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Apatite U-Pb dating at UNAM laboratories: analytical protocols and examples of its application

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ABSTRACT:

Apatite is the most common phosphate mineral in the Earth's crust and can be found in practically all magmatic and metamorphic rocks, as well as in ancient and recent sediments and in certain ore deposits. Its effective closure temperature of 450–550 °C for the U-Pb system makes apatite an important natural medium-temperature thermochronometer that can be dated by both laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) and isotope dilution thermal ionization mass spectrometry (ID-TIMS) techniques. Due to its low U content, coupled with high Pb contents incorporated during crystallization (also called common lead), apatite U-Pb dating is analytically challenging, and requires robust analytical protocols to achieve reliable ages. In this experimental study we obtained apatite U-Pb ages from six rock samples employing LA-ICP-MS (at Laboratorio de Estudios Isotópicos, Centro de Geociencias, UNAM), while one sample was also dated by ID-TIMS (at Laboratorio Universitario de Geoquímica Isotópica, UNAM). These samples have igneous emplacement or metamorphic ages broadly ranging from the Neoproterozoic to the Paleocene. The obtained apatite U-Pb ages agree well with other radioisotopic data (U-Pb on zircon and K-Ar or Ar-Ar on micas and hornblende) available for the same rock samples tested, or for the same geological units studied. These apatite U-Pb results, obtained for the first time at two main Mexican geochronology laboratories, are precise enough to be geologically meaningful and usually represent the cooling ages; though, in some cases they may also indicate the crystallization or the metamorphic age. Some advantages and disadvantages of LA-ICP-MS and ID-TIMS-based methodologies were observed and outlined. Our results validate the analytical procedures used and will serve as an important trigger towards the development or improvement of medium-temperature thermochronology techniques in Mexico.

KEYWORDS: apatite, laser ablation, isotope dilution, U-Pb dating, thermochronology, Mexico.

RESUMEN:

El apatito es el mineral de fosfato más común en la corteza de la Tierra y puede encontrarse en casi todas las rocas magmáticas y metamórficas, así como en algunos sedimentos antiguos y recientes y en ciertos depósitos minerales. La temperatura de cierre efectiva de 450–550 °C para el sistema U-Pb hace que el apatito sea un importante termocronómetro natural de temperatura media, el cual se puede fechar tanto por la técnica de ablación láser asociada a un espectrómetro de masas con fuente de plasma de acoplamiento inductivo (LA-ICPMS, por sus siglas en inglés) como por el método de dilución isotópica usando espectrometría de masas con ionización térmica (ID-TIMS, por sus siglas en inglés). Debido a su bajo contenido de U, aunado a un alto contenido de plomo incorporado durante la cristalización (denominado también plomo común), la datación U-Pb en apatitos es un proceso retador que requiere protocolos analíticos robustos para obtener edades confiables. En este estudio experimental hemos obtenido edades U-Pb en apatitos de seis muestras analizadas por LA-ICPMS en el Laboratorio de Estudios Isotópicos, Centro de Geociencias, UNAM, así como en una muestra mediante ID-TIMS en el Laboratorio Universitario de Geoquímica Isotópica, UNAM. Estas muestras tienen edades ígneas o de metamorfismo que varían en un rango amplio, desde el Neoproterozoico hasta el Paleoceno. Las edades U-Pb determinadas en

apatitos concuerdan bien con otros datos radioisotópicos (e.g., U-Pb en circones y K-Ar y Ar-Ar en micas y hornblenda) disponibles para las mismas muestras de roca analizadas, o para las mismas unidades geológicas estudiadas. Las edades U-Pb de apatitos, obtenidas por primera vez en dos principales laboratorios mexicanos de geocronología, son suficientemente precisas para tener algún sentido geológico e indican usualmente las edades de enfriamiento (tres muestras), aunque en algunos casos pueden señalar también las edades de cristalización o de metamorfismo. Algunas ventajas y desventajas entre las metodologías basadas en LA-ICP-MS e ID-TIMS fueron también observadas y señaladas. Los resultados de este estudio validan los procedimientos analíticos utilizados y servirán como un inicio importante hacia el desarrollo o mejoramiento de las técnicas de termocronología de media temperatura en México.

PALABRAS CLAVE: apatito, ablación láser, dilución isotópica, datación U-Pb, termocronología, México.

INTRODUCTION

Apatite, the most common phosphate mineral with typical natural occurrence as $\text{Ca}_5(\text{PO}_4)_3[\text{F}, \text{Cl}, \text{OH}]$, has attracted great interest as a low and medium-temperature thermochronometer (fission-track analysis, $\text{U}(\text{Th})/\text{He}$ and U-Pb dating; e.g., Wolf *et al.*, 1996; Ehlers and Farley, 2003; Hasebe *et al.*, 2004; Donelick *et al.*, 2005; Solé and Pi, 2005; Chew and Donelick, 2012; Abdullin *et al.*, 2014; Chew *et al.*, 2014; Enkelmann *et al.*, 2014; Spikings *et al.*, 2015); as a chemical indicator of petrogenetic or metallogenetic processes, as well as a novel geochemical tool in sedimentary provenance studies (Belousova *et al.*, 2002; Morton and Yaxley, 2007; Jafarzadeh *et al.*, 2014; Abdullin *et al.*, 2016a and 2016b; Bruand *et al.*, 2016); or even as an isotopic tracer (e.g., Sr isotopes, assuming that apatite does not incorporate any significant amount of initial Rb; Bizzarro *et al.*, 2003); or as a proxy of the redox state of silicic host magmas as well (Miles *et al.*, 2014).

On the other hand, zircon (ZrSiO_4) is still the most commonly used U-rich accessory mineral in U-Pb dating. Its high closure temperature for this isotopic system, greater than 800 °C, as well as its exceptional chemical and mechanical robustness (e.g., Lee *et al.*, 1997; Cherniak and Watson, 2001) allow robust igneous crystallization ages and ages of medium to high-grade metamorphism to be obtained, as well as to extract provenance signals from siliciclastic samples (e.g., Krogh, 1982; Košler *et al.*, 2002; Jackson *et al.*, 2004; Weber *et al.*, 2005, 2007, 2008; Solari *et al.*, 2014, 2015).

While zircon is relatively straightforward to date by the U-Pb method (e.g., see Faure and Mensing, 2005; Schoene, 2014; Solari *et al.*, 2015), apatite U-Pb dating is more challenging. The main issues of dating accessory minerals by LA-ICP-MS that could have incorporated Pb during crystallization are listed and described by Chew *et al.* (2014). In the case of apatite, the issues lie in the low U concentration that generally vary from 1 to 50 ppm; though, in some cases these values may be increased up to 100–300 ppm (for instance, in apatite of sample Congra-2; see details below), while apatite may incorporate high levels of lead during the magma crystallization (hereafter referred to as common Pb) (Chamberlain and Bowring, 2001; Schoene and Bowring, 2007; Chew *et al.*, 2011, 2014; Thomson *et al.*, 2012).

Another issue for apatite U-Pb dating, in particular by microanalytical instruments like LA-ICP-MS, is the lack of isotopically homogeneous and large natural apatite crystals to be used as a primary reference material. There are numerous standard zircons (Horstwood *et al.*, 2016, and references therein); meanwhile, the few apatite U-Pb standards can contain variable amounts of common Pb, which makes the U-Pb isotope ratio calculation more complex than for zircons (Chew *et al.*, 2011; Thomson *et al.*, 2012; Chew *et al.*, 2014). However, this can be corrected by applying a ^{207}Pb -based correction depending on the initial $^{207}\text{Pb}/^{206}\text{Pb}$ value of the standard apatite (Chew *et al.*, 2014; see section "LA-ICP-MS technique (LEI)" below). Ionic diffusion of radioactive and/or daughter isotopes of any geochronological system in crystal structures are assumed as dependent of ambient temperature and cooling rate, and it defines the closure temperature (e.g., Dodson, 1973; Copeland *et al.*, 1988; Villa, 1988). Apatite U-Pb is a useful thermochronometer due to the following reasons: (1) Apatite is a common accessory mineral in virtually all igneous lithologies, and it is also a common component of many metamorphic and siliciclastic and some non-clastic sedimentary rocks, as well as of certain ore deposits (e.g., Green and Watson, 1982; Piccoli and Candela, 2002; Morton and

Yaxley, 2007; Webster and Piccoli, 2015); (2) its effective closure temperature of *ca.* 450 to 550 °C for U-Pb, together with its partial retention zone of radiogenic lead (PbPRZ) of ~370–580 °C (Cherniak *et al.*, 1991; Chamberlain and Bowring, 2001; Schoene and Bowring, 2007; Cochrane *et al.*, 2014), makes apatite U-Pb dating a new, alternative technique to obtain crystallization ages (*i.e.*, of rapidly cooled shallow intrusions, volcanic and subvolcanic rocks), ages of metamorphism (Henrichs *et al.*, 2018) and of hydrothermal or mineralization processes, as well as to construct thermal histories of crystalline basement, among others; and, (3) apatite is seemingly stable during sediment transport and is extremely stable during diagenesis preserving its original composition (Morton and Hallsworth, 2007; Morton and Yaxley, 2007), which may be particularly useful for sedimentary provenance studies and to obtain approximate stratigraphic ages based on detrital apatite U-Pb results (Chew and Donelick, 2012; O'Sullivan *et al.*, 2016; Bedoya *et al.*, 2017).

In this experimental study, we report the first apatite U-Pb results obtained in Mexican geochronology laboratories, testing this technique on different igneous and metamorphic rocks, which have already been dated by conventional radioisotopic methods like U-Pb on zircons, and K-Ar or Ar-Ar dating. The chosen samples have crystallization or metamorphic ages ranging from the Mesoproterozoic to the Paleocene (for more details see Table 1). Our U-Pb results on apatite validate the used analytical procedures and will serve as a starting point for the development or improvement of medium-temperature thermochronology techniques in Mexico.

DESCRIPTION OF SAMPLES

Sample Congra-2

Congra-2 is a monzogranite from the southern Sierra Madre Occidental. For its cognate sample, collected from the same plutonic body, an emplacement age of 59.4 ± 0.5 (2 σ) Ma was proposed on the basis of zircon U-Pb results (Sample Conc-01 in Duque-Trujillo *et al.*, 2014). Hornblende and biotite concentrates from Conc-01 yielded Ar-Ar ages at 55.7 ± 0.5 (2 σ) and 56.3 ± 0.3 (2 σ) Ma, respectively, both cooling ages that were also obtained by Duque-Trujillo *et al.* (2014).

Sample OV-0421

This sample is a two mica-bearing deformed granite belonging to the Tres Sabanas pluton, which is located NW of Guatemala City, Guatemala. Torres de León (2016) reported a U-Pb zircon age of 115 ± 4 Ma (2 σ) for OV-0421, interpreted as its crystallization age. A cooling age of 102.2 ± 1.2 (2 σ) Ma, obtained by the K-Ar method, has also been reported by the same author.

Sample MCH-40

MCH-40 is an andesite of rather hypabyssal origin. This rock was sampled to the east of Jacinto Tirado town, Chiapas (Table 1) from an Early to Middle (?) Jurassic volcanic unit named informally as Andesita Pueblo Viejo (*e.g.*, Castro-Mora *et al.*, 1975; Meneses-Rocha, 2001; Godínez-Urban *et al.*, 2011). Due to its age uncertainty, MCH-40 was also dated by zircon U-Pb technique (see details below).

Sample MCH-17

This rock sample is a metamorphosed granite, collected from the Chiapas Massif near the Cintalapa de Figueroa town. According to several authors (*e.g.*, Damon *et al.*, 1981; Torres *et al.*, 1999; Meneses- Rocha, 2001; Schaaf *et al.*, 2002; Weber *et al.*, 2005), the Chiapas Massif is a Permian-Triassic continental arc and represents the basement of the southern Maya block. This unit is mainly composed of diverse plutonic rocks, most of which are deformed while some are metamorphosed. To better constrain its crystallization age, zircons from MCH-17 were also dated in the current work by employing LA-ICP-MS-based U-Pb method. This sample is the only one from which apatite were analyzed by both LA-ICP-MS and ID-TIMS techniques. This was done mainly to evaluate the U-Pb results obtained by two different analytical methods and to see eventual discrepancies (details in “Analytical Methods” section).

TABLE 1
Lithology, locality, and previously published geochronological data of selected samples.

Sample	Locality	Lithology	Published geochronological data			Notes
			Age (Ma)	$\pm 2\sigma$ (abs)	MSWD	
Congra-2 (Conc-01)	Concordia, Sinaloa (23°16'21.0"N; 106°07'01.6"W)	Monzogranite (undeformed)	59.4	0.5	0.79	Average $^{206}\text{Pb}/^{238}\text{U}$ zircon age; n=30 (Duque-Trujillo <i>et al.</i> , 2014. sample Conc-01).
			55.7	0.5		Ar-Ar hornblende age (Duque-Trujillo <i>et al.</i> , 2014).
			56.3	0.3		Ar-Ar biotite age (Duque-Trujillo <i>et al.</i> , 2014).
OV-0421	Central Guatemala (14°46'29.3"N; 90°33'08.0"W)	Deformed granite	115	4		Average $^{206}\text{Pb}/^{238}\text{U}$ zircon age; n=16 (Torres de León, 2016).
			102.2	1.2		K-Ar biotite age (Torres de León, 2016).
MCH-40	Chiapas (16°43'04.0"N; 93°45'33.0"W)	Andesitic dike				Average $^{206}\text{Pb}/^{238}\text{U}$ zircon age; n=27 (this work; see results).
MCH-17	Chiapas (16°24'46.0"N; 93°47'42.0"W)	Metagranite				Average $^{206}\text{Pb}/^{238}\text{U}$ zircon age; n=25 (this work; see results) ID-TIMS leached apatite age (this work; see results).
Gt-0340	Rabinal, Guatemala (15°06'39.0"N; 90°30'02.0"W)	Deformed granite	483	7		ID-TIMS lower intercept zircon age (Ortega-Obregón <i>et al.</i> , 2008).
OC111	Oaxaca (18°05'31.4"N; 97°20'21.0"W)	Orthogneiss	990	10		Average $^{206}\text{Pb}/^{238}\text{U}$ zircon age; n=25 (Solari <i>et al.</i> , 2014).

Sample GT-0340

This sample belongs to the undeformed Rabinal Granite, cropping out north from the Rabinal town in Baja Verapaz, Guatemala. Many zircon aliquots from GT-0340 were previously dated by U-Pb ID-TIMS, yielding an age of 482.6 ± 7 (2 σ) Ma (Concordia upper intercept) that was interpreted as its approximate crystallization age (Ortega-Obregón *et al.*, 2008). This age is similar to others obtained in different samples from the same granite (*e.g.*, Solari *et al.*, 2013).

Sample OC-111

This sample is a granulitic gneiss from the Grenville-aged Oaxacan Complex. The sample is composed by quartz, mesoperthite, rutile, garnet, and by altered mafic minerals. OC-111 was sampled *ca.* 120 km north of Oaxaca City, to the east of the tectonic contact between the Zapoteco and Mixteco tectonostratigraphic terrains (*e.g.*, see ElíasHerrera *et al.*, 2007). Previous U-Pb zircon dating yielded protolith crystallization ages at ~ 1220 Ma, although it was also demonstrated that this sample underwent “dry” granulite facies metamorphism at 990 ± 10 Ma (Solari *et al.*, 2014).

ANALYTICAL METHODS

Apatite U-Pb analyses were performed by a joint effort between the Laboratorio Universitario de Geoquímica Isotópica (LUGIS, Instituto de Geofísica, CDMX, UNAM) and the Laboratorio de Estudios Isotópicos (LEI, Centro de Geociencias, Campus Juriquilla, UNAM). Apatite U-Pb dating at LUGIS was based on ID-TIMS technique, while in LEI all the six selected samples were analyzed with LA-ICP-MS.

Preparation of samples

Sample preparation was performed at the Centro de Geociencias (Taller de Molienda and Taller de Laminación), Campus Juriquilla, UNAM. Heavy minerals were concentrated from two narrow grain-size fractions of 63–125 and 125–180 μm , using conventional methods/ instruments like crushing, sieving, WilfleyTM table, FrantzTM separator, and finally by hand-picking under a binocular microscope. For U-Pb dating by ID-TIMS, only ‘sterile’ apatite grains, without visible inclusions and other defects like cracks, were selected to be split into two aliquots with ~ 300 crystals of different size and morphology (*e.g.*, see Figure 1). Analysis by ID-TIMS was performed only for sample MCH-17. For LA-ICP-MS-based apatite U-Pb dating, ~ 200 apatite crystals extracted from each sample were mounted with EpoFixTM Struers in a 2.5 cm diameter plastic ring, and then polished with Buehler SiC MicroCutTM sandpapers P1500 ($\sim 12.5 \mu\text{m}$) and P2500 ($\sim 8.3 \mu\text{m}$), and alumina suspensions over Buehler MicroClothTM pads (see an example illustrated on Figure 2). LA-ICP-MS analyses were performed for all the six selected samples. In addition, two samples (MCH-17 and MCH-40) were also prepared for zircon U-Pb dating, from which *ca.* 100 crystals from each sample were mounted to be analyzed by LA-ICP-MS.

ID-TIMS methodology (LUGIS)

An additional observation of the selected apatite crystals using a binocular microscope, cleaning, chemical treatments, and isotopic measurements were entirely carried out at LUGIS. One of the two aliquots with ~ 300 apatite crystals was leached in 0.25M HNO₃ at ambient temperature (24 °C) for 1 min to remove any non-radiogenic Pb from their exterior surface, and then rinsed with de-ionized water. The other one, also made up of ~ 300 crystals, was cleaned only in Milli-Q water. The aliquots were dried down, transferred into TeflonTM microcapsules and weighted, then 1 ml of ultrapure 8M HNO₃ was added. The complete digestion of both aliquots (*i.e.*, leached and non-leached) was performed at 110 °C for 20 hours; then, the solutions were dried down. After evaporation, a drop of 6M HCl was added and dried to convert the produced components to chlorides. The dry residues derived from both aliquots were dissolved in 600 μl of 3M HCl, and heated at 110 °C for 7 hours. Each digested aliquot was split into two fractions; one for isotopic composition (IC) analyses, and a second for isotope dilution (ID) measurements to which a mixed ^{208}Pb – ^{238}U spike was added. U and Pb were separated by chromatographic column chemistry technique employing BioRadTM AG-1X8 resins within TeflonTM columns.

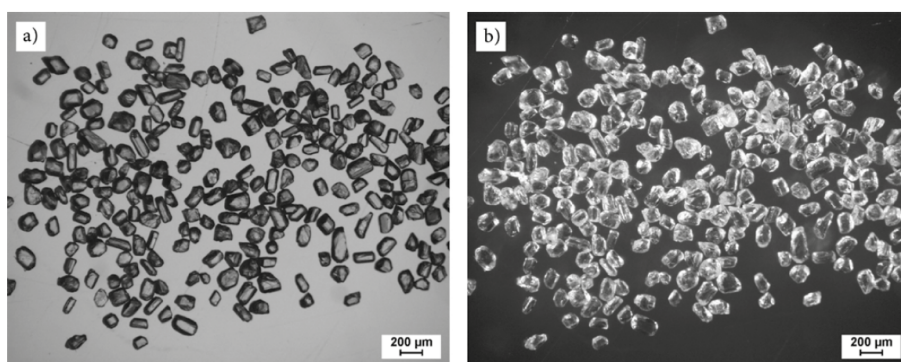


FIGURE 1

Photomicrograph of selected apatite crystals for ID–TIMS analysis from sample MCH-17 using (a) transmitted and (b) reflected light. Note that most of crystals are free of inclusions or fractures.

The prepared samples were loaded on Re filaments with the addition of a silica gel–H₃PO₄ mixture and analyzed in static mode. Pb isotopic measurements were performed in a Thermo™ TRITON Plus mass spectrometer; measuring ²⁰⁴Pb with an ion counter, whereas the remaining Pb isotopes (²⁰⁶Pb, ²⁰⁷Pb, and ²⁰⁸Pb) were measured in Faraday cups. U concentrations were determined with a Finnigan™ MAT-262 mass spectrometer. Finally, 30 and 45 isotope ratios were acquired for U and Pb, respectively. Mass fractionation was taken into account based on PbDat v. 1.24 (Ludwig, 1993) and applying a correction factor of 0.12‰ for each isotopic pair (²⁰⁶Pb/²⁰⁴Pb, ²⁰⁶Pb/²⁰⁷Pb, and ²⁰⁶Pb/²⁰⁸Pb). The values for initial common Pb were estimated at 250 Ma (Permian–Triassic Chiapas Massif; Damon *et al.*, 1981; Schaaf *et al.*, 2002; Weber *et al.*, 2005), following the well-accepted methodology of Stacey and Kramers (1975). During the experiment at LUGIS, total analytical blanks for Pb and U were obtained as 18 and 16 pg, respectively. However, in further experiments these values may change, and thus, will have to be carefully controlled again.

LA–ICP–MS technique (LEI)

Spot analyses were performed with a Resonetics RESolution™ LPX Pro (193 nm, ArF excimer) laser ablation system, coupled to a Thermo Scientific iCAP™ Qc quadrupole ICP–MS at LEI. Different analytical protocols were employed for zircon and apatite. For zircon, each spot analysis consisted in the acquisition of 15 s of background signal (gas blank), 30 s of ablation, and 15 s to allow the signal to reach the baseline again (wash-out). The spot diameter was of 23 µm, using a fluence of 6 J/cm² with a repetition rate of 5 Hz. For apatite, 35 s of ablation was chosen with the same gas blank and wash-out acquisition times as given above. In this case, a spot of 60 µm was employed, basically to obtain a larger volume of the ablated material to be analyzed with the purpose of compensate for weaker signals of U and Pb. Due to different coupling of laser energy in different materials, the laser pulse energy and repetition rate were reduced to 4 J/cm² and 4 Hz, respectively. For both zircon and apatite, 350 ml/min of He was used as a carrier gas, mixed in the cell with *ca.* 800 ml/min of Ar. Four and three ml/min of N₂ for zircon and apatite analyses were also added after the cell, respectively, to increase the sensitivity of the plasma. A “squid” signal homogenizer is located right after the ablation cell, to mix properly all the gases added before reaching the plasma. Single-spot analyses in apatite were performed in inclusion-free grains or in those parts in which no visible inclusion or other defects were present (*e.g.*, see in Figure 2).

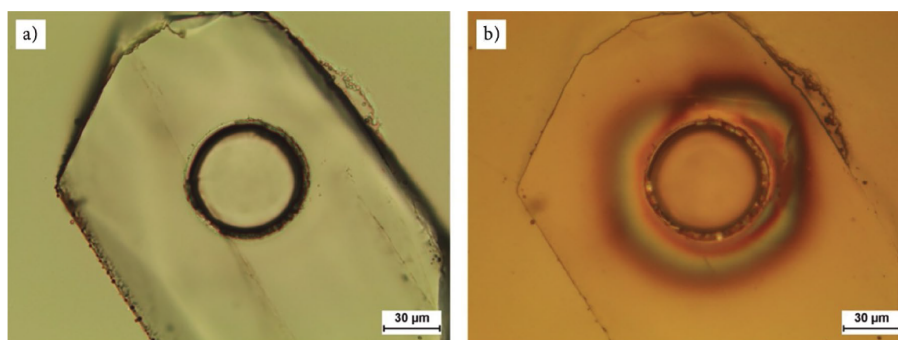


FIGURE 2

Photomicrograph of an apatite crystal after ablation in (a) transmitted and (b) reflected light. Spot diameter (center of the crystal) is 60 μm . The selected crystals for LA-ICP-MS analysis are free of inclusions or cracks. Note the concentric deposits of ablated material around the crater in reflected light (b).

Together with those isotopes needed for the U-Pb age calculation (^{206}Pb , ^{207}Pb , ^{208}Pb , ^{232}Th , and ^{238}U), LA-ICP-MS permits to sequentially detect additional elements such as REE (rare earth elements, La to Lu), Sr, Y, Mn, Mg, Cl, etc. These detailed chemical compositions are not shown because they are beyond the scope of this paper. This protocol for apatite analyses was established on the basis of numerous experiments carried out at the LEI Lab during the past years, and may be used for simultaneous U-Pb and fission-track *in-situ* double dating plus multielemental analysis (*e.g.* Abdullin *et al.*, 2016a and 2016b).

Tuning of the ICP-MS followed the parameters reported in Solari *et al.* (2010) and Ortega-Obregón *et al.* (2014). Corrected isotope ratios, ages and errors were calculated using Iolite software (Paton *et al.*, 2011) and the VizualAge data reduction scheme of Petrus and Kamber (2012). The standard zircon “91500” (Wiedenbeck *et al.*, 1995, 2004) was used as a reference mineral for zircon U-Pb analyses. In the course of this experiment, an age of 337.7 ± 1.5 (2σ) Ma was obtained for the Plešovice zircon used as a secondary standard for zircon U-Pb analyses, yielding a mean square weighted deviation (MSWD) of 1.2 from 20 single-spot U-Pb ages. This value is in line with the standard $^{206}\text{Pb}/^{238}\text{U}$ age of *ca.* 337 Ma (Sláma *et al.*, 2008).

In the case of apatite U-Pb dating, UcomPbine (Chew *et al.*, 2014) was used to model $^{207}\text{Pb}/^{206}\text{Pb}$ initial values and thus force a ^{207}Pb correction that considers the common Pb incorporated by apatite standards at the moment of their crystallization. As is described by Chew *et al.* (2014), this correction is applied immediately after baseline subtraction and previously to the “raw” standard ratios calculation in order to not compromise down-hole fractionation and mass bias corrections. Once the standard is corrected and down-hole fractionation modeled, the drift in the mass discrimination during a session is subsequently accounted by sample-standard bracketing. Tera and Wasserburg (T-W) Concordia diagrams are commonly used in apatite U-Pb dating, because results are mostly discordant. The lower intercept in T-W, is considered as a “mean” apatite U-Pb age that should have geological significance (*e.g.*, crystallization or cooling age, or the age of a metamorphic event).

For apatite U-Pb dating, the Madagascar apatite was used as a main reference material (with a mean U-Pb age of 485–475 Ma; Thomson *et al.*, 2012; Chew *et al.*, 2014), assuming that matching of the matrix is fundamental during the U-Pb dating by LA-ICP-MS (*e.g.*, Gehrels *et al.*, 2008; Solari *et al.*, 2010, 2015).

Due to limitations in the LEI Lab, no representative and wellstudied secondary U-Pb standard was used for apatite U-Pb dating. However, as was mentioned above, the sample MCH-17 was chosen to be dated by both ID-TIMS and LA-ICP-MS techniques. This sample, thus, was also used to control the final U-Pb ages obtained for all the apatite samples selected (see results below). Age and error calculations, including zircon

mean ages and intercept values and uncertainties, were carried out using Isoplot v. 3.7, written by Ludwig (2008).

RESULTS AND INTERPRETATIONS

Zircon U-Pb ages

Zircons from two of the six experimental samples (MCH-17 and MCH-40) were dated by U-Pb employing LA-ICP-MS (LEI; U-Pb results are detailed in Figure 3 and Supplementary Appendix A1). Only results within -5 to 30% of concordance were considered as reliable ages. Weighted average ages were calculated using Isoplot v. 4.15 (Ludwig, 2008) rejecting outliers (less than 95% of confidence).

Sample MCH-17

MCH-17 mainly contains prismatic and bipyramidal zircon crystals, most of which are larger than 100–150 μm length and 50–100 μm width. A detailed observation of their morphology suggests that these crystals are of igneous origin, without evidence from cathodoluminescence of possible metamorphic rims. From a total of 35 zircons dated, 27 passed the concordance filters (those within -5 to 30% of discordance) and were considered meaningful for the interpretation of the igneous crystallization age. The remaining analyses are concordant to slightly discordant (see Supplementary Appendix A1), and define a marked cluster between ~260 and ~250 Ma in the Concordia diagram (Figure 3a). The mean $^{206}\text{Pb}/^{238}\text{U}$ age of 253 ± 2.5 (2σ) Ma, displayed in Figure 3b, can be interpreted as the best approximation for the crystallization age of this granitic sample.

Sample MCH-40

Most zircon grains from the andesite sample MCH-40 are small (50–100 μm long, 30–60 μm wide) with poorly developed prism facets and pyramids. 35 zircon crystals were analyzed, 27 of which are considered reliable for age interpretation. The U-Pb ages are concordant and straddle the Concordia curve at ~195 Ma (lower Sinemurian age; Figure 3d). The mean $^{206}\text{Pb}/^{238}\text{U}$ age of 196.1 ± 1.5 (2σ) Ma is, hence, interpreted as the best crystallization age for MCH-40 (Figure 3d).

Apatite U-Pb ages

Apatite U-Pb results obtained from all the six studied rock samples are described below. Detailed information on our LA-ICP-MS U-Pb experiments is given in Supplementary Appendix A2.

Sample Congra-2

Apatite from the youngest sample Congra-2 show a well-defined linear regression in the T-W diagram, which is derived from all the 34 single-spot analyses performed with LA-ICP-MS. A lower intercept age of 60.9 ± 1.5 (2σ) Ma was detected (MSWD = 0.95; Figure 4a). Despite its young age (Paleocene; Duque-Trujillo et al., 2014), a statistically very reliable apatite U-Pb age was obtained for Congra-2, which is also supported by a MSWD that is close to 1. This is due to high U contents that vary broadly from one apatite to another (31.1 to 371.5 ppm with a mean of 108.2 ppm, standard deviation (SD) = 69.5; Supplementary Appendix A2), which was crucial to yield strongly heterogeneous isotopic ratios for the U-Pb system as well as to generate high levels of radiogenic Pb, and thus displaying a perfect order on the Discordia line (Figure 4a) with a very low age error (only 2.5% at 2σ level).

This pluton yielded a crystallization age of 59.4 ± 0.5 (2σ) Ma as it was well documented by zircon U-Pb geochronology, and then cooled to ~56 Ma as suggested by Ar-Ar ages in hornblende and biotite (Sample Conc-01; Duque-Trujillo et al., 2014). In this case, the apatite lower intercept U-Pb age of 60.9 ± 1.5 (2σ) Ma is within the analytical uncertainty of the zircon age, therefore, this apatite U-Pb date also seems to indicate

the igneous crystallization age for Congra-2 (Conc01), suggesting that the pluton was likely emplaced at a temperature above the Ar-Ar closure temperature of hornblende and biotite, and lower than the closure temperature of the apatite U-Pb system.

Sample OV-0421

For the Tres Sabanas granite OV-0421, the lower intercept in T–W yielded an apatite U-Pb date of 106.7 ± 8.7 (2σ) Ma, obtained from 42 spots (Figure 4b). These apatites have U contents of 84–177 ppm with a mean of 127 ppm and SD of 21.8 (Supplementary Appendix A2). A high MSWD of 9.89 incorporates a large uncertainty, derived from quite scattered apatite U-Pb results as well as from small age errors (Supplementary Appendix A2; Figure 4b).

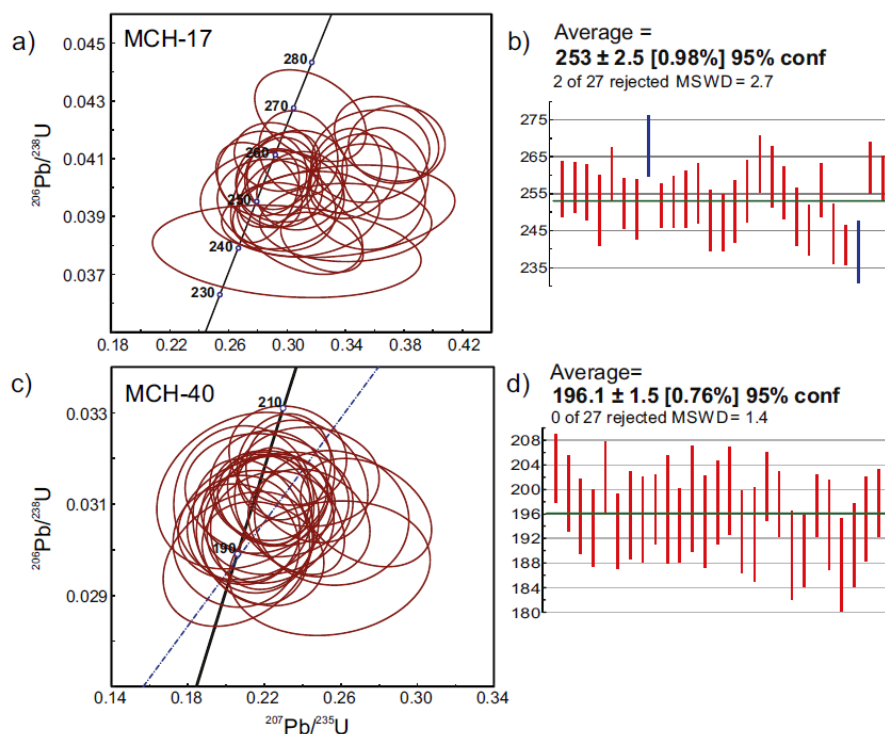


FIGURE 3

U-Pb Concordia diagrams (Wetherill, 1956) and weighted average $^{206}\text{Pb}/^{238}\text{U}$ age plots of zircons from samples MCH-17 (a and b, respectively), and MCH-40 (c and d, respectively). Blue bars correspond to rejected data.

The age determined in this work by apatite U-Pb dating, lies between the zircon U-Pb age of 115 ± 4 (2σ) and the biotite K-Ar age of 102.2 ± 1.2 (2σ) Ma, which were previously obtained for the same granite sample by Torres de León (2016). Central values decrease systematically depending on the closure temperatures for each system; from U-Pb in zircon (900 to 800 °C; Lee *et al.*, 1997; Cherniak and Watson, 2001) through U-Pb in apatite (ca. 450–550 °C; Schoene and Bowring, 2007; Chew *et al.*, 2014; Cochrane *et al.*, 2014) to K-Ar in biotite (300 to 350 °C; e.g., Faure and Mensing, 2005). On the basis of these available data, a cooling rate of ~ 41 °C/Ma can be calculated for the magmatic system from which OV-0421 was formed. This cooling rate could be significant for the whole Tres Sabanas pluton, located in Guatemala.

Sample MCH-40

Apatite crystals extracted from MCH-40 (Sinemurian andesite) fall towards the upper intercept in the T–W Concordia (Figure 4c). A total of 43 dated apatite crystals define a lower intercept of 198 ± 18 (2σ) Ma, yielding an MSWD value of 1.15. The tendency towards the upper intercept in T–W and a large age error of

9.1% at 2SE are apparently linked to very low U concentrations in those apatites, which range between 4.3 and 16.6 ppm with a mean value of 7.2 ppm and SD of 2.4 (see details in Supplementary Appendix A2).

Zircons and apatite from MCH-40 yield similar U-Pb ages of 196.1 ± 1.5 Ma and 198 ± 18 (2σ) Ma, respectively (Figures 3d and 4c). In theory, volcanic or subvolcanic samples like this andesite (or shallow plutons) should yield equivalent crystallization ages independently of the isotopic system applied. Considering that this magma cooled (crystallized) very rapidly, both those U-Pb ages can be interpreted as corresponding to the igneous crystallization age during the Sinemurian Stage. There is a large difference in the age error between the zircon U-Pb and the apatite U-Pb results, which is basically controlled by the strong differences in the abundance of their parent and daughter nuclides (i.e., zircon contains almost no common Pb and is significantly more enriched in U and radiogenic Pb when compared to apatite). It is important to note that Sinemurian igneous ages from Jurassic magmatic rocks and volcanoclastic units (Andesita Pueblo Viejo and La Silla Formation), distributed in the Chiapas Massif area, have also been detected in recent works based on U-Pb, Ar-Ar, and fission-track dating (Godínez-Urban et al., 2011; Abdullin *et al.*, 2018).

Sample MCH-17

Non-leached apatite of sample MCH-17, dated by ID-TIMS, yielded a reversely discordant $^{206}\text{Pb}/^{238}\text{U}$ age of 253.5 ± 2.7 (1σ) Ma (-8.9% of discordance; Table 2; Figure 5). The negative discordance can indicate that these crystals include non-radiogenic Pb in their outermost zones. The leached aliquot, however, yielded a concordant $^{206}\text{Pb}/^{238}\text{U}$ age of 247.1 ± 0.8 Ma with a very small 1σ -error (0.3%; Figure 5; Table 2). The results of this single exercise demonstrate that for some samples, such as MCH-17, a rapid acid leaching may be required to remove possible superficial Pb contamination of non-radiogenic origin (e.g., in apatite from weathered rocks).

Using single-spot analyses performed on 39 apatites by LA-ICP-MS, a lower intercept U-Pb age of 243 ± 12 (2σ ; 4.9%) Ma was obtained for the same sample (MSWD = 1.7; Figure 4d), a value that agrees very well with the ID-TIMS-derived concordant U-Pb age of 247.1 ± 0.8 Ma (1σ). In these apatites U vary from 11.5 to 34.7 ppm, giving a mean concentration of 18.7 ppm with a SD of 6.1 (Supplementary Appendix A2).

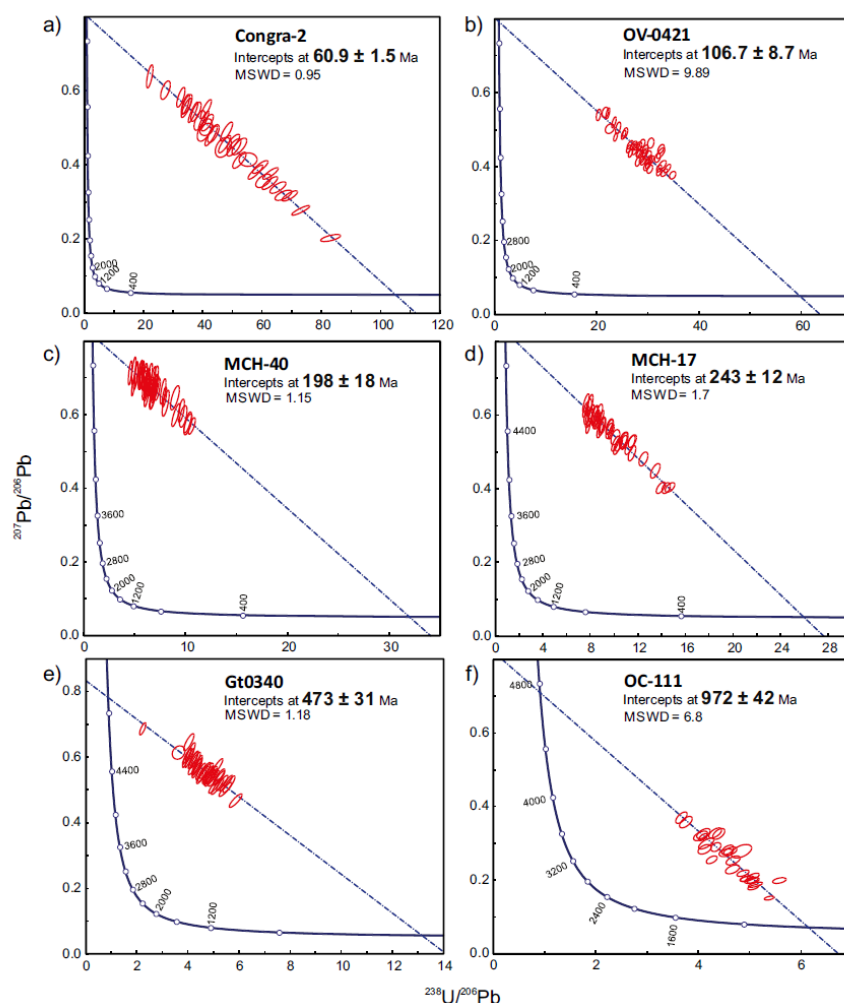


FIGURE 4

Tera-Wasserburg Concordia diagrams for U-Pb data of apatites from samples: (a) Congra-2; (b) OV-0421; (c) MCH-40; (d) MCH-17; (e) GT-0340; and, (f) OC-111.

Both apatite U-Pb dates of 247.1 ± 0.8 (1σ) Ma and 243 ± 12 (2σ) Ma obtained for sample MCH-17 by two different analytical techniques are younger than (or, within the analytical uncertainty of) the zircon crystallization age of 253 ± 2.5 (2σ) Ma (Figures 3b, 4d, and 5). In general, our U-Pb results are in close agreement with geochronological data reported for the Chiapas Massif in previous studies (Damon et al., 1981; Schaaf *et al.*, 2002; Weber *et al.*, 2005). For example, K-Ar biotite ages compiled by Torres et al. (1999) mostly lie within the Early Triassic. Triassic cooling ages from the Chiapas Massif were also detected by Rb-Sr in mica-whole rock pairs that range from 244 ± 12 to 214 ± 11 (2σ) Ma (Schaaf *et al.*, 2002). Therefore, our apatite U-Pb ages suggest that the main post-emplacement thermal cooling of the Chiapas Massif started during the earliest Triassic.

Sample GT-0340

A total of 37 apatite crystals analyzed from the Rabinal Granite GT0340 yielded a lower intercept age of 473 ± 31 (2σ ; $\sim 6.6\%$) Ma (Figure 4e), and these grains display U contents between 3.1 and 124.3 ppm with a mean of 19.6 ppm and SD of 19 (see Supplementary Appendix A2). The MSWD of 1.18 suggests that all dated apatites pertain to a single age population. The apatite U-Pb age of 473 ± 31 (2σ) Ma is identical within error to the Rabinal Granite emplacement age of 483 ± 7 (2σ) Ma obtained for the from same rock sample

(ID–TIMS-derived zircon U–Pb results, reported by Ortega-Obregón *et al.*, 2008), indicating a shallow crystallization and fast cooling.

TABLE 2
U–Pb isotopic results by ID–TIMS of leached and unleached apatite from sample MCH-17.

Sample	Corrected Isotope ratios ^(a)																	Corrected ages ^(a)					
	Weight	U	[U]	Pb _{tot}	Pb _{com}	²⁰⁶ Pb*/ ²⁰⁴ Pb	±1σ	²⁰⁶ Pb*/ ²³⁸ U	±1σ	²⁰⁷ Pb*/ ²³⁵ U	±1σ	²⁰⁷ Pb*/ ²⁰⁶ Pb*	±1σ	Rho	²⁰⁶ Pb*/ ²³⁸ U	±1σ	²⁰⁷ Pb*/ ²³⁵ U	±1σ	²⁰⁷ Pb*/ ²⁰⁶ Pb*	±1σ	Disc		
	(mg)	(ppm)	%err	(ppm)	(ppm)	(corr.)	(%)	(rad.)	(%)	(rad.)	(%)	(rad.)	(%)	6/8-7/5	(Ma)	(abs)	(Ma)	(abs)	(Ma)	(abs)	(%)		
MCH-17 Ap	2.990	7.99	0.198	2.51	2.07	28.12	0.12	0.04011	1.05	0.25769	4.03	0.04660	3.73	0.407	253.5	2.66	232.8	9.38	28.5	89.0	-8.9		
MCH-17 Ap Leached	2.170	7.10	0.216	1.79	1.40	30.81	0.22	0.03908	0.31	0.27613	3.27	0.05125	3.12	0.514	247.1	0.78	247.6	8.10	252.0	72.0	0.2		

* Denotes radiogenic lead; (a) Decay constants used $^{238}\text{U}=1.55125 \times 10^{-10}$;
 $^{235}\text{U}=9.48485 \times 10^{-10}$; $^{238}\text{U}/^{235}\text{U}=137.88$; Rho: Error correlation.

Sample OC-111

Finally, the gneiss sample OC-111 from the Oaxacan metamorphic complex, yielded an apatite U–Pb age of 972 ± 42 (2σ) Ma that was obtained from only 27 grains (MSWD = 6.8; Figure 4f). These apatites are poor in U (3.3–38.1 ppm, mean = 13.4 and SD = 6.9; Supplementary Appendix A2); but yielded a relatively precise age due to the significant ingrowth of radiogenic Pb, proportional to their “old” ages (Figure 4f). Therefore, some apatites from OC-111 display almost concordant ages, a behavior that was not observed for the younger samples tested (Figures 4a, 4b, 4c, 4d, 4e, and 4f). The high MSWD seems to be related to natural age dispersion (Supplementary Appendix A2; Figure 4f).

The apatite U–Pb age of 972 ± 42 (2σ) Ma obtained for OC-111 is within the age of the granulite facies metamorphism in the Oaxacan Complex, calculated at 990 ± 10 Ma (Solari *et al.*, 2003, 2014). On the other hand, the central age is in line with a few cooling ages reported from the northern Oaxacan Complex including a titanite U–Pb age of 968 ± 9 Ma (2σ) and a phlogopite Ar–Ar age of 945 ± 10 (2σ) Ma (Keppie *et al.*, 2004). Thus, all these cooling ages correspond to the earliest cooling period after the main metamorphic event (i.e., postmetamorphic cooling ages). It can also be proposed that the high dispersion of ages (MSWD = 6.8) may imply that some apatites actually record the age of metamorphism at 1 Ga, whereas some others may display true cooling signals (Figure 4f; see also detailed apatite U–Pb results in Supplementary Appendix A2).

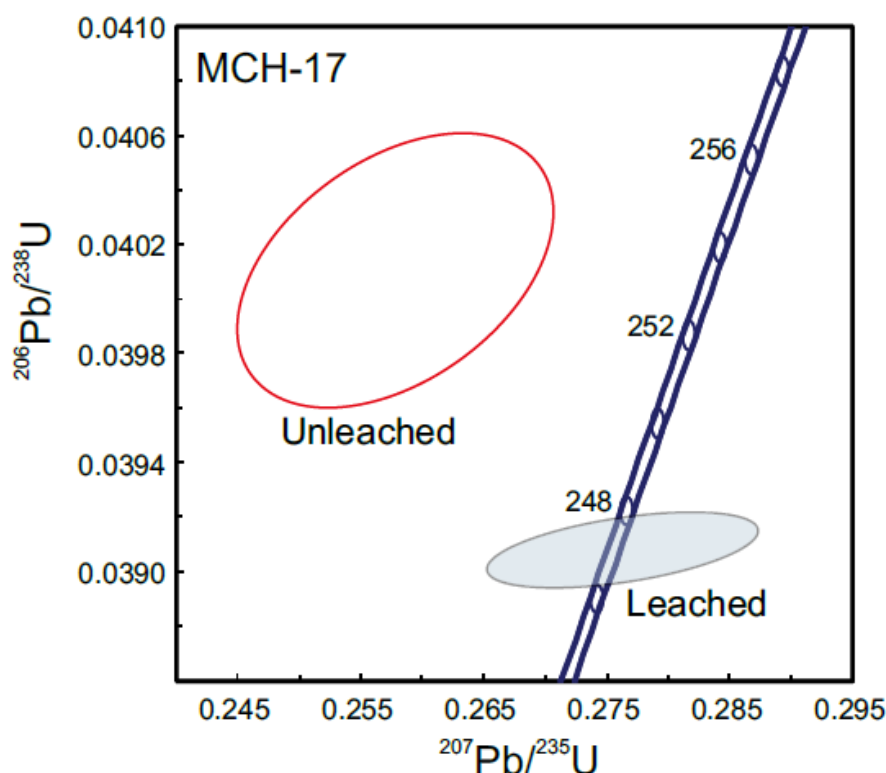


FIGURE 5

U-Pb Concordia plot (Wetherill, 1956) for U-Pb data of apatites from sample MCH-17 obtained with ID-TIMS. Leached aliquot in gray (filled) and unleached aliquot in red.

DISCUSSION

Medium-temperature thermochronology methods (*e.g.*, K-Ar, ArAr, or Rb-Sr in micas, apatite U-Pb, and titanite fission-track dating) can be employed to resolve many fundamental geological problems such as to reconstruct the thermal history of ancient crystalline basements, to obtain the ages of metamorphic events and magmatic or mineralization processes, as well as for provenance analyses (*e.g.*, Schaaf *et al.*, 2002; Keppie *et al.*, 2004; Schoene and Bowring, 2007; DuqueTrujillo *et al.*, 2014; Liu *et al.*, 2014; Bedoya *et al.*, 2017).

Some advantages and disadvantages of ID-TIMS and LA-ICPMS-based apatite U-Pb methods were observed during this study and are described in the following paragraphs. ID-TIMS technique undoubtedly yields high-precision apatite U-Pb results (MCH-17; Table 2), and this is a critical advantage over the LA-ICP-MS-based apatite U-Pb dating. ID-TIMS, thus, is very useful in certain cases; for example, during the search for new apatite U-Pb standards, for the dating of young apatite samples with low radiogenic Pb contents (*e.g.*, apatite from Rupelian-aged ore deposits and Oligocene-Miocene igneous rocks which are widespread in Mexico), as well as the dating of those rocks, in which apatite may contain extremely low U concentrations (*e.g.*, many high-pressure metamorphic or some intermediate to felsic volcanic rocks; see Henrichs *et al.*, 2018). However, there are also inconvenient points of ID-TIMS that should be outlined. First of all, ID-TIMS U-Pb is a time-consuming technique, due to numerous steps in the chemical treatment required for samples to be dated, which may generate a high analytical cost. Secondly, ID-TIMS is a totally destructive dating technique (*i.e.*, aliquots of crystals must be digested entirely; see subsection “ID-TIMS methodology, LUGIS”).

After our observations during the LA-ICP-MS experiments, for most igneous or metamorphic rock samples 40 to 60 single-spot analyses may be recommended to obtain reliable apatite U-Pb results. There are not any significant difficulties for the dating of very old apatite samples, which in any case may contain high concentrations of radiogenic lead generated over a large geological time, and a small number of grains can be sufficient to obtain quite representative U-Pb ages (*e.g.*, 20–30 spots; OC-111). It is more convenient to date isotopically highly heterogeneous apatites, which may yield a perfectly aligned trend in the T–W Discordia line with a statistically very reliable U-Pb date (*e.g.*, see results for Congra-2). To test the heterogeneity of isotopic ratios before LA-ICP-MS sessions, a simple trial can be proposed for apatite. This can be done by etching mounted and polished apatite crystals, a chemical procedure commonly applied to reveal ^{238}U spontaneous fission tracks. For either magmatic or metamorphic apatite samples with monotonic cooling histories for the fission-track system, the heterogeneity of spontaneous fission track densities is directly related to the heterogeneity of U distribution, and consequently may imply a strong heterogeneity of isotope ratios for the U-Pb system. This rapid test can be used to choose more convenient samples for apatite U-Pb dating, especially for the LA-ICP-MS-based technique.

Although apatite U-Pb ages obtained by LA-ICP-MS display larger errors than zircon U-Pb ages also obtained by LA-ICP-MS (*ca.* 2.5–9.1% for 2σ level, this study, *vs. ca.* 2% at 2σ ; *e.g.*, Horstwood *et al.*, 2016), which is mainly due to lower contents of U and thus of radiogenic Pb in apatite, they are typically accurate when there has been no subsequent open system behavior and the obtained lower intercept U-Pb ages are precise enough to be geologically meaningful. It is important to note that the statistics for apatite U-Pb dating by LA-ICP-MS may be improved by employing a larger spot diameter (80 μm or above) combined with a longer data acquisition time (*e.g.*, 55 seconds), as well as by changing other parameters of laser ablation and ICP-MS operation conditions.

A correct estimation of initial $^{207}\text{Pb}/^{206}\text{Pb}$ ratio is also very important to calculate reliable LA-ICP-MS-derived apatite U-Pb ages. There are some advantages of LA-ICP-MS over ID-TIMS. Among them, sample preparation and lower costs. The ability of LA-ICP-MS to measure also REE simultaneously with U, Th, Sr, Y, Mn, Mg, Cl, and isotopic ratios gives complementary information on the chemical composition of apatite that can be used for simultaneous double dating of single grains by U-Pb plus in-situ fission-tracks, as well as for petrogenetic, metallogenetic and provenance studies. Moreover, and this is also an important advantage, LA-ICP-MS is not such destructive technique if compared to ID-TIMS; therefore, mounted crystals already analyzed by LA-ICP-MS may be repeatedly used in further microanalytical experiments.

GENERAL CONCLUSIONS

In this experimental study, we report the first apatite U-Pb data obtained in two Mexican geochronology labs, LUGIS and LEI, both at UNAM, testing this technique on different igneous and metamorphic rocks which have already been dated by conventional radioisotopic methods like U-Pb on zircon, and K-Ar or Ar-Ar in micas and hornblende. The obtained results agree well with the isotopic ages available for the same samples as well as for the same geological units studied. The apatite U-Pb ages are precise enough to be meaningful, and in most of the studied cases they display the igneous crystallization age, suggesting rocks were emplaced at depths with temperature lower than the Pb partial retention zone. Apatite U-Pb dating has many particular advantages as compared to other medium-temperature methods. The results of our study validate the analytical procedures employed and can undoubtedly serve as a crucial starting point towards the development and improvement of medium-temperature thermochronometry tools in Mexico.

SUPPLEMENTARY MATERIAL

Supplementary Appendix A1 "Zircon U–Pb corrected isotopes and ages of samples analyzed by LA–ICP–MS" and Supplementary Appendix A2 "Apatite U–Pb corrected isotopes and ages of samples analyzed by LA–ICP–MS" can be found at the journal web site, in the table of contents of this issue.

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ADDITIONAL INFORMATION

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