Geochronological constraints on the carbonate-sulfarsenide veins in Dobšiná, Slovakia: U/Pb ages of hydrothermal carbonates, Re/Os age of gersdorffite, and K/Ar ages of fuchsite

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The abandoned deposit Dobšiná in eastern Slovakia used to be exploited for Ni–Fe–Co sulfarsenides in siderite-ankerite hydrothermal veins. These veins are locally accompanied by alteration zones of fuchsite-quartz-carbonate rocks. The Ni–Fe–Co sulfarsenides are postdated by chalcopyrite and tetrahedrite. This work aims to determine the age of these mineralizations utilizing U/Pb, Re/Os, and K/Ar dating of selected minerals. U/Pb dating of hydrothermal siderite and ankerite gave ages from 145 ± 5 Ma to 114 ± 24 Ma. They broadly coincide with the metamorphic peak in the Gemeric Unit and the formation of the Gemeric cleavage fan (CGF) during partial exhumation of this unit. Hydrothermal gersdorffite was dated (Re/Os) to 93.6 ± 0.9 Ma, the time of exhumation of the Gemeric unit. No data were extracted for the younger Cu sulfides. By correlation with other units in the Western Carpathians, they could be assigned Late Cretaceous age, corresponding to welding of the Gemeric and Veporic units and the formation of the Trans-Gemeric shear zone. The fuchsite rocks were dated (K/Ar) to 110 Ma, but this datum conflicts with textural evidence, which shows that these rocks are older than the hydrothermal carbonates. The age of 110 Ma is therefore considered to be cooling age after the reset of the geochronometer by the Alpine metamorphism, in agreement with earlier data. The geochronological results are supported by structural measurements and field observations. The main mineralized zone, investigated in one of the accessible adits, conforms with the Variscan – ?Alpine metamorphic foliation. There are also younger, less significant structures whose relevance to the ore mineralization is not yet clear.

Keywords: hydrothermal mineralization, geochronology, ore mineralogy, isotopic dating

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1. Introduction

The ore mineralizations around the small township Dobšiná in eastern Slovakia include many hydrothermal veins that contain siderite, quartz, arsenides, and sulfides of Ni, Co, Cu, Sb (Halahyjová-Andrusovová 1961, 1964; Grecula et al. 1995; Mesarčík et al. 2001; Kiefer et al. 2017). Additionally, metasomatic bodies of siderite/ankerite were mined here in the past. The Ni–Fe–Co mineralization in Dobšiná is hosted by the Gemeric Unit, one of the major thick-skinned tectonic units of the Central Western Carpathians (Fig. 1). Beside the ores in Dobšiná, the Gemeric Unit hosts more than 1200 ore bodies (Fig. 1; Grecula et al. 1995) that can be roughly divided into siderite-quartz-sulphide and quartz-stibnite assemblages. Many of the ore bodies were of significant economic importance in the past.

An excellent summary of the development of the ideas and perception of the hydrothermal Ni–Fe–Co mineralization in Dobšiná was written in the Slovak language by Mesarčík et al. (2001) and a digest thereof will be given here. The early studies of Böckh (1908) and Woldřich (1912) link the hydrothermal mineralization to the gneiss-amphibolite complex, granites, or both. Woldřich (1912, 1927) anticipated Early Carboniferous age of the veins because he assumed that the post-mineralization tectonics was of Upper Carboniferous age. Ahlburg (1913) found limonite-rich pebbles in Carboniferous conglomerates, assumed that these conglomerates represent weathered siderite ores and postulated their pre-Carboniferous age.
Papp (1919) linked the ores to the quartz porphyries and presumed their Permian age. The first views that there may be two separate stages of mineralization can be attributed to Ulrich (1928). The older stage could be connected to the basic rocks of the gneiss-amphibolite complex; the younger one is a result of remobilization. Other authors seek the source of the mineralization in the granitic rocks (Kettner 1921; Hynie 1922), but their ideas about the age of these granites diverge. Zoubek (1937), in contrast to all these studies, identified the Alpine metamorphism as the source of the mineralization. Hence, the dichotomy of the Variscan versus Alpine age of the veins dates back to the early years, although many details of geology and geochronology of the host rocks were unknown at that time.

The Alpine age of the veins was proposed and defended by Máška (1957), Varček (1962, 1985), and Bernard (1963). Some of them linked the mineralization to the granites, assuming their Cretaceous age. Rozložník (1989) insisted on the Cretaceous (or even younger) age even though at that time, the Permian age of the granites was already known. On the other hand, Ilavský (1986) maintained that the mineralizations are Variscan, connected to the voluminous granitic magmatic activity in those times.

The dichotomy persisted until today. Currently, there are two competing models that address the origin and timing of the ore mineralizations in the Gemicer Unit. Radvanec et al. (2004) linked the siderite-quartz-sulfide
ores to Permian regional fluid flow, initiated by the uplift of the Variscan metamorphic core complex. This assignment is based on ‘available indirect data’ and Radvanec et al. (2004) also stated that ‘more geochronological data both on metamorphic rocks and individual mineralization stages are needed to clarify all relationships unambiguously’. On the other hand, Hurai et al. (2006, 2015) tied the mineralizations to the deeply buried rocks complexes under the stack of nappes thrust over the Gemic Unit in Cretaceous times. The deep formation environments were confirmed also by the finds of superdense CO₂ (Urban et al. 2006) and N₂ inclusions (Hurai et al. 2008), which are incompatible with the regional fluid flow of a rock complex that is being exhumed. The Alpine green-schist facies metamorphic condition is also indicated by zircon fission-track data, which yielded ages of 115–75 Ma. Most probably, these data refer to two stages of the Gemic Unit evolution during the Alpine orogeny. The older ages can be related to thrusting or final emplacement of the Gemic Unit onto the Veporic Unit (115–100 Ma) and the younger ages are related to final exhumation of the Gemic Unit during the orogenic collapse (90–75 Ma) (cf. Plašienka et al. 2007; Králiková et al. 2016). The Alpine (mostly Cretaceous) ages were also confirmed by chemical electron-microprobe (CHIME) and laser-ablation ICP-MS dating of accessory monazite associated with the gangue minerals of the quartz-tourmaline stage. The relevance of its age, however, to the main siderite and ankerite from Dobšiná, K/Ar ages of the Cr-mica (fuchsite) closely associated with the ore veins, and Re/Os dating of gersdorffite, a major ore mineral of the hydrothermal veins studied (Halajerová-Andrusovová 1961, 1964; Kiefer et al. 2017).

### 2. Geological evolution of the studied area

The area around Dobšiná is known for its complex geological settings located in the internal zones of the Central Western Carpathians (CWC). The CWC forms the core of the Carpathian Orogen, which underwent a complex Variscan and Alpine structural and metamorphic evolution and it contains several nappe units of various orders and ages. In the study area, there are two main types of tectonic units (Fig. 1): (i) thick-skinned thrust sheets, which comprise the pre-Alpine crystalline basement along with its Upper Palaeozoic to Mesozoic sedimentary cover (the Veporic and Gemic units); and (ii) detached cover nappe systems containing Upper Palaeozoic to Mesozoic sedimentary formations with rare volcanic rocks (the Meliatic, Turnaic, and Silicic units).

The Veporic Unit was overridden by the Gemic Unit, which consists of amphibolite-facies metamorphic rocks of the Klátov Group (Spíšek et al. 1985), green-schist facies metabasites and phyllites (ophiolitic) of the Rakovec Group, and the green-schist facies volcano-sedimentary sequences of the Gelnica Group. These Gemic crystalline complexes are covered by Upper Carboniferous to Middle Triassic volcano-sedimentary succession of Variscan molasse, affected by subsequent Alpine an-metamorphism. Younger formations than the Middle Triassic are unknown in the Gemic cover sequence. The main phase of crustal shortening and nappe formation took place during the late Early Cretaceous (≈137–115 Ma) immediately after the closure of the Triassic–Jurassic Meliata Ocean (≈155–145 Ma), followed by the formation of an accretionary complex (Kozur and Mock 1973; Maluski et al. 1993; Faryad and Henjes-Kunst 1997; Árkai et al. 2003; Dallmeyer et al. 2008; Hurai et al. 2008, 2015; Vozárová et al. 2014). This earliest stage of the Cretaceous N–S convergence in CWC orogenic wedge is recorded in the Gemic Unit by the formation of Alpine metamorphic foliation mainly in incompetent lithologies partly to disjunctive foliations developed in competent rocks of the unit and probably represent a gradual transition from bedding or Variscan to Alpine structure (cf. Hroudá and Faryad 2017).

Later, the tectono-metamorphic overprint is associated with the evolution of a large-scale cleavage fan structure across the entire unit (Lexa et al. 2003). The evolution of
the Gemeric cleavage fan is associated with overthrusting of the Gemeric Unit over the underlying Veporic Unit in the north during the late Early Cretaceous (Lexa et al. 2003; Jeřábek et al. 2012; Novotná et al. 2015; Vojtko et al. 2016). The Gemeric Unit is overlain by a subduction-related melange of sediments and relics of oceanic mantle rocks of the Jurassic Meliata Ocean. The metamorphic conditions of Jurassic accretionary wedge composed of the Meliata, Turňa related rocks correspond to blueschist or greenschist facies for Alpine regional metamorphism (e.g., Faryad 1995, 1997). The Jurassic accretionary wedge and the underlying Gemeric Unit are tectonically overlain by extensive horizontally lying unmetamorphosed Silica nappe system derived most probably from the Apulian shelf (Fig. 1).

The system of siderite-type hydrothermal carbonate-quartz veins with Ni–Fe–Co-Cu ore mineralization near Dobšiná is hosted in Early Paleozoic gneiss-amphibolite complex of the Klátov group belonging to the Gemeric Unit. Minor occurrences of Ni–Fe–Co ores are known from the metasomatic bodies of siderite hosted in Carboniferous rocks (Fig. 2). The two principal vein systems were distinguished. The Zemberg-Terézia vein system is located at the northern margin of the gneiss-amphibolite complex and the Martini vein system, which is situated at the southern margin of the same complex (Fig. 2). The two principal vein systems were subdivided by the N–S trending fault to western (Zemberg) and eastern (Terézia) segments. The Zemberg segment generally consists of three (Severná, Hlavná and Južná vein) and Terézia segment of two (Terézia I., Terézia II.) veins. The Martini vein system consists of four major (Jacobi, Ezechiel, Martini,
3. Samples and Methods

Most of the samples used in this study were collected in the old mines with a precise localization (Details are included in Tab. 1). Samples of fresh hydrothermal carbonates (siderite and dolomite-ankerite series) were collected in-situ in the abandoned adits located both at the Zemberg-Terézia vein system (Karol adit-DBK1, Lower Jórémeny adit-DN1 to DN4) and Martini vein system (Remény adit – DR1 to DR4). Fuchsite samples were collected from the Martini vein system (DR3 – underground from the Remény adit and DE-1 from the dump of the Lower Ezechiel adit). Samples rich in the hydrothermal Ni-Co arsenides were collected by Halahyjová-Andrusová (1961, 1964) from both vein systems and recently re-investigated by Kiefer et al. (2017).

All samples have been examined in the form of polished sections with a Zeiss-Axio-Imager.M2m optical microscope equipped with a CCD-camera system. The chemical composition of the hydrothermal carbonates and the ore minerals was determined by electron microprobe analyses using a JEOL JXA-8230 instrument. For the carbonate minerals, the operating conditions were set to an accelerating voltage of 15 kV, a beam current of 15 nA and a beam diameter of 1 µm. The wavelength-dispersive X-ray spectrometers were used to measure the elements and X-ray lines of Mg ($K_\alpha$), Si ($K_\alpha$), Sr ($L_\alpha$), Ba ($M_\alpha$), Ca ($K_\alpha$), Fe ($K_\alpha$), and Mn ($K_\alpha$). To improve the count-rate statistics, the counting times were 40 s. The standard specimens used were: MgO for Mg, rhodonite for Si and Mn, celestine for Sr, baryte for Ba, apatite for Ca, and hematite for Fe. The detection limits, calculated from the peak and background counts, the measurement time and the standard material concentration, are 0.03 wt. % for Si, 0.02 wt. % for Mg, 0.09 wt. % for Ba, 0.09 wt. % for Sr, 0.02 wt. % for Ca, 0.05 wt. % for Fe and 0.05 wt. % for Mn. The ore minerals, mainly sulfarsenides, were analyzed with an accelerating voltage of 20 kV, a beam current of 50 nA, a beam diameter of 1 µm and counting times of 40 s. The measured elements and lines were As ($K_\alpha$), S ($K_\alpha$), Sb ($L_\alpha$), Pb ($M_\alpha$), Ni ($K_\alpha$), Co ($K_\alpha$), and Fe ($K_\alpha$). The standard specimens used were: arsenopyrite for As, pyrite for S and Fe, pure metal standards for Ni and Co, galena for Pb and stibnite for Sb. The detection limits, calculated after the same procedure as the above mentioned, were: 0.04 wt. % for Sb, 0.08 wt. % for Pb, 0.07 wt. % for As, and 0.02 wt. % for S, Co, Cu, Fe, and Ni. The elements analyzed but not listed in the corresponding tables (Pb, Cu) were always below the detection limit.

The rocks at the vast majority of exposures possess particular planar structure. In metamorphic rocks, a planar structure is known as metamorphic or tectonic foliation (cleavage, schistosity). In this paper, we use foliation as a general term for all pervasively developed planar structures found in rocks. The second type of planar structure were veins with their typical tabular shape, which were deformed during the tectonogenesis. Structures in rocks with linear, as opposed to planar, the character also occur in many forms. Observed lineations are both mineral lineations (linear alignments of minerals in metamorphic tectonites) and stretching lineations defined by strained objects with elongated, cigar-like shapes. These lineations were measured on vein surfaces to provide information on the direction of movement during or after the vein formation. All oriented data were processed in the stereographic projection (Lambert net), lower hemisphere using the FaultKin 7.5 (Marrett and Allmendinger 1990; Allmendinger et al. 2013). Uranium-Pb ages were acquired in situ in thick polished section by laser ablation-sector field-inductively coupled
plasma-mass spectrometry (LA-SF-ICPMS) at the Goethe University Frankfurt (GUF). At GUF a ThermoScientific Element 2 sector field ICP-MS is coupled to a RESOLution S-155 (Resonetics) 193 nm ArF Excimer laser (CompexPro 102, Coherent) equipped with a two-volume ablation cell (Laurin Technic, Australia). Static ablation in a helium atmosphere (0.3 l/min) used a spot size of 213 µm and a fluence of ca. 1 J cm⁻² at 6 Hz. Soda-lime glass SRM-NIST614 was used as a reference glass together with 3 carbonate standards to bracket sample analysis. Raw data were corrected offline using a macro-based in-house MS Excel© spreadsheet program (Gerdes and Zeh 2009). The $^{207}\text{Pb}/^{206}\text{Pb}$ ratio was corrected for mass bias (0.6 %/amu) and the $^{206}\text{Pb}/^{238}\text{U}$ ratio for inter-element fraction (ca. 5 %). Due to the carbonate matrix, additional correction of 9 % has been applied on the $^{206}\text{Pb}/^{238}\text{U}$, which was determined using WC-1 carbonatate reference material dated by TIMS (254 ± 6 Ma; Roberts et al. 2017). All reported uncertainties are the 2σ level. For the Re/Os geochronological analyses, acquired at the University of Alberta, ten samples of Ni–Fe–Co arsenides were selected for preliminary analyses, and only three of them contained sufficient Re to be considered further. These three samples were analyzed by the methods described in detail by Selby and Creaser (2004) and Morelli et al. (2010). Initially, the Re abundance in each of 3 mineral separates was tested by isotope dilution analysis and found to be suitable for one sample only (sample 436: 10.0 ppb Re, 12d: 0.2 ppb Re, 25b: 2.0 ppb Re). An initial full Re-Os analysis revealed little common Os was present, with $^{187}\text{Os}^*$ comprising > 99.7 % of the Os measured. As such, for this work, a mixed double spike containing known amounts of isotopically enriched $^{185}\text{Re}$, $^{190}\text{Os}$, and $^{187}\text{Os}$ analysis was then used (Markey et al. 2007). The $^{187}\text{Re}$ and $^{187}\text{Os}$ concentrations were determined by isotope dilution mass spectrometry using Carius-tube, solvent extraction, anion chromatography, and negative thermal ionization mass spectrometry techniques. The isotopic analysis used a ThermoScientific Triton mass spectrometer by Faraday collector. Total procedural blanks for Re and Os are less than <1 pico grams and 0.1 picogram, respectively, which are insignificant in comparison to the Re and Os concentrations present. The Reference Material 8599 Henderson molybdenite (Markey et al. 2007) is routinely analyzed as a standard, and during the past five years returned an average Re-Os date of 27.78±0.07 Ma (n = 15), indistinguishable from the Reference Age Value of 27.66±0.1 Ma (Wise and Watters 2011). The $^{187}\text{Re}$ decay constant used is $1.666 \times 10^{-11}$ a⁻¹ (Smoliar et al 1996).For the K/Ar dating, two portions of each sample (8–16 mg and ≈30 mg) were weighed. A small quantity of CuO (approximately 0.1 mg) was added to the lighter portion. This sample was then wrapped in an Al-foil and proceeded to radiogenic argon measurement. The heavier portion was weighted directly on platinum evaporator and used for potassium measurement. Radiogenic argon measurements were performed on Nu Instruments Noblesse multicollector noble-gas spectrometer (NG 039). The Al-foil wrapped portions was melted in a double-vacuum resistant crucible at ≈1200 °C. Pure $^{38}\text{Ar}$ produced by the Institute for Inorganic and Physical Chemistry of Bern University was used as the spike. CuO added to the samples enhances the oxidation of organic matter during this step. Titanium sponge acted as a sublimation getter and was always put to the place of the reactor where samples are heated. This was the first cleaning level of gases extracted from the samples. The final purification of argon was carried out in an isolated section of the line by a warm getter pump (D-100, SAES Getters), monitored not to release additional argon atoms (after installation it was activated for 12 h). Small portions of gas aliquot released from the sample (ca. 1 %) were then measured at least three times by opening and closing

Fig. 3a – Fuchsite-rich rocks with veinlets of siderite. b – Coarse-grained siderite (brownish) and ankerite (white) with gersdorffite (metallic grey) mineralisation. Jóremény adit. Both photographs courtesy of D. Peterec.
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Blanks measurements were performed after every sample and confirmed that released radiogenic $^{40}\text{Ar}$ for every sample was above 99.5%. Every day, the $^{40}\text{Ar}/^{36}\text{Ar}$ and $^{40}\text{Ar}/^{38}\text{Ar}$ ratios are measured for air sample aliquots, delivered from a calibrated air pipette. Based on this results $^{40}\text{Ar}/^{36}\text{Ar}$ and $^{40}\text{Ar}/^{38}\text{Ar}$ ratios were corrected for instrument mass fractionation and detector efficiencies assuming atmospheric ratios of $(^{40}\text{Ar}/^{36}\text{Ar})_{\text{air}} = 298.57$ and $(^{40}\text{Ar}/^{38}\text{Ar})_{\text{air}} = 1583.52$ (Lee et al. 2006). The amount of the original aliquot of $^{38}\text{Ar}$ spike was determined by measuring international standard GL-O glauconite, performed every day. For this series of samples $1\sigma$ for GLO was 0.7 % (13 samples). Standard age errors were calculated from the law of error propagation, also taking these values into account. The potassium contents were measured using Sherwood Model 420 flame photometer on HF dissolved samples. The maximum error of this measurement was estimated to be equal 0.03 % K. Based on six measurements, K content in GL-O glauconite was slightly higher: 6.61 ± 0.02 %, than that of the standard: 6.56 ± 0.06 % (Odin 1982). LP-6 biotite has a slightly lower amount of $^{K}$: 8.27 ± 0.022 % (4 measurements) comparing to reference value: 8.33 ± 0.03 % (Odin 1982). Both measured standards are within reference ranges assuming 0.03 % K error.

4. Results

4.1. Laboratory and field observations

4.1.1. Macroscopic observations

Samples collected for this work are either fuchsite-rich country rocks or hand specimens from ore veins, collected in old mines. Samples from both old dumps and debris were avoided as much as possible because their origin and association with the ore veins are difficult to ascertain. Various kinds of metapelites (phylite) and, less frequently, also metasandstone, with intercalations of metamorphosed mafic volcanic to volcanoclastic rocks of the Northern Gemenic Unit, are predominant host rocks of the mineralized veins.

The fuchsite-rich rocks are massive, fine-grained, distinctly green rocks that form thick (tens of meters) alteration haloes. They are not necessarily directly associated with the ore veins. If intermingled with hydrothermal siderite or ankerite, the carbonates are clearly younger than the fuchsite-rich rocks (Fig. 3a). The ore veins form tabular bodies and lenses, with a thickness of up to 1 m. Over a distance of a few meters, however, the thickness may change drastically, sometimes to only a few millimetres filled by greyish clay. The veins are usually accompanied by a splintery rim of black, carbonaceous matter. The main minerals of the veins are carbonates, either siderite or ankerite. Siderite is especially common in the lower parts of the mineralized zone (Remény adit) and occurs there together with ankerite. In the higher parts (Zemberg, Jóremény), siderite is scarce. If they are found together, siderite is older than ankerite (Fig. 4a). The Ni–Fe–Co sulfarsenides are usually dispersed in the carbonates (Figs. 4a–c), localized in small veinlets or zones near the contact of the veins with the host rocks. The Ni–Fe–Co sulfarsenides are younger than the main carbonate mass (Fig. 3b). They are hosted by both types of carbonates but, in some cases, whitish anker-
itic carbonate appears to be even younger than the ore minerals. A common, although a not volumetrically abundant component of the veins is quartz with tourmaline, chlorite, and albite. These minerals form pockets in the carbonates, but a precise temporal relationship to siderite, ankerite, and ore minerals was challenging to determine. The uppermost portions of the veins at the Kruhová (Gugl) hill contain siderite, baryte, and cinnabar. The relationship of these minerals to the main mineralization is not clear because they can be observed only as hand specimens in shallow pits.

4.1.2. Structural observations

Structural observations were carried out only in the abandoned Remény adit close to Dobšiná town because of its accessibility at the time of this study. Data for geometrical and structural analysis were obtained from the main mineralized zone emplaced in the Lower Palaeozoic metapelites in the adit. In this site, the geometry of the mineralized zone is different from other mineralized localities close to Dobšiná. In the broader area, the mineralized veins generally strike W–E with southward inclination. In the Remény adit, the mineralized structures can be roughly divided into two geometrically different groups. The main zone has the same strike as the other zone in the broader area of Dobšiná, but the dip is northwards (Fig. 5a). This relatively older structure is intersected by most probably slightly younger vein system with the NW–SE direction with the dip south-westwards (Fig. 5b).

The thickness of the main mineralized zone varies from several centimetres to several decimetres. It is a mylonite zone with shallow to medium inclination, with uniform both strike and dip in the entire accessible and observable length in the Remény adit. The measured structured along the mylonite zone refer to an extensional tectonic regime with the orientation of the principal minimal stress axis ($\sigma_3 = 185/04^\circ$) in the NNE–SSW direction and subvertical principal maximal stress axis ($\sigma_1 = 066/83^\circ$). The mineralized mylonite zone generally conforms to the Variscan–?Alpine metamorphic foliation and the slight variations in its dip are caused by younger folding. It is possible to assume that the mylonite zone was formed as a heterogeneous structure. However, a further detailed structural investigation is needed. Firstly, the mylonite zone could also be formed by the compressional tectonic regime (Early Cretaceous) and later (Mid- to Late Cretaceous) was overprinted by a younger low-angle normal fault mechanism. Unfortunately, limited access to the structure did not allow us more detailed structural research.

The younger mineralized veins are generally thinner and less significant. The veins are steeper than the older mineralized mylonite zone and were formed during an extensional tectonic regime with the orientation of the principal minimal stress axis ($\sigma_3 = 207/02^\circ$) in the NE–SW direction and subvertical principal minimal stress axis.

![Fig. 5](image-url) Stereograms of observed mineralized structures with movement directions in the Remény adit near Dobšiná. a – great circles and movement trajectories in the main mineralized zone; b – great circles and movement trajectories of the younger structures than the main mineralized zone. Note: the planar structures are drawn as the great circles, and the movement trajectories (lineation on that planes) are represented by arrows. The compressional area with calculated principal compressional stress axes ($\sigma_3$) is depicted by red contours and dots. The tensional area with calculated principal extensional axes ($\sigma_3$) is shown by blue. Stereograms use Lambert projection, lower hemisphere.
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These younger veins disrupt the Variscan–Alpine metamorphic foliation and Alpine mineralized aforementioned mylonite zone. According to the crosscut criteria, the faulting and mineralization process have to be younger than the Gemeric cleavage fan. Conditionally, it could be linked with the exhumation process in the Gemeric Unit. However, it is still not entirely clear whether the movement in these two zones is directly related to the mineralization process or it is the subsequent process of faulting in the Gemeric Unit or both.

4.1.3. Observations in optical and electron microscope

Back-scattered electron (BSE) images (Fig. 6) and WDS analyses (Tab. 2, Fig. 7a) distinguished multiple generations ofankerite (Ank I–III) and siderite (Sid I and II). Ankerite I (Ank I) is the dominant carbonate phase in all samples (Figs. 3b, 4) and shows a compositional range of 15.6–21.2 mol. % MgO, 47.0–53.3 mol. % CaO, 21.5–29.1 mol. % FeO and 3.0–5.0 mol. % MnO.

Ankerite II (Ank II) forms veinlets (hundreds of micrometres, Fig. 6a) in Ank I. The compositional range of Ank II is 25.1–29.1 mol. % MgO, 51.3–55.2 mol. % CaO, 15.0–20.2 mol. % FeO and 1.1–1.7 mol. % MnO. In general, Ank II has less Fe than Ank I (mean concentration Ank I Fe = 26.8 mol. % vs Ank II Fe = 18.1 mol. %) and more Mg (mean concentration Ank I Mg = 18.6 mol. % vs Ank II Mg = 27.7 mol. %). Some Mg/Ca-rich compositions might violate compositional rules for carbonate-nomenclature and therefore could be labelled as dolomite or Fe-dolomite. However, those compositions only represent minor portions and are texturally not isolated and therefore are part of the corresponding ankerite stage.

Ankerite III (Ank III) can be found as veins and fractures cutting through Ank I, Ank II, and siderite (Fig. 6b). The compositional range of Ank III is 19.8–20.8 mol. % MgO, 52.2–53.0 mol. % CaO, 23.5–24.4 mol. % FeO, and 2.7–3.7 mol. % MnO. This carbonate is homogeneous when compared to Ank I and II. It is comparable to carbonate chemistry of Ank I (more Fe and less Mg than Ank II).

Siderite in the investigated samples can be found as massive coarse-grained siderite in carbonate-sulfarsenide veins (Figs. 3, 4a, 6c) or as siderite in the quartz-fuchsite rocks (Figs. 3a, 6e). The main mass of vein siderite is denoted as Sid I and has a compositional range of 11.0–16.4 mol. % MgO, 0.1–1.0 mol. % CaO, 78.4–83.7 mol. % FeO, and 2.8–5.6 mol. % MnO. Siderite from the quartz-fuchsite rocks has a similar composition of 12.0–24.1 mol. % MgO, 0.2–1.4 mol. % CaO, 70.7–82.8 mol. % FeO, and 4.2–6.1 mol. % MnO. We assume that this siderite is also Sid I and was brought into these rocks during the siderite sub-stage.

Later siderite, denoted as Sid II, is found in thin veinlets populated by grains and crystals of Ni–Fe–Co sulfarsenides or as rims around these minerals (Fig. 6c). The compositional range of Sid II is 23.7–29.7 mol. % MgO, 0.8–6.4 mol. % CaO, 65.5–72.5 mol. % FeO, and 0.5–1.8 mol. % MnO. The results of all WDX spot analyses on carbonates can be seen in the electronic supplementary table S1.1.

4.1.4. Ore mineralization

Optical microscopy and BSE images suggest that the ore mineralization, represented mostly by Ni–Fe–Co sulfarsenides and arsenides from Dobšiná (in wt. %) from the samples used for geochronological dating in this work. Mineral abbreviations: gdf = gersdorffite, aspy = arsenopyrite

| Tab. 2 | Representative chemical compositions of carbonates from Dobšiná (in wt. %) in the samples used in this work. |
|-------|------------------------------------------------------------------------------------------------------------------|
|        | Sid I   | Sid II  | Siderite (disseminated) | Ank I   | Ank II  | Ank III |
| FeO    | 48.18   | 42.71   | 46.63                | 17.10   | 12.48   | 16.06   |
| MnO    | 2.18    | 0.59    | 3.16                 | 2.63    | 0.95    | 2.05    |
| MgO    | 4.52    | 9.65    | 5.31                 | 6.91    | 10.71   | 7.66    |
| CaO    | 0.19    | 1.41    | 0.35                 | 26.63   | 28.41   | 27.61   |
| BaO    | <D.L.   | <D.L.   | <D.L.                | <D.L.   | <D.L.   | <D.L.   |
| SrO    | <D.L.   | <D.L.   | <D.L.                | <D.L.   | <D.L.   | <D.L.   |
| Total  | 55.20   | 54.46   | 55.46                | 53.42   | 52.68   | 53.51   |

4.1.4. Ore mineralization

Optical microscopy and BSE images suggest that the ore mineralization, represented mostly by Ni–Fe–Co sulfarsenides and arsenides from Dobšiná (in wt. %) from the samples used for geochronological dating in this work. Mineral abbreviations: gdf = gersdorffite, aspy = arsenopyrite

| Tab. 3 | Representative chemical compositions of sulfarsenides and arsenides from Dobšiná (in wt. %) from the samples used for geochronological dating in this work. Mineral abbreviations: gdf = gersdorffite, aspy = arsenopyrite |
|-------|------------------------------------------------------------------------------------------------------------------|
|        | DR2  | DN3  | DN3  | DN2  | 436  |
| Sb     | 0.14 | <D.L. | 0.09 | 0.09 | 0.09 |
| S      | 13.94 | 19.18 | 15.56 | 11.37 | 15.74 |
| As     | 51.67 | 45.51 | 49.43 | 54.83 | 49.87 |
| Co     | 0.60 | <D.L. | 7.09 | 4.11 | 2.48 |
| Ni     | 32.17 | 0.09 | 20.82 | 26.13 | 29.47 |
| Fe     | 1.35 | 34.62 | 7.12 | 2.99 | 1.86 |

Total 99.87 99.40 100.11 99.52 99.50

<D.L. – below detection limit
sulfarsenides, is hosted by all carbonates except for the late Ank III. The ore minerals found are arsenopyrite and Ni-rich sulfarsenides, mainly gersdorffite and members of the gersdorffite-arsenopyrite solid-solution (Tab. 3, Fig. 7b). Chemical composition of ore minerals in the system Ni-Fe-Co-As-S is defined by a great chemical variability and abundant polymorphism. Such variability is also observed in the samples from Dobšiná (Kiefer et al. 2017), but it is not bound to a particular mineralization stage but rather a general observation for all samples.

Gersdorffite occurs as small disseminated crystals in Ank II (Fig. 6a), as large strongly zoned and often highly fractured assemblages in Ank I and siderite (Figs. 3b, 6c, d) and it is the most common ore mineral in the investigated samples. The compositional range of gersdorffite is 0–0.5 wt. % Sb, 10.8–19.8 wt. %

Fig. 6 Back-scattered electron (BSE) images of the samples from Dobšiná. The square holes in some images are traces after laser ablation. a – Early ankerite I with thin veinlets of ankerite II. Note that the ore mineralization is localized in Ank II. b – Early ankerite I with veinlets of the remobilized ankerite III. c – Sideritic carbonates with ore mineralization. d – Gersdorffite with oscillatory zonality. e – Intergrowth of fuchsite (fu), siderite (sid), quartz (qtz), and corroded relics of chromite (chr) in fuchsitic rocks.
S, 43.2–63.1 wt. % As, 0.3–12.7 wt. % Co, 0–0.03 wt. % Cu, 13.8–34.2 wt. % Ni, and 0.4–11.7 wt. % Fe, highlighting the variations from almost ideal stoichiometries (e.g., Ni\textsubscript{0.94}Co\textsubscript{0.04}Fe\textsubscript{0.01}As\textsubscript{1.00}S\textsubscript{1.01}) to substantial deviations, therefore (e.g., Ni\textsubscript{0.54}Co\textsubscript{0.21}Fe\textsubscript{0.28}As\textsubscript{1.04}S\textsubscript{0.94}, Ni\textsubscript{0.86}Co\textsubscript{0.07}Fe\textsubscript{0.08}As\textsubscript{1.18}S\textsubscript{0.82} or Ni\textsubscript{0.98}Co\textsubscript{0.03}Fe\textsubscript{0.02}As\textsubscript{1.52}S\textsubscript{0.45}) including some high-As compositions corresponding to rammelsbergite/krutovite-like stoichiometries. Arsenopyrite is a minor phase and only occurs as small disseminated crystals in Ank II together with gersdorffite. The mineral chemistry of arsenopyrite is relatively constant and approaches the mean stoichiometry of Fe\textsubscript{1.02}As\textsubscript{1.00}S\textsubscript{0.98}. The results of all WDX spot analyses on sulfarsenides and arsenides can be seen in the electronic supplementary table S1.2.

4.2. Geochronology

4.2.1. U/Pb LA-SF-ICP-MS dating on carbonates

Nine samples (DR1–DR4, DN1–DN4, DBK1) were selected for the U/Pb dating, but only four gave one or two statistically meaningful regression lines with sufficient variation in the U/Pb to calculate lower intercept ages. Representative Tera-Wasserburg diagrams are shown in Fig. 8. Three samples (DN1, DR2, and DN3) gave older and younger ages and one sample (DN2) only one younger age. The results of all WDX spot analyses on sulfarsenides and arsenides can be seen in the electronic supplementary table S1.2.

Fig. 7 Triangular diagrams showing a – the composition of the generations of hydrothermal carbonates and b – composition of the Ni–Fe–Co sulfarsenides in the samples used for dating.

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older isochron ages vary between 145 ± 5 Ma and 114 ± 24 Ma. All old ages have been acquired on Ank I and Sid I (Tab. 4). The three younger isochron ages vary between 20.7 and 29.5 Ma and are assigned to Ank III (Tab. 4). The U and Pb concentrations as well as the calculated isotopic ratios of the four samples DN1, DN2, DN3 and DR2 can be seen in the electronic supplementary table S2.

K/Ar dating of fuchsite

Two samples from the wall-rock alteration were chosen for K/Ar-dating. A preliminary investigation by XRD showed that both samples contained mostly quartz and fuchsite and to a lesser amount siderite and kaolinite (Tab. 5). Both samples gave similar ages averaging at 109 Ma (Tabs 4 and 5).

4.2.1. Re/Os dating of gersdorffite

Nine representative gersdorffite samples were pre-analyzed or analyzed but only the sample 436 yielded an age, with a value of 95.7 ± 1.2 Ma (Tab. 6). The age uncertainty is quoted at 2σ level and includes all known analytical uncertainty, including a ≈0.31 % uncertainty in the decay constant of $^{187}$Re. The gersdorffite sample analysis was replicated using a larger sample aliquot and yielded a more precise age of 93.6 ± 0.9 Ma, which lies within the uncertainty of the first analysis. To our best knowledge, this datum is the first Re/Os age obtained from gersdorffite.

5. Discussion

5.1. Precipitation sequence of hydrothermal minerals in Dobšiná

The precipitation sequence of hydrothermal minerals in Dobšiná was established by Halahyjová-Andrusovová (1959) on many samples from the old mines that were...
accessible during the exploration activities in the 1950s. The sequence published by Halahyjová–Andrusovová (1959) can be augmented by observations done in this work, the work of Kiefer et al. (2017), and especially correlation with siderite-sulfide deposits in the Gmeric Unit (Varček 1985; Chovan et al. 1994; Hurai et al. 2008). The sequence constructed here (Fig. 9) focuses mainly on the Ni–Fe–Co veins and neglects other important ores found in the Gmeric Unit, e.g., the quartz-sulfide phase with stibnite (see Fig. 5 in Hurai et al. 2008).

All these authors agree that fuchsite and the associated minerals represent the oldest–fuchsite stage, either in Rudňany (cf. Rojkovič 1977) or elsewhere. In our samples from Dobšiná, siderite-ankerite veinlets penetrate the fuchsitic rocks (Fig. 3a), thus confirming the observations from Rudňany. The following siderite sub-stage (Hurai et al. 2008) is mineralogically simple, with dominant siderite and subordinate ankerite. The baryte sub-stage is prominently developed at some deposits, for example in Rudňany, but not in Dobšiná. Baryte, together with siderite and cinnabar, does occur in the highest portions of the veins in Dobšiná. The hand specimens, recovered from shallow surface pits at the Kruhová hill do not allow the placement of this baryte into the precipitation sequence. We assume, however, that its position in the precipitation sequence is the same as in the Rudňany deposit.

The quartz-tourmaline stage is devoid of ore minerals and developed locally. Owing to the common occurrence of monazite, this stage has been a subject of previous geochronological work (e.g., Hurai et al. 2015) and provides an important marker that separates, in time, the older (fuchsite, siderite, baryte) and younger (sulfide) stages. Numerous observations in the old mines suggest that ankerite is generally younger than siderite. This conclusion also agrees with the earlier studies; Hurai et al. (2008) assert that the main carbonate associated with the sulfide stage is ankerite (Fig. 3b). Ni–Fe–Co arsenides are scarce minerals in the ore deposits in the Gmeric Unit, except Dobšiná and some veins at the Rudňany and Rožňava deposits (Koděra et al. 1990). If considered in the precipitation sequence, they are usually placed early in the quartz-sulfide stage (e.g., Rojkovič 1977; Chovan et al. 1994). Their chemical and mineralogical evolution in Dobšiná was described in detail by Kiefer et al. (2017), but there is no reason, either from that or the current work, to assume that they were formed in several separate hydrothermal pulses or events. The Ni–Fe–Co arsenides are often crushed and healed by younger Cu sulfides, either chalcopyrite or tetrahedrite. A simple placement of the arsenides with the later Cu sulfides in one stage, without indication of the tectonic damage, is therefore questionable. We, therefore, distinguish the Ni–Co and Cu–Sb (chalcopyrite-tetrahedrite) mineralizations as two separate substages of the quartz-sulfide stage (Fig. 9).

### Table 6: Re/Os isotopic and age data for gersdorffite from Dobšiná (sample 436)

| Sample      | Re (ppb)     | 187Re (ppb) | 187Os (ppb) | Model age (Ma) |
|-------------|--------------|-------------|-------------|----------------|
| 436         | 12.14 ± 0.10 | 7.630 ± 0.063 | 0.01218 ± 0.00011 | 95.7 ± 1.2   |
| 436 repetition | 11.40 ± 0.09 | 7.165 ± 0.056 | 0.01118 ± 0.00004 | 93.6 ± 0.9   |

**Fig. 9 Precipitation sequence of minerals from Dobšiná. Compiled from the work of Halahyjová–Andrusovová (1959), Varček 1985, Chovan et al. 1994, Hurai et al. 2008, Kiefer et al. 2017, and this work.**
5.2. Age constraints on the fuchsite stage

There is no doubt that fuchsite, together with quartz and carbonates, constitute the oldest mineralization stage in Dobšiná and Rudňany (cf. Rojkovič 1977; Hurai et al. 2008). Fuchsite-carbonate-quartz rocks have also been described from several other sites in the Gemeric Unit (Roztoky, Bindt, Stará Voda, Nálepkovo, Mlynky, Mišíkova skala, Košické Hámre, Hnilec) and interpreted as a final product of metasomatic alteration of ultrabasic or basic rocks by CO₂-rich hydrothermal fluids (Ivan 1982).

Fuchsite from the Gemeric Unit has been subjected to several K/Ar geochronological studies. The value determined from Dobšiná (Augustín quarry, Malá Vlčia dolina) is 190.8 Ma (0.83 % K₂O) (Cambel et al. 1980). This work reported K/Ar age determinations from metamorphic rocks in the broader area of Dobšiná, some of them with clear signs of retrograde metamorphism. These ages scatter considerably and are 273, 258, 227, 154 Ma. Fuchsite was also dated in Rudňany to 210±2 Ma (1.11 % K₂O) (Bagdasaryan et al. 1977), later revisited and revised to 205 Ma, with another datum of 144 Ma (Cambel et al. 1990). The significance of these scattered values in comparison with the newer geochronological data is questionable. Our K/Ar ages for fuchsite of 109 and 110 Ma (4.68–4.82 % K₂O) (Tab. 5 and Fig. 10) are distinctly younger than the previously published values and also contradict the U/Pb age of the carbonates (Tab. 4) which are texturally younger than fuchsite (Fig. 3a). Low K-amounts in previous measurements can also indicate that these dates correspond to rocks containing low amounts of fuchsite. Early on, it was noted that the K/Ar ages in this area are scattered and contradictory (Hovorka and Spišiak 1981) and may have been subject to resetting during the intensive Alpine reworking (Radvec et al. 2004). Similar ages (clustered around 100 Ma) were obtained by Rb/Sr (Kováč et al. 1986) or K/Ar (Kantor and Rybár 1979) dating of sheet silicates from the Gemeric Unit. Given that fuchsite is a Cr-rich variety of muscovite, the closure temperature of 350±40°C (Spear and Parrish 1996) also applies to this mineral. Fuchsite from Dobšiná is a product of alteration, although not too fine-grained, and its closure temperature may lie near the lower end of the interval specified by Spear and Parrish (1996). Additionally, fluid-assisted recrystallization compared to pure temperature-controlled diffusion can be of importance, as was shown, e.g. by Tartèse et al. (2011). If taken at its face value, the ≈110 Ma could be interpreted as the age of the last cooling below ≈320°C or lower, due to the influence of fluids (e.g., Villa 2015). This temperature coincides with the last hydrothermal pulses related to the crystallization of the hydrothermal carbonates (see below). This interpretation is also supported with the estimates of peak P-T conditions of the Alpine metamorphism in the Gemeric Unit of 320–350°C and 0.4–0.55 GPa (Faryad and Dianisika 1999). This event, correlated with the geochronological data of Dallmeyer et al. (1996), was assigned Cretaceous age (120–86 Ma) and the zircon fission-track data (Plašienka et al. 2007; Králiková et al. 2016). The meaning of the previously published, older ages remains questionable. Why, for example, are the two ages of fuchsite from one locality, Rudňany, 60 Ma apart (Cambel et al. 1990)? They may represent mixed ages of several minerals in the bulk samples used at that time. Some of the minerals may have been formed earlier than fuchsite and experienced partial Ar loss during the Alpine times (e.g., hornblende from amphibolites). Argon loss could be locally enhanced by the influence of hydrothermal fluids, promoting recrystallization, producing thus varying ages from a single locality.

5.3. Age constraints on the carbonate minerals

The carbonate minerals found in Dobšiná, either siderite or ankerite, formed mostly during the siderite sub-stage (Fig. 9). Among the massive vein carbonates, siderite (sid I) is usually older than ankerite (Ank I). The BSE images (Fig. 6) document repeated remobilization of carbonates, thus suggesting that their mutual relationship and assignment to a particular stage may not be straightforward.

There are no geochronological data for the hydrothermal carbonates in the Gemeric Unit. The two competing and conflicting models argue for Permian (Radvec et al. 1994a, b, 2004; Radvec and Grecura 2016) or Cretaceous (Hurai et al. 2002, 2008) age of these and associated minerals. They are based on geological arguments, application of geothermobarometry, and estimation of hydrothermal fluid properties and chemistry. All our results place the formation of the hydrothermal carbonates in Dobšiná to Cretaceous. Two ankerite samples gave ages of 145±5 and 114±24 Ma and the only siderite sample that returned an age gave 118±10 Ma. These ages mark thrusting of the Gemeric Unit onto the Veporic crystalline basement from 140 to 115 Ma (Hurai et al. 2008, 2015; Vozárová et al. 2014). This overthrust is further constrained by the cooling of the uppermost Gemeric and overlying rocks complexes of the Meliata accretionary wedge documented by zircon (U–Th)/He ages and ZFT ages (130–90 Ma; Plašienka et al. 2007; Pútíš et al. 2014; Králiková et al. 2016). Formation of the main mass of the hydrothermal siderite and ankerite appears to be confined to this crustal-scale event. The subsequent exhumation of the Gemeric Unit (115–80 Ma) was accompanied by different types of ore mineralizations, described below. Thrusting of the Gemeric Unit onto the Veporic Unit
was also accompanied by the hydrothermal remobilization of U–Mo–Cu mineralization in the northern parts of the Gemeric Unit at 130±20 Ma (Rojkovič et al. 1993) or 124±10 Ma (Rojkovič and Konečný 2005).

5.4. Age constraints on the quartz-tourmaline stage

This stage, regionally distributed in the Gemeric Unit, has been dated in the past. Monazite, an integral component of this stage, was analyzed either by electron microprobe or LA-ICP-MS. Monazite from Čučma gave ages of 120±9 and 76±12 Ma, with a weighted average of 92 Ma (Hurai et al. 2006). Later, Hurai et al. (2015) extracted a more precise age of 139±1 Ma from monazite from Rožňava. There are some data on monazite from the quartz-tourmaline stage from the Tatric and Veporic units, where the results scatter broadly around 100 Ma (Majzlan et al. 2020). Another temporal hint is the age of 131 Ma, extracted from a zircon overgrowth in the Paleozoic silicilastic metasediments (Vozárová et al. 2013).

The CHIME age of monazite from Jedľové Kostoľany is 83±9 Ma, that from Lubietová (Veporic Unit) 92±11 and 97±9 Ma (Ozdín 2008, 2015; Ozdín et al. 2016). At these localities, these authors consider the quartz-tourmaline stage with monazite to be contemporaneous with the Ni–Fe arsenide mineralization. If so, these dates would place independent constraints on the Ni–Co sub-stage, even though not in the Gemeric Unit. According to the position of this quartz-tourmaline stage in the general precipitation sequence in the Gemeric Unit, it is younger than the siderite phase and usually older than the Ni–Co sub-stage of the quartz-sulfide phase. Siderite and ankerite, perhaps components of the siderite stage, have been dated in this work to 145–114 Ma, with multiple remobilization events visible in the BSE images and geochronological data. Given that monazite could have also been repeatedly remobilized (cf. Hurai et al. 2006), we can only conclude that the available data fit roughly together. Exact differentiation of the ages of these two stages is thus complicated by multiple remobilization events.

5.5. Age constraints on the Ni–Fe–Co sulfarsenides

The carbonates Sid II and Ank II are closely associated with the Ni–Fe–Co sulfarsenides and precede their crystallization. We were not able to obtain their ages, however, because of their low U content.

In the quartz-sulfide phase in the Gemeric Unit, precipitation of the Ni–Fe–Co sulfarsenides precedes the more abundant and voluminous Cu sulfides (Fig. 9) (Varček 1985; Chovan et al. 1994). Such a position also agrees with the results from the Tatric and Veporic Unit (Ozdín 2003, 2008; Ozdín et al. 2016) where the Ni–Fe–Co sulfarsenides occur only locally and in small quantities.

There are no exact geochronological data for any opaque minerals of this phase. Our datum of 93.6±0.9 Ma is the first numerical result that documents the age of these minerals. It places constraints not only on the Ni–Fe–Co arsenides, but also on the older quartz-tourmaline stage and the younger Cu sulfides. We should note that we attempted U/Pb LA-ICP–MS dating of the gersdorffite and other Ni–Fe–Co sulfarsenides. All these attempts failed because of the high Pb content of these minerals. As mentioned above, the Cu sulfides postdate the Ni–Fe–Co sulfarsenides. The age of the Cu sulfides remains unknown; the only hint is the Ar/Ar dating of muscovite from Jedľové Kostoľany (Ozdín 2010), with the ages of 75±1 and 75±2 Ma. This muscovite was assumed to postdate the sulfide mineralization at this locality.

6. Conclusion

Geochronological data from this work place the siderite-sulfide veins from the Gemeric Unit in Cretaceous (Fig. 10), in agreement with the results and interpretations in Hurai et al. (2002, 2006, 2008). The arguments presented there are based on P–T estimates, which suggest that the mineralization formed at relatively high pressures, compatible with the overburden of crystalline rocks, Permo-Triassic cover and the Jurassic (Meliata) accretionary wedge (Hurai et al. 2008). The new geochronological data disentangle the evolution of the complex, multi-stage hydrothermal systems more precisely.

The new data can be reconciled with the models of the crustal evolution of the Gemeric Unit (Lexa et al. 2003) and its thermal history (Fig. 10). The formation of the siderite stage broadly coincides with the metamorphic peak in the Gemeric Unit and the formation of the Gemeric cleavage fan (CGF), a large arcuate fan whose origin is linked to N–S compression of the Gemeric rocks during the thrusting (Lexa et al. 2003). Continuation of the Cretaceous shortening resulted later in the formation of the Trans-Gemeric shear zone, a zone that is developed in both Gemeric and the Veporic units. Thus, at this time, these two units must have already been amalgamated. This time matches with the formation ages of some of the monazite in the quartz-tourmaline stage, the Ni–Fe–Co sulfarsenides in Dobšiná, and marginally also with the formation of the Cu sulfides. The Cu sulfides, if given the tentative Late Cretaceous age of 75 Ma, postdate the unroofing and exhumation of the Veporic Unit from beneath the Gemeric Unit. These mineralizations also date the age of the Trans-Gemeric shear zone formation (cf. Lexa et
We are very thankful for the helpful contributions and review of the results of this work and earlier related studies in relation to the tectono-thermal evolution of the Gemeric Unit. For display reasons, the data was plotted with a horizontal offset.

Fig. 10 Summary and review of the results of this work and earlier studies in relation to the tectono-thermal evolution of the Gemeric unit. For display reasons, the data was plotted with a horizontal offset.

- U/Pb on carbonates, this work, related to the siderite sub-stage and remobilization phase
- K/Ar on fuchsite, this work, related to the fuchsite stage
- Re/Os on gersdorffite, this work, related to the Ni–Co sub-stage
- K/Ar on fuchsite, Dobšiná, Rudňany (Cambel et al. 1980, 1990), related to the fuchsite stage
- U/Th/Pb on monazite, Čučma, Rožňava (Hurai et al. 2006, 2015), Lubišová, Jedľové Kostolany (Ozdín 2008, 2015, Ozdín et al. 2016), related to the quartz-tourmaline stage
- Ar/Ar on muscovite, Jedľové Koštolany (Ozdín et al. 2016), perhaps related to the Cu–Sb sub-stage

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