 0.22 \( \text{apfu} \) for a sample from the Bytíz deposit. We have found no reports on Te content in berzelianite in the literature. The empirical formula for S-poor berzelianite, \( \text{Cu}_{1.93}(\text{Se}_{1.02}\text{S}_{0.05})\Sigma_{1.07} \), is based on 4 \( \text{apfu} \) and an average of 165 analyses. The empirical formula for S-rich berzelianite, \( \text{Cu}_{1.88}\text{Fe}_{0.03}(\text{Se}_{0.69}\text{S}_{0.43})\Sigma_{1.12} \), is based on 4 \( \text{apfu} \) and an average of 52 analyses. The empirical formula for S, Te-rich berzelianite, \( \text{Cu}_{1.94}(\text{Se}_{0.85}\text{S}_{0.14}\text{Te}_{0.06})\Sigma_{1.05} \), is based on 4 \( \text{apfu} \) and an average of four analyses.

![Graph](image)

**Figure 14.** Se vs. S+Te (apfu) graph in berzelianite from various localities.

4.2.5. Brodtkorbite, \( \text{Cu}_2\text{HgSe}_2 \)

Brodtkorbite is an extremely rare selenide known only from the type locality Tumiñico in Argentina [79]. In Příbram, brodtkorbite was identified in carbonate gangue carrying abundant tiemannite, berzelianite, and other selenides [55]. It occurs as anhedral aggregates up to 150 µm across in intergrowth with tiemannite (Figure 15), clausthalite, and eucarite, and contains uraninite inclusions of a spheroidal shape. Relatively abundant lamellar intergrowth of tiemannite and umangite represents the most probable products of brodtkorbite disintegration (Figure 16).
4.2.6. Bukovite, Tl\textsubscript{2}(Cu,Fe)\textsubscript{4}Se\textsubscript{2}

In the reflected light, the mineral has properties similar to the type of material described by Paar et al. [79]. It is cream white with a light pink or bluish shade, somewhat darker than the associated tiemannite. It is strongly anisotropic, with colors under crossed polarizers changing from violet grey to yellow grey. Brodtkorbite is older than tiemannite, which replaces it along fractures.

X-ray powder diffraction data for Příbram brodtkorbite are given by Sejkora et al. [55], including refined unit-cell parameters: $a = 7.5137(5)$, $b = 4.1840(2)$, $c = 8.0183(6)$ Å, $\beta = 124.43(2)^\circ$, $V = 207.59(8)$ Å$^3$. Based on single-crystal X-ray data, Sejkora et al. [55] solved the crystal structure of the studied brodtkorbite ($R_{\text{obs}} = 0.0805$). Based on the data (416 reflections), refined unit-cell data were obtained using the CrysAlis Pro [68] program: $a = 7.505(1)$, $b = 4.1851(9)$, $c = 8.013(1)$ Å, $\beta = 124.43(2)^\circ$, with $V = 207.59(8)$ Å$^3$, $Z = 2$.

The chemical composition of Příbram brodtkorbite is relatively uniform. In addition to the main elements—Hg, Cu, and Se (Table S10)—were observed minor contents of S up to 0.02 apfu, Te to 0.01 apfu, and Ag up to 0.01 apfu. Different from the type material from Tumiñico [79], the studied brodtkorbite from Příbram does not contain trace amounts of Fe. The empirical formula of Příbram brodtkorbite (the mean of 19 analyses) based on 5 apfu is $(\text{Cu}_{1.80}\text{Ag}_{0.01})_{\Sigma1.81}\text{Hg}_{1.11}(\text{Se}_{2.07}\text{S}_{0.01})_{\Sigma2.08}$. 

**Figure 15.** Brodtkorbite grains (grey) in association with tiemannite (white) in calcite gangue. BSE photo. Příbram, a dump of shaft 16. Photo by J. Sejkora.

**Figure 16.** Brodtkorbite aggregates (grey) partly disintegrated to umangite (dark grey) and tiemannite (white). BSE photo. Příbram, a dump of shaft 16. Photo by J. Sejkora.
4.2.6. Bukovite, Tl$_2$(Cu,Fe)$_4$Se$_2$

Bukovite belongs among the rare selenides at the deposit, together with sabatierite and crookesite as other selenides containing Tl in its crystal structure. It occurs as lath-shaped crystals up to 30 µm long, intergrown with uraninite and tiemannite in association with eucairite (Figure 17). It was also identified as 20 µm euhedral crystals intergrown with sabatierite, or Zn-hakite, eskebornite, and umangite (Figure 18).

![Figure 17](image1.png)

**Figure 17.** The euhedral aggregate of bukovite (Bu) overgrowing with eucairite (Eu) and tiemannite (T) earlier spherical aggregates of uraninite (U). BSE photo. Příbram, a dump of shaft 16. Photo by J. Sejkora, P. Škácha.

![Figure 18](image2.png)

**Figure 18.** Grey bukovite growing with brownish eskebornite, violet umangite, and grey Zn-hakite. Reflected light, one polarizer. FOV 90 µm. Příbram, a dump of shaft 16. Photo by P. Škácha.

The mineral is light grey in reflected light, and weak anisotropy was observed.

The chemical composition of bukovite is relatively homogeneous (Table S11); identified were minor abundances of Ag (up to 0.20 apfu), Pb (up to 0.01 apfu), and S (up to 0.20 apfu). Specimens from the type locality [83] carry slightly increased Ag and Pb. The chemical composition of bukovite from Příbram (average of 17 analyses), based on 10 apfu, can be expressed by the empirical formula (Tl$_{1.95}$Pb$_{0.01}$)$_{1.96}$Cu$_{5.00}$Ag$_{0.04}$Fe$_{0.97}$Se$_{3.96}$S$_{0.05}$Te$_{0.01}$.$\Sigma$4.02.
4.2.7. Bytízite, Cu₃SbSe₃

Bytízite is a rare new mineral described recently from one sample from a dump of shaft 16 by Škácha et al. [47]. Bytízite forms anhedral grains up to 40 μm (in the section), growing together in aggregates up to 300 μm across in the calcite gangue. Chaméanite, bukovite, příbramite, eskebornite, giraudite, umangite, crookesite, hakite, and tetrahedrite were found in the association.

Bytízite is steel-grey in colour and has a metallic lustre. In reflected light, bytízite is grey with a yellowish hue and weak bireflectance. Pleochroism is weak. Anisotropy under crossed polarisers is strong (Figure 19), with grey to brownish rotation tints.

![Figure 19](image)

**Figure 19.** Light grey bytízite (By) forms a rim around brown chaméanite (Ch) with tiny light brown eskebornite inclusions. Dark grey giraudite (G) is in the association. Reflected light, (A) one polarizer, and (B) partly-crossed polarizers. FOV 210 μm. Příbram, a dump of shaft 16. Photo by P. Škácha.

Bytízite is steel-grey in color and has a metallic lustre. In reflected light, bytízite is grey with a yellowish hue and weak bireflectance. Pleochroism is weak. Anisotropy under crossed polarisers is strong (Figure 19), with grey to brownish rotation tints.

X-ray powder diffraction data for Příbram bytízite are given by Škácha et al. [47], including refined unit-cell parameters: a = 7.94(1), b = 10.55(3), c = 6.84(1) Å, V = 573(1) Å³. Based on the single-crystal X-ray data, Škácha et al. [47] solved the crystal structure of bytízite (R_{int} = 0.0437). Based on the data (590 reflections), refined unit-cell data were obtained using the CrysAlis Pro program [60]: a = 7.959(1), b = 10.583(1), c = 6.824(1) Å, with V = 574.8(1) Å³, Z = 4.

The chemical composition of bytízite (Table S12) corresponds very well with the formula Cu₃SbSe₃ derived from the crystal-structure study. Only weak Sb-As and S-Se substitution was observed; a similar trend was found in other Cu-Sb selenides from Příbram: permingeatite [52] and příbramite [47]. The empirical formula, based on electron-microprobe analyses (n = 68), is (Cu₃.00Fe₀.01Ag₀.01)Σ₁.03(Sb₀.97As₀.03)Σ₁.03(Se₂.94S₀.01)Σ₂.95.

4.2.8. Cadmoselite, CdSe and Greenockite, CdS

Cadmoselite is a very rare mineral identified in two polished sections prepared from one sample. In the first case, it forms anhedral to euhedral grains up to 10 μm in size in later clausthalite (Figure 20); spheroidal aggregates of uraninite are also present. In the second case, cadmoselite is intergrown with Cd-hakite and clausthalite (Figure 21) as euhedral grains up to 40 μm or as thin tabular crystals in the central part of Cd-hakite aggregates. The euhedral forms of cadmoselite and cavities in the Cd-hakite indicate that cadmoselite is a later mineral replacing earlier Cd-hakite.
In reflected light, cadmoselite is dark grey and has a rather weak anisotropy.

The cation position in the studies of cadmoselite shows a minor substitution by Hg \((0.01 \text{ apfu})\) and Cu \((0.01−0.02 \text{ apfu})\) (Table S13). The minor contents of S in the anion position do not exceed \(0.01 \text{ apfu}\). Its chemical composition is very similar to type samples from the Ust’ Uyok deposit, Russia [84]. The empirical formula (average of four spot analyses based on 2 \text{ apfu}\) is \((\text{Cd}_{0.99}\text{Hg}_{0.01}\text{Cu}_{0.01})_{\Sigma1.01}(\text{Se}_{0.98}\text{S}_{0.01})_{\Sigma0.99}\). Cadmoselite from the Tumiñico deposit [79] contains Hg up to 0.05 \text{ apfu}. The limited solid solution observed of Hg for Cd is consistent with the fact that cadmoselite and HgSe (tiemannite) have different crystal structures of, respectively, the wurtzite and sphalerite types. The diagram of Hg and Cd contents in cadmoselite, tiemannite, and greenockite are presented in Figure 22.
Cadmoselite is a very rare mineral. In addition to the type locality—a vanadium-uranium-selenium deposit in Ust’ Uyok, Siberia, Russia [84]—it is reported only from a single occurrence at the vein-type Tumiñico hydrothermal deposit by Paar et al. [79].

Greenockite occurs in the studied association as euhedral to subhedral grains up to 20 × 8 μm with euhedral chalcopyrite grains, cadmoselite, and uraninite. In reflected light, greenockite is grey and weakly anisotropic.

With regard to a minimal number of grains suitable for analysis, only two chemical analyses of greenockite were obtained; in addition to predominant Cd and S, also Hg (0.08 a 0.31 apfu) and minor Zn, Ag, and Tl were measured. The anion position shows a partial substitution Se\(\text{S}^{-1}\) with Se contents up to 0.06 apfu (Table S14). A diagram of Hg vs. Cd contents in cadmoselite, tiemannite, and greenockite are presented in Figure 22.

4.2.9. Chaméanite, (Cu,Fe)\(_4\)As(Se,S)\(_4\)

Chaméanite is a very rare selenide known only at several localities worldwide (Chaméane [77], Tumiñico [79], and Předbořice [41]). At Příbram, it was identified in a sample carrying Tl, Cu, Fe, and Sb selenides. Chaméanite forms up to 0.7 mm long subhedral grains with a very strong cleavage, in association with eskebornite, hakite, tetrahedrite, closer unidentified Tl selenides, příbramite, and bytízite (Figure 23).

In reflected light, isotropic chaméanite is grey to grey-blue with various color shades depending on the Cu/Fe ratio.

The average chemical composition of chaméanite from Příbram (Table S15) is close to the ideal formula (Cu,Fe)\(_4\)As(Se,S)\(_4\). The Cu/Fe ratio (apfu) is extremely variable between 4 and 356, due to the very low local Fe content (Figure 24). For the original Chaméane material, the Cu/Fe ratio is in the range of 6–13 [77], and from the Tumiñico locality, it is 15 [79]. The chaméanite from Příbram also contains minor Ag (0.01–0.13 apfu), Sb (0.00–0.28 apfu), Tl (up to 0.06 apfu), Hg (up to 0.05 apfu), and S (up to 0.08 apfu). The empirical formula of Příbram chaméanite (average of 100 analyses)—based on 9 apfu—is \((\text{Cu}_{3.63}\text{Fe}_{0.33}\text{Ag}_{0.03})\Sigma_{3.99}(\text{As}_{0.88}\text{Sb}_{0.08})\Sigma_{0.96}(\text{Se}_{4.03}\text{S}_{0.02})\Sigma_{4.05}\).
Clausthalite, PbSe

Clausthalite is the most abundant selenide observed at the deposit. It was described by Růžička [48] from the B4 vein and by Litochleb et al. [49] in a sample from Bytíz. Later on, clausthalite was found locally abundant in material from shafts 16 and 11A [50,51].

Clausthalite was observed in numerous mineral associations. At first, it was described with arsenic and arsenolamprite as irregular grains tens of μm long, in irregular intergrowth with arsenic [50]. Later on, its occurrence with cuprostibite was described Škácha et al. [51] as silvery-grey fine-grained and irregular aggregates up to 2 mm in dark calcite. In this mode of occurrence, it is soft, without distinct cleavage. It is white in reflected light, and it overgrows and replaces mainly an older chalcocite.

Clausthalite occurs at the deposit as fine-grained aggregates without visible cleavage, as well as perfectly cleavable aggregates up to 0.5 cm across in close association with botryoidal uraninite. It differs from megascopically-similar galena by the absence of associated sphalerite. Clausthalite is

Figure 23. Brown chaméanite aggregate with grey giraudite and light brown eskebornite inclusions in the association. Reflected light, one polarizer. FOV 210 μm. Příbram, a dump of shaft 16. Photo by P. Škácha.

Figure 24. Fe vs. Cu (apfu) graph for chaméanite from Příbram and other localities.

4.2.10. Clausthalite, PbSe

Clausthalite is the most abundant selenide observed at the deposit. It was described by Růžička [48] from the B4 vein and by Litochleb et al. [49] in a sample from Bytíz. Later on, clausthalite was found locally abundant in material from shafts 16 and 11A [50,51].

Clausthalite was observed in numerous mineral associations. At first, it was described with arsenic and arsenolamprite as irregular grains tens of μm long, in irregular intergrowth with arsenic [50]. Later on, its occurrence with cuprostibite was described Škácha et al. [51] as silvery-grey fine-grained and irregular aggregates up to 2 mm in dark calcite. In this mode of occurrence, it is soft, without distinct cleavage. It is white in reflected light, and it overgrows and replaces mainly an older chalcocite.

Clausthalite occurs at the deposit as fine-grained aggregates without visible cleavage, as well as perfectly cleavable aggregates up to 0.5 cm across in close association with botryoidal uraninite. It differs from megascopically-similar galena by the absence of associated sphalerite. Clausthalite is
integrown with the majority of other selenides and sulphides (Figure 25). It forms euhedral crystals up to 50 μm in size, rarely in association with inhomogeneous Cu-Sb oxycompound(s), eucairite, and other minerals (Figure 26). Like tiemannite, clausthalite belongs among the earlier selenides, but both minerals occur together only rarely. In that case, clausthalite grains were observed in tiemannite aggregates; thus, clausthalite is probably earlier than tiemannite. It is probable that clausthalite occurs at the locality in more than a single generation.

![Figure 25](image1.png)

**Figure 25.** Light clausthalite in form of inclusions in dark chalcopyrite. BSE photo. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.

![Figure 26](image2.png)

**Figure 26.** Euhedral clausthalite grains (white) in non-homogenous Cu-Sb oxide (dark grey). BSE photo. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.

In reflected light, clausthalite is white and isotropic.

We have observed only low contents of Ag, Tl, Zn, Sb, As, and Te, not exceeding 0.05 apfu during the study of the chemical composition of clausthalite (Table S16). Somewhat higher are minor Cu contents up to 0.12 apfu. The SeS$_{-1}$ substitution is limited in most cases; S contents up to 0.32 apfu were observed only sporadically. The limited range of SeS$_{-1}$ substitution makes the Příbram clausthalite distinctly different from members of the clausthalite-galena series from localities of the West Bohemian ore district [85], Niederschlema-Alberoda [18], Běstvina [25], or Moldava localities [26]. The empirical formula of the studied Příbram clausthalite (average of 58 analyses) based on 2 apfu is (Pb$_{0.98}$Cu$_{0.02}$)$_{2.00}$(Se$_{0.97}$S$_{0.03}$)$_{2.00}$.
4.2.11. Crookesite, Cu$_7$(Tl,Ag)Se$_4$

Crookesite has been found in only one case, as several irregular inclusions up to 10 μm. It is closely associated with berzelianite and an unnamed Cu–As selenide (Figure 27) in the specimen with bytizite and chaméanite.

![Figure 27](image-url)

**Figure 27.** Anhedral brownish grey crookesite (Cr) grain embedded in berzelianite (Bz) in association with unnamed Cu–As selenide (Un). Reflected light, one polarizer. FOV 105 μm. Příbram, a dump of shaft 16. Photo by P. Škácha.

In reflected light, it is brownish grey and is weakly anisotropic.

The chemical composition of crookesite from Příbram (Table S17) is close to the ideal formula Cu$_7$(Tl,Ag)Se$_4$; it contains only minor amounts of S (0.02 apfu) and Ag (0.01 apfu). Its empirical formula (mean of two analyses) based on 12 apfu can be expressed as Cu$_{7.12}$(Tl$_{0.93}$Ag$_{0.01}$)$_{0.94}$(Se$_{3.92}$S$_{0.02}$)$_{23.94}$.

4.2.12. Dzharkenite-Pyrite Solid Solution, FeSe$_2$-FeS$_2$

Dzharkenite is a selenium analogue of pyrite that occurs as a rare product of the youngest stage of mineralization. It forms anhedral to subhedral grains up to 60 μm across in association with ferroselite and minerals of the tetrahedrite group antimonelite and příbramite (Figure 28).

![Figure 28](image-url)

**Figure 28.** Pinkish subhedral dzharkenite grain in association with dark grey tetrahedrite enclosed light grey tiemannite inclusions. Reflected light, one polarizer. FOV 210 μm. Příbram, a dump of shaft 16. Photo by P. Škácha.
In reflected light, dzharkenite is pink and isotropic. This makes it different from ferroselite, which is distinctly anisotropic in grey or brass yellow colors. Identification of dzharkenite and ferroselite was confirmed by EBSD (Figure 29).

The chemical composition of dzharkenite from Příbram is close to the ideal formula FeSe$_2$. Copper content is 0.07 apfu at maximum and S in the range 0.01–0.18 apfu (Table S18). Its empirical formula (average of 22 analyses) based on 3 apfu is (Fe$_{0.98}$Cu$_{0.02}$)$_2$(Se$_{1.92}$S$_{0.08}$)$_2$S$_{2.00}$.

Dzharkenite is a very rare mineral on a global scale. Ferroselite is the selenium analogue of marcasite, and is more common at hydrothermal uranium deposits [4,7,86]. Dzharkenite was first described from the Suluchekinskoye Se-U deposit in Kazakhstan by Yashunsky et al. [87]. Another occurrence was reported from the Zapadno-Ozernoe deposit of Cu-Zn massive sulphides in Russia [88]; members of the solid solution pyrite-dzharkenite are here of a supergene origin. Dzharkenite occurs as minute octahedra enclosed in calcite at the Cisco locality, Utah [89]. Litochleb et al. [49] reported Se-rich pyrite (0.21–0.29 apfu Se) from the Bytíz deposit as minute subhedral grains up to 40 × 80 μm enclosed in calcite and partly rimmed by coffinite. A comparison of variation in Se and S in the dzharkenite-pyrite solid solution from various occurrences is presented in Figure 30.

**Figure 29.** Electron backscatter diffraction (EBSD) images of (A) dzharkenite and (B) ferroselite (right) from Příbram, with indexed Kikuchi bands. Photo by M. Racek.

**Figure 30.** Se vs. S (apfu) graph of dzharkenite, ferroselite, and Se-pyrite from various localities.
4.2.13. Eskebornite-Chalcopyrite Solid Solution, CuFeSe$_2$-CuFeS$_2$

Minerals of the eskebornite-chalcopyrite isomorphic series are relatively common at the deposit. Eskebornite occurs only locally, but chalcopyrite is known from many places.

Eskebornite was observed as anhedral brown 1 mm grains in association with Cu-selenides and hakite. It also forms aggregates with abundant umangite and berzelianite grains (Figure 31) or inclusions in hakite aggregates (Figure 32). Eskebornite also replaces earlier uraninite spherical aggregates (Figure 33).

![Figure 31](image1)

**Figure 31.** Brown eskebornite with inclusions of violet umangite, blue berzelianite, and dark grey uraninite. Middle grey minor tetrahedrite is in the association. Reflected light, one polarizer. FOV is 210 μm. Příbram, dump at shaft 16. Photo by P. Škácha.

![Figure 32](image2)

**Figure 32.** Eskebornite grains (dark grey) in hakite aggregates (light grey). BSE photograph. Příbram, a dump of shaft 16. Photo by J. Sejkora, P. Škácha.

In reflected light, eskebornite is light brown with a pink shade. It is strongly anisotropic under crossed polarizers.

In the course of the study of the chemical composition of eskebornite (Table S19), aside from minor contents of Te, Ag, and Tl, two groups of analyses with different S contents were observed: 0–0.17 apfu and 0.57–0.79 apfu S. The empirical formulae for these eskebornite types from Příbram are the following:
During the study of the chemical composition of chalcopyrite, only local minor contents of Ag, Pb, Tl, and Sb were observed (Table S20). The Se content still never exceeds 0.21 apfu. Figure 34 shows the S and Se contents of chalcopyrite-eskebornite solid solutions from various localities, and suggests the presence of a possible solvus gap.


Eskebornite is a relatively abundant selenide in hydrothermal uranium deposits. In the Czech Republic, it was reported from Slavkovice [90], Habři [91], and Petrovice deposits [92].

Chalcopyrite is one of the most common sulphides present in the carbonate-uraninite stage, in association with tetrahedrite, uraninite, and selenides. It forms typical golden yellow elongated aggregates up to several centimeters long. Chalcopyrite in the aggregates is mainly anhedral with some grains tending towards euhedral. A study of polished sections shows that chalcopyrite is later than tetrahedrite, and belongs to the youngest minerals of the whole mineral association (Figure 32).

In reflected light, it is strongly yellow and weakly anisotropic.

During the study of the chemical composition of chalcopyrite, only local minor contents of Ag, Pb, Tl, and Sb were observed (Table S20). The Se substitution is more significant, but the Se content still never exceeds 0.21 apfu. Figure 34 shows the S and Se contents of chalcopyrite-eskebornite solid solutions from various localities, and suggests the presence of a possible solvus gap.

**Figure 33.** Eskebornite aggregates (dark grey) partly replace earlier spherical aggregates of uraninite (white). BSE photograph. Příbram, a dump of shaft 16. Photo by J. Sejkora, P. Škácha.

**Figure 34.** Se vs. S (apfu) graph of chalcopyrite and eskebornite from various localities.
4.2.14. Eucairite, AgCuSe

Eucairite in studied samples forms abundant anhedral aggregates up to 0.5 mm, which most frequently occur in association with berzelianite and umangite (Figure 35). It is earlier than Cu-selenides, which replace eucairite (Figure 36). The eucairite inclusions in clausthalite were observed only rarely.

![Figure 36. Berzelianite (grey) replaces earlier eucairite aggregates (white). BSE photograph. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.](image)

In reflected light, it has a cream-yellow color with a pink shade. It is strongly anisotropic from blue-green to blue-grey colors.

The X-ray powder diffraction data of eucairite from Příbram (Table S21) agree very well with data calculated from the crystal structure information [95]. The refined unit-cell parameters of eucairite from Příbram are compared with the published data in Table S22.

During the chemical study of eucairite from Příbram, minor contents of Tl, As, and Te in addition to the dominant elements (Table S23) were determined. The empirical formula of eucairite from Příbram (the average of 48 analyses) based on 3 apfu is $\text{Ag}_{1.00}\text{Cu}_{1.04} (\text{Se}_{0.95}\text{Te}_{0.01})_{0.01}$.  

An interesting phase related to eucairite was observed in a single polished section. It forms rather inhomogeneous aggregates up to 200 μm, with penetrating naumannite lamellae. Both of the minerals
probably represent products of disintegration of an unknown Ag-Cu-Se mineral that is stable at higher temperatures. This mineral (after recalculation to 3 apfu) shows a strong variation in contents of Ag (1.18–1.63 apfu), Cu (0.31–0.63 apfu), and Se (1.05–1.18 apfu). Minor Hg contents (to 0.01 apfu), Te (to 0.003 apfu), and S (to 0.01 apfu) were also observed in this mineral phase (Table S24).

4.2.15. Ferroselite, FeSe₂

Ferroselite occurs only rarely in studied samples. It forms idiomorph crystals up to 15 μm (Figure 37) and their aggregates not exceeding 50 μm in calcite gangue. Dzharkenite, příbramite, antimonselite, and a new Hg-Cu-Sb-selenide were found in association.

![Figure 37. Pinkish ferroselite anhedral grain with very intensive anisotropism (A,B). Reflected light, partly-crossed polarizers. FOV 210 μm. Příbram, a dump of shaft 16. Photo by P. Škácha.](image)

In reflected light, it has a cream-yellow color with a pink shade. It is strongly anisotropic from blue-green to blue-grey. Identification of ferroselite and dzharkenite was confirmed by EBSD (Figure 29).

The chemical composition of ferroselite does not significantly differ from dzharkenite, and is close to the ideal formula FeSe₂. Copper contents up to 0.02 apfu and S contents in the range 0.01–0.09 apfu were determined (Table S25). The empirical formula of ferroselite from Příbram (the mean of 12 analyses) based on 3 apfu is (Fe1.00Cu0.01)Σ1.01(Se1.94S0.05)Σ1.99.

4.2.16. Klockmannite, CuSe

Klockmannite is very rare in the studied material. It was observed in several samples as anhedral grains up to 30 μm across composed of very thin lamellar crystals intergrown with berzelianite in association with uraninite and tiemannite (Figure 38).

It is blue in reflected light, is anisotropic, and it shows a strong bireflection.

During the study of the chemical composition of klockmannite from Příbram (Table S26), increased Ag contents (up to 0.03 apfu) were observed; the cation/anion ratio varies in the range 0.90–1.05, probably due to intergrowth with other Cu-selenides on a sub-microscopic scale. In the anion site, dominant Se is substituted by Te and S in a minor range. The empirical formula of klockmannite (average of nine analyses) on the basis of 2 apfu is (Cu0.98Ag0.02)Σ1.00(Se0.98S0.03)Σ1.01. Variation of Se vs. S in klockmannite from several localities in the Czech Republic is presented in Figure 39.
In reflected light, it has a cream-yellow color with a pink shade. It is strongly anisotropic from blue-green to blue-grey. Identification of ferroselite and dzharkenite was confirmed by EBSD (Figure 29).

The chemical composition of ferroselite does not significantly differ from dzharkenite, and is close to the ideal formula FeSe$_2$. Copper contents up to 0.02 apfu and S contents in the range 0.01–0.09 apfu were determined (Table S25). The empirical formula of ferroselite from Příbram (the mean of 12 analyses) based on 3 apfu is \((\text{Fe}^{1.00}\text{Cu}^{0.01})\Sigma^{1.01}(\text{Se}^{1.94}\text{S}^{0.05})\Sigma^{1.99}\).

4.2.16. Klockmannite, CuSe

Klockmannite is very rare in the studied material. It was observed in several samples as anhedral grains up to 30 $\mu$m across composed of very thin lamellar crystals intergrown with berzelianite in association with uraninite and tiemannite (Figure 38).

Figure 38. A euhedral tiemannite crystal (T) occurs in the cavity of uraninite aggregates (U), in the association are Cu-selenides (dark grey) including klockmannite (Kl) and berzelianite (Bz). BSE photograph. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.

It is blue in reflected light, is anisotropic, and it shows a strong bireflection. During the study of the chemical composition of klockmannite from Příbram (Table S26), increased Ag contents (up to 0.03 apfu) were observed; the cation/anion ratio varies in the range 0.90–1.05, probably due to intergrowth with other Cu-selenides on a sub-microscopic scale. In the anion site, dominant Se is substituted by Te and S in a minor range. The empirical formula of klockmannite (average of nine analyses) on the basis of 2 apfu is \((\text{Cu}^{0.98}\text{Ag}^{0.02})\Sigma^{1.00}(\text{Se}^{0.98}\text{S}^{0.03})\Sigma^{1.01}\).

Variation of Se vs. S in klockmannite from several localities in the Czech Republic is presented in Figure 39.

In the sample from Příbram described by Litochleb et al. [49], klockmannite is the predominant phase in compound grains and aggregates. The size of anhedral grains is up to 70 $\times$ 150 $\mu$m, and rarely 1 mm. In places, the aggregates enclose minute pieces of earlier umangite and athabascaite. The observed sulphur contents are up to 0.7 apfu [49].

4.2.17. Naumannite, Ag$_2$Se

Naumannite was only observed in a limited number of polished sections; however, it was relatively abundant in some cases. It forms anhedral grains up to 40 $\mu$m in association with tiemannite and inhomogenous aggregates of AgCuSe minerals (Figures 40 and 41).
In the sample from Příbram described by Litochleb et al. [49], klockmannite is the predominant phase in compound grains and aggregates. The size of anhedral grains is up to 70 × 150 μm, and rarely 1 mm. In places, the aggregates enclose minute pieces of earlier umangite and athabascaite. The observed sulphur contents are up to 0.7 apfu [49].

4.2.17. Naumannite, Ag₂Se

Naumannite was only observed in a limited number of polished sections; however, it was relatively abundant in some cases. It forms anhedral grains up to 40 μm in association with tiemannite and inhomogenous aggregates of AgCuSe minerals (Figures 40 and 41).

![Figure 40. Inhomogeneous aggregates of Ag-Cu-Se phase close to eucairite with naumannite lamellae (light grey). BSE photograph. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.](image1)

In reflected light, naumannite is grey and weakly anisotropic. The chemical composition of naumannite from Příbram is usually relatively simple; minor contents of Cd, Hg, As, and Te (Table S27) were observed locally. Naumannite aggregates with increased Cu contents up to 0.36 apfu were observed in rare cases (Figure 42). The empirical formula of naumannite (from the mean of 30 analyses) calculated on the basis of 3 apfu is \((\text{Ag}_{1.96}\text{Cu}_{0.06})\Sigma_{2.02}\text{Se}_{0.97}\).
4.2.18. Permingeatite, Cu$_3$SbSe$_4$

The occurrence of permingeatite in the studied material was described in detail by Škácha et al. [52]. The mineral occurs as anhedral grains and aggregates, in the rare case up to 1 mm across in brownish or black calcite, in association with chalcopyrite, hematite, příbramite, antimonselite, dzharkenite, ferroselite, uraninite, löllingite, clausalhalite, hakite, berzelianite, umangite, and eskebornite. A narrow clausalhalite rim was developed around some permingeatite aggregates (Figure 43). In the studied assemblage, uraninite is locally abundant as typical colloform aggregates with a strong semi-metallic lustre. Permingeatite grains show megascopic bronze-brown color with a yellow shade and a strong metallic lustre. Permingeatite is brownish pink in reflected light and its anisotropy is strong. In comparison with the data of Johan et al. [40] for material from Předbořice, the measured reflectance values for permingeatite from Příbram [52] are different, probably due to the use of different standards.

Figure 42. Ag vs. Cu (apfu) graph for naumannite.

Figure 43. Permingeatite anhedral grains (grey) rimmed by clausalhalite (white). BSE photograph. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.
WDS data for permingeatite published by Škácha et al. [52] indicate a partial Se-S substitution corresponding to the solid solution of permingeatite (Cu$_3$SbSe$_4$)-famatinite (Cu$_3$SbS$_4$). In addition to contents of Cu, Sb, Se, and S, only minor Fe contents were documented, which imperfectly correlate with Cu. The empirical formula of the studied S-rich permingeatite (an average of 10 spot analyses) on the basis of 8 $\text{apfu}$ is (Cu$_{3.01}$Fe$_{0.02}$)$_{\Sigma}$3.03(Sb$_{1.00}$As$_{0.01}$)$_{\Sigma}$1.01(Se$_{3.74}$S$_{0.26}$)$_{\Sigma}$3.95. During the study of other samples collected after the Škácha et al. [52] publication, permingeatite aggregates (Table S28) containing—in addition to S—small amounts of As (up to 0.08 $\text{apfu}$), Ag (to 0.05 $\text{apfu}$), Hg (to 0.01 $\text{apfu}$), and Tl (to 0.01 $\text{apfu}$) were detected. The empirical formula of permingeatite (the mean of 41 analyses on the basis of 8 $\text{apfu}$) can be expressed as (Cu$_{3.00}$Fe$_{0.04}$)$_{\Sigma}$3.03Sb$_{0.99}$As$_{0.01}$Se$_{3.74}$S$_{0.26}$)$_{\Sigma}$3.97. The Se vs. S diagram (Figure 44) shows a correlation with a slope of $-1$, confirming partial S-Se substitution.

![Figure 44. Se vs. S (apfu) graph for permingeatite.](image-url)

X-ray powder diffraction data for permingeatite were presented by Škácha et al. [52]. The refined lattice parameters $a = 5.6323(2)$ Å, $c = 11.2354(7)$ Å, $V = 356.41(2)$ Å$^3$ (for the space group $I4_2m$) are in good agreement with the data of Johan et al. [40] for permingeatite from the type locality Příbram, and are lower than values for a synthetic analogue of permingeatite [94]. The published data indicate that increasing contents of As and S result in lowering the volume of the unit-cell.

### 4.2.19. Příbramite-Chalcostibite Solid Solution CuSbSe$_2$-CuSbS$_2$

The occurrence of minerals of the příbramite-chalcostibite solid solution series in the Příbram uranium ore district was first mentioned in the study of antimonelite of Škácha et al. [53]. Later on, these minerals were identified in additional samples, which resulted in their detailed characterization.

The new mineral příbramite—a Se-analogue of chalcostibite—was approved in 2016 (2015-127). The paper by Škácha et al. [46] provides a detailed characteristic, including its crystal structure solution ($R_{\text{obs}} = 0.0480$; 342 unique reflections). Příbramite has been found at the dump at shaft 16 (Příbram-Háje) in two fragments of calcite gangue. In the first sample, příbramite forms locally-abundant polycrystalline aggregates up to 100 μm in size, consisting of minute lens-shaped crystals (Figure 45). The tiemannite, antimonelite, dzharkenite, hakite, ferroselite, and a new Hg-Cu-Sb selenide were identified in the association. The second sample contains anhedral příbramite grains up to 15 μm across, intergrown with bytizite in association with chaméanite, giraudite, and bukovite.
In reflected light, příbramite is grey with a yellow shade, slightly lighter than Hg-rich hakite; it is distinctly anisotropic.

Study of příbramite chemical composition (Table S29) resulted in the observation of SbAs$_{-1}$ substitution (up to 0.07 $apfu$ As) and SeS$_{-1}$ substitution (0.09–0.51 $apfu$ S). The diagram in Figure 46 indicates the range of probable příbramite-chalcostibite solid solution as supported by the present data. Copper dominates the cation position, but minor contents of Ag, Fe, Pb, Tl, and Hg (up to 0.0X $apfu$) were found locally. The empirical formula of příbramite (the mean of 53 analyses on the basis of 4 $apfu$) can be expressed as Cu$_{1.00}$(Sb$_{1.02}$As$_{0.01}$)$_{\Sigma 1.03}$(Se$_{1.81}$S$_{0.15}$)$_{\Sigma 1.96}$.

![Figure 45. Prismatic příbramite crystals embedded in calcite. Reflected light, (A) one polarizer, (B) partly-crossed polarizers. FOV 210 µm. Příbram, a dump of shaft 16. Photo by P. Škácha.](image)

Figure 45. Prismatic příbramite crystals embedded in calcite. Reflected light, (A) one polarizer, (B) partly-crossed polarizers. FOV 210 µm. Příbram, a dump of shaft 16. Photo by P. Škácha.

![Figure 46. Se vs. S ($apfu$) graph for příbramite and Se-rich chalcostibite from Příbram and China.](image)

Figure 46. Se vs. S ($apfu$) graph for příbramite and Se-rich chalcostibite from Příbram and China.

An unnamed mineral from the U-Hg-Mo deposit in the Guizhou region, China that was observed as microscopic grains in carbonate veins carrying U-Se mineralization [95] probably corresponds to příbramite because of similar chemistry and reflected light properties.

Se-containing chalcostibite forms abundant lens-shaped-to-tabular crystals up to 10 µm and their aggregates several mm across (Figure 47) in calcite gangue colored black by microscopic inclusions of sulphides and selenides. Antimonselite, luzonite, tennantite, and other minerals are present in the association. The megascopic aggregates have a pink shade and are very brittle. It is grey and strongly anisotropic in reflected light.

![Figure 47. Lenticular euhedral grains of Se-rich chalcostibite (light grey), hakite (medium), and uraninite (dark grey) aggregates in calcite gangue. Reflected light, (A) one polarizer, (B) partly-crossed polarizers. FOV 210 µm. Příbram, a dump of shaft 16. Photo by P. Škácha.](image)
Backscattered electron images of chalcostibite show distinct zoning (Figure 48), due especially to a wide S-Se\textsubscript{1−1} substitution; the analysed Se contents range from 0.07 to 0.99 \textit{apfu} (Figure 46). In the cation position, dominant Cu (Table S30) is accompanied by minor Ag, Fe, and Pb (up to 0.0X \textit{apfu}); Sb shows a minor substitution (up to 0.01 \textit{apfu}) by As. The empirical formula of Se-chalcostibite (the mean of 72 analyses on the basis of 4 \textit{apfu}) can be expressed as Cu\textsubscript{1.00}(Sb\textsubscript{0.99}As\textsubscript{0.02})\textsubscript{Σ1.01}(Se\textsubscript{1.29}S\textsubscript{0.70})\textsubscript{Σ1.99}.

Figure 48. Zonation of Se-rich chalcostibite euhedral aggregates caused by variable Se-S content. BSE photo. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.

4.2.20. Sabatierite, Cu\textsubscript{6}TlSe\textsubscript{4}

Sabatierite with the idealized formula Cu\textsubscript{6}TlSe\textsubscript{4} was first described by Johan et al. [33] from the Bukov deposit. Based on a study of powder diffraction data, Berger [96] proposed a different formula (Cu\textsubscript{4}TlSe\textsubscript{3}), but his assumption is not confirmed by chemical study. The formula and crystal structure of sabatierite are not yet known.

In the studied material from Příbram, sabatierite was observed as a single anhedral grain 30 \textmu m in size, intergrown with bukovite (Figure 49) in association with berzelianite and eskebornite. In reflected light, sabatierite is creamy white with a pinkish shade.

Figure 49. Lenticular euhedral grains of Se-rich chalcostibite (light grey), hakite (medium), and uraninite (dark grey) aggregates in calcite gangue. Reflected light, (A) one polarizer, and (B) partly-crossed polarizers. FOV 210 \mu m. Příbram, a dump of shaft 16. Photo by P. Škácha.
Figure 49. Brown-grey sabatierite is intergrown with dark grey bukovite. Reflected light, one polarizer. FOV 90 μm. Příbram, a dump of the shaft 16. Photo by P. Škácha.

Study of the chemical composition of the mineral (Table S31) shows that it contains dominant Tl, Cu, and Se, and minor Ag (0.01–0.39 apfu), Fe (0.01–0.03 apfu), and S (0.01–0.03 apfu). The empirical formula (the mean of four analyses) on the basis of 11 apfu is (Cu_{6.16}Ag_{0.12})Σ_{0.94}(Tl_{0.92}Fe_{0.02})Σ_{0.94}(Se_{3.67}S_{0.12})Σ_{3.70}; this corresponds to stoichiometry proposed by Johan et al. [33]. The elevated Ag content in the studied sample is interesting. One of the spot analyses (Table S32) indicates a possible solid solution approaching an unnamed Ag-Cu-Tl selenide described by Sejkora et al. [15] from the Zálesí deposit. The diagram of Ag vs. Cu contents (Figure 50) for both phases from Příbram and Zálesí exhibit a negative correlation of these elements.

Figure 50. Ag vs. Cu (apfu) graph for sabatierite from Příbram and an unnamed Ag-Cu-Tl-Se phase from Zálesí.

4.2.21. Tetrahedrite Group

The tetrahedrite group is the most complex isotypic series among the sulphosalts in nature. It is caused by the multiplicity of iso- and heterovalent substitutions [97]. A simplified general formula for minerals of the tetrahedrite group can be expressed according to Moëlo et al. [97] as 

\[ \text{[III]} A_{n}^{[IV]}(B,C)_{6}^{[III]}X_{4}^{[IV]}Y_{12}^{[VI]}Z_{1}, \]

where \( A \) is Cu or Ag in triangular coordination; \( B \) is Cu\(^{1+} \) or Ag in...
tetrahedral coordination, C is generally a divalent metal (typically Fe or Zn, but also Hg, Mn, Cd, Cu$^{2+}$, etc.) in the same coordination as B; X is Sb, As, Bi, or Te in trigonal pyramidal coordination; Y is S or Se in tetrahedral coordination, and Z is S or Se in special octahedral coordination. Structural studies confirmed the presence of vacancies or interstitial atoms (e.g. Cu), or heterovalent substitutions due to the incorporation of Fe$^{3+}$ or Te$^{4+}$.

The classification of the tetrahedrite group of minerals is in need of review and updating, but this has not been done to date [97,98]. Currently, the divalent substituents C in tetrahedral coordination are not generally distinguished in the IMA-approved definitions of the minerals, with the exception of Hg-dominant hakite, although varieties with different dominant C cations are distinguished in the literature [54]. However, the dominant C cation in these minerals may be Zn, Fe, Hg, Cu, or Cd, and here we distinguish these cases using the dominant-constituent rule [99]. Hakite has been originally described as an Hg variety, but for the matter of classification, Škácha et al. [54] propose the application of the dominant rule (on the atomic basis) for solid-solutions of the members of the tetrahedrite group. That is why we distinguish Zn-, Fe-, Hg-, Cu-, and Cd- members in the following text.

4.2.21.1. Giraudite-Tennantite Solid Solution, Cu$_6$[(Cu$_4$(Fe,Zn)$_2$)$_6$As$_3$Se$_3$]$_2$-Cu$_6$[(Cu$_4$(Fe,Zn)$_2$)$_6$As$_3$]$_2$S$_3$

Minerals of the giraudite-tennantite solid solution were rare, they were found in only some samples. Cu-giraudite forms rims around hakite, and it is intergrown with berzelianite (Figure 51) or its aggregates up to 400 μm long are intergrown with chaméane in association with pribramite, bytizite, and Tl-selenides. In another sample, anhedral Hg-giraudite grain about 30 μm across is replaced by later Cu-selenide (Figure 52); this is a unique case at the locality that a mineral of the tetrahedrite group is older than the Cu-selenides. Both giraudite types are of a light grey color with a brownish shade in reflected light.

![Figure 51. The core of aggregate formed by hakite (light grey) is rimmed by giraudite (grey); berzelianite (dark grey) is the youngest mineral in the association. BSE photo. Pribram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.](image)

In its chemical composition, giraudite shows typical substitutions of the tetrahedrite group. Two compositional types were identified: Cu- and Hg-giraudite (Table S32). Cu-giraudite (22 analyses) also contains Ag in the A position (up to 0.23 apfu) in addition to Cu; the B position is fully occupied by Cu. The C position is dominated by Cu (1.6–2.39 apfu), with a small Hg content (up to 0.83 apfu), Fe (up to 1.40 apfu), Zn (up to 0.11 apfu), and Cd (up to 0.03 apfu). Arsenic in the position X is substituted by Sb (up to 0.71 apfu), and locally, Te (up to 0.43 apfu). The highest S contents in the anion positions do not exceed 1.38 apfu. The empirical formula of Cu-giraudite based on 29 apfu is (Cu$_{5.82}$Ag$_{0.18}$)$_{0.00}$Cu$_{4.00}$[Cu$_{7.66}$Fe$_{0.34}$Hg$_{0.07}$Zn$_{0.03}$]$_{1.00}$As$_{12.87}$Sb$_{0.05}$Se$_{12.87}$. 

![Figure 52.](image)
Fe-tetrahedrite is fully occupied by Cu. The B position is fully occupied by Cu. The C position is filled mainly by Hg (1.3 apfu), minor Cd (0.2 apfu), and Zn (0.12 apfu). In the X position, in addition to arsenic, is a low Sb content (0.19–0.44 apfu). Sulphur contents in anion positions vary around 1.6 apfu. The empirical formula of Hg-giraudite based on 29 apfu is (Cu₅.₇₈Ag₀.₂₂)Σ₆.₀₀Cu₄.₀₀(Hg₁.₂₉Cu₀.₄₁Cd₀.₂₂Zn₀.₁₂)Σ₂.₀₄(As₃.₅₅Sb₀.₃₁)Σ₂.₃₈(Se₁₁.₄₈S₁.₆₁)Σ₁₃.₉₉.

Giraudite is a very rare member of the tetrahedrite group. It was reported from the hydrothermal Chaméane deposits [77] and Alberoda [19]. It is also mentioned from some epithermal deposits in Mexico and USA. Figure 53 shows an obvious negative correlation between As and Sb for various minerals of the tetrahedrite group, including giraudite from Příbram.

![Figure 52](image-url) Yellowish-pink giraudite anhedral grain replacing by blue-green berzelianite which is partly replaced by violet umangite. Reflected light, one polarizer. FOV 210 μm. Příbram, a dump of shaft 16. Photo by P. Škácha.

In the A position, Hg-giraudite (two analyses) contains Ag (0.22–0.23 apfu), in addition to Cu. The B position is fully occupied by Cu. The C position is filled mainly by Hg (1.3 apfu), minor Cd (0.2 apfu), and Zn (0.12 apfu). In the X position, in addition to arsenic, is a low Sb content (0.19–0.44 apfu). Sulphur contents in anion positions vary around 1.6 apfu. The empirical formula of Hg-giraudite based on 29 apfu is (Cu₅.₇₈Ag₀.₂₂)Σ₆.₀₀Cu₄.₀₀(Hg₁.₂₉Cu₀.₄₁Cd₀.₂₂Zn₀.₁₂)Σ₂.₀₄(As₃.₅₅Sb₀.₃₁)Σ₂.₃₈(Se₁₁.₄₈S₁.₆₁)Σ₁₃.₉₉.

Giraudite is a very rare member of the tetrahedrite group. It was reported from the hydrothermal Chaméane deposits [77] and Alberoda [19]. It is also mentioned from some epithermal deposits in Mexico and USA. Figure 53 shows an obvious negative correlation between As and Sb for various minerals of the tetrahedrite group, including giraudite from Příbram.

![Figure 53](image-url) Sb vs. As (apfu) graph for tetrahedrite group of minerals from Příbram.

Fe-tennantite is very rare in the studied material. It forms thin irregular rims about 30 × 10 μm around tetrahedrite aggregate (Figure 54) in association with luzonite, antimonoselite, and Se-rich chalcostibite. In reflected light, tennantite is similar to tetrahedrite. It is grey, without notable anisotropy.
Hakite and its sulphur analogue tetrahedrite are relatively common in the studied samples. In contrast to tetrahedrite, hakite is more common in microscopic form, but locally, aggregates up to several mm in size were observed. At present, Škácha et al. [54] studied in detail hakite from Příbram. They distinguish three chemical types of hakite: Hg-, Zn-, and Cd-hakite. Most detailed data were obtained for the most common Hg-hakite; its refined unit-cell parameters (space group \( I-43m \)) agree very well with the published data for Hg-hakite from Předbořice [39]. With the use of precession electron diffraction tomography (PED), the Hg-hakite crystal structure was solved and its relationship to the tetrahedrite group was confirmed.

After extending the data set used by Škácha et al. [54] by analyses of the new samples, it is possible to define five chemical types of hakite, depending on filling the \( C \) position by the respective elements: Hg-, Zn-, Cd-, Fe-, and Cu-hakite.

**Hg-hakite** is the prevailing type, which forms relatively abundant allotriomorphic aggregates together with Zn-hakite up to several mm across (Figures 13 and 55). In one case, Hg-hakite formed an aggregate of well-formed idiomorphic crystals up to 100 \( \mu m \) [54]. Both of the hakite types occur especially in association with berzelianite, or rarely, with athabascaite. In all analysed Hg-hakite samples, Cu predominates at the \( A \) site. It contains up to 1.93 \( apfu \) Ag, which is more than that observed in other types of hakite (Figure 56). The range of Ag content in the Hg-hakite is 0.04–1.93 \( apfu \) (\( n = 122 \)). The \( B \) position is fully occupied by Cu in most cases. In the \( C \) position, Hg (0.72–1.98 \( apfu \))

![Figure 54. Dark grey Fe-tennantite (Tn) overgrowing grey Fe-tetrahedrite (Te) in association with pinkish luzonite (Lu), chalcopyrite (Cpy), and uraninite (U). Reflected light, one polarizer. FOV 210 \( \mu m \). Příbram, a dump of the shaft no. 16. Photo P. Škácha.](image)
is predominant above other elements (Zn up to 0.96 apfu; Cu and Cd up to 0.71 apfu, and Fe up to 0.59 apfu). The antimony contents vary in the range 2.03–4.20 apfu; less common is As (up to 2.02 apfu). The measured S contents are 0.01–4.69 apfu, and the average Se content is 11.67 apfu. On the basis of the mean of 122 EPM analyses (Table S34), the empirical formula is based on 29 apfu and can be expressed as (Cu5.65Ag0.35)Σ6.00Cu4.00(Hg1.72Zn0.16Fe0.07Cu0.05Cd0.05)Σ2.05(Sb3.89As0.27)Σ4.16(Se11.67S1.13)Σ12.80.

![Figure 55](image_url) Homogeneous Hg-hakite (light grey) in association with uraninite (white) and Fe-tetrahedrite (dark grey). BSE photo. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.

![Figure 56](image_url) Ag vs. Cu (apfu) graph for tetrahedrite group of minerals from Příbram.

**Zn-hakite** was observed as irregular aggregates up to several mm in size in association with Hg-hakite, clausthalite, bukovite (Figure 18), and berzelianite. It also occurs together with Hg-hakite and Zn-tetrahedrite as a part of zoned idiomorphic crystals up to 100 µm [54]. Minor Ag contents not exceeding 0.62 apfu were observed in the A-site during the study of its chemical composition (Table S35); the B-site is fully occupied by Cu, Hg (up to 0.80 apfu), Fe (up to 0.59 apfu), and Cd (up to 0.47 apfu), in addition to the dominant Zn (0.60–1.88 apfu), were observed in the C site. In the X position, the dominant Sb is accompanied by As (up to 0.71 apfu), and in the anion site, Se is partly substituted by S (0.07–4.52 apfu). The empirical Zn-hakite formula (the mean of 53 analyses) on the basis of 29 apfu is (Cu5.80Ag0.20)Σ6.00Cu4.00(Zn1.46Hg0.31Cu0.12Cd0.07)Σ1.96(Sb3.78As0.38)Σ4.16(Se11.94S0.91)Σ12.85.

**Cd-hakite** forms very rare allotriomorphic aggregates up to 50 µm in size, intergrown with clausthalite (Figure 21). Its aggregates are locally replaced by lath-shaped cadmoselite crystals; the
replacement starts from the center of aggregates. According to chemical analyses (Table S36), the A site in Cd-hakite contains minor amounts of Ag (0.06–0.59 apfu) in addition to the dominant Cu. The B site is fully occupied by Cu. In the C position, Cd in the range of 1.05–1.54 apfu is accompanied by Zn (up to 0.89 apfu) and Hg (up to 0.57 apfu). Sb shows a substitution by As (up to 0.37 apfu), and in the anion group, the dominant Se is locally accompanied by S contents in the range 0.67–4.54 apfu. The chemical composition of Cd-hakite (mean of 14 analyses) corresponds to the empirical formula (Cu5.60Ag0.14)Σ5.74Cu4.00(Cd1.37Hg0.45Zn0.15)Σ1.97(Sb3.99As0.32)Σ4.31(Se9.95S3.05)Σ13.00.

**Fe-hakite** has been found very rarely as allotriomorphic aggregates up to 300 × 30 µm in size, in association with chalcopyrite, Cu-hakite, chaléanite, giraudite, permingeatite, tetrahedrite, bytízite, and příbramite (Figure 57). During the study of the chemical composition (Table S37), dominant Cu contents were found in the A position accompanied by Ag in the range 0.01–0.29 apfu and, exceptionally, by Tl (up to 0.02 apfu). The B position is fully occupied by Cu, and in the C position, in addition to the dominant Fe (0.50–1.33 apfu), and mainly Zn (0.15–0.72 apfu), and Hg (0.00–0.52 apfu) occur. In the X position is Sb (2.61–3.86 apfu), partly replaced by As (0.36–1.37 apfu). Sulphur contents in the Y+Z positions are low (0.04–0.47 apfu). The empirical formula of Fe-hakite based on 29 apfu (the mean of 27 analyses) is (Cu5.87Ag0.13)Σ6.00Cu4.00(Fe0.92Cu0.08Zn0.45Hg0.21)Σ2.13(Sb3.63As0.47)Σ4.10(Se10.66S2.11)Σ12.77.

![Figure 57. The complicated aggregate of dark grey Fe-hakite (Fe-Hk), Cu-hakite (Cu-Hk) and Fe-tetrahedrite (Te), bytízite (Bt), and bukovite (Bu). BSE photo. Příbram, a dump of shaft 16. Photo by J. Sejkora and P. Škácha.](image)

**Cu-hakite** forms very rare, small allotriomorphic aggregates intergrown with bytízite in association with Cu-hakite, chaléanite, giraudite, bytízite, and příbramite (Figure 57). In its chemical composition (Table S38), the dominant Cu in the A position is accompanied by Ag (0.14–0.39) and, rarely, Tl (up to 0.015 apfu). The B position is fully occupied by Cu, and in the C position, minor Fe (0.06–0.82 apfu) and Zn (0.03–0.23 apfu) accompany the dominant Cu (0.67–1.44 apfu). In the X position, the dominant Sb (2.61–4.02 apfu) is—in part—replaced by As (0.36–1.37 apfu). Isomorphic substitution by S in the Y+Z positions is low (0.04–0.47 apfu). The empirical formula of Cu-hakite on the basis of 29 apfu (the mean of eight analyses) is (Cu5.72Ag0.28)Σ6.00Cu4.00(Cu1.02Hg0.38Fe0.45Zn0.19)Σ2.04(Sb3.10As0.11)Σ4.11(Se12.74S0.12)Σ12.86.

**Tetrahedrite** is abundant in the studied material. It is a common component of hypidiomorphic to idiomorphic aggregates several mm in size (Figure 57), sometimes intergrown with later chalcopyrite. Tetrahedrite was also observed as zoned granular aggregates with hakite (Figure 58) or as rims around chalcocite (Figure 59); in these aggregates, tetrahedrite usually forms the outer parts of grains. In reflected light, it is grey, isotropic, and slightly darker than hakite.
Two types of tetrahedrite were encountered in the course of the study of the chemical composition: Zn-tetrahedrite and Fe-tetrahedrite. Fe-tetrahedrite was observed in the association with antimonellite, Se-chalcostibite, Fe-tennantite, luzonite, chalcopyrite, or with chaméanite, bytízite, příbramite, giraudite, bukovite, crookesite, new As-Cu selenide, Fe-hakite, and chalcopyrite. Zn-tetrahedrite occurs as rims around chalcocite euhedral grains with clausthalite, in association with antimonellite, příbramite, new Hg-Cu-Sb selenide, tiemannite, and other minerals. Tetrahedrite in Příbram forms a continuous solid solution series with hakite (Figure 60) as previously reported by, e.g., Förster and Rhede [19] and Förster et al. [21].
A similar phase was already described as an unnamed mineral with a proposed formula, 

\[ \text{Cu}^{0.97-1.81} \text{apfu} \]

up to 5.21 \text{apfu}. The empirical formula of Zn-tetrahedrite (the mean of 15 analyses) on the basis of 29 \text{apfu} is

\[ (\text{Cu}^{5.99}\text{Ag}^{0.01})_{2.32}\text{Cu}_{4.00}(\text{Zn}^{1.42}\text{Fe}^{0.45}\text{Cd}^{0.02}\text{Hg}^{0.07}\text{Pb}^{0.02})\Sigma_{1.98}(\text{Sb}^{3.91}\text{As}^{0.11})\Sigma_{4.01}(\text{S}^{11.51}\text{Se}^{1.40})\Sigma_{12.91}. \]

The A and B positions of the crystal structure in Zn-tetrahedrite are dominantly occupied by Cu, but low Ag contents below 0.03 \text{apfu} occur in the A position (Table S39). The dominant Zn (0.97–1.81 \text{apfu}) is accompanied by Fe (0.06–0.99 \text{apfu}), Cu (up to 0.24 \text{apfu}), Hg (up to 0.32 \text{apfu}), Cd (up to 0.32 \text{apfu}), and Pb (up to 0.17 \text{apfu}) in the C position. In the X position is the dominant Sb, replaced in part by As in the range of 0.02–1.32 \text{apfu}. Sulphur is partly substituted by selenium (up to 4.27 \text{apfu}). The empirical formula of Zn-tetrahedrite (the mean of 15 analyses) on the basis of 29 \text{apfu} is

\[ (\text{Cu}^{5.99}\text{Ag}^{0.01})_{2.32}\text{Cu}_{4.00}(\text{Zn}^{1.42}\text{Fe}^{0.45}\text{Cd}^{0.02}\text{Hg}^{0.07}\text{Pb}^{0.02})\Sigma_{1.98}(\text{Sb}^{3.91}\text{As}^{0.11})\Sigma_{4.01}(\text{S}^{11.51}\text{Se}^{1.40})\Sigma_{12.91}. \]

The A and B positions in Fe-tetrahedrite are—like in Zn-tetrahedrite—occupied dominantly by Cu (Table S40), and Ag in the A position is below 0.03 \text{apfu}. The C position contains—in addition to the dominant Fe (up to 2.35 \text{apfu})—Zn (0.01–1.12 \text{apfu}), Pb (up to 0.16 \text{apfu}), Cd (up to 0.10 \text{apfu}), Cu (up to 0.73 \text{apfu}), and Hg (up to 0.14 \text{apfu}). Antimony is slightly substituted by As (up to 0.59 \text{apfu}) and S by Se (up to 5.21 \text{apfu}). The empirical formula of Fe-tetrahedrite (the mean of 27 analyses) on the basis of 29 \text{apfu} is

\[ (\text{Cu}^{5.99}\text{Ag}^{0.01})_{2.66}\text{Cu}_{4.00}(\text{Fe}^{1.22}\text{Zn}^{0.32}\text{C}^{0.47}\text{Pb}^{0.02}\text{Hg}^{0.01})\Sigma_{0.01}(\text{Sb}^{3.69}\text{As}^{0.31})\Sigma_{4.01}(\text{S}^{7.74}\text{Se}^{3.16})\Sigma_{12.90}. \]

Tetrahedrite is a very abundant and characteristic mineral, especially in the neighboring Březové Hory base-metal district [100]. Its occurrences in the uranium district are most frequently tied to the base-metal mineralization of the siderite-sulphide stage. In the calcite-sulphidic stage, it belongs to locally-common minerals, often associated with chalcopyrite.

4.2.21.3. Unnamed Hg-Cu-Sb Selenide from the Tetrahedrite Group

A probable new Hg-Cu-Sb selenide was discovered in a single sample with dzharkenite, ferroselite, příbramite, antimonelite, and Hg-hakite (Figure 61). This phase contains up to 30.73 wt % Hg (Table S41). A similar phase was already described as an unnamed mineral with a proposed formula 

\[ (\text{Cu}, \text{Hg})_{1-x}\text{Sb}_x(\text{Se},\text{S}), \]

from Guizhou Province, China, by Chen et al. [95].
The A and B positions in Fe-tetrahedrite are—like in Zn-tetrahedrite—occupied by Fe and Sb in almost equal amounts. The A and B positions in Fe-tetrahedrite are—like in Zn-tetrahedrite—occupied by Fe and Sb in almost equal amounts. An analogous shift has been reported as a consequence of S-Se substitution in Kharbish et al. [101]. Hooke’s Law [102] predicts significant shifts to lower wavenumbers for Sb-Se vibrations relative to Sb-S; this can be seen if the spectra of Figure 64A,B are compared. An analogous unnamed selenide from China [95] is isotropic with a white color and yellowish tint. The analogous unnamed selenide from China [95] is isotropic with a white color and yellowish tint. The relation of the crystal structure of the unnamed Hg-Cu-Sb selenide from Příbram to minerals of the tetrahedrite group was checked with EBSD, Raman spectroscopy, and single-crystal diffraction. A zoned aggregate of this mineral overgrown by Hg-hakite and tetrahedrite was selected for the EBSD study. The fit of all three sets of EBSD data in the studied aggregate not only supports them having

**Figure 61.** The core of the zonal aggregate is formed by new Hg-Cu-Sb selenide and Hg-hakite (indistinguishable from each other in the one microscope); the rim is grey tetrahedrite. Bluish-grey uraninite and white anhedral grains of tiemannite are the oldest phases in the association. Reflected light, one polarizer. FOV 210 μm. Příbram, a dump of shaft 16. Photo by P. Škácha.

The relation of the crystal structure of the unnamed Hg-Cu-Sb selenide from Příbram to minerals of the tetrahedrite group was checked with EBSD, Raman spectroscopy, and single-crystal diffraction. A zoned aggregate of this mineral overgrown by Hg-hakite and tetrahedrite was selected for the EBSD study. The fit of all three sets of EBSD data in the studied aggregate not only supports them having
the tetrahedrite type (Figure 63), but also the identical crystallographic orientation of all three phases. Raman spectra were very similar for the unnamed selenide and coexisting Hg-hakite (Figure 64a), the main differences being a band at 261 cm\(^{-1}\) which was observed only for hakite, strong splitting of the stretching modes at 210–230 cm\(^{-1}\), and a shift to a lower wavenumber of the lattice modes near 70 cm\(^{-1}\) for the unnamed selenide. Bands were assigned based on the data of Kharbish et al. [101]. Hooke's Law [102] predicts significant shifts to lower wavenumbers for Sb-Se vibrations relative to Sb-S; this can be seen if the spectra of Figure 64A,B are compared. An analogous shift has been reported as a consequence of S-Se substitution in permigateite-famatinite/luzonite [52]. Single-crystal X-ray diffraction experiments proved that the studied extracted grains of the unnamed Hg-Cu-Sb selenide are cubic, with \(a = 10.9636(12) \text{Å}\) and \(V = 1317.83(8) \text{Å}^3\), \(Z = 2\).

![Figure 62](image1.png) ![Figure 63](image2.png)

**Figure 63.** EBSD images of (A) new Hg-Cu-Sb selenide, (B) Hg-hakite, and (C) tetrahedrite from Příbram with indexed Kikuchi bands. These photos were taken from the zonal aggregate depicted in Figure 62. Photo by M. Racek.

![Figure 64](image3.png)

**Figure 64.** Raman spectra of (a) the new Hg-Cu-Sb selenide and Hg-hakite and (b) coexisting tetrahedrite. The vertical shift of spectra was done for clarity.

The interpretation of chemical composition of an unnamed Hg-Cu-Sb selenide is somewhat complicated; observed stoichiometry (especially low Cu content) indicates a presence of vacancies
at the A or B-C positions. That is why the composition of this phase was calculated on the basis \((\text{Sb+As}) = 4 \text{ apfu}\). In the A position of unnamed Hg-Cu-Sb selenide from Příbram, minor Ag (0.18–0.34 apfu) and occasionally Tl contents (up to 0.05 apfu) were found. The range of AgCu$_{1-1}$ substitution of the unnamed phase is distinctly lower than in the case of coexisting Hg-hakite (Figure 65). Copper at the A and B positions (4.79–7.17 Cu apfu) is partly substituted by Hg (Figure 66). In the available literature for minerals of the tetrahedrite group sensu stricto, including synthetic phases (e.g. [103–108]), Hg contents above 2 apfu connected with substitution Hg$^{2+}$-Cu$^{1+}$ were not observed, including tetrahedrites from Hg deposits [100]. On the other hand, the Hg$^{2+}$-Cu$^{1+}$ substitution was observed in some minerals structurally related to tetrahedrite (e.g., galkhaite, aktashite) [109], or related to polybasite-fettelite [110]. The observed Hg contents (2.73–4.09 apfu) positively correlate (Figure 67) with calculated vacancies (1.73–3.06 pfu); the presence of vacancies also provides a charge balance to the formula. In contrast to coexisting Hg-hakite, the contents of Zn (up to 0.14 apfu), Fe (up to 0.12 apfu), and Cd (up to 0.10 apfu) in the unnamed phase are significantly lower (Figure 68). In the X position, Sb is slightly substituted by As in the range 0.12–0.28 apfu. The determined range of S-Se$_{1-1}$ substitution in the unnamed Hg-Cu-Sb selenide is distinctly more limited (0.67–2.02 S) than in the coexisting Hg-hakite (Figure 69).

**Figure 65.** Ag vs. Cu (apfu) graph for unnamed Hg-Cu-Sb selenide and Hg-hakite.

**Figure 66.** Hg vs. Cu (apfu) graph for unnamed Hg-Cu-Sb selenide and Hg-hakite.
Minerals 2017, 7, 91

Figure 67. Hg (apfu) vs. vacancies pfu graph for unnamed Hg-Cu-Sb selenide and Hg-hakite; vacancies were calculated as 12-(Cu¹⁺Ag⁺Tl⁺Hg⁺Fe⁺Zn⁺Cd).

Figure 68. Hg vs. Fe + Zn + Cd (apfu) graph for unnamed Hg-Cu-Sb selenide and Hg-hakite.

Figure 69. Hg vs. S (apfu) graph for unnamed Hg-Cu-Sb selenide and Hg-hakite.
The empirical formula of a new Hg-Cu-Sb selenide (mean of 62 analyses) on the basis of (Sb + As) = 4 $\text{apfu}$ is $\text{Cu}_{5.83}\text{Ag}_{0.28}\square_{2.40}\text{Hg}_{3.43}\text{Fe}_{0.01}\text{Zn}_{0.03}(\text{Sb}_{3.88}\text{As}_{0.12}\text{Se}_{10.83}\text{S}_{1.29})_{2.12}$.2\text{Se}_{13}$ and the expected ideal formula is $\text{Cu}_6\text{Hg}_4\text{Sb}_4\text{Se}_{13}$.

4.2.21.4. Trends in the Chemical Composition of Minerals of the Tetrahedrite Group at Příbram

Minerals of the tetrahedrite group viewed from the perspective of element substitution represent some of the most complex phases. The complex compositional variations in S-dominant members of the tetrahedrite group have been studied by numerous authors (cf. [100,103,105–108]), and particularly by Velebil et al. [100]. In contrast, information on possible substitutions in Se-dominant members is rather limited. Papers by numerous authors [11,15,19,39,111–114] present related information.

In the frame of the study of minerals of the tetrahedrite group from Příbram were data obtained for co-existing Se- and S-dominant members of this group. Increased Ag contents (1–2 $\text{apfu}$) in the $A$ position were observed only in Hg-hakite; analysed contents in all remaining members of this group do not exceed 0.5–0.7 $\text{apfu}$ (Figure 56). Ag content is lower in the case of tetrahedrite; conversely, Ag content is higher in hakite—especially in Hg-hakite (Figure 70). In the $C$ position of tetrahedrite is a preference for Zn and Fe, in contrast to closely-associated hakite with a preference for Hg, Zn, and Cd. A larger ionic radius of selenium atoms with respect to sulphur or the local geochemical conditions may be responsible for the situation. The rare undersubstituted Fe-tetrahedrite was found in association with selenides from Příbram (Figure 71) [54]. The selenides contain larger voids in the structure, which may more easily host larger cations such as Hg, Ag, or Cd. A change in fluid chemistry over the time span of crystallization can be an additional factor [54].

**Figure 70.** Ag vs. Se ($\text{apfu}$) graph for tetrahedrite group of minerals from Příbram.
4.2.22. Tiemannite, HgSe

In the studied association, tiemannite is one of the relatively common minerals. It forms metallic grey anhedral grains up to 500 μm across in calcite gangue and is accompanied by other selenides. Tiemannite is intergrown with brodtkorbite, eucairite, and minerals of the tetrahedrite group (Figure 72) or příbramite. It is light grey in reflected light, and is isotropic.

The chemical composition of the studied tiemannite (Table S42) contains minor Ag (up to 0.03 apfu), Cd (up to 0.14 apfu), As (to 0.01 apfu), and Cu (to 0.05 apfu) in addition to the dominant Hg. In the anion position, besides dominant Se, S (up to 0.02 apfu) occurs locally. Its empirical formula (based on 93 analyses) on the basis of 2 apfu is \( \text{(Hg}_{0.98}\text{Cd}_{0.02}\text{Cu}_{0.01})_{\Sigma1.01}(\text{Se}_{0.96}\text{S}_{0.1})_{\Sigma2.99} \). An anomalous Te-rich tiemannite was found in a single sample (Table S43). In addition to elevated Te contents (0.10–0.24 apfu), minor Cu contents (up to 0.12 apfu) were also observed. The empirical formula of studied Te-tiemannite (the average of 38 analyses) on the basis of 2 apfu is \( \text{(Hg}_{0.95}\text{Cd}_{0.01}\text{Cu}_{0.05})_{\Sigma1.01}(\text{Se}_{0.76}\text{Te}_{0.18})_{\Sigma2.97} \).
Minor contents of Cu (up to 0.17 wt %) and Cd (up to 0.15 wt %) are reported for tiemannite from the Tumiñico deposit, Argentina [79]. The distinctly negative correlation of Se and Te (Figure 73) confirms the existence of at least a partial solid solution between isostructural tiemannite (HgSe) and coloradoite (HgTe). No information on significant contents of Te in tiemannite was found in the literature.

![Figure 73. Se vs. Te (in apfu) graph for tiemannite and Te-rich tiemannite from Příbram.](image)

4.2.23. Umangite, Cu₃Se₂

In the studied samples, umangite is relatively abundant as minute anhedral aggregates with a typically violet color. The size of the aggregates is rarely up to 3 mm. The mineral is frequently associated with berzelianite and eucairite, and less commonly with hakite, tiemannite, and brodtkorbite. S-rich umangite was found in association with chalcocite and S-rich berzelianite.

In reflected light, the mineral has a red-pink color (Figure 74); it is strongly anisotropic, with red-orange and yellow colors.

![Figure 74. Lamellae of violet umangite replace bluish berzelianite. White eucairite is in the association. Reflected light, (A) one polarizer; (B) crossed polarizers. FOV 210 µm. Příbram, a dump of the shaft 16. Photo by P. Škácha.](image)

The study of the chemical composition of umangite (Table S44) confirmed—in the cation position—minor contents of Ag (up to 0.28 apfu) and Fe (up to 0.20 apfu), in addition to the dominant
Cu. In the anion position, besides the dominant Se, there is some S (up to 0.36 apfu), and locally, Te (up to 0.01 apfu). The average empirical formula of the studied umangite from Příbram (the mean of 70 analyses) on the basis of 5 apfu is (Cu$_{3.00}$Ag$_{0.02}$Fe$_{0.02}$)$_{Σ3.04}$(Se$_{1.91}$S$_{0.05}$)$_{Σ1.96}$.

One sample contained S-rich umangite (Table S45). The dominant selenium in the anion position (Se 1.18–1.45 apfu) is significantly substituted by S (0.61–0.85 apfu). Elevated sulphur contents in umangite (Figure 75) were reported from the Tilkeroide uranium deposit, with values up to 13.5 wt % S by Wallis [115]. The chemical composition of S-rich umangite from Příbram (the mean of eight analyses) corresponds with the empirical formula (Cu$_{2.88}$Fe$_{0.14}$Ag$_{0.01}$)$_{Σ2.98}$(Se$_{1.25}$S$_{0.75}$)$_{Σ2.02}$. Te-rich umangite was found in the case of the only point analysed (Table S46). Its chemical composition is Cu$_{2.98}$(Se$_{1.86}$Te$_{0.10}$S$_{0.06}$)$_{Σ2.02}$.

![Figure 75. Se vs. S (apfu) graph for umangite from Příbram and other localities.](image)

Umangite is a relatively abundant selenide in hydrothermal uranium deposits. It occurs in association with other Cu selenides. It is often formed via disintegration of berzelianite, which changes to umangite at temperatures below 112 °C [116]. Litochleb et al. [49] identified umangite in a historical sample from Bytíz. In addition to the major elements, it contains minor contents of Pb and Hg.

4.2.24. Unnamed Cu-As Selenide, CuAsSe

Very tiny inclusions of an unnamed selenide with a maximum size of 15 μm, with chemistry corresponding to CuAsSe were found, showing extremely rare association with crookesite, chameánite, giraudite, and other selenides (Figure 27).

It is grey in the reflected light, and anisotropy was not observed.

Except for the main elements, it contains a small amount of Fe, Ag, and Tl (up to 0.004 apfu). Only weak substitution SSe$_{−1}$ (up to 0.013 apfu) and SbAs$_{−1}$ (up to 0.003 apfu) was observed (Table S46). The chemical composition of unnamed Cu-As selenide from Příbram on the basis of 3 apfu (mean of two analyses) corresponds with the empirical formula Cu$_{1.04}$As$_{0.95}$(Se$_{0.99}$S$_{0.01}$)$_{Σ1.00}$.

Synthetic CuAsSe was explored by Whitfield [117]. Crystals of this phase have a structure with a simple polytypic relationship to the lautite (CuAsS) structure [118]. Both structures are ordered superstructures of the diamond arrangement, with Cu-As, As-(Se,Se), and Cu-(S,Se) bonds.

5. Succession of Crystallization

In evaluating a succession of selenide mineralization in the studied ore district, it is important to note that all succession data were obtained from the dump material without knowing its original vein or its geological situation. The limited joint occurrence of some minerals, and mutual intergrowth relations, impose limits on the definition of a full succession. As all of the samples were collected at
dumps, it is difficult to say if an unusual association (e.g., native arsenic, arsenolamprite, clausthalite, and berzelianite) represent certain evolutionary stages, or whether the minerals derived from a single vein—possibly affected by local lithology—are confined to a particular locality.

The presented scheme of mineral relations of the selenide and sulphide mineralization is based on a study of more than 250 polished sections. The possible or probable presence of some selenides or sulphides belonging to different generations (e.g., clausthalite, hakite, or chalcocite) would affect the accuracy of such a scheme. A general succession is proposed based on partial or local successions, but information is missing for some minerals of uncertain paragenetic position:

Eucairite, brodtkorbite, hessite → clausthalite → tiemannite → berzelianite → umangite → eskebornite → hakite, löllingite → giraudite, Se-tetrahedrite → tetrahedrite → Se-chalcopyrite → permingeatite, bytizite, příbramite, Se-chalcostibite, luzonite, antimonelite.

6. Discussion of the Formation Conditions

The main part of the uraninite at this ore district formed in the course of the III. calcite-uraninite mineralization stage (5th–6th import stage) [59]. The same author classified selenides in the later calcite-sulphide stage corresponding to the 7th or 8th import stage. The relatively later age of selenides in relation to uraninite is in agreement with our observations. Other mineralization stages in the uranium district, and in the Březové Hory ore district, carry no selenide occurrences or sulphides with increased selenium content [119–121].

This situation follows from differences in Se and S properties and from specific conditions of the selenium mineralization. Similar to the situation at other deposits in the Czech Republic, selenides in the Příbram area formed at temperatures near 100 °C. The abundance of calcite indicates a neutral to a weakly-alkaline environment [122,123]. Data on fluid inclusions in carbonates of the uraninite-antraxolite stage indicate crystallization temperatures of 80–140 °C [62]. The selenium thermometers confirm relatively low temperatures. Berzelianite—one of the most abundant selenides in the Příbram area—forms at temperatures exceeding 112 °C, and it alters to umangite at lower temperatures. Both the minerals are relatively abundant in the studied samples. Very rare is bellidoite, which separates from berzelianite at temperatures lower than 123 °C. However, at higher selenium concentrations, berzelianite can coexist with bellidoite at still lower temperatures [124]. The local athabascaite occurrence indicates temperatures below 100 °C [68]. Similar data based on a study of selenide mineralization and fluid inclusions were obtained at the Tumiñico selenium deposit, Argentina, by Paar et al. [79].

The solutions involved in selenide precipitation were of low salinity, and partly of meteoritic origin [49,62]. A strong hematitic alteration is characteristic for the veins, although not evident in all samples.

The selenides in our samples crystallized before most of the sulphides of the same mineralization stage, in contrast to the observations of Litochleb et al. [49].

Selenides of the vein hydrothermal deposits formed under conditions of \( f_{O2} \) above the hematite-magnetite buffer, probably in the range of 5.8 log units (hematite-ferroselite univariant reaction) to 7 log units (anglesite-galena buffer), above the hematite-magnetite buffer [125]. Such high oxygen fugacity values result in geochemical separation of selenium from sulphur in the hydrothermal fluids, a high \( \frac{Se}{S} \) fugacity ratio, and in the deposition of various selenide minerals. The deposition of the various selenide minerals, as well as the extent and nature of the association formed depend on the concentration of available Se in solution [124].

The mineralogy of the selenide association changed through the paragenetic sequence as a function of the changing environmental conditions. The first ore mineral to crystallize was the first generation of chalcocite, accompanied by subordinate bornite. Next, colloform uraninite precipitated as spherical aggregates and vein fillings. Then, selenides crystallized, often replacing uraninite.

Clausthalite and berzelianite form an independent association with arsenic and arsenolamprite. The dominant role of arsenic in the oldest phases of Ag-rich associations is known from the neighboring
Minerals 2017, 7, 91 of 56

shaft 21, Příbram-Haje [126]. The local crystallization of both arsenic polymorphs resulted in a lower As/Sb fugacity, and only a limited quantity of arsenic entered the selenide structure during the main period of their crystallization. With regard to the strongly-limited association of both arsenic polymorphs with selenides, it is not possible to determine their relative age in relation to the main stage of selenide crystallization.

The large volume of precipitated selenides is evidence of a high Se$_2$ / S$_2$ fugacity ratio. During the earlier part of the selenide crystallization stage, tiemannite, clausthalite, eucairite, and Cu selenides precipitated, while minerals of the tetrahedrite group formed later. The presence of Cu or Cu-Sb in selenides from Příbram is typical. Crystallization of these minerals lowered the Se fugacity sufficiently to cause the transition into the calcite-sulphidic crystallization stage, with the formation of minerals such as tetrahedrite and chalcopyrite. Similar relations were also observed at the Tumiñico locality [79].

During the later phase of selenide mineralization, there is evidence of repeated partial dissolution of older selenides, with replacement by the youngest Se-bearing phases and the production of significant textural complexity. This stage features considerable mineralogical complexity, even if it is of a limited occurrence. Variation in S and Se contents or other elements at the micrometer scale indicates highly variable fugacities in time and space of the respective low-temperature processes. Antimonselite, permingeatite, Se-rich chalcostibite, part of hakite, and two new minerals for the mineral system—příbramite [46] and bytzite [47]—belong to minerals of this regeneration stage of mineralization in particular. Rare As-dominant selenides and sulphides—in contrast to local common native arsenic occurrence—are concentrated in this later phase. Arsenic was probably supplied by hydrothermally-altered older minerals.

The variability of selenium minerals of the calcite-sulphidic stage derives primarily from a compositional variation of hydrothermal fluids supplying elements for selenide crystallization. Variation in Se/S fugacity contributed to the occurrence of sulphide and selenide pairs, such as Se-chalcostibite-příbramite, chalcopyrite-eskebornite, hakite-tetrahedrite, giraudite-tennantite, and others. Selenides containing Cd and TI are very rare. Minerals containing increased concentrations of these elements were not found, with the exception of Cd substitution in sphalerite in the Příbram ore district.

The selenide mineralization in the Příbram uranium district shows variations in Se$_2$ and S$_2$ fugacities (−12 to −18 in the case of Se, and −17 to −24 of S, respectively), which are similar to those published in detail by Simon et al. [125] for “selenide-bearing nonconformity-related uranium deposits”. Considering the absence of krut’aite in the association and according to the stability field diagram for the temperature 100 °C [125], we can expect the origin of the selenium mineralization under the krut’aite stability field. This corresponds to a log fSe$_2$ lower than −12.

A characteristic feature of selenide association in the Příbram uranium district is the fine-grained character of this mineralization. The individual selenide grains rarely exceed the limit of 2 mm. The occurrences include particularly abundant selenides of Cu, Sb, and Hg. In contrast, no selenides of Co, Ni, Bi, Au, and other elements have been found, although these occur at the Předbořice deposit, not far from the Příbram area. The presence of minute and rare inclusions of hessite is surprising. The origin of tellurium contained in this mineral is unlikely in hydrothermal alterations of an older gold-bearing quartz gangue, which typically contains tellurides. However, characteristic quartz gangue was absent in veins carrying selenides.

Hessite in association with selenides is known from the Frederik VII mine in Southern Greenland [127]. In uranium deposits, it was found only at Niederschlema by Förster [17]. The accompanying selenides in Příbram—with the exception of tiemannite, and occasionally berzelianite and umangite—are free of notably-elevated Te, as at the Niederschlema deposit.

7. Conclusions

The Příbram uranium and base-metals district contains important occurrences of selenide mineralization. Twenty-six mineral species were identified, including two new approved minerals
(příbramite and bytizite) and two possible new mineral species (Hg-Cu-Sb and Cu-As selenide). The minerals were characterized by chemical and optical data, in some cases by powder or single-crystal X-ray diffraction. The detailed study confirmed the occurrence of several minerals which are rare on a worldwide scale (e.g., antimonselite, brodtkorbite, chaméanite, giraudite, cadmoselite, permingeatite), or a wide range of substitutions (especially of tetrahedrite-group minerals).

Crystallization of the selenides reflects a high \( \text{Se}_2/\text{S}_2 \) fugacity ratio, but the phases formed changed through time as conditions evolved. Simple selenides of Pb, Hg, Ag, and Cu formed first, with tetrahedrite-group minerals later, and finally, some very late minerals as replacements of earlier selenides. Micron-scale variation of \( \text{S}/\text{Se} \) ratios and contents of other elements reflect conditions that varied rapidly in time and space.

The origin of selenide mineralization in the Příbram uranium district occurred between the calcite-uraninite and carbonate-sulphidic stage. Selenides—always later than uraninite—crystallized at temperatures near 100 \(^\circ\)C in a neutral-to-weakly-alkaline environment from solutions with high oxygen fugacity and a high \( \text{Se}_2/\text{S}_2 \) fugacity ratio.

Supplementary Materials: The following are available online at http://www.mdpi.com/2075-163X/7/6/91/s1.

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