Provenance and Variscan low-grade regional metamorphism recorded in slates from the basement of the (SW Hungary)

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Abstract
A metapelitic sequence with Silurian protolith from the Horváthertelend Unit (Tisza Mega-unit, Hungary) has a K-white mica + chlorite + quartz + albite + anatase ± tourmaline mineral assemblage. Moderately developed disjunctive and well-developed continuous foliations are present. Geochemical results reflect a dominantly felsic source of the protoliths, suggesting an intermediate to acidic volcanic arc in the provenance area. Metasandstone pebbles in the metaconglomerate indicate a recycled sedimentary source. Raman spectroscopy of carbonaceous material indicates a ~350–370 °C peak metamorphic temperature. The Kübler Index (KIBasel) values of the phengitic K-white mica indicate epizonal metamorphism (0.22 ± 0.04 Δ²θ). The Chlorite ‘Crystallinity’ Index (ChCCIS) suggests metamorphic alteration near to the anchizone–epizone boundary (0.31 ± 0.06 Δ²θ). K–Ar ages of K-white mica are interpreted as a result of Variscan metamorphism (c. >310 Ma) and post-Variscan uplift (c. 290 Ma). The predominance of lydite and slate of Llandoverian age and the overlying coarse-grained metagreywacke and metaconglomerate beds of the Szalatnak Slate Formation show strong lithological similarities with the proximal Silurian sequences in the Małopolska Massif (Kielce Region, Holy Cross Mountains, Poland). The original position of the Horváthertelend Unit is presumably to the northeast from the Bohemian Massif, next to the Upper Silesian Block (Moravo-Silesian Zone) and the Małopolska Terrane.

Keywords Carbonaceous material · K-white mica · Provenance · Variscan metamorphism · Palaeogeography · Tisza Mega-unit

Introduction
The European Variscan and Alpine mountain chains are collisional orogens and are built up of pre-Variscan basement blocks originated at the Gondwana Palaeo-tethyan margin (von Raumer et al. 2003; Novo-Fernández et al. 2016). Pre-Variscan domains, such as the Iberian Massif, Bohemian Massif and French Massif Central, are essential components in the Variscan basement areas in Western and Central Europe (von Raumer et al. 2003; Kroner et al. 2008). Reconstructions of the Variscan Orogen illustrate a complex evolution with different tectonic units, including a variety of (par)autochthonous sections suffered metamorphism (e.g., Novo-Fernández et al. 2016). The Variscan orogenic phase resulted in superimposed structures and juxtaposition of pre-Variscan and Variscan blocks (von Raumer et al. 2003; Kroner et al. 2008; Kříbek et al. 2009). The Bohemian Massif represents the eastern termination of the European Variscan Orogen. Further Variscan continental crust fragments to the southeast are, however, partly hidden and/or moderately to strongly overprinted by the subsequent Alpine Orogeny (Fig. 1a).

The southern part of the Bohemian Massif belongs to the Moldanubian Zone (sensu stricto) which is a key part of the Variscan Internides (Kroner et al. 2008 and references therein). This internal zone is characterized by individual
complexes which preserve regional low- to high-grade metamorphism as well as synorogenic sedimentation and magmatism dominated by felsic rocks (Kroner et al. 2008; Kříbek et al. 2009). According to Buda et al. (2000, 2004) and Klötzli et al. (2004), the Carboniferous Mórágy Granite Complex of the Tisza Mega-unit, Carpathian–Pannonian area, Hungary (Figs. 1b and 2), also originated from the southern part of the Moldanubian Zone and could represent a shear-zone-bounded coherent block in allochthonous position.

The Tisza Mega-unit is a large composite structural unit, which is actually an exotic terrane of European Plate origin in the south-eastern part of the basement of the Pannonian Basin (Haas and Péró 2004). It is made up of Variscan crystalline complexes, molasse-type post-Variscan formations and Alpine overstep sequences with variable evolutionary history. Based on the Variscan and early Alpine tectonostratigraphic characteristics, the Tisza Mega-unit was located at the southern margin of the European Plate, east to the Bohemian Massif, prior to a rifting period in the Middle Jurassic (Csontos and Vörös 2004; Haas and Péró 2004; Klötzli et al. 2004; Varga et al. 2007). The existing palaeo-geographic reconstructions, however, are quite generic and sometimes contradicting, so further accurate and reliable correlation studies on the Palaeozoic records of the Tisza Mega-unit are needed.

In the Tisza-Mega-unit a marine Silurian formation occurs in the subsurface in southern Transdanubia (Oravecz 1964; Kozur 1984; Árkai et al. 1995; Szederkényi 1996). The Szalatnak Slate Formation composed of low-grade organic-rich metapelite, metagreywacke, and metaconglomerate is known from numerous boreholes in the northwestern marginal zone of the Tisza Mega-unit (Fig. 2). Due to petrographic characteristics as well as metamorphic and deformational evolution this metasedimentary sequence has a considerable importance for correlation of the Tisza Mega-unit.

Normally, these very fine-grained rocks consist of high amounts of phyllosilicates, dominantly K-white mica and chlorite, with lesser amounts of quartz and feldspar which are stable in a wide pressure-temperature range (Miyashiro 1994; Frey and Robinson 1999). Using X-ray powder diffraction (XRPD) methods, illite and chlorite ‘crystallinity’ (IC and ChC, respectively) are expressed by indices and the most common ones are the Kübler index (KI) for IC and Árkai index (AI) for the ChC (Kübler 1967; Árkai 1991; Árkai et al. 1995; Kübler and Jaboیدoff 2000). These indices are widely applied for determining the grade of diagenesis and low temperature metamorphism of (meta)pelitic rocks. Additionally, the ‘b’ cell dimension of K-white mica estimates the pressure conditions (Sassi 1972; Sassi and Scolari 1974; Guidotti and Sassi 1986; Kisch et al. 2006).

A tool to study the thermal alteration of organic-rich metasedimentary rocks is Raman spectroscopy (e.g., Yui et al. 1996; Beyssac et al. 2002; Rahl et al. 2005; Rantitsch and Judik 2009; Aoya et al. 2010; Wiederkehr et al. 2011; Kouketsu et al. 2014; Lünsdorf et al. 2014; Mori et al. 2016; Lünsdorf et al. 2017). Several authors published empirical geothermometers based on the Raman spectral properties of carbonaceous material (CM), determining peak metamorphic temperatures in the low-grade (Lahfid et al. 2010; Kouketsu et al. 2014) or low- to high-grade (Beyssac et al. 2002; Rahl et al. 2005; Aoya et al. 2010) temperature range.

The major aim of this study is to evaluate the metamorphic conditions and provenance of the Palaeozoic organic-rich metasedimentary Szalatnak Slate Formation from southern Transdanubia, Tisza Mega-unit (Horváthertelend Unit), Hungary (Fig. 2). For this purpose, a multidisciplinary approach including petrography, XRPD, Raman spectroscopy, K–Ar isotopic geochronology and electron microprobe analysis (EMPA) of white mica has been used. Whole-rock geochemistry of the samples has been determined to interpret the provenance area.

Geological setting

The Tisza Mega-unit forms the basement of the Pannonian Basin south of the Mid-Hungarian (Zagreb–Zemplin) Lineament (Fig. 1b) (Csontos and Vörös 2004; Haas and Péró 2004; Schmid et al. 2008). On the north the SW–NE trending Zagorje–Mid-Transdanubian Shear Zone (Pamić and Tomljenović 1998; Haas et al. 2000; Vozár et al. 2010; Fig. 2) comprises a 100 km wide and 400 km long unit between the Periadriatic–Balaton Lineament and the Zagreb–Zemplin Lineament.

The pre-Cenozoic basement of the Hungarian part of the Tisza Mega-unit crops out in the Mecsek and Villány Mts, S Transdanubia. The crystalline complexes and the overlying Palaeozoic and Mesozoic sequences show heterogeneous lithological and metamorphic characteristics (Haas and Péró 2004; Szederkényi et al. 2012). In the studied area, the Palaeozoic formations are predominated by continental sequences. Marine Silurian deposits are present as well. During the Pennsylvanian a molasse-type siliciclastic and locally coal-bearing sequence was deposited, while the Permian is represented by continental red beds with volcanoclastic rocks (Vozár et al. 2010; Szederkényi et al. 2012). The studied Silurian Szalatnak Slate Formation is known in the relatively small and isolated Horváthertelend
and Szalatnak Units (Fig. 2). In the surroundings gneiss, and mica schist with amphibolite intercalations as well as Variscan granitoids (durbachites) of the Mórágy Granite Complex form the basement (Szederkényi 1996; Buda et al. 2004). The Szalatnak Slate was penetrated by boreholes (e.g., Horváthertelend-1 and Szalatnak-3; Árkai et al. 1995; Szederkényi 1996) and this is the oldest fossil-bearing formation in the area (Szalatnak Unit; Fig. 3). Despite its regional importance, only sporadic information is available about the metamorphism of the Horváthertelend Unit (Szederkényi et al. 2012). The Szalatnak Slate is composed of black slate, metagreywacke and polymictic metaglomerate with rare fossiliferous lydite intercalations with a maximum thickness of ~ 500 m (Árkai et al. 1995). According to Oravecz (1964), this formation contains Silurian Hystri-chosphaerida (acritarch) remains and graptolite fragments. Muellerisphaeridae microfauna, radiolarians and conodonts were recovered from the siliceous rocks (Kozur 1984; Fülöp 1994), corresponding to the Pterospathodus amorphognathoides Zone (Llandovery–Wenlock transition).

Based on the phyllosilicate characteristics and vitrinite reflectance of CM, Árkai et al. (1995) proved an anchi–epizonal metamorphic alteration of the Szalatnak Slate. These authors presumed a contact metamorphic overprint caused by a subvolcanic syenite body, belonging to the Mórágy Granite Complex (Fig. 3). The K–Ar isotopic age of the < 2 µm biotite-rich fraction ranges between c. 200 Ma and c. 170 Ma. Árkai et al. (1995) presumed a Variscan metamorphic age with a subsequent thermal overprint.

**Samples and methods**

Detailed macroscopic core investigations from the borehole Horváthertelend–1 (Hh–1) were carried out (Fig. 3). Petrographic studies were conducted on hand specimen (n = 50) and thin sections (n = 75). A multi-method approach, involving metamorphic petrology, microstructural analysis (Paschier and Trouw 2005), whole-rock and mineral chemistry, Raman CM thermometry, mineralogy using XRPD analysis and K–Ar illite geochronology was applied. Mineral abbreviations follow Whitney and Evans (2010).

**Whole rock major and trace element chemistry**

A total of six representative slate samples were selected for chemical analysis (Fig. 3). Whole rock major (ICP-ES method) and trace (ICP-MS method) element analyses were performed at the Bureau Veritas Mineral Laboratories (AcmeLabs), Vancouver, Canada. As a quality control, duplicate analyses were performed on selected samples. The accuracy and analytical precision of the analytical methods were also verified against a standard internal (in house) reference material (standard STD SO-19).

In this study, major element ratios are used to characterize the sedimentary protolith of the Horváthertelend metapelite samples, following the chemical classification schemes of Cox et al. (1995), Herron (1988) and Pettijohn et al. (1972). Additionally, selected major (e.g., Ti) and trace (e.g., La, Th, Hf) element parameters and discriminatory diagrams.
(Bhatia and Crook 1986; Floyd and Leveridge 1987; Floyd et al. 1989) are used to determine the provenance and tectonic setting.

Raman CM thermometry

Eight samples were selected from the borehole Hh-1 to prepare thin sections normal to the foliation for the Raman analysis. It is well known that CM is sensitive to the polishing process which leads to erroneous Raman spectra (Lünsdorf 2016 and references therein). Lünsdorf (2016) proved that the fine polishing (0.05 µm) influences the Raman spectra of the CM significantly, while all preparation steps before, using coarser slurry, do not. To avoid erroneous measurements, abrasion was carried out in three steps using P800, P1000, and P1200 SiC-abrasive powder. Afterwards the samples were polished in a single step using diamond slurry with a grain size of 6 µm. The Raman spectra of 175 CM

Fig. 3 Schematic lithological columns of the two main sections of the Silurian Szalatnak Slate Formation (modified after Árkai et al. 1996 and Mészáros et al. 2016) showing the positions of the investigated cores within the hole Horváthertelend-1
grains were measured using a THERMO Scientific DXR Raman spectrometer. The laser beam was focused beneath the surface of CM grains. The measurements were carried out using an X100 objective lens, a 50 µm pinhole aperture and a 532 nm wavelength Nd-YAG laser with an irradiation power of 2 mW. Every measurement was made using a 900 line/mm grating and the exposure time was 100 s on each individual grain.

The Raman spectrum of CM was fitted by several distinct peaks using peak-fitting software (Peak Fit 4.12 ver.; Sea-Solves Software Inc., MA, USA) with a Voigt function, following the decomposition procedures for Raman CM spectra presented by Kouketsu et al. (2014). For the samples in this study, five distinct bands are identified in the range of 1000–1750 cm⁻¹: ~1580 cm⁻¹ (G-band), 1350 cm⁻¹ (D1-band), 1620 cm⁻¹ (D2-band), 1510 cm⁻¹ (D3-band) and 1245 cm⁻¹ (D4-band). Both calibration procedures of Beyssac et al. (2002) and Kouketsu et al. (2014) were used to provide two independent datasets. Geothermometers proposed by Beyssac et al. (2002), Rahl et al. (2005) and Kouketsu et al. (2014) were applied, respectively.

**XRPD and phyllosilicate characterization**

The whole rock mineralogical composition and characterization of the separated <2 µm grain size (clay) fraction were estimated by X-ray powder diffractometry (XRPD). Unaltered, macroscopically homogeneous rock chips were grounded and homogenised in an agate mortar (< 2 min. grinding time per sample). Grain size separation for clay fraction analysis was achieved by repeated ultrasonic defocclusion and gravitational settling. A total of 19 samples were measured by a Rigaku Ultima IV X-ray diffractometer using Bragg–Brentano geometry, CuKα radiation, graphite monochromator, proportional counter, divergence and detector slits of 2/3°.

For whole rock analysis, random powder mounts were made using ~0.04 g rock powder on a Si single crystal sample holder to determine the mineralogical composition and to characterize mica polytypes. The specimen were scanned at 50 kV/40 mA from 3 to 70°2θ with a goniometer step rate 1°/min and data acquisition steps of 0.05°. The qualitative evaluation of the XRPD spectra was made by Rigaku PDXL 1.8 software using the ICDD (PDF2010) database. Semi-quantitative mineralogical composition was estimated based on reference intensity ratio (RIR) method.

For clay fraction analysis, highly oriented XRPD slides with 3 mg/cm² density were prepared by repeated sedimentation of the separated clay fraction. Both air-dried and ethylene–glycol solvated preparations were scanned at 45 kV/35 mA, from 3 to 50°2θ with goniometer step rate 1°/min and step-width 0.1°. For determination of illite and chlorite ‘crystallinity indices’ and calculation of crystallite size, a triplicate scan of the same slides were run at 40 kV/30 mA, from 3 to 14°2θ with goniometer step rate 0.6°/min and step-width 0.01°.

The determination of the phyllosilicate ‘crystallinity’ was made using the Crystallinity Index Standards (CIS) of Warr and Rice (1994) and the Küberl index (KI_Basel) calculated after Warr and Mählmann (2015).

For standardization of the Küberl index (KI_Basel) and the chlorite ‘crystallinity’ index (ChC_CIS), the Crystallinity Index Standards (CIS) scale was utilized after instructions of Warr and Rice (1994). Instrumental line broadening was determined using an in-house muscovite standard. The narrowest FWHM value measured on the muscovite standard was 0.064° and the sharpest reflection measured for a sample sedimented on a slide was 0.114°. Sample treatment and instrumental conditions for the calibration procedure were strictly the same as during experimental determination of illite and chlorite ‘crystallinity indices’ of the studied sample set. Analytical quality of the standardization was controlled by 10 subsequent parallel measurements of the same slide. Standard deviation is ~2.5% in the case of FWHM10Å = 0.195 and FWHM2Å = 0.198 (n = 10). A good reproduction of the measured values together with a significant correlation were found between CIS_Heidelberg (given by the calibration card and Warr and Mählmann 2015) and CIS_Szeged using FWHM values of ~10 Å reflections (n = 6):

\[ \text{CIS}_\text{Heidelberg} = 1.096 \times \text{CIS}_\text{Szeged} + 0.144, \quad r^2 = 0.960. \]  \hspace{1cm} (1)

A less good correlation was arisen for chlorite ‘crystallinity’ using FWHM values of ~7 Å peaks (n = 4):

\[ \text{CIS}_\text{Heidelberg} = 0.618 \times \text{CIS}_\text{Szeged} + 0.199, \quad r^2 = 0.833. \]  \hspace{1cm} (2)

The conversation of the CIS scale to KI_Basel scale was made according to Warr and Mählmann (2015).

For differentiation between K-, Na- and Ca-rich white micas, the XRPD (00,10) basal reflections around 2 Å were checked using the highly oriented slides (see Frey and Niggli 1972; Árkai et al. 2003, 2004). For a better resolution and precise determination of position of the 00,10 K-white mica peaks, triplicate analyses were made at 50 kV/40 mA, from 40 to 50°2θ with a goniometer scan speed of 0.5°/min and step-width of 0.02°. K-white mica ‘b’ cell dimension was measured on random powder mounts made of the separated clay fraction to eliminate the detrital contamination after the proposal of Padan et al. (1982). Triplicate analyses were made at 50 kV/40 mA from 58 to 53°2θ with a goniometer step rate 0.333°/min and data acquisition steps of 0.03°.
Mineral chemistry

K-white mica composition analysis was made with an energy-dispersive and a wavelength-dispersive equipped JEOL JSM-6310 microprobe. The instrument operated at an acceleration voltage of 15 kV, a focal distance of 15 mm and a beam current of ~6 nA. Mineral standards were used: adularia (Si, Al, K), garnet (Mg, Fe), rhodonite (Mn), titanite (Ca, Ti), jadeite (Na).

K–Ar dating

K–Ar isotope geochronology was carried out on four slate samples containing a large proportion of K-white mica. For every slate sample, both the whole-rock and the <2 µm size fraction were measured. These samples do not contain any other K-bearing mineral phase in detectable amount. After crushing in an agate mortar, 1 g of every bulk samples was isolated. The <2 µm size fraction was separated from aqueous suspension after ultrasonic disaggregation.

Samples were analysed following the procedure of Balogh (1985) at the K–Ar laboratory of the Institute for Nuclear Physics, Hungarian Academy of Sciences, Debrecen. The potassium content was measured on 50 mg sample aliquots after dissolution by HF and HNO₃ using a Sherwood–400 type flame spectrophotometer with an accuracy better than ±1.5%. Separated mineral sample splits were subjected to heating at 100 °C for 24 h under vacuum to remove atmospheric Ar contamination that adsorbed on the surface of mineral particles during sample preparation. Argon was extracted from the minerals by fusing the samples by high frequency induction heating at 1300 °C. The released gases were cleaned in two steps in a low-blank vacuum system by St-700 and Ti-getters. The isotopic composition of the spiked Ar was measured by a Nier-type mass spectrometer. The atmospheric Ar ratio was analysed each day during the measurement period and averaged 295.9 ± 1.85 (1σ) for 60 independent determinations. This value is not significantly different from the theoretical one (295.5; Nier 1950). All isotope measurements were corrected by the atmospheric ⁴⁰Ar/³⁶Ar ratios determined on the day of the analysis.

The accuracy and reproducibility of the isotope ratio measurements were periodically controlled by the Rodina 2/65 internal standard for which the radiogenic ⁴⁰Ar content averaged 13.79 ± 0.12 (2σ) × 10⁻⁶ cm³ g⁻¹ STP after five independent determinations. The recommended value is 13.71 ± 10⁻⁶ cm³ g⁻¹. The decay constants recommended by Steiger and Jäger (1977) were used for age calculation with an overall error of ±2%.

Results

Mineralogical composition and rock structure

The studied samples are composed of dark grey–black slate with grey metasiltstone and metasandstone (metagreywacke) intercalations and red metasandstone (metaarkose) clasts up to several cm in diameter (Fig. 4). The lower part of the section (below the depth of 769 m is, however, predominated by well-foliated pale green–grey slate and monomictic matrix-supported metaconglomerate which contains metasandstone pebbles embedded into a well-foliated fine-grained matrix. The lower part of the metaconglomerate section contains rhyolite pebbles. The samples have a well-developed pressure solution cleavage. The cleavage surfaces are covered by limonitic coatings.

In some samples quartz veins cut through the cores (Figs. 4 and 5). The axial planes of these folded veins coincide with the cleavage planes of the rock. These veins are apparently sheared by pressure solution seams (Fig. 5a). In less-deformed domains of the slate, the original sedimentary lamination is observed. The metasandstone intercalations and clasts are lenticular, often sigmoidal, in shape and occasionally form boudinage structures (Figs. 4 and 5b). Regarding the three principal axes of the clasts, their average axial ratios are X/Z: ~2.8, Y/Z: ~2.7 and X/Y: ~1.0 where X indicates the maximal elongation and Z the maximal shortening direction so the longest axis is parallel to the foliation.

Based on the semi-quantitative XRPD analysis (Supplementary Table 1), the bulk slate samples have a highly variable mineralogical composition with 10–40 mass% K-white mica, 20–50 mass% chlorite, 10–40 mass% quartz and 5–20 mass% albitic plagioclase together with a relatively high amount of amorphous material (up to ~5 mass%, dominantly organic matter and limonite). The matrix of the slate has a moderately developed continuous foliation with oriented sericitic K-white mica and chlorite bands. Occasionally, the continuous foliation associates with an anastomosing pressure solution cleavage, having carbonaceous material and limonite (presumably after pyrite) (Fig. 5c). Both in the matrix and in the quartz veins all studied samples contain randomly oriented needle-shaped anatase crystals (~50 µm), reflecting prekinematic idio- and hypidioblasts (Fig. 5d). Additionally, both the lenticular metasandstone clasts and the prekinematic anatase grains have chlorite + quartz pressure shadows parallel to the foliation. In the pale-grey slate, these deformation structures have a slightly monoclinal symmetry. The folded preterctic quartz veins are dynamically recrystallized (Fig. 5d) and the quartz grains have a strongly undulose extinction and subgrain microstructure. Along the serrated grain-boundaries small undeformed neoblasts were formed. In some samples euhedral tourmaline needles
mica and chlorite content increases. Margins of the clasts, plagioclase disappears while K-white rite and opaque minerals. Additionally, the relatively large grains. Detrital biotite is intensively degraded to chlo-

erate composition (~ 20 vol%) and detrital plagioclase (~ 10 vol%) (Fig. 4e–h). The metaarkose clasts are well-sorted, quartz (~70 vol%), volcanic rock fragments with intermedi-
tely active environment (e.g., Cox et al. 1995). Relatively cate geochemically immature deposits, suggesting a tectoni-
dering diagram (Fig. 7), the samples display systematic light REE enrichment trends. Apart from the sample at the maximum depth, the metapelites show quite uniform patterns with a negative Eu anomaly and near-flat heavy REE patterns. These features reflect their derivation from typical fraction-
ated (mature) upper continental crust (Taylor and McLennan 1985; McLennan 1989). On the other hand, the unique sample has the highest ΣREE value and is more enriched in light REEs. These features together with a fractionated heavy REE pattern reflect its higher content in accessories (e.g., zircon).

Geochemical results indicate a dominantly felsic (silicic) source area and point to an intermediate to acidic volcanic arc terrane as an important component of the provenance area. High amounts of metasandstone pebbles in the meta-
conglomerate, however, indicate that a recycled quartz-rich sediment source is also possible.

Raman spectroscopy

CM flakes are randomly dispersed in the rocks but they are enriched in the pressure solution seams and in some meta-

greywacke samples. The graphitic material often associates with anatase and altered pyrite, which generally is trans-
formed to goethite. In addition to the autochthonous grain population, a highly ordered, probably allochthonous, gra-
phitic carbon grain population is also present. This highly ordered population has a well-rounded grain-shape and small grain-size (10–15 µm) with a distinctive, ordered Raman spectra, showing strong G band and very weak D1 and D2 bands. This population was excluded from the thermometry procedure because of its assumed detrital origin. The grain population located in microlithons was used in this study for RSCM thermometry due to the lack of shearing deformation.

The autochthonous CM population is a low-grade disor-
dered graphitic material (Table 2). Its spectra display a sym-
metric D1 band and a negligible D3 band with an average R1 (D1intensity/Gintensity, Beyssac et al. 2002) ratio of ~1.5. D2 defect band forms a distinct shoulder on the G band (Fig. 8). The Raman spectra suggest that the metamorphism took place between ~330 and ~400 °C maximum metamorphic temperature (Beyssac et al. 2002; Aoya et al. 2010; Lahfid et al. 2010; Kouketsu et al. 2014; Beyssac et al. 2016).

Using the calibration of Beyssac et al. (2002), RSCM thermometry shows a normal distribution with a mode at 351 °C whereas the arithmetic mean of the estimated temperature is 356 ± 21 °C. The modified thermometer of Rahl
et al. (2005) shows a mode of 362 °C and a mean temperature of 349 ± 35 °C. Obviously, the mode provides a better estimation of the expected value of the metamorphic temperature than does the arithmetic mean because of the skewness of the distribution. The applied thermometers suggest an estimated peak metamorphic temperature of ~ 350–370 °C.

**X-ray powder diffraction examination of the < 2 µm fraction**

The mineralogical composition of the < 2 µm fraction is K-white mica + chlorite + quartz ± albite ± kaolinite (Supplementary Table 1). Kaolinite-bearing samples were excluded from the chlorite ‘crystallinity’ determinations because of the resulted interference of 001 peak of the kaolinite and 002 peak of the chlorite at ~ 7 Å.

Characteristic d spacings of the K-white mica (3.74 Å, 3.00 Å, 2.80 Å) indicate a predominance of the 2M₁ polymytype. The position of the 00,10 reflection estimates a 1.995–2.001 Å d spacing. There is no shoulder observable on any sides of this peak, suggesting a near-theoretical ‘muscovite’ composition for the K-white mica without any Na- or Ca-substitutions referring ‘mixed’ K-Na-mica (Livi et al. 1997). According to the ethylene glycol treatment of the samples, neither K-white mica nor chlorite contains any interstratified smectitic swelling phase.

Esquevin index of the K-white mica, i.e., intensity ratio of the 5 Å and 10 Å peaks (Esquevin 1969), shows a phengitic composition for most of the studied samples (Fig. 9a, Table 3). After Dunoyer de Segonzac (1970) and Árkai et al. (1995), only samples with > 0.25 Esquevin index were used to determine the KI (Kübler 1964; Kübler and Jaboyedoff 2000). The KI_Basel values cover a range from 0.19 to 0.38...
Δ^2θ with a mean of 0.22 ± 0.04 Δ^2θ (Fig. 9b; Table 3). The ChC_CIS values range from 0.27 to 0.53 Δ^2θ with a mean of 0.31 ± 0.06 Δ^2θ (Fig. 9c, Table 3). Between the KI_base and ChC_CIS values a moderate correlation (R = 0.77) is observed (Fig. 10). The mean crystallite size of the K-white mica calculated applying the Scherrer-equation (Klug and Alexander 1974; Merriman et al. 1990) falls between 288 ± 14 Å and 1293 ± 54 Å. Estimated mean crystallite size based on the chlorite 002 reflection fluctuates from 157 ± 4 Å to 1043 ± 43 Å. The calculated mean crystallite size values, excluding the anomalous values of the altered samples (745.1b, 779.6 and 787), are 890 ± 198 Å for K-white mica and 692 ± 175 Å for chlorite (Table 3).

Because of the K-white mica + chlorite + quartz + albite ± anatase mineral assemblage and the absence of paragonite, pyrophyllite or chloritoid indicative for Al-saturated metapelites (Franceschelli et al. 1989), the method and the scale of Guidotti and Sassi (1986) are applied. Due to the high amount of CM reducing (low fO2) conditions are presumed during metamorphism, and primary origin of hematite or goethite are ruled out. The measured ‘b’ cell dimension values range from 9.011 to 9.029 Å with a mean of 9.019 ± 0.004 Å (Fig. 9d).

**Mineral chemistry**

The Si content of the K-white mica ranges between 6.26 and 6.73 apfu. Fe^{2+} content varies between 0.28 and 0.38 and the Mg ranges between 0.34 and 0.57. The calculated Na/(Na + K) ratios fluctuate between 0.02 and 0.08 (Supplementary Table 2).

**Geochronology**

The K–Ar radioisotopic age data of the whole rock samples and those of the <2 μm grain-size fraction fall in the range
| Major elements (in mass%) | Samples | 765.5 | 776.2 | 777 | 777.2 | 778 | 779 |
|---------------------------|---------|-------|-------|-----|-------|-----|-----|
| SiO₂                      | 61.59   | 62.48 | 60.14 | 62.87| 63.61 | 64.18 |
| Al₂O₃                     | 17.93   | 16.60 | 17.17 | 17.59| 17.01 | 18.87 |
| Fe₂O₃                     | 7.00    | 7.72  | 7.99  | 6.19 | 6.02  | 4.00  |
| MgO                       | 2.81    | 3.19  | 3.68  | 2.92 | 2.92  | 2.05  |
| CaO                       | 0.46    | 0.28  | 0.26  | 0.30 | 0.29  | 0.08  |
| Na₂O                      | 1.06    | 2.86  | 2.36  | 2.71 | 1.79  | 1.01  |
| K₂O                       | 3.95    | 2.19  | 2.45  | 2.69 | 3.03  | 4.73  |
| TiO₂                      | 0.77    | 0.76  | 0.74  | 0.76 | 0.72  | 0.71  |
| P₂O₅                      | 0.30    | 0.12  | 0.11  | 0.13 | 0.16  | 0.05  |
| MnO                       | 0.09    | 0.07  | 0.10  | 0.08 | 0.05  | 0.03  |
| LOI                       | 3.80    | 3.50  | 4.80  | 3.60 | 4.20  | 4.10  |
| Total                     | 95.96   | 96.27 | 95.00 | 96.24| 95.60 | 95.71 |
| ICV                       | 0.90    | 1.03  | 1.02  | 0.89 | 0.87  | 0.67  |
| Trace elements (in ppm)   |         |       |       |     |       |      |
| Ba                         | 543     | 418   | 430   | 469 | 488   | 681   |
| Ni                         | 33      | 52    | 51    | 41  | 57    | 25    |
| Sc                         | 21      | 21    | 20    | 21  | 20    | 17    |
| Be                         | 4.0     | 6.0   | <1    | 1.0 | 2.0   | 3.0   |
| Cr                         | 13      | 15    | 14    | 15  | 16    | 18    |
| Co                         | 18.2    | 23.5  | 22.8  | 20.2| 23.0  | 8.4   |
| Cs                         | 4.3     | 2.8   | 4.6   | 3.9 | 3.5   | 6.5   |
| Ga                         | 26.6    | 18.5  | 23.4  | 21.5| 17.8  | 21.6  |
| Hf                         | 4.4     | 4.2   | 3.3   | 3.9 | 4.4   | 6.9   |
| Nb                         | 10.3    | 8.1   | 8.7   | 10.5| 8.0   | 12.4  |
| Rb                         | 120.9   | 71.2  | 75.7  | 84.4| 94.1  | 155.6 |
| Sn                         | 3.0     | 1.0   | 3.0   | 2.0 | 2.0   | 3.0   |
| Sr                         | 64      | 156   | 121   | 148 | 94    | 89    |
| Ta                         | 0.7     | 0.6   | 0.6   | 0.7 | 0.9   | 0.8   |
| Th                         | 9.6     | 8.1   | 8.1   | 8.7 | 8.7   | 29.0  |
| U                          | 2.2     | 2.9   | 2.4   | 2.4 | 2.4   | 7.5   |
| V                          | 128     | 141   | 125   | 134 | 124   | 122   |
| W                          | 2.9     | 3.0   | 2.2   | 2.7 | 2.2   | 4.3   |
| Zr                         | 144     | 146   | 122   | 139 | 149   | 240   |
| Y                          | 28.9    | 20.5  | 20.8  | 24.2| 20.7  | 23.7  |
| La                         | 24.8    | 25.4  | 25.5  | 23.9| 29.4  | 61.3  |
| Ce                         | 57.4    | 53.5  | 53.7  | 53.9| 60.7  | 117.2 |
| Pr                         | 6.9     | 6.1   | 6.4   | 6.0 | 7.0   | 12.2  |
| Nd                         | 26.1    | 23.1  | 23.6  | 22.3| 27.7  | 43.4  |
| Sm                         | 5.47    | 4.17  | 4.90  | 4.58| 4.91  | 6.68  |
| Eu                         | 1.14    | 0.93  | 1.09  | 0.99| 1.13  | 1.71  |
| Gd                         | 4.77    | 3.96  | 4.53  | 4.46| 4.31  | 6.03  |
| Tb                         | 0.75    | 0.59  | 0.69  | 0.68| 0.65  | 0.84  |
| Dy                         | 5.21    | 3.85  | 4.18  | 4.36| 4.05  | 4.80  |
| Ho                         | 1.12    | 0.77  | 0.79  | 0.93| 0.77  | 0.82  |
| Er                         | 3.02    | 2.25  | 2.54  | 2.84| 2.57  | 2.46  |
| Tm                         | 0.51    | 0.36  | 0.36  | 0.39| 0.37  | 0.36  |
| Yb                         | 3.03    | 2.47  | 2.40  | 2.69| 2.57  | 2.21  |
| Lu                         | 0.46    | 0.39  | 0.41  | 0.43| 0.41  | 0.34  |
of 302–319 Ma and 288–300 Ma, respectively (Table 4). The K–Ar isochron method (Shafiqullah and Damon 1974) is used to demonstrate extraneous argon or argon loss in a co-genetic suite of samples of different K-contents or the presence of detrital minerals. Two arrays of data can be distinguished on the diagram and the fitted lines intersect the 40Ar/36Ar axis close to the atmospheric value at 295.5. This indicates that the two sample sets are co-genetic and post-genetic thermal processes did not result in argon loss or gain from the samples. Whole rock data yield an isochron age of c. 312 Ma and the separated <2 µm fraction samples yield an age of c. 297 Ma (Fig. 11).

Discussion

Metamorphic conditions and age

Thermometers of Beyssac et al. (2002) and Kouketsu et al. (2014) give temperature estimations in the range of errors of RSCM methods. For the Szalatnak Slate, the peak temperature of metamorphism is estimated ~ 350–370 °C and shows homogeneous distribution along the studied Horváthertelend section. The XRPD-based phyllosilicate 'crystallinity' data suggest a metamorphic grade corresponding to the ~ 350 °C temperature.

In most of the samples, KI values indicate a K-white mica formation during epizonal metamorphic conditions (Kübler and Jaboyedoff 2000; Warr and Rice 1994) and suggest a low-grade metamorphism. Based on the I 5 Å/l 10 Å ratios (Esquevin 1969) and EMPA data, these K-white mica populations are phengitic in composition which is common in low-grade epizonal conditions (Erns 1963; Frey and Robinson 1999; Massonne and Schreyer 1987; Velde 1965). The CIS calibrated Árkai indices, however, indicate anchizonal metamorphic conditions (zone boundaries after Kübler and Jaboyedoff 2000 correlated with ChC₀₀₂ in Árkai 1991). Nevertheless, the correlation between the KI_base and ChC_CIS values indicates a genetic relationship between the phengitic K-white mica and chlorite in the studied rocks. The

Table 1 (continued)

| Samples | 765.5 | 776.2 | 777 | 777.2 | 778 | 779 |
|---------|------|------|-----|------|-----|-----|
| ΣREE    | 141  | 128  | 131 | 128  | 147 | 260 |

Table 2 Estimated metamorphic temperatures of the studied samples based on Raman spectroscopy of the carbonaceous material

| Depth (m) | n   | R1         | R2         | T (°C)α  | T (°C)β  | FWHM D1 (cm⁻¹) | FWHM D2 (cm⁻¹) | T (°C)γ  | T (°C)δ  |
|-----------|-----|------------|------------|----------|----------|----------------|----------------|----------|----------|
| 753.0     | 28  | 1.42 ± 0.25| 0.63 ± 0.06| 360 ± 26 | 352 ± 43 | 52.3 ± 10.9    | 24.7 ± 3.5     | 364 ± 24 | 367 ± 28 |
| 772.3     | 23  | 1.65 ± 0.25| 0.64 ± 0.06| 358 ± 25 | 356 ± 30 | 51.9 ± 6.6     | 26.4 ± 3.6     | 366 ± 14 | 356 ± 24 |
| 777.0     | 22  | 1.40 ± 0.26| 0.64 ± 0.04| 356 ± 20 | 340 ± 26 | 57.6 ± 12.5    | 27.7 ± 3.2     | 354 ± 27 | 347 ± 22 |
| 779.6     | 19  | 1.47 ± 0.21| 0.64 ± 0.05| 356 ± 23 | 348 ± 42 | 59.9 ± 14.2    | 28.1 ± 3.3     | 349 ± 31 | 344 ± 23 |
| 780.3     | 20  | 1.42 ± 0.17| 0.63 ± 0.04| 360 ± 19 | 354 ± 34 | 56.8 ± 13.3    | 27.3 ± 2.6     | 356 ± 29 | 350 ± 18 |
| 780.0     | 18  | 1.50 ± 0.15| 0.64 ± 0.05| 353 ± 22 | 344 ± 44 | 59.5 ± 12.7    | 27.0 ± 6.1     | 350 ± 28 | 352 ± 41 |
| 781.0     | 25  | 1.64 ± 0.15| 0.63 ± 0.02| 357 ± 9  | 363 ± 17 | 53.5 ± 8.7     | 26.0 ± 3.5     | 363 ± 19 | 359 ± 24 |
| 790.0     | 15  | 1.45 ± 0.20| 0.63 ± 0.04| 357 ± 20 | 349 ± 32 | 61.2 ± 10.1    | 26.3 ± 2.4     | 346 ± 22 | 357 ± 17 |
| Sum       | 170 | 1.50 ± 0.26| 0.27 ± 0.04| 356 ± 21 | 349 ± 35 | 56.1 ± 11.5    | 26.6 ± 3.7     | 365 ± 25 | 355 ± 25 |

FWHM, full width at half maximum
Temperature estimation using thermometer of αBeyssac et al. (2002); βRahl et al. (2005); Kouketsu et al. (2014) based on γD1 band and δD2 band
calculated crystallite sizes of the white mica and the chlorite are also consistent with this observation.

The "b" cell dimension values of 9.00 Å–9.04 Å of the K-white mica in the < 2 µm grain size fraction are characteristic of a medium pressure geodynamic environment (Fig. 12) during the formation of K-white mica in the ms + chl + qz + ab ± ant assemblage (Guidotti and Sassi 1986; Sassi F 1972; Sassi and Scolari 1974). The estimated 2.5–4 kbar pressure supports this conclusion.

The characteristic deformational microstructures, such as the pervasive continuous foliation, the moderately developed pressure solution cleavage and chl + ms + qz mica beards in pressure shadows, indicate diffusional mass transfer. Deformed vein quartz with subgrain structure and neoblasts along its margins are indicative of a low-temperature grain boundary migration recrystallization (Paschier and Trouw 2005; Stipp and Kunze 2008; Stipp et al. 2002). These features correspond to epizonal thermal conditions during the deformation as well (Paschier and Trouw 2005; Stipp et al. 2002). The appearance of the deformational microstructures indicates flattening strain based on the principal axis of the clasts (Flinn 1962; Ramsey and Huber 1983).

The whole rock and the separated <2 µm fraction mean ages slightly differ (310.1 ± 8.1 Ma and 293.1 ± 7.7 Ma, respectively). The K–Ar age increases with the increasing grain-size which is considered as detrital effect or multi-phase K-white mica crystallization (Clauser and Chaudhuri 1995; Clauser and Weh 2014). Based on the petrographic and XRPD examinations, only the K-white mica is a stoichiometrically K-bearing mineral phase. The K-white mica is regarded as product of a single, progressive metamorphic event. Therefore, both studied grain-size fractions suffered resetting near the closure temperature of phengitic K-white mica (~350 °C in the case of coarse-grained crystals; Hunziker 1986; Hunziker et al. 1986). Consequently the K–Ar radioisotopic age of <2 µm size fraction with lower closure temperature (~260 °C; Hunziker 1986; Hunziker et al. 1986) is considered as a cooling age (Glasmacher et al. 2001). The relatively small K–Ar age difference of the grain-size fractions supports the cooling age model rather than the detrital effect. Because the estimated peak metamorphic temperature is higher than the closure temperature a resetting of the K–Ar isotopic system is assumed (Leitch and McDougall 1979). The K–Ar data of the < 2 µm suggest post-Variscan (c. 290 Ma) uplift while the whole rock age data show a Variscan (c. > 310 Ma) metamorphism.

Regional correlation

The Tisza Mega-unit is a large lithosphere block of complex internal structure. From the point of view of the palaeogeographic reconstruction, this exotic terrane was often neglected, or its role in the regional correlation was not carefully considered (Haas and Péro 2004). Verniers et al. (2008) provided an extensive review of the available literature for outcrop areas and subsurface presence of the Silurian in Central Europe, including some Slavonian occurrences in the Tisza Mega-unit (Mt. Psunj, Mt. Papuk and Mt. Krndija; Fig. 2). Unfortunately, however, there was no mention about Hungarian records in that article.

The most intensively studied metasediments of the Tisza Mega-unit are located in the Slavonian Mts. (Croatia) (e.g., Balen et al. 2015; 2018). The Radlovac Complex (Fig. 2) consists of slates, phyllices, metagreywackes and metaconglomerates with local (meta)basic intrusions (Pamić and Jamičić 1986; Jamičić 1988). In spite of some lithological similarity, however, the Radlovac metasedimentary rocks significantly differ from those of the Horváthertelend Unit. The possible presence of Late Silurian slates, belonging to the Noric–Bosnian Terrane of peri-Gondwana (Verniers et al. 2008), was mentioned by Jerenić et al. (1994). Nevertheless, in the Radlovac Complex Pennsylvanian and Permo triassic rocks are dominantly exposed with Alpine low-T regional metamorphism (Biševac et al. 2010, 2011, 2013).
Fig. 9  a Esquevin index versus KI_{Basel} values plot (after Esquevin 1969) suggests dominantly phengitic K-white mica composition of most samples. b KI_{Basel} plot indicates epizonal metamorphism of the formation. Zone boundaries are indicated after Kübler and Jaboyedoff (2000) and Warr and Mählmann (2015). c ChCCIS plot shows chlorite ‘crystallinity’ values of the studied samples. Note: empty circle indicates the presence of kaolinite in the studied sample. d Esquevin index versus ‘b’ cell dimension diagram of the samples. The highlighted area indicates samples suitable for pressure characterization.

Table 3  XRPD parameters of the 10 Å and 14 Å mineral phases of the <2 µm fraction

| Sample (m) | I(5 Å)/I(10 Å) | d_{0010} (Å) | KI_{Basel} (Δ^2θ) | ChCCIS (Δ^2θ) | K-white mica crystallite size (Å) | Chlorite_{002} crystallite size (Å) | ‘b’ cell dimension (Å) |
|------------|----------------|-------------|-------------------|--------------|----------------------------------|-----------------------------------|----------------------|
| 745.1a     | 0.41           | 2.000       | 0.226±0.003       | 0.417±0.045  | 739±24                           | 250±48                            | 9.007±0.006          |
| 745.1b     | 0.50           | 2.001       | 0.264±0.15       | 0.529±0.009  | 528±61                           | 157±14                            | 9.009±0.007          |
| 749.2      | 0.50           | 2.000       | 0.242±0.005      | 0.317±0.003  | 625±30                           | 483±14                            | 9.017±0.005          |
| 752.6      | 0.44           | 1.999       | 0.231±0.000      | 0.315±0.001  | 701±3                            | 497±4                            | 9.029±0.005          |
| 759        | 0.36           | 1.998       | 0.219±0.002      | 0.308±0.001  | 804±18                           | 537±10                            | 9.023±0.002          |
| 765.6      | 0.31           | 1.999       | 0.224±0.001      | 0.309±0.004  | 757±10                           | 531±24                            | 9.018±0.003          |
| 768.1      | 0.30           | 1.998       | 0.211±0.004      | 0.318±0.002  | 894±45                           | 478±10                            | 9.015±0.003          |
| 772.3      | 0.25           | 1.997       | 0.209±0.005      | 0.291±0.002  | 926±65                           | 669±16                            | 9.012±0.006          |
| 775.6      | 0.29           | 1.997       | 0.208±0.001      | 0.278±0.000  | 941±13                           | 853±6                             | 9.022±0.003          |
| 777        | 0.24           | 1.997       | 0.205±0.002      | 0.276±0.002  | 1061±75                          | 897±52                            | 9.022±0.002          |
| 777.2a     | 0.31           | 1.997       | 0.218±0.001      | 0.290±0.001  | 813±14                           | 682±9                             | 9.020±0.003          |
| 777.2b     | 0.31           | 1.997       | 0.218±0.003      | 0.299±0.001  | 817±28                           | 598±7                             | 9.015±0.005          |
| 779.6      | 0.31           | 1.998       | 0.196±0.001      | 0.359±0.010  | 1193±37                          | 341±23                            | 9.015±0.002          |
| 780.3      | 0.27           | 1.999       | 0.217±0.011      | 0.290±0.002  | 825±11                           | 683±15                            | 9.015±0.003          |
| 780.4      | 0.32           | 1.997       | 0.205±0.002      | 0.280±0.002  | 992±41                           | 809±30                            | 9.025±0.002          |
| 786.7      | 0.31           | 1.997       | 0.206±0.003      | 0.277±0.003  | 979±55                           | 876±48                            | 9.018±0.000          |
| 786.8      | 0.29           | 1.997       | 0.198±0.001      | 0.270±0.003  | 1144±36                          | 1043±82                           | 9.013±0.005          |
| 787        | 0.10           | 1.995       | 0.379±0.014      | 0.280±0.001  | 288±14                           | 813±18                            | 9.027±0.005          |
| 796        | 0.26           | 1.996       | 0.192±0.001      | 0.285±0.003  | 1293±54                          | 745±38                            | 9.022±0.003          |
In the surrounding area of the Tisza Mega-unit, the graptolite-bearing Silurian deposits in the Medvednica Mountain (Croatia; Fig. 2) belong to the distal sedimentary successions on peri-Gondwana, Eastern Alps (Verniers et al. 2008) which passed through Alpine prograde metamorphism as well (Judik et al. 2008; Rantitsch and Judik 2009). Further west, the Silurian of the Carnic Alps (Fig. 13), corresponding to the Proto-Alps Terrane, displays a northern Gondwana appearance. Sequences consist typically of either black shales or calcareous shales and limestones with a low terrigenous influx (Verniers et al. 2008). Consequently, the abovementioned Silurian rocks have no direct importance for palaeogeographic correlation of the study area with a conglomerate-bearing proximal succession.

Previous works on correlation of the Tisza Mega-unit (Buda et al. 2000) pointed out that there are similarities in lithology and in Variscan evolutionary history of the southern Transdanubian area and the Moldanubian Zone of the Bohemian Massif. The Variscan Mórágy Granite known from the SW Tisza Mega-unit was correlated with the Variscan durbachitic granitoid plutons of the central and eastern Bohemian Massif (Gy et al. 2004; Klötzli et al. 2004; Varga et al. 2003) highlighted petrographic and geochemical similarities between the Pennsylvanian continental Téseny Sandstone, Transdanubia (Hungary), and the Cracow Sandstone, Upper Silesian Coal Basin (Poland). Furthermore, the protolith and metamorphic evolution of the sporadic serpentinite occurrences of the SW Tisza Mega-unit (Gyód and Helesfa Serpentinite) reflect a close relationship with the serpentinite bodies of the Góry Sowie Massif in the W Sudetes (Kovács et al. 2009, 2016).

In the Bohemian Massif, numerous occurrences of Silurian sedimentary sequences are known (Fig. 13). The most characteristic territories are the Barrandian Basin (Prague Basin, corresponding to the Silurian Perunica Terrane), the ‘Islet Zone’, the Železné Hory Mts and the Hlinsko and the Lužice Regions (Suchý et al. 2002, 2015; Štorch 1999; Štroch and Kraft 2009). The Silurian succession of the Prague Basin was only slightly deformed (Verniers et al. 2008). In contrast, the Silurian successions of the ‘Islet Zone’ passed through contact and regional metamorphism. In the Moravo-Silesian Zone of the Brunovistulicum, only a unique Silurian occurrence is known in the Drahany Highland near the village of Stínava (Kalvoda et al. 2008, Verniers et al. 2008).

It is important to note, however, that the Silurian sediments of the Bohemian Massif belong to the distal

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![Kübler index versus chlorite ‘crystallinity’ index (ChCكس) plot of the studied samples from the borehole Horváthertelend-1](image1)

**Fig. 10** Kübler index versus chlorite ‘crystallinity’ index (ChCكس) plot of the studied samples from the borehole Horváthertelend-1

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![40K/36Ar versus 40Ar/36Ar isochron plot of the analysed whole rock (solid line) and separated < 2 μm grain-size fractions (dashed line)](image2)

**Fig. 11** $^{40}\text{K}/^{36}\text{Ar}$ versus $^{40}\text{Ar}/^{36}\text{Ar}$ isochron plot of the analysed whole rock (solid line) and separated < 2 μm grain-size fractions (dashed line)

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**Table 4** Results of the K–Ar analysis of the whole rock and the < 2 μm grain-size fraction

| Sample | Depth (m) | Size fraction | K (%) | $^{40}\text{Ar}_{\text{rad}} \times 10^{-6}$ (cm$^3$/g) | $^{40}\text{Ar}_{\text{rad}}$ (%) | Age (Ma + 1σ) |
|--------|-----------|---------------|-------|---------------------------------|----------------|--------------|
| 1-01-B  | 777.2–778.0 | Whole rock     | 1.98  | 26.14                           | 68.8           | 311.8 ± 7.9 |
| 1-01-2 M | 777.2–778.0 | < 2 μm         | 3.04  | 37.70                           | 70.6           | 293.5 ± 7.9 |
| 1-02-B  | 777.2–778.0 | Whole rock     | 2.19  | 29.77                           | 79.0           | 319.4 ± 8.2 |
| 1-02-2 M | 777.2–778.0 | < 2 μm         | 3.29  | 40.20                           | 72.2           | 289.8 ± 7.7 |
| 1-03-B  | 777.2–778.0 | Whole rock     | 2.55  | 33.32                           | 76.9           | 308.0 ± 8.0 |
| 1-03-2 M | 777.2–778.0 | < 2 μm         | 3.63  | 44.13                           | 74.1           | 288.5 ± 7.6 |
| 1-04-B  | 777.0      | Whole rock     | 3.16  | 40.52                           | 78.1           | 302.8 ± 7.8 |
| 1-04-2 M | 777.0      | < 2 μm         | 4.32  | 54.92                           | 86.1           | 300.5 ± 7.6 |

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sedimentary successions on northern Gondwana and peri-Gondwana (Verniers et al. 2008 and references therein). The Silurian sequences are black shales or calcareous shales and basinal limestones (Fig. 13). Additionally, the sedimentation was influenced by volcanic activity with deposition of lava flows, tuffs, volcanioclastics as well as tuffaceous limestones (Verniers et al. 2008).

Eastwards from the Bohemian Massif, in the foreland of the Variscan orogen, widespread occurrences of more proximal non-metamorphosed Silurian sequences are known (Łysogóry Region and Małopolska Massif, Holy Cross Mountains, Poland), belonging to the Baltica Terrane (Verniers et al. 2008). In the northern Łysogóry Region (N Holy Cross Mts), the Silurian begins with black graptolite shale (Llandovery–Wenlock) which turns into sandy shale, siltstone and greywacke in the Ludlow–Pridoli. In this sequence carbonate rocks are known in subordinate amounts forming interbeds and lenses (Kozłowski 2008).

To the south from the Łysogóry Region, in the Kielce Region (N Małopolska Massif, S Holy Cross Mts) the Silurian shows similarities to the Łysogóry occurrences. Here the Llandovery–Wenlock is characterized by dark graphite-shales with lydite intercalations and by calcareous black shales in the Wenlock–Ludlow. Contrarily, Ludlow coarse-grained greywacke and the Upper Ludlow–Pridoli siltstone and fine-grained sandstone predominates the Upper Silurian sediments in the Kielce Region (Kozlowski 2008; Kozlowski et al. 2014; Malec 1993).

The Ludlow greywackes in the Holy Cross Mts indicate that they have a reworked continental island arc provenance (Kozlowski et al. 2014). The transport directions reflect location of this area westwardly of the Holy Cross Mts. Older sedimentary and metasedimentary rocks with high amounts of cherts and variously differentiated volcanic rocks, andesitic to dacitic in composition, were the source of detritus. The rapid deposition of the sediments, composed of immatured material, was related to turbidite sedimentation (Kozłowski 2008; Kozłowski et al. 2014). Conglomerate-bearing coarse-grained formations are known only from the Małopolska Massif (Nida Region and Kielce Region; Malec et al. 2016; Verniers et al. 2008). In the Nida Region, the Silurian strata are composed of graphitic shale with lydite and carbonate intercalations in the Llandovery–Ludlow and siltstone, greywacke and fossil-free conglomerate (Miedziana Góra Conglomerate; Malec 1993) in the Pridoli (Moldinski and Szymanski 2001). The locally preserved conglomerate bodies are composed of clasts of Ordovician sandstones and Cambrian quartzites (Kozłowski et al. 2014). The Carpathian Foreland possibly represents the easternmost part of the Małopolska Massif and indicates a similar but more variable depositional environment to that in the Nida Region (Verniers et al. 2008). In the Kielce Region, greywackes with mudstone interbeds and conglomerates of Ludlow age above graphitic claystones are exposed in the Holy Cross Mts at Niestachów (Malec et al. 2016). Petrographic examination of Malec et al. (2016) indicates that these greywacke conglomerates and sandstones are composed of acidic–intermediate volcanic and sedimentary rock fragments, with subordinate metamorphic and scarce plutonic clasts.

In the Horváthertelend Unit, the absence of calcareous successions together with the dominance of coarse clastics...
suggested a more proximal palaeogeographic position. The Horváthertelend metapelite samples are texturally and geochemically immature. Petrographic and geochemical features described above indicate recycled orogenic signatures with older quartz-rich sedimentary rocks and with a distinct intermediate to acidic magmatic arc component. Based on the presence of thick polymictic conglomerate intercalations with pebbles up to 20–25 cm in diameter, it can be assumed that the Silurian sediments of the Horváthertelend Unit had source areas located on the orogen side of the depositional basin.

The Szalatnak Slate Formation in the northern part of the SW Tisza Mega-unit has some remarkable similarities to the Silurian fossil-free marine, coarse clastics of the Holy Cross Mts, especially of the southern part of the Malopolska Massif near to the Upper Silesia terrane. In both depositional areas carbonate intercalations are present in subordinate amounts; additionally, these areas are predominated by proximal lithologies (greywacke, conglomerate) with geochemical signatures of active continental margin and volcanic arc tectonic settings in the upper part of the sections. In accordance with the more proximal character of the Horváthertelend foreland basin in comparison to the Kielce and Łysogóry Regions, we conclude that the Horváthertelend Unit is maybe located to the west of the Malopolska Massif (present coordinates). This reconstruction is independently confirmed by the similar framework composition of the Pennsylvania Téseny Sandstone (SW Tisza Mega-unit) and the Cracow Sandstone (Upper Silesia) (Varga et al. 2003).

It is important to note, however, that the Silurian and the Upper Palaeozoic of the Malopolska Massif did not pass through Variscan metamorphism. The easternmost occurrence of the Variscan metamorphic Silurian and Upper Palaeozoic in the Bohemian Massif is known from the Moravo-Silesian Zone characterized by 340–320 Ma uplift ages (Dallmeyer et al. 1992; Schulmann et al. 2014; Štípská et al. 2015). In the Silesian Block of the Moravo-Silesian Zone, K-white mica K–Ar and monazite U–Th–Pb ages of 300–280 Ma are related to the post-orogenic felsic magmatism which overprinted a Barrovian type regional metamorphism reported from the NE part of the Moravo-Silesian Zone (Schulmann et al. 2014). Similar post-orogenic granodiorites with zircon U–Pb ages of ~300 Ma are known from the border region between the Malopolska Massif and the Upper Silesian Massif along the Kraków–Lubliniec Fault zone (Żelaźniewicz et al. 2008). The post-orogenic Variscan K–Ar ages (c. 290 Ma) of clay-size K-white mica in the

Horváthertelend Unit (Tisza Mega-unit) shows very similar ages like the overprint of the post-orogenic magmatism in the NE part of Moravo-Silesian Zone and S Malopolska Massif.

Based on the lithological and geochemical features together with mica cooling ages, the Silurian Szalatnak Slate Formation of the Tisza Mega-unit seems to be referred to the Silurian of the S Malopolska Massif (Baltica) and the NE part of the Moravo-Silesian Zone.

Conclusions

The Silurian Szalatnak Slate Formation from the Horváthertelend Unit, NW part of the Tisza Mega-unit (Pannonian Basin, Hungary) consists of slates, volcanic lithic-rich metagreywackes and metaconglomerates with a combined felsic volcanic arc and recycled-orogen (e.g., quartz-rich metasediments) provenance. Both petrographic and geochemical features reflect their derivation from fractionated (mature) upper continental crust. The appearance of coarse-grained lithologies in the studied section suggests a proximal palaeogeographic position during sedimentation.

The slate samples have a highly variable mineralogical composition with a K-white mica, chlorite, quartz and albite plagioclase assemblage together with a relatively high amount of amorphous material. The matrix of the slate has a moderately developed continuous foliation with oriented sericitic K-white mica and chlorite bands. In some cases, it associates with an anastomosing pressure solution cleavage, enriched in CM and limonite.

The K-white mica in the clay fraction has a phengitic composition and shows a strong genetic relationship with chlorite. KL базел of the K-white mica and ChC_C16 values of the chlorite suggest an epizonal metamorphic alteration. This is in accordance with the results of RSCM thermometry which show ~350–370 °C temperature for the peak metamorphism and contemporaneous ductile deformation. The ‘b’ cell dimension of K-white mica in the samples suggests a regional medium pressure character of the metamorphism.

The peak T condition is higher than the closure temperature of K-white mica so the K–Ar ages are interpreted as cooling ages. Whole rock isochron age with individual age data suggests Variscan peak metamorphism older than c. 310 Ma while data from the clay fraction show post-Variscan cooling ages of c. 290 Ma.

In accordance with the proximal character of the Silurian Horváthertelend foreland basin, we conclude that the studied Horváthertelend Unit, exotic with respect to the surrounding units in Hungary, could not belong to the Moldanubian Zone of the Bohemian Massif. The original position of this unit within the framework of the Variscan mountain chain is maybe to the northeast from the Bohemian Massif (present
coordinates), showing many similarities with the Silurian sequences of the Upper Silesian Block (Moravo-Silesian Zone) and the Małopolska Terrane (Kielec Region).

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