LA-ICP-MS U-Pb apatite dating of Lower Cretaceous rocks from teschenite-picrite association in the Silesian Unit (southern Poland)

KRZYSZTOF SZOPA1, ROMAN WŁODYKA1 and DAVID CHEW2

1Faculty of Earth Science, University of Silesia, Będzińska Str. 60, 41-200 Sosnowiec, Poland; krzysztof.szopa@us.edu.pl; roman.wlodyka@us.edu.pl
2Department of Geology, Trinity College Dublin, Dublin 2, Ireland; chewd@tcd.ie

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Abstract: The main products of volcanic activity in the teschenite-picrite association (TPA) are shallow, sub-volcanic intrusions, which predominate over extrusive volcanic rocks. They comprise a wide range of intrusive rocks which fall into two main groups: alkaline (teschenite, picrite, syenite, lamprophyre) and subalkaline (dolerite). Previous 40Ar/39Ar and 40K/40Ar dating of these rocks in the Polish Outer Western Carpathians, performed on kaersutite, sub-silicic diopside, phlogopite/biotite as well as on whole rock samples has yielded Early Cretaceous ages. Fluorapatite crystals were dated by the U-Pb LA-ICP-MS method to obtain the age of selected magmatic rocks (teschenite, lamprophyre) from the Cieszyn igneous province. Apatite-bearing samples from Boguszowice, Puńców and Lipowa yield U-Pb ages of 103 ± 20 Ma, 119.6 ± 3.2 Ma and 126.5 ± 8.8 Ma, respectively. The weighted average age for all three samples is 117.8 ± 7.3 Ma (MSWD = 2.7). The considerably smaller dispersion in the apatite ages compared to the published amphibole and biotite ages is probably caused by the U-Pb system in apatite being less susceptible to the effects of hydrothermal alteration than the 40Ar/39Ar or 40K/40Ar system in amphibole and/or biotite. Available data suggest that volcanic activity in the Silesian Basin took place from 128 to 103 Ma with the the main magmatic phase constrained to 128–120 Ma.

Key words: geochronology, U-Pb dating, Outer Western Carpathians, Cieszyn magmatic province, apatite.

Introduction

The Early Cretaceous alkaline volcanic region, with the teschenite-picrite association (TPA) of volcanic rocks, is unique to the western part of the Outer Western Carpathians. This magmatic province (~1500 km²) is 15–25 km wide and extends in a NE direction for over 100 km from Hranice in Moravia, Czech Republic, to Cieszyn and Bielsko-Biała in southern Poland (Fig. 1). Teschenite is named after the original German name for the Teschen locality that is now divided into two parts: Těšín (in the Czech Republic) and Cieszyn (in Poland). The term “teschenite” (originally teschinite) was used for the first time by Hohenegger (1861) to describe all granular rocks from the Moravo-Silesian Beskids Mountains. Tschermak (1866) distinguished melanocratic olivine-rich rocks (picrite), confining the term teschenite to olivine-free granular rocks. The studies of Smulikowski in Cieszyn (Smulikowski 1929, 1930) and the region as a whole (Smulikowski 1930) along with the work of Pacáč (1926), Mahmood (1973), and Kudlášková (1987) have provided basic knowledge on the chemistry and petrography of these rocks.

The main products of volcanic activity in the TPA are shallow, sub-volcanic intrusions, which predominate over eruptive volcanic rocks. In the Moravian part of the Cieszyn magmatic province they form submarine lava flows, sills and dykes, while in the Polish sector, they chiefly comprise sill complexes and more rarely dykes with thicknesses varying from a few centimeters to 40 meters (Konior 1963; Lemberger 1971). Minor amounts of submarine volcanism in Cieszyn (Puńców and Zamarski) were described by Gucwa et al. (1971). The teschenite-picrite association contains a wide range of intrusive rocks which belong to two main groups: alkaline (teschenite, picrite, syenite, lamprophyre) and subalkaline (typically dolerite) (Fig. 2). Based on modal relationships between plagioclase and alkaline feldspars three varieties of teschenites can be distinguished: theralitic, essexitic and monzonitic (Smulikowski 1929, 1930). The nepheline syenites occur only as small irregular bodies with sharp boundaries in the upper part of the teschenite sills or as veins cross-cutting the upper or lower chilled margins. They constitute the final product of the extensive fractional crystallization in individual teschenite sills and do not form independent intrusions. Dolerites are not very common rocks within the TPA. They contain high concentrations of SiO₂; some of the dolerite samples are quartz and hyperstene normative while others are nepheline normative using a CIPW normative mineralogy calculation (Fig. 2). Monchiquite, sannaite and camptonite represent the alkaline lamprophyre group in the TPA (Smulikowski 1929, 1930). They are very common throughout the region and form sills up to 4-6 meters thick. According to Wieser (1971), contact metamorphic assemblages indicate that temperatures of 400–500 °C were reached in the aureoles (diopsid-sandine hornfels facies) of the thickest sills.

The volcanic activity was sited in a zone parallel to the axis of the Proto-Silesian Basin (Hovorka & Spišiak 1988). It was confined to isolated tensional fissures (Hovorka & Spišiak 1988) and the upwelling magma was emplaced in an extensional horst-graben system mainly as sills into uncon-
solidated Cretaceous sediments. Sometimes, on reaching the sea-floor, they flowed laterally to form local lava piles. This initial rifting phase never resulted in sea-floor spreading (Nemčok et al. 2001). The major- and trace-element patterns, and also the Nd-Sr isotopic values, indicate that the parental magma of the differentiated rocks series of the TPA was likely to have been the product of partial melting in an enriched, HIMU OIB-like upper mantle (Narębski 1990; Dostal & Owen 1998; Harangi et al. 2003; Włodyka 2010). The generation of magma is inferred to have occurred at depths of 70–80 km (Włodyka 2010). Spišiak et al. (2011) interpreted the Cretaceous alkaline volcanism in the Alpine-Carpathian-Pannonian realm as a result of partial melting of Sub-Continental Lithospheric Mantle (SCLM) on the peripheries of upwellings of asthenospheric mantle confined to slow-spreading ridges of the Alpine Tethys. Most researchers agree on an Early Cretaceous age for the TPA, with the exception of Konior (1977) who proposed two main magmatic phases during the Cretaceous and Miocene in the Silesian Basin. 40Ar/39Ar and 40K/40Ar dating (see Table 1) on amphibole, pyroxene, phlogopite and biotite from a variety of rock types within the TPA (Lucińska-Anczkiewicz et al. 2002; Grabowski et al. 2003; Harangi et al. 2003) have yielded Early Cretaceous ages. Available geochronological and biostratigraphic data indicate that Cretaceous alkaline volcanism in the Western Carpathians (in various tectonic units of the central and external zones) started during the earliest Cretaceous (at ca. 140 Ma) and culminated during the Aptian and early Albian (from 125 to 100 Ma — Spišiak et al. 2011). In Central Europe, Lower Cretaceous alkaline rocks which exhibit close genetic and tectonomagmatic relationships to the TPA occur in the Mecsek-Alföld Igneous Province (southern Hungary, Harangi et al. 2003). K-Ar age
data range between 130 and 110 Ma (Harangi & Árva-Sós 1993). Mesozoic teschenites are rare elsewhere although occurrences have been documented in Georgia (Lebedev et al. 2009), the French Pyrenees (Azambre et al. 1992; Storetvedt et al. 1999) and Russia (Transbaikalia — Metelkin et al. 2004; Stupak et al. 2004) where volcanic activity associated with teschenite emplacement occurred during the Late Cretaceous (from 86 to 110 Ma).

In this paper LA-ICP-MS U-Pb data of apatite separated from teschenite and monchikite rocks from the TPA are presented. Of the five main types of magmatic rocks in the TPA (Fig. 2), three (from Lipowa, Boguszowice and Puńców) yielded sufficient high quality apatite mineral separates suitable for LA-ICP-MS analysis.

### Geological setting and sampling

The Outer Carpathians are sub-divided into the Subsilesian, Silesian, Fore-Magura and Magura Units (Nappes) (Oszczypko 2006) which are the structural remnants of several basins developed on the margin of the European Platform that were incorporated later into the Tertiary Carpathian accretionary wedge (Fig. 1). The Proto-Silesian Basin (Waskowska et al. 2009) developed during Late Jurassic times as a rift and/or back-arc basin. It existed until Late Cretaceous times when it was tectonically compartmentalized into the Silesian and Skole basins. The oldest sediments in the western part of the Proto-Silesian Basin are represented by marly shales of the Vendryně Shale Formation, previously termed the Lower Cieszyn Shales of Oxfordian-Tithonian age. This formation passes upwards into turbiditic limestones and marls of the Cieszyn Limestone Formation (Tithonian—Valanginian). The overlying Valanginian—Aptian Hradiště Formation comprises calcareous shales with intercalations of calcareous sandstones. Within this formation two members are distinguishable: a shaly facies termed the Cisownica Member (formerly termed the Upper Cieszyn Shales) and a sandstone facies termed the Piechowka Member. The Hradiště Formation is overlain by Aptian—Albian black shales of the Veřovice Formation which are in turn overlain by Albian—Early Cenomanian black shales of the Lhôt Formation.

During Late Jurassic to Aptian times the development of the Outer Carpathian basins was controlled by normal faulting and syn-rift subsidence which was associated with alkaline volcanism in the Western Carpathians. This was followed by post-rift thermal subsidence, resulting in the Albian—Cenomanian expansion of deep-water facies (Némčok et al. 2001; Poprawa et al. 2002). The sills of alkaline rocks occur mainly in the lower part of the Hradiště Formation (the Cisownica Member) and sporadically within the underlying Cieszyn Limestone Formation and the Vendryně Shale Formation. Redeposited fragments of these rocks have been recognized in Albian sediments of the Lhôt Formation (Geroch et al. 1972). The TPA rocks encountered at Stara Wies (Nowak 1978), Barachowice (Gucwa & Wieser 1985) and boreholes in the Skoczów area (Konior 1959) represent detached blocks in Miocene deposits and are believed to have been derived from the Cieszyn Beds. Heavy mineral studies from the Hradiště Formation (Cisownica Member) next to the top of a tethalite teschenite sill in Rudów (Szczurowska 1961) showed the presence of detrital diopside and kaersutite grains in shale. Their origin is interpreted as the result of a disaggregation of these early teschenite sills in submarine conditions. It has been assumed that the age of these submarine eruptions was contemporaneous with the deposition of the Upper Cieszyn Shales at Rudowo (Late Valanginian — Grabowski et al. 2003). The above stratigraphic data suggest the duration of the alkaline volcanism ranged from the Valanginian through to the Aptian.

### Analytical techniques

Apatite crystals were separated using standard techniques, including: crushing, hydrofracturing, washing, Wilfley shaking table, Frantz magnetic separator and handpicking. The separation was undertaken at the Institute of Geological Sciences, Polish Academy of Sciences, Cracow.

The morphology and chemical homogeneity of apatite crystals were investigated using a scanning FET Philips 30 electron microscope (15 kV and 1 nA) equipped with an EDS (EDAX) detector at the Faculty of Earth Sciences, University of Silesia, Sosnowiec, Poland.

Apatite analyses (major/minor elements) were carried out from teschenite and monchikite rocks from the TPA. The apatite analyses have been performed on the Inter-Institution Laboratory of Microanalyses of Minerals and Synthetic Substances, Warsaw (CAMECA SX-100 electron microprobe; 1 kV, 2 nA).

### Table 1: Radiometric age data for the Early Cretaceous alkaline rocks of the TPA

<table>
<thead>
<tr>
<th>Locality</th>
<th>Rock type</th>
<th>Analyzed fraction</th>
<th>Age (Ma)</th>
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<tbody>
<tr>
<td>Boguszowice</td>
<td>teschenite</td>
<td>amphibole</td>
<td>122.0±1.5, 122.4±1.1</td>
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<td></td>
<td>monchikite</td>
<td>apatite</td>
<td>103±20</td>
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<tr>
<td>Międzyrzeczec</td>
<td>picrite</td>
<td>flogopite</td>
<td>126.4±1.8, 133.4±1.8</td>
</tr>
<tr>
<td>Lipowa</td>
<td>monchikite</td>
<td>apatite</td>
<td>126.5±2.8</td>
</tr>
<tr>
<td>Puńców</td>
<td>teschenite</td>
<td>amphibole</td>
<td>111.7±1.8, 97.0±1.8, 99.4±1.6, 89.9±3.5, 96.3±3.7</td>
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<tr>
<td></td>
<td>biotite</td>
<td></td>
<td>134.9±2.0, 137.9±2.0</td>
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<tr>
<td></td>
<td>apatite</td>
<td></td>
<td>119.6±3.2</td>
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<tr>
<td>Rudów</td>
<td>teschenite</td>
<td>amphibole</td>
<td>122.2±0.9, 112.5±1.6</td>
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<tr>
<td>Świętoszówka</td>
<td>dolerite</td>
<td>pyroxene</td>
<td>122.7±4.7</td>
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<tr>
<td>Horni Bładowice</td>
<td>basalt</td>
<td>pyroxene</td>
<td>122.4±6.4</td>
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<tr>
<td>Nový Jičín</td>
<td>lamprophyre</td>
<td>amphibole</td>
<td>109.8±4.2</td>
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<tr>
<td>Markov</td>
<td>teschenite</td>
<td>amphibole/biotite</td>
<td>109.8±4.0</td>
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<tr>
<td>Stronik</td>
<td>canptonite</td>
<td>amphibole</td>
<td>113.6±4.4</td>
</tr>
<tr>
<td>Zilina</td>
<td>canptonite</td>
<td>amphibole</td>
<td>128.3±5.6</td>
</tr>
<tr>
<td>Zivotice</td>
<td>lamprophyre</td>
<td>biotite</td>
<td>106.1±4.4</td>
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</table>

* — 40Ar/39Ar, Lucińska-Anczkiewicz et al. (2002); ** — 40K/40Ar, Grabowski et al. (2003); *** — 40K/39Ar, Harangi et al. (2003); **** — Apatite U-Pb data from this study.
according to the ideal chemical formula of apatite: \( A_2(BO_3)_6(X_2) \) where the A site is occupied by Ca, Fe, Mn, Mg, Th, REE, Y and Na, the B site by P (substituted by S, Si) and the X site by F, Cl and OH\(^-\) ions. The hydroxyl content was calculated by normalization assuming ideal stoichiometry (i.e. no vacancies in the X site so that F\(^+\)+Cl\(^-\)+OH\(^-\) = 2).

Apatite U-Pb data were acquired using a Photon Machines Analyte Exite 193 nm ArF Excimer laser-ablation system coupled to a Thermo Scientific iCAP Qc at the Department of Geology Trinity College Dublin. Twenty-eight isotopes (\(^{31}\)P, \(^{35}\)Cl, \(^{42}\)Ca, \(^{54}\)Mn, \(^{86}\)Sr, \(^{89}\)Y, \(^{130}\)La, \(^{140}\)Ce, \(^{141}\)Pr, \(^{146}\)Nd, \(^{147}\)Sm, \(^{152}\)Eu, \(^{154}\)Gd, \(^{159}\)Tb, \(^{163}\)Dy, \(^{166}\)Ho, \(^{169}\)Er, \(^{172}\)Yb, \(^{175}\)Lu, \(^{206}\)Pb, \(^{206}\)Pb, \(^{207}\)Pb, \(^{208}\)Pb, \(^{232}\)Th, \(^{238}\)U and mass 248(\(^{232}\)Th\(^{16}\)O) were acquired using a 50 µm laser spot, a 4 Hz laser repetition rate and a fluence of 3.31 J/cm\(^2\). A ca. 1 cm sized crystal of Madagascar apatite which has yielded a weighted average ID-TIMS concordia age of 473.5±0.7 Ma (Thomson et al. 2012; Cochrane et al. 2014) was used as the primary apatite reference material in this study. McClure Mountain syenite apatite (the rock from which the \(^{40}\)Ar/\(^{39}\)Ar hornblende standard MMhb is derived) was used as a secondary standard. McClure Mountain syenite has moderate but reasonably consistent U and Th contents (~23 ppm and 71 ppm — Chew & Donelick 2012) and its thermal crystallization age (weighted mean \(^{207}\)Pb/\(^{235}\)U age of 523.51±2.09 Ma) and initial Pb isotopic composition \(^{206}\)Pb/\(^{204}\)Pb = 15.47±0.04 are known from high-precision TIMS analyses (Schöene & Bowring 2006). Durango apatite was analysed in this study as a secondary standard. Durango apatite is a distinctive yellow-green fluorapatite widely used as a mineral standard in apatite fission-track and (U-Th)/He dating and apatite electron micro-probe analyses. It is found as large crystals within an open pit iron mine at Cerro de Mercado, Durango, Mexico. The apatite formed between the eruptions of two major ignimbrites which have yielded a sandine-anorthoclase \(^{40}\)Ar/\(^{39}\)Ar age of 31.44±0.18 Ma (McDowell et al. 2005). NIST 612 standard glass was used as the apatite trace element concentration reference material.

The raw isotope data were reduced using the “VizualAge” data reduction scheme of Petrus & Kamber (2012) within the freeware IOLITE package of Paton et al. (2011). User-defined time intervals are established for the baseline correction procedure to calculate session-wide baseline-corrected values for each isotope. The time-resolved fractionation response of individual standard analyses is then characterized using a user-specified down-hole correction model (such as an exponential curve, a linear fit or a smoothed cubic spline). The data reduction scheme then fits this appropriate session-wide “model” U-Th-Pb fractionation curve to the time-resolved standard data and the unknowns. Sample-standard bracketing is applied after the correction of down-hole fractionation to account for long-term drift in isotopic or elemental ratios by normalizing all ratios to those of the U-Pb reference standards. Common Pb in the apatite standards was corrected using the \(^{207}\)Pb-based correction method using a modified version of the VizualAge DRS that accounts for the presence of variable common Pb in the primary standard materials (Chew et al. 2014). Over the course of two months of analyses, McClure Mountain apatite \(^{207}\)Pb/\(^{235}\)U TIMS age of 523.51±1.47 Ma — Schöene & Bowring 2006) yielded a U-Pb Tera-Wasserburg concordia lower intercept age of 524.5±3.7 Ma with an MSWD=0.72. The lower intercept was anchored using a \(^{207}\)Pb/\(^{206}\)Pb value of 0.88198 derived from an apatite ID-TIMS total U-Pb isochron (Schöene & Bowring 2006).

### Results

**Petrography and apatite chemistry**

In the Polish part of the Outer Western Carpathians outcrops of TPA alkaline and sub-alkaline rocks are scarce. The best outcrops are found in several abandoned quarries, but even at these localities the state of preservation is often very poor due to common alteration by hydrothermal fluids and/or chemical weathering. Apatite crystals were separated from rocks from three localities (Fig. 1); two within the lower part of the Hradiště Formation (Puńców, Boguszowice) and one within the Cieszyn Limestone Formation (Lipowa).

Samples collected in Puńców come from an abandoned quarry, ca. 1.5 km north of the church in Puńców village (No. 1, N 49°43′ 49.6774″ and E 18°40′ 1.9152″) and represent the central parts of a theralite-teschenite sill. The samples dated by Lucinska-Anzchkiewicz et al. (2002), Grabowski et al. (2003) and Harangi et al. (2003) come from the same location. They were probably sampled from the bottom portion of the same teschenite sill which is also visible in a small outcrop by the right inflow of the Puńcówka creek. The typical medium-grained theralite-teschenite from Puńców (Figs. 3A, 4) is formed of elongated plagioclase laths sub-ophically intergrown with purplish-red, sector-zoned sub-silicic diopside of “fassaitic” composition. In these rocks intergrowths of clinopyroxene and amphibole (kaersutite) occur, indicating simultaneous crystallization. Sometimes the intergrowth comes along a surface corresponding to a crystal face of pyroxene. During the late-stages of melt crystallization kaersutite superseded growth of pyroxenes. We can then observe sub-silicic diopsides variably replaced by late magmatic brown kaersutite (Fig. 4). It can be attributed to a reaction between pyroxene and melt at temperatures below 1050 °C (Yagi et al. 1975). The rock also contains titanomagnetite as the dominant spinel phase which is oxidized to titanomagnhemite (Haraničyzk et al. 1971) during extensive sub-solidus, low-temperature alteration. The feldspars are the most altered mineral phases and are replaced mainly by a mixture of zeolites (alun�me, natrolite, thomsonite, mesolite) and chlorite. Individual crystals of diopside (Fig. 3B) often have rims of mica (anne-siderophyllite), Apatite crystals sometimes form inclusions in kaersutite and diopside but occur mainly in secondary mesositas (i.e. late stage interstitial material) after decomposed plagioclase. The apatite crystals are thin (up to 0.3 mm) and forms accicular crystals up to 2 mm.

The second sample (No. 2) comes from a small closed quarry found on the bank of the Kalembianka stream in Boguszowice Valley near Cieszyn (N 49°46′ 16.39″ and E 18°37′ 25.0095″). At this locality the central and top por-

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tions of a thin (~6 m) monchiquite sill are visible. The fine-grained central portion of the sill contains phenocrysts of sector-zoned, sub-silicic diopside and biotite (Fig. 3C). Green aegirine, locally overgrowing Al-Ti diopside is present in accessory amounts. Opaque minerals (titanomagnetite) are equally distributed throughout the rock matrix. Microcrysts of diopside, biotite, kaersutite are common in a cryptocrystalline groundmass in addition to lesser amounts of titaniferous magnetite and alkaline feldspars which are replaced by variable amounts of analcime, natrolite, calcite.

Fig. 3. Photographs of the investigated samples. Photographs A, C and E show polished rock slabs while photomicrographs B, D and F are thin sections of samples from Boguszowice, Lipowa and Puńców, respectively. Ap — apatite, Bt — biotite, Cal — calcite, Chl — chlorite, Di — diopside, MtTi — Ti-magnetite, Mt — magnetite, Kae — kaersutite, Eg — aegirine, At — annite, Sdf — siderophyllite (abbreviation according to Whitney & Evans (2010)).
and chlorite. Biotite, pyroxene and the groundmass encloses prismatic needles (up to 2.5 mm long) of apatite.

The third (No. 3) sample was collected from an abandoned quarry near Nadkościół village (N 49° 40‘ 51.6271“ and E 19° 5‘ 18.834”), to the west of Lipowa in the district of Żywiec. The apatite crystals, which take the form of long prisms up to 2–3 mm, were separated from an altered monchiquite sample (Fig. 3E,F). They occur in a groundmass of primary alkali feldspars and pyroxenes altered to an analcime-calcite-chlorite-quartz mixture. A characteristic feature of these rocks is the presence of a significant amount of secondary pyroxene of aegirine and aegirine-augite compositions. They form irregular rims on diopside crystals. Their secondary origin was confirmed by a fluid inclusion study by Dolníček et al. (2010).

All of the dated apatite crystals from the different magmatic rocks of the Cieszyn igneous province can be classified as fluorapatite with 1.6–3.7 wt. % F [ca. 1.4 atoms per formula unit] (Table 2; Fig. 5). The crystals range from 0.1 to 0.8 mm long and are 0.1 to 0.2 mm wide (Fig. 6); in general stubby apatite dominates over acicular varieties in the studied populations. Apatite crystals in all the samples show high but variable Sr contents 2580–3218 ppm (mean value 2980 ppm), 145–184 ppm Y (mean value 171 ppm) and low Mn contents (230–540 ppm). Their chondrite (Cl)-normalized REE patterns are dominated by very strong LREE fractionation (La/LaN = 7.8–73.6 — Puńców, 68–77.1 — Lipowa and 67.1–72.5 — Boguszowice), minor positive Eu anomalies (1.04–1.15) for apatite from Puńców and slightly negative Eu anomalies for Lipowa (0.93–0.95) and Boguszowice (0.91–0.93) (Fig. 6; Table 3).
Table 2: Representative electron microprobe analyses of apatite (EMPA) and number of ion p.f.u. calculated on the basis of 24 O

<table>
<thead>
<tr>
<th></th>
<th>Lipowa</th>
<th>Boguszowie</th>
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None of the investigated apatite crystals show any internal zonation (Fig. 7) in either optical or BSE images.

**Apatite dating**

Tera-Wasserburg concordia plots of the LA-ICPMS U-Pb apatite data from three magmatic samples from the TPA (a monchique from both Boguszowie and Lipowa and a theralite-teschenite sill from Puńców) are plotted in fig. 8 and the analytical data are listed in Table 4. All the analyses from the three samples are characterized by relatively high amounts of common Pb (206Pb/204Pb values typically between 0.4 and 0.8). This is a function of the relatively young age of the samples and the relatively low uranium contents (3.54–6.44 ppm — Boguszowie; 2.47–5.43 ppm — Lipowa and 2.99–5.89 ppm — Puńców).

The Tera-Wasserburg concordias were anchored using initial Pb isotopic ratios calculated using the Stacey & Kramers (1975) terrestrial Pb evolution model. The apatite U-Pb age was used as a starting estimate for the Stacey &
Kramers (1975) model and the initial Pb isotopic composition was calculated using an iterative approach (cf. Chew et al. 2011). The anchored concordia lower intercept ages are 103 ± 20 Ma (MSDW = 3.5) for Boguszowice (Fig. 8A), 126.5 ± 8.8 Ma (MSDW = 1.4) for Lipowa (Fig. 8B), and 119.6 ± 3.2 Ma (MSDW = 1.4) for Puńców (Fig. 8C). Taking into account the age uncertainties, the U-Pb data cluster around 120 Ma yields a weighted average age of 117.8 ± 7.3 Ma (MSWD = 2.7). This age is consistent with the 120 Ma 40Ar/39Ar and 40K/40Ar pyroxene and amphibole

Fig. 7. BSE images of the dated apatites. A–C — Outer crystal surfaces, D–E — an examples of polished grain interiors which were used for EMP and LA-ICP-MS analysis.

Fig. 8. Tera-Wasserburg concordia plots for LA-ICP-MS U-Pb apatite analyses from the Cieszyn igneous province (A — Boguszowice, B — Lipowa, C — Puńców). Data-point error ellipses are 2 σ.
All of the calculated ratios have been done after normalization to chondrite-C1 (Sun & McDonough 1989). LOD — limit of detection for concentration of analysed elements.
Table 4: Representative LA-ICP-MS U-Pb apatite data for rock samples from Boguszowice, Puńców and Lipowa.

<table>
<thead>
<tr>
<th>Locality</th>
<th>206Pb (ppm)</th>
<th>207Pb (ppm)</th>
<th>208Pb (ppm)</th>
<th>206Pb/207Pb</th>
<th>235U/206Pb</th>
<th>238U/206Pb</th>
<th>232Th/208Pb</th>
<th>208Pb/238U</th>
<th>207Pb/235U</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lipowa</td>
<td>1.27 ± 0.07</td>
<td>1.19 ± 0.07</td>
<td>1.23 ± 0.07</td>
<td>1.07 ± 0.07</td>
<td>0.70 ± 0.07</td>
<td>0.73 ± 0.07</td>
<td>0.72 ± 0.07</td>
<td>1.40 ± 0.07</td>
<td>2.30 ± 0.07</td>
</tr>
<tr>
<td>Boguszowice</td>
<td>1.31 ± 0.08</td>
<td>1.19 ± 0.08</td>
<td>1.23 ± 0.08</td>
<td>1.07 ± 0.08</td>
<td>0.70 ± 0.08</td>
<td>0.73 ± 0.08</td>
<td>0.72 ± 0.08</td>
<td>1.40 ± 0.08</td>
<td>2.30 ± 0.08</td>
</tr>
<tr>
<td>Puńców</td>
<td>1.30 ± 0.09</td>
<td>1.19 ± 0.09</td>
<td>1.23 ± 0.09</td>
<td>1.07 ± 0.09</td>
<td>0.70 ± 0.09</td>
<td>0.73 ± 0.09</td>
<td>0.72 ± 0.09</td>
<td>1.40 ± 0.09</td>
<td>2.30 ± 0.09</td>
</tr>
</tbody>
</table>

Discussion

Closure temperature of apatite

Experimental determination of the diffusion parameters of Pb in apatite by high-temperature annealing experiments (Watson et al. 1985) and ion implantation with Rutherford backscattering techniques (Cherniak et al. 1991) imply closure temperatures between 450 °C and 550 °C for typical apatite grain dimensions and crustal cooling rates. These estimates are consistent with field-based studies (Chamberlain & Bowring 2000; Schoene & Bowring 2006). They are also compatible with those predicted on the basis of crystal chemistry and ionic porosity (Dahl 1997) and the empirical estimates based on U-Pb dating of large apatites from pegmatites (Krogstad & Walker 1994). Apatite is chemically stable in middle-amphibolite facies conditions (i.e. temperatures above its closure temperature). Under such conditions it is believed the U-Pb systematics of apatite are controlled predominantly by volume diffusion rather than by new growth or re-crystallization (Chamberlain & Bowring 2000).

The ages of the TPA magmatic rocks range from 90 to 138 Ma (Table 4 — Lucińska-Anzekiewicz et al. 2002, 120—122 Ma; Harangi et al. 2003, 96—128 Ma; Grabowski et al. 2003, 90—138 Ma) which is longer than the time span of magmatic activity in the Mecsek-Alföld Igneous Province in
Hungary and comparable to the range of Cretaceous alkaline volcanism documented from various tectonic units in the Western Carpathians (ca. 140 to 125–100 Ma — Spišiak et al. 2011). The very small age range between ca. 122 Ma (teschenite) and 120 Ma (nepheline syenite) emplacement documented by Łucińska-Anczkiewicz et al. (2002) is interpreted as recording the time of the solidification of teschenite sills by fractional crystallization (FC) process which is a result of crystallization of the later interstitial melts as a small irregular bodies or cross-cutting veins of nepheline syenites in the upper and lower part of the sill.

Much of the teschenite geochronological data is often somewhat contradictory from data as the same silt at an individual locality often show significant age dispersion. This is likely due to low-temperature alteration (chloritization) of the dated mafic mineral phases, and the late magmatic crystallization of amphibole. This situation is documented in Puńców where the same theralite-teschenite sill was dated by four different authors (see Table 1). The younger ages (<100 Ma) were obtained by the K-Ar amphibole dating (Grabowski et al. 2003; Harangi et al. 2003) and Ar-Ar amphibole dating (Łucińska-Anczkiewicz et al. 2002). Petrographic studies reveal that this sill shows evidence for intensive late magmatic crystallization of amphibole (Fig. 4) which followed crystallization of pyroxene with amphibole commonly mantling pyroxene. In contrast, the oldest ages (>130 Ma) were obtained by K-Ar dating of biotite (Puńców) and phlogopite (Międzyrzecze) (Grabowski et al. 2003).

The results obtained in this study yield similar ages (ca. 120 Ma) to those from the studies of Łucińska-Anczkiewicz et al. (2002) and Harangi et al. (2003). The apatite population is magmatic and EMP analyses and BSE observation imply no significant alternation which is common in magmatic rocks from the Cieszyn magmatic province. An analysis of the available robust geochronological data, including the new U-Pb apatite data from this study, suggest that volcanic activity in the Proto-Silesian Basin took place from 128 to 103 Ma and most likely peaked between 128 and 120 Ma. The U-Pb apatite dating technique has clear potential to constrain the emplacement and evolution of the Cieszyn magmatic province, as apatite is the main U-bearing phase suitable for U-Pb geochronology and the U-Pb apatite system appears unaffected by secondary alteration that has historically plagued K-Ar and $^{40}$Ar-$^{39}$Ar dating studies on these rocks.

**Conclusions**

1. The weighted mean LA-ICP-MS U-Pb age for all three samples is 117.8±7.3 Ma and is similar to previously published K-Ar and Ar-Ar ages.
2. The considerably smaller dispersion in the U-Pb apatite age data compared to K-Ar (whole rock) and $^{40}$Ar-$^{39}$Ar amphibole and biotite ages is likely because the U-Pb apatite system is unaffected by alteration compared to the K-Ar and $^{40}$Ar-$^{39}$Ar systems.
3. The probable time of volcanic activity in the Silesian Basin took place from 128 to 103 Ma and most likely peaked between 128 and 120 Ma.
4. A lack of primary magmatic zircon, monazite or xenotime makes apatite the most suitable phase for U-Pb dating of the igneous rocks from the Cieszyn magmatic province.

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