The paragenesis of silver minerals in the Pb-Zn Stan Terg deposit, Kosovo: an example of precious metal epithermal mineralization

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Abstract: This study reports silver mineral association found recently in the Stan Terg lead and zinc mine, located in the Vardar zone (in northern Kosovo). The described mineralization comprises pyrargyrite (Ag₃SbS₃), freieslebenite (AgPbSbS₃), high-Ag bearing tetrahedrite and freibergite ((Ag⁴⁺₂ₓCu²⁻₂ₓ)(Cu,Ag)₄(Fe,Zn)₂Σ₆Sb₄S₁₂S₁⁻ₓ with (0 < x < 1)); as well as native compounds (Electrum, composition of those minerals was confirmed by the electron microprobe. The freibergite from native silver is native antimony). The Ag-minerals occur in vuggs and cracks in a massive galena ore and have signs of the latest minerals, which precipitated in the deposit. The chemical of the Stan Terg deposit reveals zonality and contains between 13.91–20.28% of Ag. The high concentration of Ag in solutions is also indicated by relatively high silver content in Au-Ag alloy (electrum), which is between 47.02% and 73.19% of Ag. The Ag association is supposed to be an epithermal equivalent of precious metal mineralization, which could be located in the external part of the Stan Terg hydrothermal system. This association occurs in low temperatures, below 200°C. The Ag-minerals can be a part of epithermal veins from the external part of the Stan Terg hydrothermal system. Similarly to the other known Pb-Zn-Ag hydrothermal systems, the Ag association is related to the formation of the rhodochrosite banded ore and Ag-Au-Sb dominated mineralization.

Keywords: Ag-minerals, pyrargyrite, freieslebenite, freibergite, Kosovo, Trepça mineral belt, Vardar zone

INTRODUCTION

Silver minerals are common ore minerals in various types of ore mineralizations and deposits primarily in epithermal vein systems, but they can be found also in skarn-, sedimentary-exhalative-, sediment-hosted-, porphyry-types of mineralization (e.g. Pearson et al. 1988, Grossou-Valta et al. 1990, Püttmann et al. 1991, Höller & Gandhi 1995, Schalamuk et al. 1997, Zeng et al. 2000, Cheilletz et al. 2002, Shalaby et al. 2004, Seifert & Sandmann 2006). Silver in the ore deposits occurs primarily in solid solutions with galena in the form of Bi(Sb) + Ag vs 2Pb substitution (lillianite type). In high temperature conditions, silver could form various groups of Ag-Pb-Bi(Sb) sulphosalts. In epithermal mineralization, silver commonly forms its own sulphide minerals; e.g. pyrargyrite, stephanite, proustite, argentite and/or native silver.

Stan Terg is one of numerous Pb-Zn deposits in the Vardar zone, located in the central part of the Balkan Peninsula (Southern Europe). In ancient times, mineral occurrences in the Balkan Peninsula were the main sources of Au and Ag. Mineralization is associated with Tertiary volcanism and is related to hydrothermal activity. Silver
mineralization is known from different base metal vein districts and deposits, for example the Podrinje Metallogenic District (Bosnia and Herzegovina), mainly from Srebrenica, where Ag-bearing tetrahedrite, stephanite, polybasite, pyrargyrite, argyrodite and native silver have been described (Radosavljević et al. 2016). The Zletovo mining district (Macedonia), freibergite, proustite, and pyrargyrite were described in sulphosalts paragenesis (Serafimovski et al. 2006). In the territory of Kosovo, pyrrargyrite was reported by Smejkal (1960), and Smejkal & Rakić (1957), in the Stan Terg, Hajvalia, Kizhnica and Kopaonik deposits. Berjaktarevic (1995) described the “Glama silver” prospect, near Gjilan, with a base metal ore with increased Ag, Au and Sn contents.

Silver in the Stan Terg deposit is linked mostly with galena ore, as isomorphic substitution in a lattice with Bi (Kołodziejczyk 2012), as well as in the form of its own minerals (Smejkal & Rakić 1957, Smejkal, 1960, Kołodziejczyk 2012). The Ag content in ore from the Stan Terg mine is 70 g/Mg on average, with Pb + Zn content around 10% (Féraud & Deshamps 2009, Trepça Mining Company internal documents). According to these works, most of the silver is linked to galena, and a minor part of it may belong to separate Ag-associations.

The purpose of this study is to report the electron microprobe data of the epithermal, Ag-bearing mineral association from Stan Terg in the external part of the deposit. This association may indicate the presence of an epithermal stage of evolution in the Stan Terg hydrothermal system.

**GEOLOGICAL SETTING**

The Stan Terg deposits and its hydrothermal system are located in Kosovo, within the Šumadija-Kopaonik ore district, known also as the Trepça mineral belt. It is a central part of the Vardar zone, in the vicinity of the Kopaonik granite massive (Hyseni et al. 2010). Stan Terg is the main Pb-Zn deposit, surrounded by numerous smaller occurrences, of the same hydrothermal origin (Fig. 1).

The major lithologies in the area are fragments of Paleozoic crystalline schists and phyllites, with unconformable overlying Triassic sediments, phyllites, volcanoclastic rocks and Upper Triassic carbonates. In the broad surrounding of the deposit, ultrabasic rocks and serpentinites of Jurassic age are exposed on the surface. In addition a Cretaceous complex occurs with carbonates, clastics, serpentine, volcanics and volcanoclastic rocks of basaltic composition. Tertiary (Oligocene-Miocene) volcanics of andesite, trachyte and latite composition are common in the area, and the surface is covered by felsic pyroclastic rocks (Hyseni et al. 2010, Strmić Palinkaš et al. 2013).

Mineralization in the Stan Terg deposit is generally hosted in the Mesozoic carbonates, at the contact with schists and the volcanic rocks of calc-alkaline affinity (Hyseni et al. 2010). The orebodies have an elongated shape of long mantos, or huge veins (with diameters up to few hundreds of meters). The orebodies stretch parallel to the conduit as columns. They split and join together in a form of a huge stockwerk.

The Stan Terg deposit has a large hydrothermal system with various mineralization styles, such as skarns, skarn-free carbonate-replacements, veins and breccias (Féraud & Deshamps 2009, Hyseni et al. 2010, Strmić Palinkaš et al. 2013). Skarn minerals are observed primarily close to the central part of the deposit. According to Dangić (1993), part of the ore mineralization could occur during skarn formation; mainly magnetite, pyrite, pyrrhotite, chalcopyrite, sphalerite and galena. The skarns at Stan Terg are of Ca-type and consists of typical contact metamorphic assemblages, with garnet, hedenbergite, ilvaite and actinolite. Within the skarn mineralization, Bi minerals were identified (Kołodziejczyk et al. 2015). The Bi-assemblage is also related to breccia mineralization, close to the central volcanic rocks. The main mineralization at Stan Terg was formed during the hydrothermal stage and comprises several generations of pyrite, galena, sphalerite, chalcopryrite, arsenopyrite, marcasite and lead-antimony sulfosalts (Féraud & Deshamps 2009). The gangue minerals are various Ca-Mn-Fe-Mg carbonates and quartz. Small veins and veinlets which run offset from the main orebodies are observed in the external part of the deposit. The hydrothermal phase was divided into four sub-stages: “Zn + Pb” stage I, “Cu” stage II, “Ag” stage III and “Sb” stage IV (Kołodziejczyk 2012), where galena and sphalerite precipitated mostly during the first stage. According to this classification, Ag-mineralization is postponed only by the “Sb” stage comprising of boulangerite and heteromorphite.
ANALYTICAL METHODS

Silver minerals were found in samples collected from the central part of 149e orebody. This orebody is localized in the external, northern part of the deposit. All of the described phases were found in massive, fine-grained galena hand specimens, with minor pyrite and siderite in the vuggs.

Polished sections were examined by optical microscopy, and the chemical composition of the minerals was determined by a JEOL JXA-8230 Super Probe electron microprobe in the Critical Elements Laboratory at the Faculty of Geology, Geophysics and Environmental Protection, AGH University of Science and Technology in Krakow.

The operating conditions for Ag-sulphosalts and native elements were as follows: accelerating voltage 20 kV, current 10 nA, peak time 10 s, background time 5 s, beam diameter was 3 μm. The following wavelengths were used: TeLα, SKα, PbMα, CuKa, BiMa, AgLa, SbLa, SeLa, AsLa. Natural mineral standards (PbS, FeS₂) and synthetic compounds (Bi₂Te₃, Cu, Ag, Sb, Se, InAs) were used for the calibration.

For freieslebenite and tetrahedrite, operating conditions were: accelerating voltage 20 kV, beam current 20 nA, peak time 20 s, background time 10 s and a beam diameter between 0 and 5 μm. The following wavelengths were used: PbMα, TeLa, BiMa, SbLa, SeLa, CdLa, InLa, ZnKa, CuKa, SKa, SnLa, HgMa, FeKa, MnKa, AgLa, GaLa, GeLa, AsLa. Natural mineral standards (PbS, ZnS, FeS₂, Sb₂S₃, MnS) and synthetic compounds (InAs, Cu, Bi, CdS, PbTe, Ag, HgTe, SnS, Se, Ga) were used for the calibration.

Fig. 1. Location and geological setting of the Stan Terg deposit (Kołodziejczyk et al. 2015)
RESULTS

The ore character of the 149e orebody is a little bit different from ore bodies in different parts of the deposit. The main ore mineral is a finely crystallized galena, with numerous microscopic boulangerite inclusions. Macroscopically visible crystals of boulangerite are rare. The minor minerals in the ore body are pyrite and pyrrhotite, and they occur especially at the contact with the host rocks. Sphalerite occurs in minor quantities, together with chalcopyrite and usually is observed only by using a microscope. The ore in this paragenesis is massive, with fine vuggs filled up by siderite and, less commonly, quartz. Galena has various textures, ranging from very fine (pure galena ore) up

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**Fig. 2.** BSE images of silver minerals from Ag-association at the Stan Terg deposit: A) pyrargyrite (pg) grains overgrown on galena (gn); B) pyrargyrite (pg) grains overgrown on carbonates and galena (gn); C) freieslebenite (fr) crystal overgrown on galena (gn), sph – sphalerite; D) zonal freibergite (frb) and electrum grain (Au) in the vugg in galena (gn); E) zonal electrum crystal (Au) overgrowing galena (gn); numbers in bracket indicate Au content in atomic %; F) native antimony (Sb) grain in a vugg, blg – boulangerite; numbers in brackets indicate Ag content (apfu) in freibergite, and Au content in atomic % in electrum

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to coarse-grained (as mixture with other ore minerals in the contact zone).

The Ag minerals seem to be the youngest phases in the ore paragenesis. They occur usually as free, rounded crystals overgrowing galena, or they fill the fine cracks or vuggs in the ore (Fig. 2A–F). Paragenetic observations suggest that pyrargyrite and pyrostilpnite are younger than chalcopyrite and tetrahedrite. The position of native elements with other Ag minerals is still questionable, because they are not occurring together.

Back-scattered electron images (BSE) provided information about the zonality of the analyzed phases, observed mainly in tetrahedrite and electrum phases.

**Pyrargyrite/pyrostilpnite**

Pyrargyrite is the most abundant Ag mineral in the described association. It occurs as separate grains, overgrowing galena in the vuggs, or replacing chalcopyrite and tetrahedrite. No zonation is observed in these crystals (Fig 2A, B).

Microscope observations indicate the presence of different colours of internal reflections in the phases corresponding to Ag₅SbS₃ composition. According to Uytenbogaardt & Burke (1985), this phenomenon may separate pyrargyrite from pyrostilpnite that may be distinguished with red or brownish internal reflections, respectively. Hence, we can conclude that in the Stan Terg deposit both phases occur, as a result of the internal reflections of both colours.

The chemical composition of measured grains is close to the ideal Ag₅.0₂Sb₀.94S₃.00, and is presented in Table 1, and plotted on diagram (Fig. 3). No arsenic was detected, and there is no sign of the presence of a pyrargyrite-proustite solid solution.

![Fig. 3. Ag-Sb-Pb tertiary plot (at. %) for Ag-minerals from the Stan Terg deposit](image-url)
Fereislebenite

Fereislebenite is rare and occurs similarly as the pyargyrite, overgrowing galena. Single grains are up to 20 μm and reveal no zonation (Fig. 2C). The chemical composition is close to the ideal, AgPbSbS3. Representative analyses are presented in Table 1 and are plotted on the diagram (Fig. 3). No As was detected in the analyses.

Ag-rich tetrahedrite and freibergite

The status of freibergite as a separate mineral is discussed. Moëlo et al. (2008) summarized reports concerning Ag-rich tetrahedrite and freibergite. Following Rozhdestvenskaya et al. (1993) the actual structural formula of freibergite may be represented as (Ag4+2xCu2−2x)[(Cu,Ag)4(Fe, Zn)2]Σ6Sb4S12S1−x with (0 < x < 1). Hence, the Ag content in freibergite is between 4–6 apfu (atoms per formula unit). The phases with Ag below 4 apfu should be called Ag-rich tetrahedrite.

Tetrahedrite group minerals found together with Ag phases are represented by Ag-rich tetrahedrite and freibergite. Representative chemical analyses of these minerals are presented in Table 2. The minimal Ag content in Ag-rich tetrahedrite is 10.65%, what corresponds to 3.09 apfu. In freibergite from Stan Terg, the Ag content varies between 13.91–20.28% (4.02–5.68 apfu) and the x value varies between 0.01 and 0.84. There is a strong correlation between Ag and Cu content (Fig. 4A). The freibergite shows zonality in the BSE images which is due to differentiated Ag content (Fig. 2D). The Fe and Zn contents are variable and do not depend on Ag content. Most of the results indicate 6% of Fe and 1% of Zn, what corresponds to about 1.7 Fe apfu, and 0.3 Zn apfu (Fig. 4B). The Ag-rich tetrahedrite and freibergite from Stan Terg contain up to 0.03 apfu of Pb.

Native elements

Numerous minute grains of electrum have been found in the described Ag-association. The electrum occurs as separate, rounded grains overgrown on galena, or within freibergite. The maximum dimension of the electrum is up to 20 μm. Chemical analyses of the electrum show variable Au vs Ag contents (Tab. 3). The electrum from the Stan Terg mine is characterized by a high content of Ag, which at its maximum could reach 73.19%. Practically all analyses are silver dominant, which indicates the presence of Ag-rich solutions.

Table 1

Chemical composition of pyrargyrite (1–3) and freieslebenite (4–6) from the Ag-association at the Stan Terg deposit (wt. %); b.d. – below detection limits, n.a. – not analyzed

<table>
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<td>Pb</td>
<td>0.07</td>
<td>b.d.</td>
<td>b.d.</td>
<td>39.75</td>
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<td>Fe</td>
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<td>n.a.</td>
<td>n.a.</td>
<td>0.43</td>
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<tr>
<td>Cu</td>
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<td>b.d.</td>
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<td>59.52</td>
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<td>20.03</td>
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<td>Sb</td>
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<td>21.65</td>
<td>22.02</td>
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<td>Bi</td>
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<td>b.d.</td>
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<td>b.d.</td>
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<td>17.60</td>
<td>17.61</td>
<td>18.08</td>
<td>18.13</td>
<td>18.07</td>
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<tr>
<td>Total</td>
<td>99.62</td>
<td>99.03</td>
<td>98.89</td>
<td>100.05</td>
<td>100.98</td>
<td>101.63</td>
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</table>

* Chemical formula based on 4 cations for pyrargyrite and 4 cations for freieslebenite.
Table 2
Chemical composition of Ag-rich tetrahedrite (1–4) and freibergite (5–7) from the Ag-association at the Stan Terg deposit (wt. %); b.d. – below detection limit

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<tr>
<td>S</td>
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<td>22.69</td>
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<td>22.41</td>
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<tr>
<td>Cu</td>
<td>22.02</td>
<td>21.31</td>
<td>20.81</td>
<td>20.72</td>
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<td>17.96</td>
<td>14.45</td>
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<tr>
<td>Fe</td>
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<td>0.54</td>
<td>7.15</td>
<td>6.50</td>
<td>8.25</td>
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<td>5.26</td>
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<td>Zn</td>
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<td>b.d.</td>
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<td>b.d.</td>
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<td>0.08</td>
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<td>100.23</td>
<td>100.58</td>
<td>100.41</td>
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Chemical formula on the basis of 16 cations

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<td>3.60</td>
<td>3.91</td>
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<td>5.68</td>
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<td>Cu</td>
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<td>0.18</td>
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<td>b.d.</td>
<td>b.d.</td>
<td>0.01</td>
<td>0.01</td>
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</table>

As, Se, Bi, Hg – were below detection limits.

Fig. 4. Composition of high Ag-tetrahedrite from Ag-mineralization at the Stan Terg deposit: A) binary plot Ag vs. Cu (at. %); B) binary plot Zn vs. Fe (at. %)

Table 3
Chemical composition of native elements from the Ag-association at the Stan Terg deposit (at. %); 1–4 electrum, 5–7 native antimony, b.d. – below detection limits, n.a. – not analyzed

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<tr>
<td>Hg</td>
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<td>0.11</td>
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<tr>
<td>Au</td>
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<td>33.43</td>
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<td>24.97</td>
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<tr>
<td>Fe</td>
<td>0.03</td>
<td>0.64</td>
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<tr>
<td>Sb</td>
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<td>0.05</td>
<td>0.10</td>
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As, Bi, Cu – were below detection limits.
A strong correlation between Ag and Au (Fig. 5) is observed, with Ag content between 47.02% and 73.19% (Au content 52.77% and 24.97%, respectively).

Native antimony was found as a separate, irregular grain in a vugg of about 15 µm size (Fig. 2F). Composition of the native Sb is presented in Table 3. The analyses indicated small Fe impurities; however they could be caused by the surrounding siderite.

DISCUSSION

This paper presents the electron microprobe (EPMA) geochemical results for pyrargyrite, pyrostilpnite, freieslebenite, Ag-rich tetrahedrite and freibergite. In the first stage of research, the authors carried out SEM-EDS analyses that additionally indicated the presence of stephanite and polybasite (Pršek et al. 2012). Unfortunately, due to the small size of crystals and difficult analytical conditions, we were not able to repeat these measurements by the means of the electron microprobe. Those analyses of stephanite indicated its elevated antimony content (2.4–2.5 apfu). The identified polybasite had an irregular form and occurred within chalcopyrite, together with Ag-bearing tetrahedrite and freibergite. The polybasite had a chemical composition with an increased content of copper (12%) and a reduced content of silver (39%) in comparison to the theoretical one.

Epithermal character of Ag-mineralization

The described mineral association is common in epithermal systems around the world. Similar associations occurring as tiny crystals in the fractures were described by Gemmell et al. (1989) from the Fresnillo district in Mexico. The mineral from Fresnillo has an increased As vs Sb and Se vs S substitution, which indicated the additional presence of proustite and selenian polybasite. Lynch (1989) described the association of pyrargyrite, acantite, polybasite, stephanite, native Ag in the polymetallic veins form Yukon.

Pyrargyrite from Stan Terg has been assigned by Kepuska (1998) to the mesothermal phase of the hydrothermal stage of mineral precipitation in this deposit, whereas according to Topalović (1971), pyrargyrite occurred in the epithermal stage together with jamesonite, antimonite, boulangerte, falkmanite, melanikovite, aragonite, calcite, dolomite, rhodochrosite and barite.

Ag-sulphosalts are believed to precipitate in low temperatures, late in the paragenetic sequence. Keighin & Honea (1969) determined 197°C as the maximum temperature of stephanite formation. The mineral breaks down to pyrargyrite and argentite during heating above this temperature. Toulmin (1963) analyzed precipitation conditions of the pyrargyrite-proustite solid solution. They indicate it exists only up to 300°C. Below this is a miscibility gap. Comparison to conditions of such association in the other deposits worldwide, may suggest that Ag-minerals in Stan Terg have precipitated in low temperatures from hydrothermal fluids, below 200°C.

In the presented results, the content of silver in the electrum is relatively high, and it is around 48–74%. The other generation of electrum found in the Stan Terg deposit has 12–17% of Ag (authors’
unpublished data). The gold dominant electrum was found in the skarn paragenesis, together with Bi-bearing minerals.

In comparison to most of the know epithermal mineralizations, Ag content in the described electrum is relatively high (Fig. 6). Comparing to the mineralogical composition of these occurrences, the high Ag content in electrum is usually related to a rhodochrosite-banded ore with predominant sulphides, such as pyrite, sphalerite, galena, chalcoprite, tetrahedrite, Ag-minerals and stannite. Rhodochrosite from the Stan Terg has been characterized by Tomić (2003), where the assay of trace elements indicated the presence of Ag in this carbonate. We can assume that silver was transported and originated from the same hydrothermal solutions.

**Fig. 6.** Binary plot Ag vs. Au of electrum (at. %) from Stan Terg and various epithermal systems

### Application in exploration

Tetrahedrite may be of great interest in exploration. Wu & Petersen (1977) noticed that silver content in tetrahedrite is increasing with distance from the center of volcanic activity in complex hydrothermal systems. They found Ag-content variations in a district scale in a vein, base metal system, in Casapalca, Peru. Lynch (1989) analyzed the tetrahedrite from the Keno Hill Ag-Zn-Pb district in Yukon and had similar conclusions. In the external parts of this system, freibergite dominates over tetrahedrite. Another epithermal mineral indicator is cinnabar, reported often together with Ag-mineralization (e.g. Cheilletz et al. 2002).

Stan Terg deposit is the largest known base metal deposit in northern Kosovo. Since the early 20th century, it is well known that the deposit is surrounded by numerous different ore occurrences, such as Zijaqa, Melenica, Mazhiq, Rashan and Trstena (e.g. Tmava et al. 2003). These areas have been investigated by the Trepça Mining Company, but never excavated. The only signs of mining activity are ancient tunnels and shafts scattered over a wide area. The mineralization in this area has various characters (veins, stock work, replacement, disseminated) and is trapped in various lithostratigraphic unites. Previous investigations in the Stan Terg area also indicated the presence of
epithermal mineralization. Tiny cinnabar im-
pregnations and stringers, mostly within the over-
laying tuffs in the neighboring area, were reported
by Titcom et al. (1936). Pyrargyrite was reported
previously by Smejkal (1960) and Topalovič (1971),
without any precise position.

The presence of freibergite and Ag-rich tetra-
hedrite in the northern, external part of Stan Terg,
as well as previous reports of cinnabar presence
in the vicinity could mean that besides medium
temperature stages with base metals (main eco-

nomic interest) and high temperature associations
with Bi and Au (Kołodziejczyk et al. 2015), there is
also an epithermal overprint in the external parts
of the Stan Terg hydrothermal system, probably
dominated by Ag-Au-Sb±Hg mineralization.

CONCLUSIONS

1. Silver in the Stan Terg deposit occurs as
a substitution in the lattice of galena, as well
as in the form of its own minerals. The chem-
ical composition of these phases has been
confirmed with the use of an electron micro-
probe.

2. Paragenesis of silver minerals comprises pyr-
argyrite, pyrostilpnite, Ag-rich tetrahedrite,
freibergite, freieslebenite, electrum and native
antimony. These minerals occur within gale-
na-dominated ore, with minor occurrences
of pyrite, sphalerite and stannite. This kind of
mineralization is surrounded by banded-rho-
dochrosite.

3. The Stan Terg deposit is a significant Pb-Zn hy-
drothermal system, with high temperature par-
agogeneses, and low temperature phases, among
which, Ag-bearing minerals have been recently
found. The mode of Ag-association occurrence is
following main ore deposition and indicates late precipitation. Comparing to other known
data, the Ag-mineralization at the Stan Terg
mine could occur at around 200°C.

4. Similar geochemical zonality in the distribu-
tion of minor elements to hydrothermal sys-
tems formed elsewhere has been confirmed,
and further exploration in the vicinity may
lead to the discovery of new epithermal vein
systems.

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REFERENCES

Bahna B. & Chovan M., 2001. Low-sulfidation type of ep-
ithermal Au-Ag mineralization near Pukanec (Cen-
tral Slovakia Neogene Volcanic fields). GeoLines,
13, 11–17.

Berjakarevic D., 1995. Polymetallic mineral phenome-
on of Glama silver near by Gnjlane. [in:] Geološka
i metalogenija Kopaonika, 19–22 jun 1995, Beograd,
326–333.

Cheilletz A., Levresse G., Gasquet D., Azizi-Samir M., Zyadi
silver deposit: Neoproterozoic epithermal mineraliza-
tion in the Anti-Atlas, Morocco. Mineralium Deposita,
37, 8, 772–781.

Dangić A., 1993. Tetriary lead-zinc ore deposits and cal-
co-alkaline magmatism of the serbo-macedonian prov-
ince: metallogenic and geochemical characteristics, hy-
drothermal systems and their evolution. Geološki anal-
ski Balkanskoga poluostrva (Annales géologiques de la pén-
insule Balkanique), 57, 1, 257–285.

Féraud J. & Deschamps Y., 2009. French scientific coopera-
tion 2007–2008 on the Trepça lead-zinc-silver mine and
the gold potential of Novo Brdo/Artana tailings (Kosovo).
BRGM Report No RP-57204-FR.

Gemmell J.B., Zantop H. & Birnie R.W., 1989. Silver sulfos-
als of the Santo Nino vein, Fresnillo District, Zacatecas,

Grossou-Valta M., Adam K., Constantinides D.C., Prevoo-
steau J.M. & Dimou E., 1990. Mineralogy of and poten-
tial beneficiation process for the Molai complex sulphide
orebody, Greece. [in:] Gray P.M.J., Bowyer G.J., Castle
J.F., Vaughan D.J. & Warner N.A. (eds.), Sulphide depos-
its – their origin and processing, The Institution of Min-
ing and Metallurgy, 119–133.

Höller W. & Gandhi S.M., 1995. Silver bearing sulfoslates
from the metamorphosed Rampura Agucha Zn-Pb-(Ag)
deposit, Rajasthan, India. Canadian Mineralogist, 33,
1047–1057.

Hyseni S., Durmishaj B., Fetahaj B., Shala F., Berisha A. &
Large D., 2010. Trepça Ore Belt and Stan Terg mine –
geological overview and interpretation, Kosovo (SE Eu-
rope). Geologija, 53, 1, 87–92.

Keighin C.W. & Honea R.M., 1969. The system Ag-Sb-S
from 600°C to 200°C. Mineralium Deposita, 4, 157–171.

Kěpuska H., 1998. Distribušni i elementeva šperndarëse
dve mikroelemente përcjellëse ne mineralea kry-
sore xeheroformjonëse në vendburimi e plumb – zanuk
“Trepça”. Universiteti i Prishinës, Fakulteti Xehetarišë –
Metalurgišë [doctoral dissertation].

deposit, Kosovo. Faculty of Geology, Geophysics and
Environmental Protection, AGH University of Science and Technology, Cracow [MSc thesis, unpublished].

Lynch J.G., 1989. Large scale-hydrothermal zoning reflected concerning the major ore minerals and sulphosalts from the Eastern Erzgebirge, Germany.

Lynch J.G., 1989. Large scale-hydrothermal zoning reflected concerning the major ore minerals and sulphosalts from the Eastern Erzgebirge, Germany.


Mining Geology, 38, 211, 369–373 [in Japanese].


Economic Geology, 83, 8, 1737–1759.


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Ore Geology Reviews, 6, 6, 563–579.


Ore Geology Reviews, 72, 253–268.


Mineralogicheskii Zhurnal, 15, 2, 9–17.


Ore Geology Reviews, 28, 1, 1–31.


Mineralium Deposita, 39, 5–6, 608–621.


Geology Reviews, 12, 3, 173–186.


Vesnik ZGGL, 14, 7–28.


Mining and Metallurgy, 17, 9, 424–426; 17, 10, 481–484; 17, 11, 514–518 and 527; 17, 12, 584–585.


Buletini i Shkencave Gjeologjike, 2, 26–31.


Mining and Metallurgy, 17, 9, 424–426; 17, 10, 481–484; 17, 11, 514–518 and 527; 17, 12, 584–585.


Buletini i Shkencave Gjeologjike, 2, 26–31.


