A geochemical study of gersdorffite from the Trepça Mineral Belt, Vardar Zone, Kosovo

Sławomir MEDERSKI1*, Marcin WOJSŁAW1, Jaroslav PRŠEK1, Juraj MAJZLAN2, Stefan KIEFER2, Burim ASLLANI3

1. Introduction

Gersdorffite, with the nominal formula NiAsS, is a common Ni-bearing sulfarsenide occurring in various hydrothermal and magmatic sulfide systems.

Minerals of the gersdorffite-ullmannite series (GUS) belong to the pyrite structure group. Three crystal structure variants of gersdorffite have been described: an ordered structure (space group P213, ullmannite subgroup) (Bayliss and Stephenson 1967), a disordered structure (space group Pa3, pyrite supergroup) (Bayliss 1982) and distorted a disordered structure P1 (Bayliss and Stephenson 1968).

Gersdorffite incorporates Ni, Co and Fe in varying proportions (Klemm 1965) within the ternary gersdorffite–cobaltite–arsenopyrite solid solutions. The Ni and Co end-members of the series exhibit complete miscibility above 550°C, but the Ni–Fe substitution is not complete (Yund 1962; Hem and Makovicky 2004a, 2004b). Moreover, the As and S content may deviate from the nominal molar 1:1 ratio, thus forming solid solutions towards krutovite (NiAs2) and vaesite (NiS2) (Yund 1962). In addition, the extensive As–Sb substitution (towards ullmannite) results in an additional dimension of solid-solution formation (Bayliss 1969).

Geochemical data of gersdorffite in hydrothermal systems were already provided by many authors (e.g., Grapes and Challis 1999; Fanlo et al. 2004, 2006; Belkin and Luo 2008; Voudouris et al. 2008; Ahmed et al. 2009; Harms 2010; Gervilla et al. 2012; Emam and Zoheir 2013; Mikuš et al. 2013; Qiu and Zhu 2015; Radosavljević et al. 2015; Ferenc et al. 2016; Kiefer et al. 2017; Voudouris et al. 2018; Rezazadeh et al. 2020; Števko and Sejkora 2020). In addition to those hydrothermal systems, gersdorffite appears in the so-called five-element (Ni–Co–As–Ag–Bi) association, which was extensively investigated (e.g., Watkinson et al. 1975; Kissin 1992; Markl et al. 2016; Moroni et al. 2019; Scharer et al. 2019).

Nickel mineralization in the magmatic-hydrothermal systems occurring in the Vardar Zone is rare and only limited data are available. The mineralization with gersdorffite–ullmannite is part of Pb(Ag)–Zn deposits and was described from the Rogozna ore field (ROF) at the boundary between Serbia and Kosovo. It is widespread in the Kaludjer deposit, in less amount at Plakaonica, whereas it was not reported in the largest central deposit Crnac (Radosavljević et al. 2015; Boroević Šoštarić et al. 2011, 2013). Nickel mineralization in the Vardar zone

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contains sulfides, arsenides, and sulfarsenides and occurs in listvenite and vein-type mineralization. Other Ni-mineralization occurrences within the Vardar Zone were reported from several localities in Serbia: Pb–Zn–Sb Rajićeva Gora deposit (Janković and Zarić 1980), Ni–Fe–Cu(Pb, Zn, Sb) mineralization of the Asanovo Selo ore field (Stajević and Zarić 1984), the Rudnik Pb–Zn–Cu(Bi, W, Ag) ore field (Cvetković et al. 2011; Stojanović et al. 2018), the Lece Pb–Zn–(Au) deposit (Radosavljević et al. 2011), where Ni sulfides, Ni sulfantimonides, as well as Ni arsenides were reported. Additionally, Serafimovski et al. (2010) described occurrences of Ni sulfides (bravoite and polydymite) in the Cu–Au porphyry deposits of the Buchim–Damjan–Borov Dol Ore District in the territory of North Macedonia. Furthermore, the presence of GUS was described from the epithermal mineralization at Kamariza Mining District (Clemence Mine and “km3” locality) from the Lavrion ore field in Greece (Voudouris et al. 2008, 2018).

Earlier publications about the Stan Terg deposit and Kizhnica–Hajvalia–Badovc ore field did not indicate the presence of Ni minerals in the hydrothermal system (Dangić 1993; Féraud and Deschamps 2009; Kołodzieczyk et al. 2016b). New research performed on various small occurrences in the broad Stan Terg area and the Kizhnica–Hajvalia–Badovc ore field shows that Ni mineralization seems to be fairly common in the hydrothermal systems in the Vardar Zone (Bal et al. 2019; Mederski et al. 2019; Wojsław et al. 2019).

The aim of this paper is the description of spatial and textural relations on the microscopic scale and chemical variability of gersdorffite from the Ni–As–Sb–S–(Pb–Zn–Cu) assemblages in various small occurrences in the broad surrounding of Stan Terg deposit and Kizhnica–Hajvalia–Badovc ore field.

2. Geological setting

The hydrothermal field of the Pb–Zn–Ag Stan Terg deposit is located in Kosovo, within the central part of Vardar Zone in the vicinity of Kopaonik granite massif, between the Serbo–Macedonian Massif to the east and

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Fig. 1 – Simplified geological map of Vardar Zone and Trepça Mineral Belt (TMB) with marked metallogenic zones (I–III) (modified after Hyseni et al. 2010b; Strmić Palinkaš et al. 2013). I – Batlava–Artana Zone, II – Belo Brdo–Stan Terg–Hajvalia Zone, III – Crnac Zone, a – Stan Terg area, b – Hajvalia–Kizhnica–Badovc ore field.
the Dinarides to the west (Fig. 1) (Hyseni et al. 2010b). Together with the Kizhnica ore field, it is a part of the Trepça Mineral Belt (TMB) (Fig. 1). The main deposit in the area is Stan Terg, surrounded by numerous small occurrences and deposits (Fig. 2a). All deposits and occurrences have the same hydrothermal-metasomatic origin and Oligocene–Miocene age, related to the post-collisional magmatic activity (Strmić Palinkaš et al. 2013). The main lithologies in the TMB are represented by schists and phyllites of the Paleozoic crystalline basement, Triassic metasediments, and volcanoclastic rocks, Upper Triassic carbonates, as well as Jurassic ultramafic rocks (Elezaj 2009) (Fig. 1). Additionally, the Cretaceous complex is composed of series of flysch, carbonates, as well as volcanic and volcanoclastic rocks. Tertiary (Oligocene–Miocene) volcanic rocks are represented by andesites, trachytes, latites, and felsic pyroclastic rocks (Hyseni et al. 2010b, Strmić Palinkaš et al. 2013, 2016). The mineralization in the TMB area is hosted by Mesozoic carbonates at the contact with schists and the volcanic rocks of calc-alkaline affinity (at Stan Terg and Gjidoma) (Hyseni et al. 2010b) (Fig. 2a). It also occurs at the contact of serpentinites and volcanic rocks (at Vllahi, Selac, Melenica, Mazhiq, Kizhnica–Badovc) (Fig. 2b), or fills veins and veinlets in highly to moderately hydrothermally altered rocks – Mazhiq, Gumnishte, Badovc, Kizhnica. The mineralization forms elongated manto-like replacement orebodies, veins or is disseminated in the country rocks. The Stan Terg deposit and its broad surrounding is part of an extensive hydrothermal system with various mineralization styles, such as skarns, skarn-free carbonate-replacements, veins and breccias (Féraud and Deshamps 2009; Hyseni et al. 2010b; Strmić Palinkaš et al. 2013, 2016; Kołodziejczyk et al. 2015). The mineralogy of the deposits and occurrences is related to the mineralization styles and consists of various Ca–Mn–Mg–Fe carbonates, quartz, skarn silicates, magnetite, sulfides and native elements (Đangić 1993; Féraud and

![Fig. 2a – Simplified geological map of Stan Terg area (modified after Hyseni et al. 2010b).](image)

![Fig. 2b – Simplified geological map of Kizhnica–Hajvalia–Badovc ore field (modified after Hyseni et al. 2010a).](image)
Sulfides are represented mainly by pyrite, arsenopyrite, sphalerite, galena, pyrrhotite, marcasite, tetrahedrite, and chalcopyrite. Additionally, Bi sulfosalts, Bi sulfotellurides, Sn minerals, gold, Ag minerals and Pb–Sb sulfosalts were described recently (Kołodziejczyk et al. 2015, 2016a, b, 2017; Mederski et al. 2018; Węgrzynowicz et al. 2019). A similar Pb–Zn–Sb mineralization was observed in the Kizhnica–Hajvalia–Badovc ore field (Hyseni et al. 2010a; Kołodziejczyk et al. 2016b; Mederski et al. 2019), including various Pb–Sb sulfosalts, as well as accessory minerals such as Bi sulfosalts, Sn minerals and gold (Kołodziejczyk et al. 2016b; Mederski et al. 2019).

3. Methods

Ore samples for mineralogical and geochemical studies were collected from the five occurrences related to polymetallic Pb–Zn ores from the Trepça Mineral Belt: Kizhnica–Badovc, Melenica, Mazhiq, Vllahia and Selac. Samples from each locality were examined macroscopically and more than 100 thin and polished sections were prepared for mineralogical studies, including textural relationships using reflected and transmitted light microscopy. Seventeen samples with gersdorffite from all investigated mineralized rocks were selected for further studies using electron microprobe analyses (EPMA). GUS members were analyzed using a JEOL Super Probe 8230 in the Laboratory of Critical Elements at Faculty of Geology Geophysics and Environmental Protection, AGH-UST, Kraków Poland. The following operating conditions and standards were used: accelerating voltage 20 kV, beam current 20 nA and a beam diameter up to 5 µm. The following X-ray emission lines were used: CuKα, SKα, FeKα, AsLα, SbLα, BiMα, NiKα, CoKα. Natural mineral standards (FeS2) and synthetic compounds (Sb2S3, CuFeS2, Bi, Ni, Co, GaAs) were used for calibration. All interferences between the element emission lines were calculated using autocorrections based on the standard materials. The detection limits for analyzed elements in gersdorffite were as follows: Ni – 0.02 wt. %, Fe – 0.03 wt. %, Co – 0.03 wt. %, S – 0.012 wt. %, As – 0.07 wt. %, Sb – 0.02 wt. %, Bi – 0.03 wt. %, Cu – 0.025 wt. %. All analyses of GUS were calculated on the basis of the sum of cations (Fe + Ni + Co) = 1.

4. Investigated material

Samples for this study were collected from the various outcrops of the mineralized listvenite rocks in the broad Stan Terg and Kizhnica areas (Fig. 1). Listvenites are widespread in this area and were formed by hydrothermal alteration of the serpentinites in the tectonic zones or at the contact of serpentinites with volcanic rocks or the Triassic metamorphic series. They are represented by highly silicified and carbonatized rocks with macroscopically visible sulfide mineralization (Fig. 3a, b, c, d). Listvenites studied from each location differ mainly in polymetallic mineralization grades, as well as a degree of silicification and carbonate geochemistry. Relics of chromite, as well as green micas, are common in silicified rocks. Sulfides usually form veinlets, crosscutting listvenites, filling the voids, or are disseminated in highly altered rock. The most common sulfides in the veins are galena,
sphalerite, pyrite and chalcopyrite. Rarely, tetrahedrite, various Bi sulfosalts, Pb–Sb sulfosalts, arsenopyrite, marcasite and Ni minerals were observed. Disseminated sulfide mineralization in listvenites is mainly formed by pyrite and arsenopyrite with minor galena, gersdorffite and other Ni minerals. A brief description of the samples containing GUS members used for geochemical studies is shown in Tab. 1.

### 4.1. Kizhnica–Badovc

Representative samples from the Kizhnica–Badovc area were collected from the old dumps and listvenite outcrops on the transitional contact of serpentinites and andesites. Various ore types (massive – vein – banded ore and listvenite disseminated type of ore) can be found here. Massive ore with base metal mineralization consists of Fe–Mn carbonates, quartz, sphalerite, galena, Pb–Sb sulfosalts (boulangerite, senseyite, chovanite and jamesonite), pyrite, marcasite, stibnite and berthierite (Mederski et al. 2019). The Pb–Zn–Sb–Ni mineralization forms veinlets, lenses and disseminations in listvenite (Fig. 3a, b). It consists of Fe–Mn carbonates and quartz with sphalerite, galena, pyrite, chalcopyrite, tetrahedrite, GUS members (gersdorffite, Sb-rich gersdorffite and ullmanite), Pb–Sb sulfosalts (boulangerite and senseyite), with minor arsenopyrite, pyrrhotite, hematite, chromite, native gold, nickel sulfides (millerite and pentlandite) and nickel thiospinels (polydymite and violarite) (Mederski et al. 2019) (Fig. 4a, b). Individual ore samples differ mainly in Pb–Zn–Sb mineralization grades. Seven samples containing GUS members from all 45 investigated samples from this locality were selected for EPMA analyses.

### 4.2. Melenica

Samples from Melenica were collected from a 20 m × 50 m outcrop of hydrothermally altered rocks. The mineralization was formed on the contact of Jurassic basalts with the Miocene andesites. Listvenites from Melenica are highly silicified and are characterized by low-grade base-metal mineralization. At Melenica, sulfide mineralization in listvenites is mainly formed by pyrite and arsenopyrite with minor galena, gersdorffite and other Ni minerals. A brief description of the samples containing GUS members used for geochemical studies is shown in Tab. 1.
of quartz, Fe–Mn carbonates, pyrite, marcasite, galena, sphalerite, chalcopyrite, tetrahedrite, bournonite and minor Ni mineralization: GUS members (gersdorffite, Sb-rich gersdorffite and ullmannite), millerite, siegenite and violarite (Bal et al. 2019; Wojsław et al. 2019). Baryte veins are younger than those mentioned above. Two samples from 8 investigated listvenites from Melenača were selected for EPMA analyses.

4.3. Mazhiq

Samples from Mazhiq were collected along a mineralized body over a distance of 900 m on the contact of serpentinites or phyllites with volcanic rocks as well as from the surrounding listvenites. Samples are either massive, from the vein-type of mineralization, or disseminated, collected from the highly silicified rocks. Listvenite-hosted ores are silicified and characterized by the presence of disseminated pyrite crystals and aggregates of arsenopyrite, chalcopyrite, galena, pyrite, sphalerite, tetrahedrite. Massive ore consists of Fe–Mn carbonates, quartz, massive aggregates of pyrite, arsenopyrite, gersdorffite, needle-like crystals of Bi sulfosalts: bismuthinite and cosalite; galena, sphalerite and minor chalcopyrite, marcasite, tetrahedrite and native gold (Fig. 4c, d) (Bal et al. 2019; Wojsław et al. 2019; Węgrzynowicz et al. 2019). Two samples containing GUS members from all 17 investigated samples from Mazhiq were selected for EPMA analyses.

4.4. Vllahia VL4 + VLX

Samples from Vllahia were collected from two outcrops with macroscopically visible mineralization. The first one is gossan VL4 which was formed on listvenite at the contact of serpentinite and volcanic rocks. It is characterized by the presence of rare relics of base metal + nickel minerals. Only two samples with visible mineralization in typical listvenite rocks were found there. The second place VLX is a tectonic zone with hydrothermal massive sulfide vein on the contact of volcanic rocks and serpentinites. The mineral association contains galena, gersdorffite, millerite, pyrite, violarite, chalcopyrite, tetrahedrite, sphalerite, marcasite and rammelsbergite (Fig. 4c). Listvenite in gossan is rich in Ni mineralization and mineral association contain galena, tetrahedrite, Pb-Sb sulfosalts (semseyite, boulangerite and chovanite), sphalerite, gersdorffite–ullmannite, löllingite and Sb-bearing löllingite, rammelsbergite and Sb-bearing rammelsbergite, nickelene and breithauptite (Fig. 4f–h) (Bal et al. 2019; Wojsław et al. 2019; Węgrzynowicz et al. 2019). Four samples containing GUS members from all 20 investigated samples from both localities were selected for EPMA analyses.

4.5. Selac

The samples from Selac were collected from a mineralized tectonic zone in a serpentinite outcrop. Most of the nickel mineralization in Selac is hosted by veins and small lenses within dolomite veins up to 1 m in thickness. Additionally, disseminated mineralization is visible in the surrounding silicified and carbonatized serpentinites. The main gangue minerals here are dolomite, quartz, as well as magnesite and fuchsite. Mineralized zones consist of galena, sphalerite, nickeline, maucherite, gersdorffite, rammelsbergite, polydymite, marcasite, pyrite, millerite and chromite (Bal et al. 2019). In addition, the presence of Ni mineralization with a higher proportion of base-metal minerals is observed on the prolongation of the tectonic zone: galena, sphalerite, pyrite, arsenopyrite, marcasite, tetrahedrite with minor Ni phases: nickelene and gersdorffite. One sample from 10 investigated mineralized rocks from Selac was selected for EPMA analyses.

5. Results

Representative results of the chemical analyses of GUS members from each locality are shown in the supplementary electronic table. Additionally, average EPMA data are presented in Tab 2.

5.1. Kizhnica–Badovc

Members of GUS from the Kizhnica area occur as disseminated separate euhedral or subhedral crystals up to 150 μm as well as aggregates with galena and nickel minerals (Fig. 4a). Additionally, these crystals often occur as numerous accumulations at the edges of galena and sphalerite aggregates (Fig. 4b). Some of the gersdorffite crystals show patchy zonation (Fig. 5a). Other crystals are characterized by typical oscill-
Tab. 2. Average EPMA data of gersdorffite, ullmannite and arsenopyrite from Trepça Mineral Belt (in wt. %).

| Locality         | Mineral                      | Ni | Co | Fe | Sb | As | Apfu |
|------------------|------------------------------|----|----|----|----|----|------|
| Gersdorffite     |                              |    |    |    |    |    |      |
| Melenica         | Gersdorffite                 | 57 | 32 | 42 | 16 | 4.1 | 9.15 |
| Vllahia VLX      | Sb-bearing gersdorffite      | 35 | 53 | 3 | 8.4 | 22.4 | 83.2 |
| Vllahia VL4      | Arseneopyrite                | 29 | 28 | 62 | 5.5 | 4.4 | 22.4 |
| Kizhnica         | Gersdorffite                 | 14 | 73 | 7 | 3.5 | 4.1 | 19.1 |
| Selac            | Ullmannite                   | 15 | 33 | 66 | 0.1 | 0.01 | 0.2 |
| Melenica         | Gersdorffite                 | 8  | 28 | 62 | 0.02 | 0.04 | 0.01 |
| Kizhnica         | Ullmannite                   | 3  | 31 | 71 | 0.02 | 0.04 | 0.01 |

| Mineral          | Apfu                          | Ni  | Co  | Fe  | Sb  | As   | (As+Sb)/S |
|------------------|-------------------------------|-----|-----|-----|-----|------|------------|
| Gersdorffite     |                               | 57  | 32  | 42  | 16  | 4.1  | 9.15       |
| Sb-bearing gersdorffite |                   | 35  | 53  | 3   | 8.4 | 22.4 | 83.2       |
| Arseneopyrite    |                               | 29  | 28  | 62  | 5.5 | 4.4  | 22.4       |
| Gersdorffite     |                               | 14  | 73  | 7   | 3.5 | 4.1  | 19.1       |
| Ullmannite       |                               | 15  | 33  | 66  | 0.1 | 0.01 | 0.2       |
| Gersdorffite     |                               | 8   | 28  | 62  | 0.02 | 0.04 | 0.01      |
| Ullmannite       |                               | 3   | 31  | 71  | 0.02 | 0.04 | 0.01      |

| Locality         | Mineral                      | Ni  | Co  | Fe  | Sb  | As   | (As+Sb)/S |
|------------------|------------------------------|-----|-----|-----|-----|------|------------|
| Melenica         | Gersdorffite                 | 57  | 32  | 42  | 16  | 4.1  | 9.15       |
| Vllahia VLX      | Sb-bearing gersdorffite      | 35  | 53  | 3   | 8.4 | 22.4 | 83.2       |
| Vllahia VL4      | Arseneopyrite                | 29  | 28  | 62  | 5.5 | 4.4  | 22.4       |
| Kizhnica         | Gersdorffite                 | 14  | 73  | 7   | 3.5 | 4.1  | 19.1       |
| Selac            | Ullmannite                   | 15  | 33  | 66  | 0.1 | 0.01 | 0.2       |
| Melenica         | Gersdorffite                 | 8   | 28  | 62  | 0.02 | 0.04 | 0.01      |
| Kizhnica         | Ullmannite                   | 3   | 31  | 71  | 0.02 | 0.04 | 0.01      |

As-bearing gersdorffite crystals are more Fe-rich (Fig. 5b). Dissolution and replacement of some GUS zones by younger hydrothermal galena and quartz are common (Fig. 5c,d). Corroded rounded pyrite crystals sometimes form the cores of gersdorffite aggregates (Fig. 4b; 5e). This pyrite is enriched in Co (up to 4 wt. %) and Ni (up to 2.4 wt. %). Gersdorffite end-member compositions are the most widespread in the Kizhnica GUS population. Gersdorffite composition from this site can be expressed by the formula (Ni0.05-0.15Fe0.66–0.99)0.84–0.91Sb0.00–0.09As0.84–1.03S0.87–1.15 (Tab. 2). Two types of substitution can be observed: As–Sb, as well as Ni–Fe. The first one is manifested by the variable As–Sb content, from low to those belonging to ullmannite (Fig. 7a). The formula of Sb-rich gersdorffsite can be expressed as (Ni0.95–1.05Fe0.00–0.05)0.83–1.00Sb0.00–0.09As0.89–1.05S0.88–1.00 Most of the measurements of Sb-rich gersdorffite were carried out within the oscillatory zoned crystals (Fig. 5b). One of the GUS aggregate from the quartz-galena breccia is characterized by the presence of a zoned aggregate composed of GUS end members, as well as Sb-bearing gersdorffite (Fig. 5e). The crystallization sequence in this case is: gersdorffite I > ullmannite > Sb-enriched gersdorffite > gersdorffite II. The contact between the earlier gersdorffite and ullmannite is sharp, while that between ullmannite and later gersdorffite shows gradual transition or fluctuations. The Ni–Fe substitution (Fig. 8) is caused by secondary enrichment related to the migration of younger fluids, responsible for base-metal mineralization and observed within the irregular zones often near crystal cracks (Fig. 5a,b). Therefore, the most Fe-enriched gersdorffite is Sb-poor and is found within the highest-grade listvenite-type Pb–Zn ore. Gersdorffite is mostly Fe- and Co-poor. Arsenopyrite from the Kizhnica area occurs as euhedral crystals with sizes up to 200 µm in the quartz–palierite breccia. The formula for arsenopyrite is Fe(As0.82–0.95Sb0.00–0.02)0.84–0.93S0.87–1.11 (Tab. 2). Arsenopyrite is enriched in S (1.07–1.11 apfu)
Gersdorffite from the Trepça Mineral Belt, Vardar Zone, Kosovo
and depleted in As (0.82–0.93 apfu). Limited As–S substitutions were observed, and the As/S ratio is 0.8–1.05 in gersdorffite, 0.60–0.89 in Sb-bearing gersdorffite and 0.76–0.87 in arsenopyrite.

5.2. Melenica

The gersdorffite in the Melenica samples is disseminated in listvenites and hard to identify by reflected-light microscopy due to the small crystal size (up to 20 μm). The Melenica GUS members occur as anhedral aggregates together with millerite, violarite and siegenite. GUS members are frequently replacing the boundaries of Ni–Co sulfide crystals and are replaced by younger Pb–Zn–Cu sulfide minerals (Fig. 5f,g,h; 6a). Chemical analyses identified a gersdorffite–ullmannite series (Fig. 7a). The average chemical formulae of individual GUS populations divided by their As/(As + Sb) ratios can be presented as: (Ni$_{0.73–0.99}$Co$_{0.00–0.21}$Fe$_{0.00–0.06}$)$_{∑1.00}$As$_{0.71–1.08}$Sb$_{0.00–0.11}$S$_{0.97–1.09}$ for gersdorffite, (Ni$_{0.94–0.99}$Co$_{0.00–0.05}$Fe$_{0.00–0.01}$)$_{∑1.00}$As$_{0.51–0.74}$Sb$_{0.23–0.45}$S$_{0.96–0.98}$ for Sb-bearing gersdorffite and Ni(Sb$_{0.88}$As$_{0.09–0.1}$)$_{∑}$S$_{0.97–1.09}$S$_{0.98–0.99}$ for ullmannite (Tab. 2). The measured Co concentrations in gersdorffite are the highest within the Trepça Belt (Fig. 8). The Sb-rich gersdorffite and ullmannite are characterized by low Co and Fe contents. The chemical data indicate limited cation substitution with the increasing Sb content. Sb-rich GUS and ullmannite are intergrown with the As-end member gersdorffite and are related to the irregular Sb-rich zones. Limited As–S substitution was observed; the As/S ratio of gersdorffite is 0.66–1.05 and in Sb-bearing gersdorffite 0.52–0.74.

5.3 Mazhiq

In Mazhiq, gersdorffite is one of the main ore minerals with crystals up to a few mm in size. It occurs as a) irregular masses rimming crystals of arsenopyrite, pyrite I, or pyrite II (Fig. 4c; 6b) and b) intergrowth of euhedral to subhedral crystals with oscillatory zoning (Fig. 4d). Gersdorffite is often fractured parallel to ore zonation and is commonly porous, especially in the outer parts of crystals. Free spaces between consecutive gersdorffite generations are filled with quartz (Fig. 6c).
Arsenopyrite is often found in oscillatory zoned gersdorffite and forms a sharp boundary with its host (Fig. 6c, d). Moreover, arsenopyrite that is crystallized in open cavities appears as euhedral crystals up to 500 μm in size, commonly showing corroded boundaries, overgrown by gersdorffite and Sb-rich gersdorffite. BSE images show zonation caused by enrichment in Fe as well as the As–Sb substitution (Fig. 6c–e). The average formula of gersdorffite is (Ni$^{0.77–0.99}$Fe$^{0.00–0.26}$Co$^{0.00–0.03}$)$^{1.00}$(As$^{0.84–1.03}$Sb$^{0.00–0.14}$)$^{0.96–1.05}$S$^{0.89–0.97}$ (Tab. 2). On the cation position, only Fe–Ni substitution is observed (Fig. 8). Gersdorffite enriched in Fe usually composes the outer part of the zoned crystals and is assumed to be formed by late fluids responsible for the base-metal mineralization (Fig. 6c). An intermediate As–Sb member, with the composition (Ni$^{0.84–0.98}$Co$^{0.02–0.12}$Fe$^{0.00–0.04}$)$^{1.00}$(As$^{0.97–1.04}$Sb$^{0.01–0.05}$)$^{0.98–1.09}$S$^{0.91–1.04}$ (Tab. 2), was found as zones within inner, non-porous part of oscillatory zoned GUS aggregates (Fig. 6c, d). Sb-bearing gersdorffite tends to be Fe- and Co-poor. On the ternary diagram of the system Ni–Co–Fe, gersdorffite extends directly towards arsenopyrite composition (Fe trend). However, there is a miscibility gap (Fig. 8). Gersdorffite from this site has the highest content of Fe (0.26 apfu) among the occurrences from the Trepça Mineral Belt. The As/S ratio is 0.90–1.16 in gersdorffite and 0.78–0.9 in Sb-bearing gersdorffite. The formula for arsenopyrite from Mazhiq is (Fe$^{0.85–0.98}$Ni$^{0.00–0.13}$Co$^{0.00–0.07}$)$^{1.00}$(As$^{0.87–0.99}$Sb$^{0.00–0.02}$)$^{2.09–1.07}$S$^{0.95–1.07}$ (Tab. 2), with only slight Ni and Co enrichment. Arsenopyrite occurs as tiny core crystals (up to 10 μm) within oscillatory zoned gersdorffite crystals (Fig. 6d) or as euhedral crystals (up to 30 μm) overgrowing masses of GUS (Fig. 6b). Moreover, it is associated with chalcopyrite and pyrite and is often replaced by sulfsalts.

5.4. Vllahia (Vlx)

Gersdorffite occurs here as tiny (up to 20 μm in size) anhedral to subhedral crystals or aggregates together with rammelsbergite, all enclosed in galena. Gersdorffite frequently replaces rammelsbergite and may form horseshoe-shaped zonal aggregates (Fig. 4e; 6f). The formula of gersdorffite from Vllahia can be expressed as (Ni$^{0.84–0.98}$Co$^{0.02–0.12}$Fe$^{0.00–0.04}$)$^{1.00}$(As$^{0.97–1.04}$Sb$^{0.01–0.05}$)$^{0.98–1.09}$S$^{0.91–1.04}$ (Tab. 2). The chemical composition of gersdorffite from Vllahia is nearly stoichiometric, with only a slight Co enrichment (Fig. 8).

5.5. Vllahia (Vl4) gossan

Gersdorffite forms subhedral to euhedral intergrowths up to 50 μm in size and is overgrown by younger galena and Pb–Sb sulfsalts (Fig. 4f, h) or occurs as aggregates together with rammelsbergite disseminated in silicified listvenite (Fig. 6g). Gersdorffite edges and cores are
frequently corroded and replaced by quartz, galena, pyrite, or Pb–Sb sulfosalts (Fig. 4h). This phenomenon led to the development of skeletal textures, i.e., caries texture (Fig. 4f) and atoll texture (Fig. 4g). Gersdorffite is associated with rammelsbergite, löllingite, Sb-bearing löllingite, breithauptite and ullmannite. The chemical composition of gersdorffite can be expressed as 

\[
\left( \text{Ni}^{0.95–0.99} \text{Co}^{0.00–0.03} \text{Fe}^{0.00–0.04} \right) \sum 1.00 \left( \text{As}^{1.23–1.43} \text{Sb}^{0.04–0.15} \right) \sum 1.30–1.52 \text{S}^{0.53–0.67}.
\]

The substitution is limited at the cation position. On the other hand, gersdorffite is enriched in As (1.23–1.67 apfu) and depleted in S (0.32–0.68 apfu) in comparison with stoichiometric gersdorffite (Fig. 7a, b). The Vllahia (VL4) gersdorffite represents the As-richest composition within the Trepça Belt, approaching rammelsbergite/krutovite (Fig. 7a, b). The As/S ratio varies from 1.86 to 2.67. Chemical analyses (EPMA) identified one ullmannite crystal with the composition Ni(Sb\^{0.87–1.19}As\^{1.01–0.03})\sum 1.01\text{S}^{0.81–0.64} (Tab. 2). It is characterized by stoichiometric content of Ni, significant enrichment of As (1.15–1.33 apfu) and depletion of S (0.64–0.81 apfu) (Fig. 7a, b). Gersdorffite from Selac, together with Vllahia VL4 has the As-richest composition among all localities. There is no correlation among cations and the ratio As/S is 1.44 to 2.07, suggesting that As content is independent of the metal content.

6. Discussion

6.1. Paragenetic position of gersdorffite

Polymineral aggregates are corroded, possess cellular textures and are indicative of the high intensity of replacement processes (Fig. 6h). Gersdorffite from Selac is associated with sphalerite and tennantite and has the composition Ni(As\^{1.33–1.15}Sb\^{0.01–0.03})\sum 1.33–1.18\text{S}^{0.81–0.64} (Tab. 2). It is characterized by stoichiometric content of Ni, significant enrichment of As (1.15–1.33 apfu) and depletion of S (0.64–0.81 apfu) (Fig. 7a, b). Gersdorffite from Selac, together with Vllahia VL4 has the As-richest composition among all localities. There is no correlation among cations and the ratio As/S is 1.44 to 2.07, suggesting that As content is independent of the metal content.
tained results, the two main types of paragenetic sequence with nickel mineralization and GUS are distinguished.

The first one is characterized by crystallization of Ni sulfarsenides and sulfantimonides after Ni–Co–Fe sulfides (millerite, violarite, siegenite, polydymite and bravoite) precipitation and was observed in Kizhnica–Badovc and Melenica localities. At Melenica, Ni–Co–Fe sulfides were transformed into GUS, which forms typical outer rims (Fig. 5f). Most GUS from Kizhnica occurs in the form of euhedral crystals, but rare mineral aggregates are observed, where it grows around earlier sulfides – millerite, polydymite and violarite. There are no Ni arsenides in the paragenetic sequence at these locations.

On the other hand, the Ni sulfarsenides from Vllahia (VLX and VL4) and Selac were formed before Ni–Fe arsenide and diarsenide (nickelene, rammelsbergite and maucherite) precipitation. However, sulfarsenides from Vllahia VL4 and Selac have a non-stoichiometric composition with As > S. Gersdorffite from Selac is rare and forms residues after arsenides replacement (nickelene–rammelsbergite–maucherite massive aggregates). Additionally, the replacement by rammelsbergite and löllingite is the characteristic feature for gersdorffite from Vllahia VL4. Sulfarsenides from Vllahia VL4 and Selac (with non-stoichiometric composition, where As > S) seem to crystallize from primary, intermediate As-rich solution to become unstable and further replaced into arsenides, representing an environment with increasing As activity. The opposite trend was reported by Radosavljević et al. (2015) in Kaludjer from Rogozna Ore Field in Kosovo, where nickelene was replaced by sulfantimonides or sulfarsenides. Vllahia VLX confirms the trend described by Radosavljević et al. (2015), where the stoichiometric gersdorffite replaces rammelsbergite to form a banded aggregate, which suggests variable As activity. A transformation of arsenides to sulfarsenides was described in hydrothermal Ni–Co–Fe–As veins in Dobšiná (Kiefer et al. 2017) and the Zemberg-Terézia vein system near Dobšiná (Števko and Sejkora 2020). Additionally, paragenetic sequences from Vllahia VLX and Selac are characterized by the presence of Ni sulfides (millerite, polydymite and violarite) at the last stage of the Ni crystallization sequence.

However, gersdorffite is the only nickel phase in the paragenetic sequence of polymetallic Cu–Bi–Ni–Au mineralization from Mazhiq. GUS does not coexist with both Ni–Co–Fe sulfides and arsenides. The co-existence of gersdorffite with hydrothermal Bi–Au association is raised in a subsequent paragraph.

6.2. Chemical composition of GUS

Gersdorffite recognized at the studied localities within Trepça Mineral Belt comprises two populations concerning As/S ratio, which is a hint of two different hydrothermal fluids. The low As/S ratio of Kizhnica–Badovc (0.60–1.06), Mazhiq (0.79–1.16), Melenica (0.52–1.05), Vllahia (VLX) (0.98–1.14) indicate decreasing As concentrations in the hydrothermal fluids in relation to the fluids from Vllahia (VL4) (1.86–2.67) and Selac (1.44–2.07) (Tab. 2). The former localities are characterized by relatively homogenous compositions of gersdorffite in comparison to Vllahia (VL4) and Selac. This reveals the fluctuation of the compositions of the hydrothermal fluids during crystallization. Harms (2010) found more clear and distinguishable populations between post-Variscan vein deposits and strata-bound mineralization concerning the variability of As/S ratio and homogeneity of hydrothermal fluids.

However, a complete GUS solution (gersdorffite–ullmannite) is found in the Trepça Mineral Belt (Radosavljević et al. 2015; this study). Otherwise, Harms (2010) show only limited content of Sb up to 0.05 apfu. Stoichiometric ullmannite is rare in studied samples from TMB and is observed only in samples from Kizhnica–Badovc, Melenica and Vllahia VL4. Ullmannite was also described by Radosavljević et al. (2015) in the Rogozna Ore Field. Additionally, As–Sb substitution is observed within the oscillatory zoned crystals (Fig. 5b, c) from typical listvenite type of mineralization from Kizhnica, as well as in massive gersdorffite aggregates from Mazhiq. The majority of Sb-bearing gersdorffite in Mazhiq occurs within the inner part of crystals, which implies that As–Sb substitution is present mainly at an early stage, with an environment with a high Sb concentration. As–Sb exchange of gersdorffite–ullmannite solid solution is not continuous; there is a gap between ullmannite and Sb-bearing gersdorffite (Fig. 7a). Analogous behavior was reported by Radosavljević et al. (2015) and the opposite by Voudouris et al. (2018).

In addition to that, Harms (2010) gathered reliable evidence that a small extent of substitution on cation sites together with a narrow range of As/S ratio (0.88–1.01) of post-Variscan vein deposits imply thermochemical As reduction and mixing (homogenization) of the hydrothermal fluids leading to ore precipitation far away from the site of ore deposition. Meanwhile, the Co–Fe trend and broader As/S ratio (1.06–1.21) in the strata-bound mineralization evoke the mixing of the hydrothermal fluids close to the site of ore deposition. It seems to be with disagreement with the data collected within the Trepça Mineral Belt, where only Kizhnica–Badovc and Mazhiq are assumed to occur in proximity to the site of ore deposition concerning an appearance as massive ore. The latter author also suggests that the variability in the As/S ratios could be a consequence of disequilibrium crystallization, which might be the case for Selac where the replacement process is well-developed. What is more, the relationship in Selac Ni-rich association might be regarding low-temperature re-equilibrium and replacement of preceding gersdorffite into nickelene, maucherite and rammelsbergite.
Melenica and Vllahia (VL4) display a Co + Fe substitutional trend on the ternary Ni–Co–Fe diagram (Fig. 8), which is a result of a coupled substitution of Ni by Co and Fe. According to Fanlo et al. (2004), Co + Fe substitution is found in mineral associations in an environment of high As activity expressed as a presence of skutterudite, diarsenides, or sulfarsenides. On the other hand, only Vllahia (VL4) samples are enriched with arsenides (rammelsbergite and nickeline), while Melenica, which presents a stronger Co + Fe trend (Fig. 8), none of those minerals were detected. Additionally, Selac gersdorffite, which is replaced by maucherite and later by galena, is depleted in Co. Kizhnica–Badovc and Mazhiq, concerning the lack of Co, are characterized by a smooth Fe trend (Fig. 8). Gersdorffite enriched in Fe from Kizhnica–Badovc and Mazhiq usually occurs in the outer part of the crystals (Fig. 6c) or forms the irregular zones, often near crystal cracks (Fig. 5a, b). It is assumed to be formed by late fluids responsible for base-metal mineralization.

**6.3. Formation temperature**

Based on the geochemical data obtained in this study, it may be virtually impossible to directly determine the crystallization temperatures of the GUS in each locality from the TMB. So far, scientific studies on gersdorffite crystallization temperatures (Yund 1962; Klemm 1965; Hem and Makovicky 2004a, 2004b) have been done on dry systems with unrealistic temperatures encountered in hydrothermal conditions. Therefore, the geochemical data of other mineral phases (arsenopyrite and sphalerite) from studied occurrences were used to determine the formation conditions. An arsenopyrite geothermometer (Kretschmar and Scott 1976; Sharp et al. 1985) was used for listvenites from Mazhiq and Kizhnica–Badovc. The formation temperatures of sphalerite in Kizhnica were estimated based on a GGMFis geothermometry (Frenzel et al. 2016).

Arsenopyrite is formed simultaneously with gersdorffite in Mazhiq samples as well as gersdorffite in Kizhnica. In both occurrences, arsenopyrite is found together with pyrite as the only sulfide phase. To estimate the formation temperature of arsenopyrite, the As content can be used (Kretschmar and Scott 1976; Sharp et al. 1985). Three out of 19 and 2 out of 3 chemical analyses of arsenopyrite from Mazhiq and Kizhnica, respectively, were selected, which satisfy the conditions: 1) Ni + Co + Sb ≤ 1 wt. %, 2) the total does not deviate more than 2 wt. % from 100 wt. %. Thus, the As values of arsenopyrite fall in the "pyrite + arsenopyrite" zone of the buffered assemblages involving arsenopyrite in the Fe–As–S system.

The temperature is constrained from 440 °C to 495 °C and –6 to –4.5 log $f(S_2)$ for arsenopyrite from Mazhiq (Fig. 10). Arsenopyrite appears as the first phase together with gersdorffite and pyrite in the paragenetic sequence in

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**Fig. 10** – Activity of log $f(S_2)$ and temperature (T) projection of the stability field of arsenopyrite (after Kretschmar and Scott 1976, Sharp et al. 1985).
Mazhiq. Temperatures obtained for arsenopyrite seem to be consistent with the fact that Mazhiq assemblage was formed close to the deposition site based on As/S ratio in gersdorffite. A similar range of formation temperatures (400–450 °C) of arsenopyrite was estimated by Voudouris et al. (2018) in the Clemence deposit from Kamariza Mining District in Greece. The authors suggest the possible overestimation of temperatures of arsenopyrite from the Ni–Bi–Au assemblage from the Clemence deposit by comparing the results from fluid inclusion studies.

On the other hand, the temperature is constrained from 274 °C to 370 °C and yielded –12 to –8 log f(S₂) for arsenopyrite from Kizhnica (Fig. 10). Measured arsenopyrite (sample KZH2_12) forms euhedral crystals which overgrow sphalerite aggregates. Additionally, GGIMFis geothermometry (“Ga, Ge, In, Mn and Fe in sphalerite”) (Frenzel et al. 2016) was used to determine the crystallization temperature of the same sphalerite from the quartz-sphalerite breccia and it yielded temperatures from 330 °C to 370 °C (Mederski, unpublished). Therefore, it is assumed that arsenopyrite crystallizes with the decreasing temperatures of the hydrothermal fluids and is connected with the younger, base-metal stage of precipitation. Moreover, sphalerite from listvenites (e.g., samples SHA02_02) was formed at similar temperatures (340–390 °C) (Mederski, unpublished). Comparable temperatures have been estimated for sphalerites coexisting with GUS from Rogozna Ore Field by Radosavljević et al. (2015). The crystallization temperature of the main sulfide association is estimated at 390–430 °C (Crnac) and 305–440 °C (Plakaonica) (using the distribution of cadmium and manganese between coexisting galena and sphalerite) (Radosavljević et al. 2015).

### 6.4. Bi mineralization related to GUS occurrences (post-Ni mineralization)

The occurrence of GUS from Mazhiq is significantly different from other studied GUS locations from TMB, mainly due to the presence of bismuth sulfosalts (bismuthinite and cosalite) in post-Ni mineralization. The co-existence of the GUS, as well as the Bi minerals, is known for many locations in the Balkan area (Voudouris et al. 2008, 2018; Vavelidis and Melfos 2004; Tombros et al. 2007, 2008; Radosavljević et al. 2016). The Cu–Bi–Ni–Au mineral assemblage occurring in Mazhiq shows many similarities with polymetallic Ni–Bi–Au assemblage from the Kamariza Mining District (Clemence Mine and “km3” locality) from the Lavrion ore field in Greece (Voudouris et al. 2008, 2018). Voudouris et al. (2018) described, e.g., pyrite, arsenopyrite, chalcopyrite, native Au, bismuthinite, Ag-sulfosalts and oscillatory zoned GUS. The authors proposed a mixed hydrothermal–magmatic origin of the mineralization in the Kamariza region. Furthermore, the Bi–Ag–Cu Kapsalina occurrence in Thasos islands, Northern Greece with gersdorffite was described by Vavelidis and Melfos (2004). The Bi–Ag–Cu mineralization hosted in quartz veins consists of chalcopyrite, pyrite, bismuthinite, Bi–Ag enriched tetrahedrite group minerals, and two types of gersdorffite (Fe-poor and Fe-rich). The Ni–Fe substitution observed in gersdorffite from Kapsalina caused by secondary enrichment is a common feature for occurrences in Vardar Zone (Mazhiq and Kizhnica). In addition, as in Mazhiq, tetrahedrite replaced gersdorffite. On the other hand, the authors described in Kapsalina that crystallization of bismuthinite took place before gersdorffite, while in Mazhiq, bismuthinite crystallized during the last hydrothermal stage.

As mentioned above, the mineral association in Mazhiq is characterized by the presence of native Au and Bi sulfosalts (bismuthinite and cosalite). A similar Bi–Au rich mineralization was reported from Stan Terg Pb–Zn–Ag deposit (Kolodziejczyk et al. 2015, 2017). Kolodziejczyk et al. (2015, 2017) described Bi-sulfotelurides associated with Pb–Bi–Sb sulfosalts and galena in pyrite–pyrrhotite–galena-rich, in both skarn and hydrothermal ore bodies. The Stan Terg association exhibits a lack of nickel minerals in the system. Moreover, on the east side of Mazhiq locality, the occurrence of Bi–Au rich ores (with Bi sulfosalts: cosalite, bismuthinite, aikinite; native Bi and native Au; without Ni minerals) were reported (Mederski et al. 2018; Węgrzynowicz et al. 2019). Therefore, both Mazhiq localities could be assumed as an N–NE continuation of the Bi–Au rich ore bodies of the Stan Terg hydrothermal system. However, within Kizhnica–Hajvalia–Badovc ore field, a similar Bi–Au system is observed in the eastern part of the ore field, although it is not linked to listvenitization (Mederski, unpublished data).

Additionally, members of GUS are widespread within other types of polymetallic occurrences within the Balkan region, e.g., in Panormos Bay Au–Ag–Te mineralization (Tombros et al. 2007) or Čumavići Sb–Zn–Pb–Ag deposit (Radosavljević et al. 2016), however, with no particular mineralogical studies about the geochemistry of gersdorffite.

### 7. Conclusions

The previously unknown Ni mineralization was described in hydrothermally altered serpentinites (listvenites) from the broad Stan Terg area and Kizhnica–Hajvalia–Badovc ore field from Trepça Mineral Belt. Gersdorffite is the dominant Ni phase in the Ni–As–Sb–S association (Kizhnica–Badovc, Mazhiq, Vllahia VLX, Vllahia VL4, except Melenica and Selac) and forms disseminated crystals or aggregates with other Ni phases and base metal sulfides in listvenite rocks.
The two main types of paragenetic sequence with nickel mineralization and GUS are distinguished in TMB. The first one is characterized by the sequence: 1) Ni–Fe–Co sulfides; 2) Ni-sulfarsenides and sulfantimonides; and is observed in Kizhnica–Badovc and Melenica. The second type of sequence includes 1) Ni-sulfarsenides; 2) Ni–Fe arsenide and diarsenide; 3) ± Ni sulfides. Second type of sequence occurs in Vllahia VLX, Vllahia VL4 and Selac. However, gersdorffite is the only nickel phase in the paragenetic sequence from Mazhiq.

Gersdorffite recognized in surroundings of Stan Terg area and Kizhnica–Hajvalia–Badovc ore field is characterized by three substitutional trends: Fe + Co/Ni (all localities, except Selac); As/Sb (Kizhnica, Mazhiq, Vllahia VLX) and As + Sb/S (Selac and Vllahia VL4). Variability among the cations, As–Sb, as well as As + Sb/S, indicates the fluctuation of As, Sb and S activity in hydrothermal fluids.

The As/S ratio differentiates two hydrothermal fluids: narrow range and low values (Kizhnica, Mazhiq, Melenica, Vllahia VLX) suggest depressed As activity, mixing in the proximity of deposition site (Kizhnica, Mazhiq), broad range and elevated values: increased As activity, more inhomogeneous fluids and disequilibrium crystallization (Selac, Vllahia VL4).

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