

Secondary arsenic minerals from the Złoty Stok As-Au abandoned mine (SW Poland)

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Secondary arsenic minerals (SAM) formed recently in abandoned adits of the former Au-As mine at Złoty Stok (SW Poland) constitute two assemblages. The first consists of two types of scorodite, pitticite, kaňkite, hörnesite, picropharmacolite and minor amounts of jarosite and gypsum. Formation of the Fe arsenates took place under acidic conditions (pH ~3–4) as a result of löllingite, arsenopyrite and pyrite oxidation. Hörnesite and picropharmacolite crystallized as products of interactions between acidic arsenic-rich pore solutions with Mg-Ca carbonates from rocks that surround the ore mineralisation. The interaction of carbonates with acid pore solutions caused a rapid increase in pH that reached neutral or weakly alkaline values. The chemical compositions of hörnesite and picropharmacolite correspond well to their ideal compositions: $(Mg_{3.17}Ca_{0.07})_{23.24}(AsO_4)_{1.90} \cdot 8H_2O$ and $Ca_{4.31}Mg_{0.92}(HAsO_4)_{1.91}[(AsO_4)_{1.99}(SO_4)_{0.01}]_{22.00} \cdot 11H_2O$, respectively. The second assemblage of SAM comprises exclusively the Mg-enriched erythrite $[(Co_{1.66}Mg_{1.03}Ni_{0.28}Ca_{0.05}Zn_{0.02})_{23.03}(AsO_4)_{1.99} \cdot 8H_2O)]$ annabergite $[(Ni_{1.48}Mg_{0.94}Co_{0.66}Ca_{0.12}Fe_{0.01}Zn_{0.01})_{23.20}(AsO_4)_{1.92} \cdot 8H_2O]$ series. These minerals crystallized from slightly acidic $(pH \sim 5-6)$ to neutral media. Dissolution of SAM and other secondary phases (e.g., schwertmannite) causes the release of arsenate and sulphate ions into mine waters. These ions can be reduced under anaerobic conditions by different strains of bacteria. The product of this process is orpiment.

Key words: erythrite-hörnesite-annabergite series, picropharmacolite, secondary arsenic minerals.

INTRODUCTION

Arsenic is an example of toxic element released during decomposition of its primary minerals. Secondary arsenic minerals (SAM) are among the main As sinks in the supergene environment. Their crystallisation strongly depends on the physicochemical properties of the weathering zones of As-rich deposits (e.g., Drahota and Filippi, 2009). Changes in these conditions lead to the dissolution of SAM. This process releases arsenic into underground and surface waters, thus leading to their contamination (Marszałek and Wąsik, 2000). Particular species of SAM are well-studied because of arsenic toxicity, which strongly depends on the oxidation state of this metalloid. Thus, understanding SAM stability plays an important role from the environmental point of view. Secondary phases from the Złoty Stok deposit have been poorly recognized in previous research. Only several weathering minerals are known from this locality (Kubisz, 1964; Budzyńska, 1971). This paper is the first to presents the results of detailed study

of recently forming SAM found within abandoned adits of the Au-As mine at Złoty Stok. Attention is paid to the chemical diversity of SAM and their relationships to the known crystallochemical solid solutions. We also reconstruct the crystallisation conditions of the SAM under study.

GEOLOGY

The Złoty Stok Au-As deposit is located in the central part of the Lower Silesian Block (Żelaźniewicz et al., 2011) at the contact between the Złoty Stok-Skrzynka Shear Zone and the Kłodzko-Złoty Stok granitoid pluton. The Złoty Stok-Skrzynka Shear Zone developed during Variscan deformation of different metamorphic rocks (Cymerman, 1996; Mazur et al., 2006). It is composed of blastomyllonites, gneisses, mica schists and leptinites intercalated with marbles, amphibolites and serpentinites; the latter coexisting with minor nephrites (Kowalski, 1969; Smulikowski, 1979; Gil et al., 2015; Fig. 1). The Kłodzko-Złoty Stok granitoid pluton consists of rocks ranging from granodiorite to monzonite (Mikulski et al., 2013; Mikulski and Williams, 2014). Different types of hornfelses, skarns and skarnoids have been formed in the granitoid contact zone due to the thermal and metasomatic influence of the pluton (Wierzchołowski, 1976). Metasomatic events related to the Variscan magmatism

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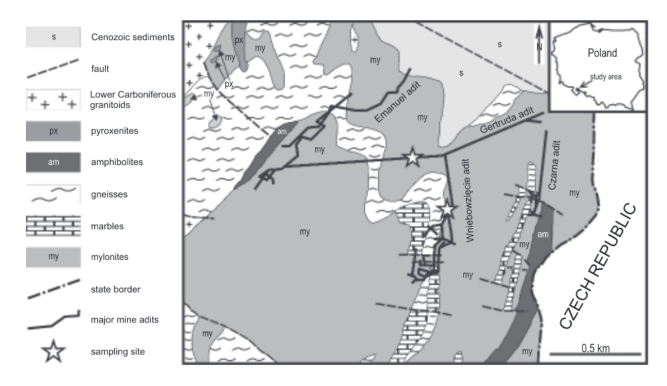


Fig. 1. Geological map of the Złoty Stok area, modified after Cwojdziński (1974); location of adits is based on old mining maps

were also responsible for the main ore mineralization in the Złoty Stok deposit (Mikulski and Speczik, 2008, 2016).

Three types of ore mineralisation occur in the Złoty Stok Au-As deposit. The first type contains löllingite-arsenopyrite mineralisation associated with diopside skarns and serpentinites surrounding dolomitic marbles. The second type is a pyrrhotite-magnetite ore with minor löllingite and arsenopyrite residing within diopside-tremolite rocks. The third variety is a disseminated löllingite-arsenopyrite ore that occurs inside black serpentinite. Sub-microscopic and microscopic gold occurs in all the ore types (Mikulski and Speczik, 2016). Other ore minerals, such as pyrite, galena, sphalerite, chalcopyrite, scheelite, glaucodot, cobaltite, cubanite, marcasite, bismuthinite, bismuth, arsenic, and Ni-Fe-Co sulphoarsenides occur as small aggregates of no economic significance (Mikulski, 1996; Muszer, 2011).

The Złoty Stok deposit has a long mining history. The first information about gold mines dates back to 1341 in this area (Dziekoński, 1972). Production of arsenic and synthetic arsenic pigments started at the beginning of the 17th century. Due to the poor quality of the arsenic ores, mining and arsenic-gold smelter activity ceased in 1961 (Muszer, 2011).

METHODS AND SAMPLES

Mineral samples were collected from the Gertruda and Wniebowzięcie adits of the former Au-As Złoty Stok mine (Fig. 1). The most diverse supergene mineralisation was found in the Gertruda adit (~842 m from the entrance). A small, up to 30 cm in size, ore lens surrounded by barren rocks rich in calcite and dolomite, was found there. The primary minerals of this lens (löllingite, arsenopyrite and pyrite) have been almost completely decomposed by weathering, leading to the formation of several characteristic zones with different mineral compositions

(Fig. 2). The first, central zone is dominated by powdery scorodite that contains relicts of the ore minerals. In the marginal parts of this scorodite type occurs its second form represented by botryoidal or spherical green aggregates. It is followed by pitticite, next to which the kaňkite accumulation zone is found. Segregations of Mg and Ca arsenates occur in the barren rocks surrounding the ore lens. In this zone, large amounts of biofilms with colonies of microorganisms are seen. The other mineral samples were taken from the Wniebowziecie adit, which is situated to the south of the Gertuda adit. Co-Mg-Ni arsenates form small aggregates on the surface of the weathering Co-rich löllingite. In Wniebowzięcie adit, schwertmannite was also found; however, it does not occur with SAM. The samples vary from 0.5 to 10 cm in diameter. To avoid humidity changes, all mineral samples were placed in tightly-sealed plastic containers. The samples were transported to the laboratory and stored at a temperature of about +10°C.

Quantitative chemical data were acquired using a Cameca SX 100 electron microprobe of the Inter-Institute Analytical Complex from Minerals and Synthetic Substances, University of Warsaw. The following standards, analytical lines and crystals were used: $V_2O_5 - V$ (K α , LIF), rhodochrosite - Mn (K α , LIF), Fe₂O₃ – Fe (Kα, LIF), chalcopyrite – Cu (Kα, LIF), ZnS – Zn (K α , LIF), albite – Na (K α , TAP), diopside – Mg (K α , TAP), orthoclase – Al (K α , TAP), diopside – Si (K α , TAP), GaAs – As $(L\alpha, TAP)$, apatite – P $(K\alpha, PET)$, orthoclase – K $(K\alpha, PET)$, diopside – Ca (K α , PET), NiO – Ni (K α , LIF), and metallic Co – Co (Kα, LIF). Analyses were conducted using the accelerating voltage of 10 kV and beam current of 5 nA. The beam diameter was 5-10 µm. ZAF corrections were applied. Elevated analytical totals for minerals with hydroxyl groups or crystallisation water are generally caused by evaporation of water under high vacuum conditions or by its evaporation due to heating of the analysed area by the electron beam. Lower totals are due to the porous nature of some minerals. The presence of all minerals was con-

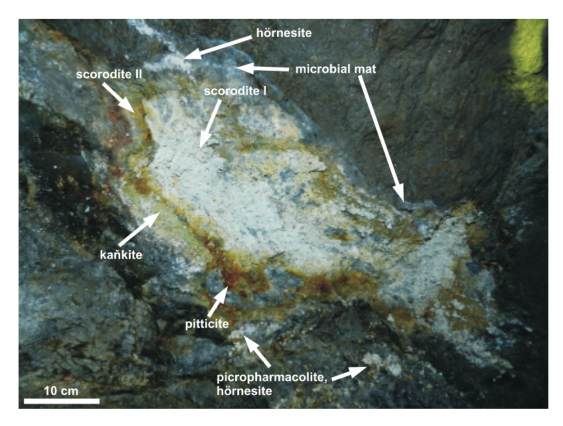


Fig. 2. Secondary Fe and Mg-Ca arsenates on the wall of the Gertruda adit

firmed by powder X-Ray diffraction (PXRD). Identification of those minerals was performed using a powder X-ray diffractometer X`Pert PRO MPD by the DSH method. The parameters of the X-ray beam were as follows: $\text{CoK}\alpha$ wavelength, finally monochromatized by parabolic X-rays mirrors, emitted from the X-Ray tube with 40 mA and 40 kV current applied. X-ray patterns were recorded in 2.5 to 75.99 °20 range, with a step size of 0.02 °20. The results were processed using the X-ray analysis software X'Pert Plus HighScore (ver. 2.2e) and ICDD PDF-2 database. Morphology of the samples was studied at the Faculty of Geology, University of Warsaw, using a FE-SIGMA VP (Carl Zeiss Microscopy GmbH) scanning electron microscope with an energy-dispersive (EDS) detector (Quantax XFlash 310, Bruker Nano GmbH). IR absorption spectra were recorded with a Nicolet Magna 550 specrometer from 4000 to 400 cm⁻¹, using KBr pellets (Faculty of Chemistry, Warsaw University). Raman spectra were recorded on a Jobin spectrometer -Yvone-Horiba coupled with a confocal microscope Olympus BX40 (AGH University of Science and Technology). The spectra were collected from mineral samples at room temperature, using laser with a wavelength of 514.5 nm in the range of 4000-50 cm⁻¹ at a resolution of 3.5 cm⁻¹ and a measurement time from 30 to 360 s.

SECONDARY MINERALS FROM THE GERTRUDA ADIT

SCORODITE FeAsO $_4 \cdot 2H_2O$

In the Gertruda adit, this mineral forms two types of aggregate. The first is powdery and grey-green and occurs in the cen-

tral part of the strongly oxidized ore lens, almost completely replacing the primary löllingite-arsenopyrite-pyrite aggregates. The diameter of these type-I scorodite aggregates reaches tens of centimetres. Even by applying very large magnifications, no crystals of this mineral can be observed. The chemical composition of the type-I scorodite does not strongly differ from the theoretical composition (Table 1). The iron content varies from 0.84 to 0.99 apfu. This element is replaced by a small amount of aluminum (0.01-0.08 apfu) and magnesium (up to 0.02 apfu). In the tetrahedral site the arsenate ion (0.66-0.99 apfu) is replaced by sulphate anion (0.02-0.28 apfu) accompanied by a phosphate ion (0.01 apfu). The calculated empirical formula of the type-I scorodite (based on four oxygen atoms per formula unit) is $(Fe_{0.92}AI_{0.02})_{\Sigma 0.94}[(AsO_4)_{0.75}(SO_4)_{0.22}(PO_4)_{0.01}]_{\Sigma 0.98} \cdot 2H_2O$. The second type of scorodite (scorodite II) forms green spherical aggregates up to 100 µm in diameter (Fig. 3A). They appear outside the earthly scorodite and coexist with pitticite. The identification of the scorodite II was based only on the PXRD method (Fig. 4A).

KAŇKITE FeAsO₄ · 3.5H₂O

Kaňkite is another iron arsenate. It forms semi-crystalline green coatings on the barren rock surface. These encrustations are built of very fine, thin tabular crystals up to 20 μm in size (Fig. 5A). The chemical composition of kaňkite is shown in Table 2. It shows a relatively small variation in iron content (from 0.88 to 0.98 apfu). This element is accompanied by aluminum (from 0.04 to 0.14 apfu) and magnesium (from 0.01 to 0.03 apfu). The arsenate content varies from 0.86 to 0.97 apfu. This anion is substituted to a small extent by sulphate (from 0.01 to 0.04 apfu) and phosphate (0.01 apfu) ions. Based on mi-

Table 1

Chemical composition [wt.%] of type-I scorodite from the Gertruda adit

2 3 4 5 6 7 8 9 10

Analysis number	1	2	3	4	5	6	7	8	9	10
Fe ₂ O ₃	29.73	30.56	36.01	36.93	37.12	35.02	36.26	34.04	34.63	35.13
Al_2O_3	1.78	1.80	0.16	0.17	0.19	0.20	0.19	0.20	0.18	0.22
MnO	0.00	0.00	0.03	0.00	0.00	0.14	0.00	0.00	0.00	0.00
MgO	0.39	0.37	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
As ₂ O ₅	49.68	50.80	38.40	37.68	35.49	37.21	38.01	39.00	39.04	38.51
SO ₃	1.34	0.89	9.75	10.24	10.44	10.65	9.93	10.51	10.25	10.22
P ₂ O ₅	0.38	0.35	0.15	0.16	0.15	0.14	0.18	0.16	0.13	0.17
SiO ₂	0.00	0.04	0.12	0.09	0.08	0.06	0.11	0.07	0.08	0.08
H ₂ O∗	15.90	16.11	17.04	17.21	16.88	16.94	17.08	17.08	17.08	17.08
Total	99.20	100.92	101.66	102.47	100.36	100.35	101.76	101.06	101.40	101.40
					apfu					
Fe ³⁺	0.84	0.86	0.95	0.97	0.99	0.93	0.96	0.90	0.91	0.93
Al ³⁺	0.08	0.08	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01
Mn ²⁺	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Mg ²⁺	0.02	0.02	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Σ A side	0.94	0.96	0.96	0.98	1.00	0.95	0.97	0.91	0.92	0.94
AsO ₄ ³⁻	0.98	0.99	0.71	0.69	0.66	0.69	0.70	0.72	0.72	0.71
SO ₄ ²⁻	0.04	0.02	0.26	0.27	0.28	0.28	0.26	0.28	0.27	0.27
PO ₄ ³⁻	0.01	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.01
SiO ₄	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Σ T side	1.03	1.03	0.97	0.96	0.94	0.98	0.97	1.00	0.99	0.98
H ₂ O	2.00	2.00	2.00	2.00	2.00	2.00	2.00	2.00	2.00	2.00

^{*}H₂O was calculated from the ideal content in the formula H₂O = 2; Ca, Ni, Co, V below detection limit

cro-area analyses, the empirical formula of kaňkite (calculated based on four oxygen atoms) is $(\text{Fe}_{0.93}\text{Al}_{0.08}\text{Mg}_{0.02}\text{Ca}_{0.01})_{\Sigma}$ $_{1.05}[(\text{AsO}_4)_{0.93}(\text{SO}_4)_{0.03}(\text{PO}_4)_{0.01}]_{\Sigma 0.97} \cdot 3.50\text{H}_2\text{O}$. The presence of kaňkite was confirmed by the PXRD method (Fig. 4B).

PITTICITE

Pitticite appears between scorodite II and kaňkite. It occurs as honey-yellow or brownish crust-like aggregates with a distinctive conchoidal fracture and vitreous to greasy luster (Fig. 3B). It is also very brittle and strongly cracked. The chemical composition is shown in Table 3. The iron content varies from 33.15 to 36.92 wt.% Fe₂O₃. Small amounts of aluminum (from 0.20 to 0.31 wt.% Al₂O₃) and magnesium (up to 0.09 wt.% MgO) were also detected. Arsenic, ranging in content from 36.72 to 39.95 wt.% As₂O₅, is accompanied by sulphur (10.45 to 11.36 wt.% SO₃), phosphorus (from 0.09 to 0.26 wt.% P₂O₅) and silicon (up to 0.06 wt.% SiO₂).

HÖRNESITE Mg₃(AsO₄)₂ · 8H₂O

Hörnesite crystallises on barren rock surfaces (Mg-enriched marble) on the outside of the kaňkite accumulations zone. It forms very fine, white, spherical aggregates up to 4 mm in size (Fig. 3C). Snowball-like aggregates of hörnesite are built of automorphic lath-like crystals. The size of the crystals is up to

20 μm. At large magnifications, a mosaic structure of the crystal terminations is visible (Fig. 5B). Hörnesite aggregates are usually surrounded by mucous products of microbiological metabolism processes (Fig. 5C). The chemical composition of hörnesite from the Gertruda adit is also close to the theoretical value (Table 4). There is only a small amount of calcium. The chemical analysis of hörnesite yields the formula (Mg_{3.17}Ca_{0.07})₂ _{3.24}(AsO₄)_{1.90} · 8H₂O (calculated based on eight oxygen atoms). The Raman spectra of hörnesite from the Gertruda adit are shown in Figure 6A. The results of the Raman analyses are reported in Table 5. Bands at 3055, 3044 and 3480 cm⁻¹ are associated with O-H stretching vibrations in the water. The position of these bands is very similar to those reported by Frost et al. (2003). Several broad bands at 2817-1141 cm⁻¹ are observed. Very strong bands at 876 and 807 cm⁻¹ are assigned to the v₃ and v₁ anti-symmetric and symmetric starching modes at arsenate groups, respectively. The band at 559 cm⁻¹ results from the Mg-O stretching vibrations. The Raman spectrum of the low-wavenumber region includes several sharp bands. Their presence is linked to v₄, v₂ modes and AsO bending and lattice modes (Makreski et al., 2015). The presence of hörnesite was confirmed by the PXRD method (Fig. 7A).

PICROPHARMACOLITE Ca₄Mg(HAsO₄)₂(AsO₄)₂ · 11H₂O

This mineral coexists with hörnesite. It appears as acicular crystals (Fig. 5D) forming spherical aggregates on barren rock

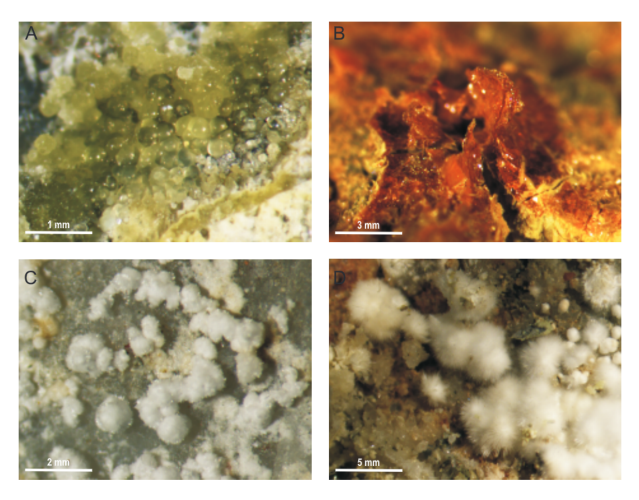


Fig. 3. Formation of secondary arsenic minerals from the Gertruda adit

 $\label{eq:local_problem} \textbf{A}-\text{green aggregates of scorodite II, } \textbf{B}-\text{accumulations of pitticite, } \textbf{C}-\text{white aggregates of h\"{o}rnesite,} \\ \textbf{D}-\text{sprays of picropharmacolite}$

surfaces (Fig. 3D). The investigated picropharmacolite is characterized by high chemical purity (Table 6). Only small amounts of sulphur (up to 0.03 apfu) were found, replacing the arsenate anion. The quantitative chemical analysis of picropharmacolite gives the following empirical formula based on 16 oxygen atoms: $\text{Ca}_{4.31}\text{Mg}_{0.92}(\text{HAsO}_4)_{1.91}[(\text{AsO}_4)_{1.99}(\text{SO}_4)_{0.01}]_{\Sigma 2.00} \cdot 11\text{H}_2\text{O}$. The presence of picropharmacolite was confirmed by the PXRD method (Fig. 4C).

JAROSITE KFe₃(SO₄)₂(OH)₆

Yellow, fine-grained, powdery aggregates (up to 0.2 mm in diameter) of jarosite were identified at the boundaries of the kaňkite and Mg-Ca arsenate zones. Jarosite accumulations are composed of automorphic crystals up to 10 μ m in size. Results of the chemical analysis are given in Table 7. Site A is dominated by potassium, varying from 0.79 to 0.90 apfu. It is accompanied by small amounts of sodium (from 0.02 to 0.03 apfu) and the hydronium ion (from 0.08 to 0.19 apfu). In the tetrahedral site, a minor substitution of the sulphate ion (1.85 to 2.04 apfu) by the phosphate ion is observed (from 0.02 to 0.04 apfu). The calculated empirical formula of jarosite (based on 14 oxygen atoms per formula unit; the amount of OH and H₃O $^+$ ions was cal-

culated based on the charge balance) is: $(K_{0.83}H_3O^+_{0.15}Na_{0.02})_{\Sigma}$ $_{1.00}Fe_{3.01}[(SO_4)_{1.97}(PO_4)_{0.03}]_{\Sigma2.00}(OH)_{6.00}$. This mineral was also identified by the PXRD method (Fig. 4D).

GYPSUM CaSO₄ · 2H₂O

This mineral occurs as small plate-like crystals, up to 20 μm in length, forming spherical aggregates. They were identified at the contact of the kaňkite and Mg-Ca arsenate zones. The presence of gypsum was confirmed only by PXRD.

ORPIMENT

In the Gertruda adit, unidentified arsenic sulphide is quite common. It occurs as impregnations on wooden mining supports submerged in water. It comprises bright yellow earthy accumulations. The aggregates are formed of thin crystals up to 3 μm in diameter (Fig. 5E), which coexist with bacterial colonies. The chemical composition is shown in Table 8. The analysed material shows high variability of arsenic and sulphur content (57.71–64.55 wt.% As, 21.23–31.67 wt.% S). The presence of orpiment was confirmed by the PXRD method (Fig. 4E).

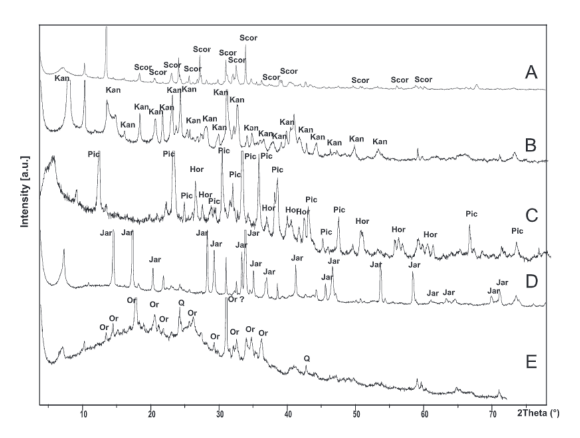


Fig. 4. PXRD patterns of secondary minerals from the Gertruda adit

A – scorodite II, B – kaňkite, C – picropharmacolite, D – jarosite, E – orpiment

SUPERGENE MINERALS FROM THE WNIEBOWZIĘCIE ADIT

ERYTHRITE $Co_3(AsO_4)_2 \cdot 8H_2O - H\ddot{O}RNESITE Mg_3(AsO_4)_2 \cdot 8H_2O$ - ANNABERGITE $Ni_3(AsO_4)_2 \cdot 8H_2O$ SOLID SOLUTION

In the Wniebowzięcie adit, two types of spherical aggregates of Co-Ni-Mg arsenates were found. The first type is purple and consists of plate-like crystals up to 30 μm in size. Purple aggregates contain higher amount of cobalt and accumulate on the weathering surface of Co-rich löllingite (average content of cobalt is 4.5 wt.%). The second type of Co-Ni-Mg arsenate is light pink in colour and formed on gersdorffite aggregates associated with löllingite. The chemical composition of crystals that build the purple aggregates is shown in Table 9. Their composition is variable. The octahedral site is dominated by cobalt and magnesium, with the contents varying from 1.39 to 2.01 apfu and from 0.61 to 1.28 apfu, respectively. These elements are accompanied by small amounts of nickel (from 0.23 to 0.34 apfu), calcium (from 0.03 to 0.07 apfu), and zinc (up to 0.04 apfu). The analysed crystals belong to the erythrite-hörnesite series with a slight enrichment in nickel (Fig. 8). The empirical formula of this phase was calculated based on eight oxygen atoms to be $(Co_{1.66}Mg_{1.03}Ni_{0.28}Ca_{0.05}Zn_{0.02})_{3.03}(AsO_4)_{1.99} \cdot 8H_2O$. In the case of the second type of aggregates, the dominant role in the cation site is played by nickel, whose content varies from 1.38 to 1.62 apfu (Table 10). In the structure, it is partially replaced by magnesium (from 0.87 to 1.01 apfu) and cobalt (from 0.60 to 0.70 apfu). They are accompanied by small amounts of calcium (from 0.09 to 0.15 apfu), zinc (up to 0.02 apfu) and iron (up to 0.01 apfu). The empirical formula of the analysed phase is (calculated based on eight oxygen atoms per formula unit) (Ni_{1.48}Mg_{0.94}Co_{0.66}Ca_{0.12}Fe_{0.01}Zn_{0.01}) $_{3.20}$ (AsO₄)_{1.92} · 8H₂O. Thus, the phase represents an Mg- and Co-enriched variety of annabergite (Fig. 8).

The Raman spectrum of Mg-enriched erythrite from the Wniebowzięcie adit (Fig. 6B) differs slightly from that of erythrite from Bou Azzer (Fig. 6C). The analysed sample shows the characteristic presence of three strong bands with maxima at 3456, 3228 and 3068 cm⁻¹. Compared to the Raman spectrum of pure hörnesite from the Gertruda adit, bands in this range are more visible. The bands observed at 856 and 796 cm⁻¹ appear from the anti-symmetric and symmetric As-O₄ vibrations. The positions of these bands are associated with the cation type at position A (Frost et al., 2003). In pure hörnesite, they appear at higher wave numbers (Table 5) than in pure erythrite. The position of the same bands for Mg-enriched erythrite from the Wniebowzięcie adit is intermediate and similar to the positions noted by Dumańska-Słowik et al. (2018). This shift is due to substitution between Mg²⁺ - Co²⁺ and Ni²⁺ ions in the cation site.

GOETHITE FEOOH AND FERRIHYDRITE Fe₁₀O₁₄(OH)₂

These minerals coexist with each other and form small stalactites and drapes of up to several centimetres in length. In the natural humidity state, they are soft and become brittle after drying. The presence of these minerals has been confirmed only by PXRD.

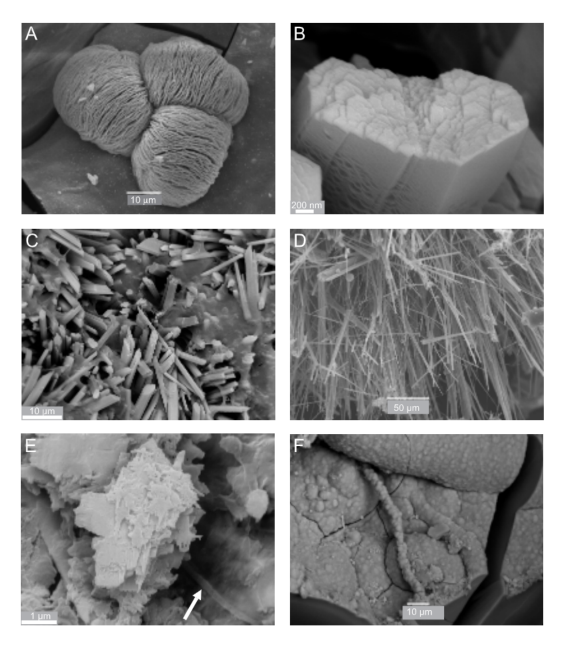


Fig. 5. SEM images of secondary arsenic minerals from the Gertruda adit

A – spherical aggregates of kaňkite on pitticite; B – mosaic termination of a hörnesite crystal; C – hörnesite crystals surrounded by a microbial mat; D – needle-like crystals of picropharmacolite; E – tabular aggregates of an unidentified arsenic sulphide from an old wooden mining construction, the arrow indicates bacterial colonies; F – microbial colonies mineralised by pitticite

SCHWERTMANNITE Fe₁₆(OH, SO₄)₁₂₋₁₃O₁₆·10-12H₂O

Schwertmannite occurs only in narrow zones (up to several centimetres in size) associated with strongly oxidized pyrite mineralisation. It forms small, up to 0.5 cm, stalactites. At large magnifications, typical thin, needle-like crystals of this mineral are visible. The presence of this mineral has been confirmed by PXRD.

DISCUSSION

The mineral compositions of SAM parageneses are strictly dependent on the conditions of their crystallisation (Drahota

and Filippi, 2009). The main factors that affect the crystallisation are the chemical composition of primary ore mineralisation and surrounding barren rocks, the humidity, pH and Eh, and the biological activity of microorganisms. Variability of the SAM crystallisation conditions can be observed in the oxidized ore lens exposed in the Gertruda adit. Arsenides and sulphides of iron (löllingite, arsenopyrite and pyrite) are unstable under supergene conditions. In the presence of oxygen and water, they undergo decomposition (Walker et al., 2006). This process is rapidly accelerated by Fe³⁺ ions – a strong oxidizing agent (Moses et al., 1987; Holmes and Crundwell, 2000). These ions are among the products of microbial activity of some bacteria strains (e.g., Schippers et al., 1996; Jones et al., 2003, Devasia and Natarajan, 2010; Vera et al., 2013; Tu et al., 2017). The presence of such bacteria was also con-

Table 2 Chemical composition [wt.%] of kaňkite from the Gertruda adit

Analysis number	1	2	3	4	5	6	7	8	9	10
Fe ₂ O ₃	30.16	31.32	29.06	29.47	29.16	29.51	30.36	30.00	31.99	30.90
Al ₂ O ₃	1.96	2.87	1.94	1.63	2.66	1.58	1.20	1.04	1.28	0.90
CaO	0.41	0.15	0.18	0.15	0.12	0.17	0.12	0.10	0.24	0.18
MgO	0.35	0.54	0.35	0.38	0.47	0.29	0.24	0.16	0.17	0.19
ZnO	0.08	0.22	0.34	0.17	0.06	0.11	0.00	0.00	0.00	0.05
K ₂ O	0.00	0.00	0.02	0.00	0.00	0.09	0.01	0.03	0.00	0.06
As ₂ O ₅	42.42	40.53	45.56	43.59	42.53	43.26	46.24	43.82	44.17	43.03
SO ₃	1.10	1.32	0.82	1.28	1.36	0.79	0.87	0.41	0.34	0.55
P ₂ O ₅	0.23	0.30	0.32	0.32	0.26	0.22	0.22	0.23	0.40	0.28
SiO ₂	0.15	0.08	0.02	0.06	0.01	0.22	0.08	0.07	0.11	0.06
H ₂ O [*]	25.51	25.79	26.06	25.62	25.63	25.21	26.22	24.89	25.83	24.98
Total	102.36	103.11	104.67	102.67	102.25	101.46	105.57	100.76	104.53	101.18
					apfu					
Fe ³⁺	0.93	0.96	0.88	0.91	0.90	0.92	0.91	0.95	0.98	0.98
Al ³⁺	0.09	0.14	0.09	0.08	0.13	0.08	0.06	0.05	0.06	0.04
Ca ²⁺	0.02	0.01	0.01	0.01	0.01	0.01	0.01	0.00	0.01	0.01
Mg ²⁺	0.02	0.03	0.02	0.02	0.03	0.02	0.01	0.01	0.01	0.01
Zn ²⁺	0.00	0.01	0.01	0.01	0.00	0.00	0.00	0.00	0.00	0.00
K⁺	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
Σ A side	1.07	1.14	1.01	1.02	1.06	1.03	0.99	1.02	1.06	1.04
AsO ₄ ³⁻	0.91	0.86	0.96	0.93	0.91	0.94	0.97	0.97	0.94	0.95
SO ₄ ²⁻	0.03	0.04	0.02	0.04	0.04	0.02	0.03	0.01	0.01	0.02
PO ₄ ³⁻	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01	0.01
SiO ₄	0.01	0.00	0.00	0.00	0.00	0.01	0.00	0.00	0.00	0.00
Σ T side	0.96	0.92	1.00	0.99	0.96	0.98	1.00	0.99	0.97	0.98
H ₂ O	3.50	3.50	3.50	3.50	3.50	3.50	3.50	3.50	3.50	3.50

 $^{^{\}star}$ H₂O was calculated from the ideal content in the formula H₂O = 3.5; Mn, Ni, Co, V below detection limit

Table 3

Chemical composition [wt.%] of pitticite from the Gertruda adit

Analysis number	1	2	3	4	5	6	7	8	9	10
Fe ₂ O ₃	34.14	33.6	33.94	33.15	35.03	34.44	36.92	34.94	36.43	36.28
Al ₂ O ₃	0.2	0.22	0.3	0.28	0.31	0.31	0.22	0.24	0.28	0.25
MgO	0.04	0.05	0.00	0.04	0.00	0.08	0.09	0.00	0.00	0.00
As ₂ O ₅	37.23	37.67	36.72	38.34	39.95	39.88	38.42	38.38	38.64	38.11
SO ₃	10.67	10.45	10.57	10.6	10.84	10.85	11.36	10.62	10.96	11.13
P ₂ O ₅	0.12	0.09	0.20	0.17	0.23	0.14	0.15	0.18	0.20	0.26
SiO ₂	0.02	0.03	0.03	0.04	0.06	0	0.06	0.01	0.04	0.05
Total*	82.42	82.11	81.76	82.62	86.42	85.70	87.22	84.37	86.55	86.08

^{*} Excluding water content; Mn, Ca, Ni, Co, V below detection limit

Chemical composition [wt.%] of hörnesite from the Gertruda adit 2 3 4 5 6 7 8 9 24.94 25.71 26.70 26.66 26.38 25.58 24.35 24.31 0.08 0.03 0.00 0.06 0.01 0.00 0.00 0.00 0.22 0.00 0.00 0.00 0.00 0.00 0.00 0.00 0.98 0.93 0.79 0.84 0.69 0.26 1.21 0.52 0.06 0.03 0.03 0.00 0.00 0.00 0.00 0.03 44.55 43.89 42.44 43.82 43.80 44.36 44.74 45.89 29.00 28.84 29.18 28.91 29.01 29.01 29.38 28.81 99.83 99.60 98.80 100.78 100.05 99.11 99.72 99.11 apfu 3.08 3.17 3.31 3.25 3.23 3.17 3.02 3.00 0.01 0.00 0.00 0.00 0.00 0.00 0.00 0.00

0.00

0.07

0.00

3.32

1.87

0.00

0.06

0.00

3.29

1.88

8

Table 4

0.00

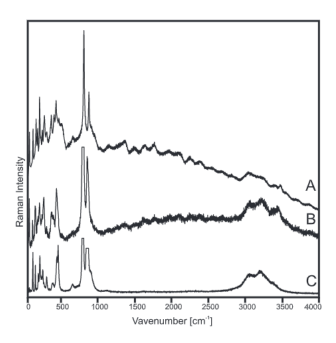
0.07

0.00

3.38

1.85

8



Analysis

number MgO

CoO

ZnO

CaO

FeO

 As_2O_5

H₂O^{*}

Total

Mg²⁺

Co2+

Ni²⁺

Ca2+

Fe²⁺

AsO₄³

 H_2O

 Σ A side

1

27.06

0.07

0.00

1.04

0.00

43.33

29.50

100.99

3.28

0.00

0.00

0.09

0.00

3.38

1.84

8

0.00

0.09

0.00

3.17

1.93

8

0.00

0.08

0.00

3.26

1.90

8

Fig. 6. Raman spectra of hörnesite from the Gertruda adit (A), erythrite-hörnesite from the Wniebowzięcie adit (B) and erythrite from Bou Azzer (Marocco) (C)

firmed in the microbial mats found in the Gertruda adit (e.g., Drawniak et al., 2010). Decomposition of primary arsenic minerals leads to the release of soluble forms of arsenic (such as arsenate and arsenite species). Arsenate ions dominate in en-

vironments with high Eh potentials. Arsenite ions can be rapidly oxidized by Fe³⁺ ions or during microbiological processes (Nicholas et al., 2003; Drewniak et al., 2012) to the AsO₃⁻¹ species (Mandl and Vyskovsky, 1994; Wiertz et al., 2006; Qin et al., 2013). In the course of iron sulphide oxidation, large amounts of sulphate ions are released into the environment, thus triggering a significant reduction in pH.

0.00

0.02

0.00

3.19

1.92

8

0.00

0.11

0.00

3.13

1.95

0.00

0.05

0.00

3.04

1.98

8

Under low pH conditions and very high activity of Fe³⁺ and AsO₄ ions, type-I scorodite is formed. This mineral replaces arsenic ore minerals in situ. Only small relicts of primary minerals are present within the scorodite accumulations. Scorodite is stable in the relatively narrow pH range: from 3 to 4 (Krause and Ettel, 1989; Bluteau and Demopoulos, 2007; Paktunc and Bruggeman, 2010). As the pH decreases, scorodite undergoes congruent dissolution which leads to the formation of Fe³⁺ and AsO₄ -rich pore solutions. These solutions migrate outside the type-I scorodite accumulations. Scorodite II crystallises from these solutions. Spherical aggregates of this mineral are typical of scorodite which crystallises slowly from pore solutions rich in the As(V) and Fe(III) species (Haffert and Crow, 2008). Situations of decreasing pH, leading to dissolution of scorodite I, are likely insignificant and short-term. The absence of zýkaite and bukovskýite confirms this observation, as these minerals are characteristic of low pH environments (2.5-3.5) (Gas'kova et al., 2008; Majzlan et al., 2012b, 2015).

The scorodite II-dominated zone contacts with pitticite accumulations. Pitticite is an amorphous hydrated sulphoarsenate of iron. It is a typical product of weathering of primary arsenic minerals which takes place in acidic environments (e.g., Kocourková et al., 2011b; Parviainen et al., 2012). The presence of highly mineralised bacterial colonies within the pitticite accumulations (Fig.

^{*} H₂O was calculated from the ideal content in the formula H₂O = 8; Al, Mn, Ni, Co, S, V, P, Si below detection limit

Table 5

Position and assignment of the bands [cm⁻¹] in the Raman spectra of Mg and Co arsenates from Złoty Stok, and comparison with literature data

Hörnesite from the Gertruda adit	Hörnesite (Frost et al., 2003)	Hörnesite (Makreski et al., 2015)	Erythrite from the Wniebowzięcie adit	Erythrite from Bou Azzer (this work)	Erythrite (Frost et al., 2003)	Band assignment
68			68	70		
118	117		113	113		Lattice mode
		138				Lattice mode
			149	149		Lattice mode
160	159	158			162	Lattice mode
181	180	180		189	188	Lattice mode
205	206	205	206			Lattice mode
				211	209	Lattice mode
222	218				223	Lattice mode
					234	
241	243	242	240	245		Lattice mode
				250	249	Lattice mode
	262	262	263	262	263	Lattice mode
273	273	271				Lattice mode
304	303	301	303	302	301	v ₂ (AsO ₄)
365	363	365				v ₂ (AsO ₄)
			377		378	v ₂ (AsO ₄)
			389	383		v ₂ (AsO ₄)
					391	
404	403	403				v ₄ (AsO ₄)
430	429	430				v ₄ (AsO ₄)
			437	442	439	v ₄ (AsO ₄)
	446					
				459	457	v ₄ (AsO ₄)
467	467	468			467	v ₄ (AsO ₄)
508						
657	660			651	652	
					727	
			796	793	792	v ₁ (AsO ₄)
807	807	808				v ₁ (AsO ₄)
			856	852	852	v ₃ (AsO ₄)
875	875	875				v ₃ (AsO ₄)
	907			906	902	
2817-1141			2280-1140 poor band intensity			
3055	3030		3068	3065	3052	OH stretchin
	3166					
			3228	3202	3200	OH stretchin
					3337	
3404						OH stretchin
			3456			OH stretchin
3480	3479					OH stretchin

5F) indicates the involvement of microorganisms in the precipitation of this phase. In relation to crystalline iron arsenates, the amorphous hydrated sulphoarsenate of Fe is unstable. Maturation of pitticite leads to the formation of crystalline Fe arsenates (Paktunc et al., 2008). On the other hand, pitticite can dissolve due to a slight decrease in arsenate or sulphate ions concentration (Chukhlantsev, 1956; Robins, 1987; Langmuir et al., 2006). For this reason, it may be a precursor to the coexisting kaňkite. Kaňkite crystallises primarily in post-mining or underground mining excavations (Majzlan et al., 2014). No direct crystallisation of kaňkite has been observed on the weathering surfaces of ar-

senic ore minerals. This observation may indicate that crystallisation of kaňkite is related to migration of arsenate-rich porous solutions. Furthermore, kaňkite is found only in very highly humid environments (Kato et al., 1984; Haffert et al., 2010; Majzlan et al., 2014). Kaňkite from the Gertruda adit is a younger phase in relation to scorodite and pitticite (Fig. 5A). This confirms the observations of other authors who related crystallisation of this mineral to dissolution of previously formed secondary iron arsenates (Haffert et al., 2010; Kocourková et al., 2011a). Kaňkite coexists with a small amount of jarosite, which is an indicator of low pH. Experimental data show that kaňkite is metastable with

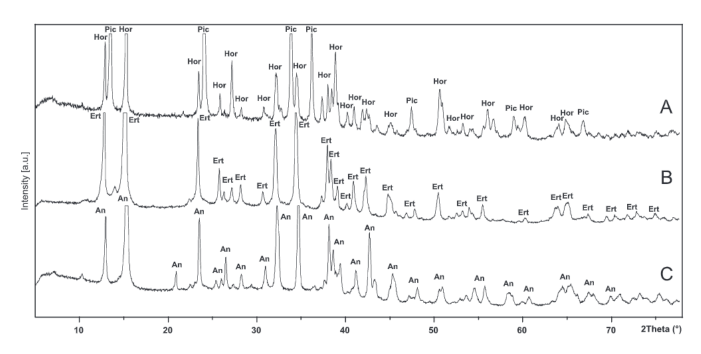


Fig. 7. PXRD patterns of Mg, Co, Ni arsenates from Złoty Stok

A - hörnesite (from the Gertruda adit), B - erythrite (from the Wniebowzięcie adit), C - annabergite (from the Wniebowzięcie adit)

Table 6

Chemical composition [wt.%] of picropharmacolite from the Gertruda adit

Analysis number	1	2	3	4	5	6	7	8	9	10
CaO	24.28	28.03	25.10	25.16	25.13	25.29	25.97	25.76	27.20	27.71
MgO	3.89	3.51	4.12	4.09	3.68	3.87	4.16	4.16	4.11	4.29
As ₂ O ₅	47.37	49.56	47.89	46.62	45.00	46.52	50.10	50.51	48.83	48.90
SO ₃	0.12	0.00	0.14	0.11	0.20	0.28	0.00	0.20	0.00	0.13
H ₂ O [*]	22.56	23.80	23.26	22.68	22.19	22.69	23.70	23.97	23.62	23.85
Total	98.22	104.89	100.52	98.65	96.21	98.65	103.93	104.59	103.75	104.87
					apfu					
Ca ²⁺	4.15	4.54	4.16	4.28	4.37	4.30	4.22	4.14	4.44	4.48
Mg ²⁺	0.93	0.79	0.95	0.97	0.89	0.91	0.94	0.93	0.93	0.97
Σ A side	5.07	5.33	5.11	5.24	5.26	5.21	5.17	5.07	5.37	5.44
AsO ₄ ³⁻	1.99	2.00	1.98	1.99	1.98	1.97	2.00	1.98	2.00	1.98
SO ₄ ²⁻	0.01	0.00	0.02	0.01	0.02	0.03	0.00	0.02	0.00	0.02
(HAsO ₄) ²⁻	1.97	1.92	1.89	1.88	1.84	1.89	1.98	1.99	1.89	1.87
Σ T side	3.97	3.92	3.89	3.88	3.84	3.89	3.98	3.99	3.89	3.87
H ₂ O	11.00	11.00	11.00	11.00	11.00	11.00	11.00	11.00	11.00	11.00

^{*} H₂O was calculated from the ideal content in the formula H₂O = 11 and charge balance; Fe, Al, Mn, Ni, Co below detection limit

respect to liquid water or water vapour (Majzlan et al., 2012a). Decomposition of this mineral may be the source of the AsO₄³⁻¹ ions for other secondary arsenates to form. The formation of Hörnesite and picropharmacolite is associated with the reaction of acidic arsenic-rich pore solutions with carbonates from barren rocks. These rocks contain variable amounts of calcite and dolo-

mite, which are the source of Mg²⁺ and Ca²⁺ ions. The interaction of carbonates with acid pore solutions causes a rapid increase in pH. Based on observations of crystallisation conditions of Mg-Ca arsenates in southern France (Juillot et al., 1999) it can be assumed that the corresponding process in the Gertrude adit takes place under neutral or weakly alkaline pH conditions. Sulphate

Table 7

Chemical composition [wt.%] of jarosite from the Gertruda adit

Analysis number	1	2	3	4	5	6
K ₂ O	6.63	7.20	6.95	7.12	7.83	7.61
Na ₂ O	0.16	0.16	0.13	0.09	0.12	0.15
Fe ₂ O ₃	43.54	44.01	43.12	46.54	45.20	44.40
SO ₃	24.92	31.14	30.57	30.04	28.85	30.09
P ₂ O ₅	0.37	0.53	0.40	0.33	0.35	0.34
H ₂ O [*]	9.30	10.66	10.45	10.67	10.14	10.35
Total	84.92	93.70	91.62	94.79	92.49	92.94
			apfu			
K ⁺	0.84	0.80	0.79	0.79	0.90	0.86
Na⁺	0.03	0.03	0.02	0.02	0.02	0.03
H ₃ O ⁺	0.13	0.18	0.19	0.19	0.08	0.11
Σ A side	1.00	1.00	1.00	1.00	1.00	1.00
Fe ³⁺	3.24	2.88	2.88	3.05	3.06	2.96
SO ₄ ²⁻	1.85	2.03	2.04	1.96	1.95	2.00
PO ₄ ³⁻	0.03	0.04	0.03	0.02	0.03	0.03
Σ T side	1.88	2.07	2.07	1.98	1.97	2.02
OH ⁻	6.93	5.45	5.48	6.14	6.20	5.80

^{*}H₂O content was calculated based on charge balance; Al, Mn, As below detection limit

Chemical composition [wt.%] of orpiment from the Gertruda adit

Analysis number	1	2	3	4	5	6	7	8	9
S	22.08	21.23	31.67	23.07	25.85	27.78	25.63	23.44	20.72
As	59.87	57.71	64.55	56.41	62.44	60.44	63.54	59.53	49.73
Total	81.95	78.94	96.22	79.48	88.29	88.22	89.17	82.97	70.45

ions that accompany arsenate ions are bound in gypsum crystals that co-exist with hörnesite and picropharmacolite. A similar process of the formation of Mg-Ca arsenates has been described from abandoned mine galleries in Jachymov, Czech Republic (Ondruš et al., 1997a, b). Crystallisation of hörnesite (and the accompanying picropharmacolite) under neutral or alkalescent pH conditions is also confirmed by the existence of hörnesite in soils with high arsenic content (Voigt et al., 1996; Foster et al., 1997). It is also worth noting that hörnesite aggregates are often associated with colonies of microorganisms (Fig. 5C). Their role in the formation of this mineral has not been fully established. There is a possibility that As and Mg²⁺ concentrations within such colonies somehow increase to the value at which crystallisation of hörnesite is possible.

Supergene Co-Mg-Ni arsenates of the erythrite-hörnesite--annabergite solid solution occur beyond the above-described ore lenses at the Wniebowzięcie adit. Their crystallisation is related to the decomposition of arsenides containing cobalt and nickel. Weathering of these minerals does not cause a significant reduction in pH. This is because the decomposition of Co-Ni arsenides does not release sulphuric acid but only poorly dissociated arsenic acid. The arsenic acid reacts with the released Co2+ and Ni2+ ions, which in turn leads to the appearance of secondary minerals directly at the surface of the ores. Mg²⁺ ions are supplied to the system from dolomite-containing barren rocks. The formation of the erythrite-hörnesite-annabergite solid solution is very similar to the mechanism described by Markl et al. (2014) to occur in supergene zones of various German polymetallic mines. According to data obtained by Langmuir et al. (1999) and Mahoney et al. (2007) these arsenates crystalised from slightly acidic media (pH ~5-6). These data were confirmed by Yuan et al. (2005), who proved that the solubility of annabergite decreases with increasing pH (minimal at pH = 9). The same relationship was confirmed for erythrite (Zhu et al., 2013). In several parts of the Wniebowziecie adit, schwertmannite was found. Its occurrence is limited to small zones containing disseminated pyrite mineralisation (without arsenopyrite and löllingite) which appears inside blastomyllonites. Moreover, some SAM were found with schwertmannite. This mineral is an indicator of acidic environmental conditions

Table 8

Table 9

Chemical composition [wt.%] of erythrite from the Wniebowzięcie adit

Analysis number	1	2	3	4	5	6	7	8	9	10
CoO	25.93	25.30	24.35	21.71	21.33	21.14	20.30	19.10	19.03	19.18
MgO	4.26	4.77	5.02	8.47	7.15	7.22	9.09	8.97	9.11	9.43
NiO	3.91	3.47	3.54	3.53	3.59	4.47	3.50	3.75	3.06	3.43
ZnO	0.31	0.40	0.17	0.00	0.18	0.38	0.16	0.14	0.00	0.63
CaO	0.47	0.28	0.30	0.44	0.59	0.40	0.38	0.44	0.59	0.68
FeO	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00	0.00
As ₂ O ₅	39.51	39.14	39.80	39.61	39.99	39.79	40.68	40.27	40.32	42.02
H₂O [*]	26.40	26.19	26.75	26.52	26.78	26.43	26.92	26.61	26.62	26.48
Total	100.78	99.55	99.93	100.27	99.61	99.82	101.02	99.28	98.73	101.85
				•	apfu					
Co ²⁺	2.01	1.98	1.90	1.64	1.63	1.61	1.51	1.44	1.44	1.39
Mg ²⁺	0.61	0.69	0.73	1.19	1.02	1.02	1.26	1.26	1.28	1.27
Ni ²⁺	0.30	0.27	0.28	0.27	0.28	0.34	0.26	0.28	0.23	0.25
Zn ²⁺	0.02	0.03	0.01	0.00	0.01	0.03	0.01	0.01	0.00	0.04
Ca ²⁺	0.05	0.03	0.03	0.04	0.06	0.04	0.04	0.04	0.06	0.07
Σ A side	3.00	3.01	2.95	3.13	3.00	3.05	3.07	3.04	3.02	3.02
AsO ₄ ³⁻	2.00	2.00	2.02	1.95	2.00	1.98	1.97	1.98	1.99	1.99
H ₂ O	8.00	8.00	8.00	8.00	8.00	8.00	8.00	8.00	8.00	8.00

^{*} H₂O was calculated from the ideal content in the formula H₂O = 8; Mn, S, P, V, Si below detection limit



Fig. 8. Ternary plot of the chemical composition of Mg-Co-Ni arsenates from the Gertruda and Wniebowzięcie adits (in atoms per formula units)

 H_2O

Analysis 2 3 4 5 7 8 9 1 6 number 18.51 18.80 19.96 19.92 19.30 21.06 NiO 19.58 19.25 18.19 CoO 8.97 9.01 7.96 8.55 8.47 9.20 7.76 8.70 9.23 MgO 7.07 7.27 6.12 7.11 6.11 6.52 6.11 6.67 6.95 ZnO 0.22 0.04 0.00 0.00 0.17 0.00 0.00 0.30 0.00 CaO 1.38 1.53 1.02 1.15 0.85 1.29 0.91 1.21 1.38 0.00 0.01 0.07 0.04 FeO 0.07 0.16 0.14 0.08 0.01 39.23 38.65 38.56 38.50 39.16 40.05 38.42 37.83 38.64 As_2O_5 H₂O 25.70 25.67 24.99 25.53 25.18 25.93 25.05 24.93 25.38 101.14 102.43 99.39 100.11 Total 100.96 98.76 100.75 99.59 98.60 apfu Ni²⁺ 1.41 1.62 1.39 1.54 1.51 1.50 1.44 1.49 1.38 Co²⁺ 0.67 0.68 0.61 0.64 0.65 0.68 0.60 0.67 0.70 Mg²⁺ 1.01 0.88 0.87 0.87 0.96 0.98 0.98 1.00 0.90 Zn²⁺ 0.01 0.00 0.00 0.00 0.01 0.00 0.00 0.00 0.02 Ca2+ 0.15 0.09 0.09 0.14 0.14 0.10 0.12 0.13 0.12 Fe²⁺ 0.01 0.00 0.01 0.00 0.01 0.01 0.01 0.00 0.00 Σ A side 3.20 3.26 3.15 3.26 3.12 3.16 3.19 3.24 3.23 AsO₄³ 1.91 1.89 1.94 1.89 1.95 1.94 1.92 1.90 1.91

Ta ble 10

Chemical composition [wt.%] of annabergite from the Wniebowzięcie adit

8

8

8

8

(pH = 3–4) and high activity of sulphate ions (Parafiniuk and Siuda, 2006). Schwertmannite is a metastable mineral, susceptible to environmental changes (Yu et al., 2002). In humid periods of the year, an increased flow of meteoric waters of neutral pH results in transformation of schwertmannite into ferrihydrite and goethite. This causes the schwertmannite-bound sulphate ions to be released into mine waters.

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Arsenate and sulphate ions of mine waters can be reduced under anaerobic conditions. Bacteria play a key role in this process (Nicholas et al., 2003). Arsenic sulphides of various As:S ratios stand for the products of such bacterially-driven reduction (e.g., Newman et al., 1997; Ledbetter et al., 2007; Battaglia-Brunet et al., 2012; Rodriguez-Freire et al., 2014, 2016). Different strains of bacteria, which reduce As(V) to As(III) were identified in the microbial mats from the Gertruda adit (Drewniak et al., 2010, 2012; Tomczyk-Żak et al., 2013, Cłapała et al., 2016). In this process, the arsenic and sulphate ions are electron acceptors, whereas organic matter from the old wooden mining construction acts as an electron donor. Then, arsenic sulphide is formed and precipitates only in the interstices of wood impregnated with the bacterial colonies, which clearly indicates that this process is related to organic matter and microbial activity.

CONCLUSIONS

Secondary arsenic minerals (SAM) from the As-Au mine at Złoty Stok are a unique assemblage of supergene minerals in Poland. These minerals record changes in environmental conditions in the abandoned mine galleries. Under oxidizing conditions, the dominant solid phases are arsenates. Iron arsenates occur only in areas of low pH and high activity of Fe³⁺ ions. Such

zones develop when the primary arsenic minerals and pyrite are weathered. The first mineral to crystallise is scorodite I. This mineral is partly dissolved and recrystallized as scorodite II. Arsenate and ferric ions released in this process are bound up in pitticite and kaňkite. The barren rocks, which surround the weathering ore minerals, contain calcite and dolomite that are a source of Ca²⁺ and Mg²⁺ species.

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Hörnesite and picropharmacolite are the products of the reaction between Ca²⁺, Mg²⁺ and AsO₄³⁻ ions. These minerals crystallise under neutral or slightly alkaline pH conditions, with increasing pH values resulting from carbonates dissolution. Arsenates of the erythrite-annabergite-hörnesite solid solution are produced by decomposition of primary Co-Ni arsenides. This process does not cause a significant pH reduction. For this reason, these minerals crystallise under slightly acidic pH conditions. Under anoxic conditions arsenate and sulphate ions present in the mine waters undergo bacterial reduction. The arsenic sulphides form in this way. Microorganisms take part in the formation of secondary arsenic minerals. They accelerate both reactions: the oxidation of reduced arsenic forms and the reduction of arsenate ions. The abandoned adits of the Złoty Stok As-Au mine are the first documented place in Poland where the SAM record the full circulation of arsenic in a supergene environment.

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^{*} H₂O was calculated from the ideal content in the formula H₂O = 8; Mn, S, P, V, Si below detection limit

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